# The Lifetime Prevention Schedule 

## Establishing Priorities among Effective Clinical Prevention Services in British Columbia

Summary and Technical Report October 2019 Update

An update of clinically preventable burden and cost-effectiveness estimates for all services reviewed to date.

Update completed by H. Krueger \& Associates Inc.

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## Establishing Priorities among Effective Clinical Prevention Services in British Columbia: 2019 Update

## Executive Summary

## Background

The report, A Lifetime of Prevention, was published by the Clinical Prevention Policy Review Committee (CPPRC) in December of 2009. ${ }^{1}$ A key goal of the CPPRC was to determine which clinical prevention services are worth doing in British Columbia (BC), culminating in a proposed Lifetime Prevention Schedule (LPS). Clinical prevention services were included on the LPS if they were considered to be effective, had a significant positive impact on population health and were cost-effective.

Clinical prevention services (CPS) are defined as:
Manoeuvres pertaining to primary and early secondary prevention (i.e., immunization, screening, counselling and preventive medication/device) offered to the general population (asymptomatic) based on age, sex and risk factors for disease and delivered on a one-provider-to-one-client basis, with two qualifications:
(i) the provider could work as a member of a care team or as part of a system tasked with providing, for instance, a screening service; and
(ii) the client could belong to a small group (e.g. a family, a group of smokers) that is jointly benefiting from the service.

This definition does not refer to the type of provider or the type of funding. This allows for the evaluation of the appropriate implementation of the service as a separate program planning matter.

Since 2009, a total of 29 CPS have been reviewed by the Lifetime Prevention Schedule Expert Committee (LPSEC) for potential inclusion in the LPS. Three new reviews were concluded in 2019, namely, screening for depression in children and youth, screening for osteoporosis to prevent fractures in older women and screening for abdominal aortic aneurysms in older men.

Note that this document has a companion document, the Reference and Key Assumptions Document, in which all key model assumptions are recorded in one location.

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## CPS Intervention Rate

Table ES-1 provides a one-page summary of the 29 CPS reviewed by the LPSEC to date. Included on the table are the relevant cohort and the frequency with which the service is to be provided. In addition, an estimated rate of coverage for the service in British Columbia and the best rate in the world is provided. The three new reviews completed in 2019 are highlighted in yellow.

For example, the best available evidence suggests that screening for colorectal cancer is effective in the general asymptomatic population ages 50 to 74 (the relevant cohort). Ideally, screening should take place every 2 years using a fecal occult blood test (FOBT) or every 10 years using sigmoidoscopy (frequency). An estimated $50 \%$ of the relevant cohort in BC are currently receiving screening at this frequency (rate of coverage in BC). International evidence suggests that this rate could be improved to $76 \%$ (best rate in the world).

Table ES1: Potential Clinical Prevention Services in B.C.
Summary of the Applicable Cohort, Service Frequency and Coverage
Clinical Prevention Services

## Summary of the Clinically Preventable Burden and Cost-Effectiveness

Table ES-2 also provides a one-page summary of the 29 CPS reviewed by the LPSEC to date. Included on this table, however, is information on the clinically preventable burden (CPB) and cost-effectiveness (CE) associated with each of the 29 maneuvers.

CPB is defined as the total quality-adjusted life years that could be gained if the clinical preventive service were delivered at recommended intervals to a BC birth cohort of 40,000 individuals over the years of life that a service is recommended. CE is defined as the average net cost per QALY gained in typical practice by offering the clinical preventive service at recommended intervals to a BC birth cohort over the recommended age range.

The $C P B$ columns identify the clinically preventable burden (in terms of quality adjusted life years or QALYs) that is being achieved in BC based on current coverage, and the potential CPB if the best coverage rate in the world ( BiW ) is achieved. For example, if coverage for colorectal cancer screening were as high as the BiW (76\%), we would expect a CPB of 1,189 QALYs. Since BC's coverage is at $50 \%$, a CPB of 703 QALYs is being achieved. This is 486 QALYs short of the potential 1,189 QALYs achievable based on BiW coverage, as identified in the Gap column.

Note that coverage rates in BC are unknown for 22 of the 29 maneuvers.
The $C E$ columns identify the cost-effectiveness ratio associated with a service stated in terms of the cost per QALY. The ratio is given based on the use of a $1.5 \%$ and a $0 \%$ discount rate. For example, the cost/QALY associated with colorectal cancer screening in BC is estimated at $\$ 47,265$, based on a discount rate of $1.5 \%$. If a $0 \%$ discount rate is used, then the cost/QALY would be reduced to $\$ 44,213$.

## Table ES2: Potential Clinical Prevention Services in B.C.

## Summary of the Clinically Preventable Burden and Cost-Effectiveness

|  | CPB(2) (0\% Discount) |  |  | CE(3) (\% Discount) |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Clinical Prevention Services | B.C. | 'BiW'(1) | Gap | 1.5\% | 0\% |
| Screening for Asymptomatic Disease or Risk Factors - Children/Youth (C/Y) |  |  |  |  |  |
| Vision screening for amblyopia | 23 | 23 | 0 | \$546,597 | \$240,992 |
| Screening for depression (ages 12-18) | Unknown | 222 |  | \$28,215 | \$27,331 |
| Behavioural Counseling Interventions - Children/Youth (C/Y) |  |  |  |  |  |
| Interventions to support breastfeeding | Unknown | 5,002 |  | $(\$ 9,021)$ | $(\$ 11,966)$ |
| Screening for obesity and referral to comprehensive, intensive behavioral intervention to promote improvement in weight status | Unknown | 80 |  | \$77,441 | \$46,302 |
| Preventing tobacco use (school-aged children \& youth) | Unknown | 4,123 |  | $(\$ 7,349)$ | $(\$ 9,538)$ |
| Preventive Medication / Devices - Children |  |  |  |  |  |
| Fluoride varnish | Unknown | 150 |  | \$43,038 | \$43,038 |
| Dental sealants | Unknown | 157 |  | $(\$ 24,690)$ | (\$29,320) |
| Screening for Asymptomatic Disease or Risk Factors - Adults |  |  |  |  |  |
| Screening for breast cancer | 703 | 1,189 | 486 | \$19,720 | \$18,326 |
| Screening (cytology-based) for cervical cancer | 1,153 | 1,471 | 318 | \$25,542 | \$26,980 |
| Addition of HPV-based cervical cancer screening | 0 | 655 | 655 | $(\$ 21,556)$ | $(\$ 19,264)$ |
| Screening for colorectal cancer | 1,141 | 1,734 | 593 | \$47,265 | \$44,213 |
| Screening for lung cancer | Unknown | 1,745 |  | \$2,240 | \$2,080 |
| Screening for hypertension | Unknown | 11,587 |  | \$15,254 | \$10,760 |
| Screening for cardiovascular disease risk and treatment (with statins) | Unknown | 9,370 |  | \$3,223 | \$1,392 |
| Screening for type 2 diabetes mellitus (T2DM) | Unknown | 3,494 |  | (\$3,121) | $(\$ 3,453)$ |
| Screening for depression in general adult population | Unknown | -8 |  | Domi | nated |
| Screening for depression in pregnant and postpartum women | Unknown | 109 |  | \$23,042 | \$10,140 |
| Screening for osteoporosis | Unknown | 91 |  | (\$29,412) | $(\$ 34,145)$ |
| Screening for abdominal aortic aneurysm | Unknown | 340 |  | \$11,995 | \$9,973 |
| Screening for Sexually Transmitted Infections and Blood Borne Pathogens - Adults |  |  |  |  |  |
| Screening for human immunodeficiency virus | Unknown | 360 |  | \$16,434 | \$16,434 |
| Screening for chlamydia and gonorrhea | Unknown | 143 |  | \$57,174 | \$53,410 |
| Screening for hepatitis C virus | 2,695 | 3,920 | 1,225 | \$3,427 | \$2,810 |
| Behavioural Counseling Interventions - Adults |  |  |  |  |  |
| Prevention of sexually transmitted infections (STIs) | Unknown | 3,285 |  | \$10,267 | \$10,267 |
| Counselling and interventions to prevent tobacco use | 3,730 | 5,944 | 2,214 | $(\$ 1,863)$ | $(\$ 3,344)$ |
| Alcohol misuse screening and brief counseling | Unknown | 2,175 |  | \$23,607 | \$16,611 |
| Screening for and management of obesity | Unknown | 2,287 |  | \$12,160 | \$11,140 |
| Preventing falls | Unknown | 429 |  | \$35,213 | \$35,213 |
| Preventive Medication / Devices - Adults <br> Routine aspirin use for the prevention of cardiovascular disease (CVD) and colorectal cancer | Unknown | 1,098 |  | \$2,302 | \$411 |
| Folic acid supplementation for the prevention of neural tube defects | Unknown | 95 |  | \$195,379 | \$113,155 |
| (1) 'BiW' = best in world; (2) CPB = clinically preventable burden; (3) CE = cost-effectiveness |  |  |  |  |  |

## Comparison by Clinically Preventable Burden

Figure ES-1 provides a summary of the CPB associated with each service. Results are displayed based on a $0 \%$ discount rate. Results based on a $1.5 \%$ discount rate are available in the body of the text. Using a $1.5 \%$ discount rate tends to reduce the CPB. The results are organized from left to right based on the services with the highest to lowest potential CPB. For example, full implementation of the service hypertension screening and treatment (Hypertension) (i.e., achieving levels that are comparable to the best in the world) would result in a CPB of 11,587 QALYs, the highest of any service reviewed.

For the seven services for which BC coverage rates are known, we have indicated (by the darker bar insert) what proportion of the potential BiW rate is currently being achieved in BC .

The black bars associated with each service represent a potential range in CPB based on oneway sensitivity analysis. That is, the range is based on varying (over a plausible range) the one assumption that has the largest effect on the results generated by the model.
Simultaneously varying more than one assumption would increase the potential range. A larger range suggests a higher sensitivity to the assumptions used.


Note that the labels on the horizontal axis in Figures ES-1 and ES-2 refer to the CPS included in Table ES-1. The ' A ' refers to adults, the ' C ' to children, the ' $\mathrm{C} / \mathrm{Y}$ ' to children/youth and the ' Ca ' to cancer.

## Comparison by Cost-Effectiveness

Figure ES-2 provides a summary of the CE associated with each service. Results are displayed based on a $1.5 \%$ discount rate. Results based on a $0 \%$ discount rate are available in the body of the text. Using a $0 \%$ discount rate tends to improve the CE. Furthermore, the results are organized from left to right based on the services with the best to worst potential CE, including a plausible range for each service based on sensitivity analysis. Screening for osteoporosis in women $65+$ has the best CE result of any service reviewed. That is, this service is considered to be cost-saving, with a cost per QALY of -\$29,412 (with a potential range from - $\$ 43,257$ to $\$ 38,997$ ).


The base models include an estimate of costs associated with a person's time used in accessing the preventive service. The most significant effect of these inclusions/exclusions is seen in services that require frequent contact with health care providers, such as behavioural counselling to prevent alcohol misuse. For this service, the cost/QALY is reduced from $\$ 23,607$ to $\$ 4,572$ if patient time costs are excluded.

## Combined Comparison Using CPB and CE

The results for CPB and CE are combined in Figure ES-3. CPB is on the vertical axis, ranging from 0 to 12,000 QALYs. CE is on the horizontal axis, ranging from $\$ 100,000 / \mathrm{QALY}$ at the intersection of the x - and y -axis to $-\$ 50,000$ at the far right of the x axis. By arranging CPB and CE in this manner, the most positive results are on the upper right of the chart and the least positive results are in the lower left of the chart. We also divided CPB into three equal segments as follows; 0 to 4,000 QALYs, 4,001 to 8,000 QALYs and 8,001 to 12,000 QALYs. CE was also divided into three equal segments as follows: $\$ 100,000$ to $\$ 50,000$ per QALY, $\$ 50,000$ to $\$ 0$ per QALY and $\$ 0$ to $-\$ 50,000$ per QALY.

The resulting nine equivalent segments are shown in Figure ES-3. Services in the upper right segment have the most favourable combination of CPB and CE while services in the lower left segment have the least favourable combination of CPB and CE.


In Figures ES-4 to ES-6, we have incorporated visual information on plausible ranges (based on one-way sensitivity analysis) with the point estimates for each service. To avoid overcrowding the above figure (ES-3), we have separated the services into three figures. Figure ES-4 includes services specific to children and youth, Figure ES-5 includes screening services and Figure ES-6 includes the remainder of the services reviewed.



## List of Abbreviations

AAA - Abdominal Aortic Aneurysm
AABR - Automated Auditory Brainstem Response
ABR - Auditory Brainstem Response
ACC - American College of Cardiology
AD - Anti-Depressant(s)
AD - Atopic Dermatitis
ADAM - Aneurysm Detection and Management
AHA - American Heart Association
apoB - Apolipoprotein B
ASA - Acetylsalicylic Acid
ASCVD - Atherosclerotic Cardiovascular Disease
AOAE - Automated Otoacoustic Emissions
AUD - Australian Dollars
AUDIT - Alcohol Use Disorders Identification Test
AUGIB - Acute Upper Gastrointestinal Bleeding
BC - British Columbia
BCEHP - British Columbia Early Hearing Program
BD - Biotinidase Deficiency
BDI - Beck Depression Inventory
BiW - Best in the World
BFHI - Baby Friendly Hospital Initiative
BMD - Bone Mineral Density
BMI - Body Mass Index
BMT - Bone Marrow Transplant
CAD - Canadian Dollars
CAGE - Cut Down, Annoyed, Guilty, Eye-Opener
CBT - Cognitive Behavioural Therapy
CCHD - Critical Coronary Heart Disease - also used for Critical Congenital Heart Defects
CCHS - Canadian Community Health Survey
CCS - Canadian Cardiovascular Society
CE - Cost-Effectiveness
CHD - Coronary Heart Disease
CI - Confidence Interval
CIN - Cervical Intraepithelial Neoplasia
CLEM - Cardiovascular Life Expectancy Model

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CMG - Case Mix Group
CPB - Clinically Preventable Burden
CPS - Clinical Prevention Service
CRC - Colorectal Cancer
CSS - Canadian Cardiovascular Society
CSVS - Canadian Society for Vascular Surgery
CTFPHC - Canadian Task Force on Preventive Health Care
CUD - Carnitine Uptake Disorder
CV - Cardiovascular
CVD - Cardiovascular Disease
dB - Decibels
DSM - Diagnostic and Statistical Manual of Mental Disorders
DXA - Dual-Energy X-ray Absorptiometry
ES - Executive Summary
ETS - Environmental Tobacco Smoke
EVAR - Endovascular Aneurysm Repair
FASD - Fetal Alcohol Spectrum Disorder
FDA - Food and Drug Administration (US)
FIT - Fecal Immunochemical Test
FOBT - Fecal Occult Blood Test
FRS - Framingham Heart Study Risk Score
FTE - Full Time Equivalent
gFOBT - Guaiac Fecal Occult Blood Test
GBD study - Global Burden of Disease study
GI - Gastrointestinal
GP - General Practitioner
HDL-C - High-Density Lipoprotein Cholesterol
HMO - Health Maintenance Organization
HPV - Human Papillomavirus
HR - Hazard Ratio
ICD - International Classification of Diseases
IR - Intermediate Risk
IQ - Intelligence Quotient
ISH - Intentional Self-Harm
LEEP - Loop Electrosurgical Excision Procedure
LDL - Low-Density Lipoprotein
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LDL-C - Low-Density Lipoprotein Cholesterol
LHA - Local Health Areas
LRTI - Lower Respiratory Tract Infection
LPS - Lifetime Prevention Schedule
LPSEC - Lifetime Prevention Schedule Expert Committee
MASS - Multicentre Aneurysm Screening Study
MAST - Michigan Alcoholism Screening Test
MDD - Major Depressive Disorder
MEA - Middle Ear Analysis
MSP - Medical Service Plan
NHANES - National Health and Nutrition Examination Survey
NICE - National Institute for Health and Clinical Excellence
NSAID - Nonsteroidal Anti-Inflammatory Drug
NSDUH - National Survey on Drug Use and Health
NTD - Neural Tube Defect
OM - Otitis Media
OME - Otitis Media with Effusion
OR - Odds Ratio
OAE - Otoacoustic Emissions
PCHI - Permanent Childhood Hearing Impairment
PCI - Percutaneous Coronary Intervention
PCP - Primary Care Provider
PDC - Proportion of Days Covered
PHQ-A - Patient Health Questionnaire for Adolescents
PHSA - Provincial Health Services Authority
POS - Pulse Oximetry Screening
PSBC - Perinatal Services British Columbia
QALY - Quality-Adjusted Life-Year
QoL - Quality of life
RCT - Randomized Controlled Trial
RR - Relative Risk
SCID - Severe Combined Immune Deficiency
SF-36 - Short Form (Health Survey) with 36 items
SIDS - Sudden Infant Death Syndrome
TC - Total Cholesterol
TEOAE -Transient Evoked Otoacoustic Emissions

TG - Triglycerides
TREC - T-cell Receptor Excision Circles
UK - United Kingdom
UKSAT - United Kingdom Small Aneurysm Trial
UNHS - Universal Newborn Hearing Screening
US - United States
USD - United States Dollars
USPSTF - United States Preventive Services Task Force
WHO - World Health Organization

## Clinical Prevention in Children and Youth

## Screening for Asymptomatic Disease or Risk Factors

## Vision Screening for Amblyopia

## United States Preventive Service Task Force Recommendations (2011)

Approximately $2 \%$ to $4 \%$ of preschool aged children have amblyopia, an alteration in the visual neural pathway in the developing brain that can lead to permanent vision loss in the affected eye. Amblyopia usually occurs unilaterally but can occur bilaterally. Identification of vision impairment before school entry could help identify children who may benefit from early interventions to correct or to improve vision.
The USPSTF recommends vision screening for all children at least once between the ages of 3 and 5 years, to detect the presence of amblyopia or its risk factors (grade B recommendation).

The USPSTF concludes that the current evidence is insufficient to assess the balance of benefits and harms of vision screening for children <3 years of age (I statement). ${ }^{2}$

## Canadian Task Force on Preventive Health Care Recommendations (1990)

In the 1990 publication on well-baby care in the first 2 years of life, the CTFPHC recommended that there was good evidence to include repeated examination of the eyes and hearing during the first year of life in the periodic health examination. This was given an ' A ' recommendation. ${ }^{3}$ Based on this information, vision screening was included in the BC Lifetime Prevention Schedule. ${ }^{4}$

## Canadian Task Force on Preventive Health Care Recommendations (1994)

Once detected, simple refractive errors affecting visual acuity are readily treatable with eye glasses. However, evidence for the treatment of amblyopia is more controversial and inconclusive. It is widely held that for any potential benefit to be realized, amblyopia must be detected during the "sensitive" period, i.e. between birth and about the seventh year.

Systematic screening for visual deficits has been found to decrease prevalence later.
Fair evidence for inclusion in periodic health examination (B Recommendation). ${ }^{5}$

## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with screening all children at least once between the ages of 3 and 5 years, to detect the presence of amblyopia or its risk factors.

[^1]In modelling CPB, we made the following assumptions:

- $99.57 \%$ of individuals in a birth cohort of 40,000 would survive to age 4 , based on data from the BC life tables for 2010 to 2012.
- Estimates of the prevalence of amblyopia ('lazy eye') range from $2.9 \%{ }^{6}$ to $4.8 \%$. $^{7}$ We used the mid-point of this range ( $3.85 \%$ ) for the base case (Table 1, row $c$ ) and the range in sensitivity analysis.
- We assumed that $70 \%$ of children with amblyopia would be asymptomatic. That is, $30 \%$ would be symptomatic and would thus be detected without the need for screening (Table 1, row e). ${ }^{8}$
- We assumed an average life expectancy for a 4 year-old of 78.6 years (Table 1, row $g$ ), based on data from the BC life tables for 2010 to 2012.
- The annual incidence of permanent visual impairment or blindness attributable to loss of vision in the non-amblyopic eye has been estimated at .00004 (.00001 to 0.00006 ) during the ages of 5 to 15 years, 0.00005 ( 0.00004 to 0.00007 ) for ages 16 to 64 and $0.00046\left(0.00039\right.$ to 0.00052 ) for ages $65+{ }^{9}$ (Table 1 , row $h, i$ and $j$ ). In screening a cohort of 40,000 , we would expect to find 1,073 four-year olds with amblyopia. Of these, approximately 10 would be expected to have permanent visual impairment or blindness attributable to loss of vision in the non-amblyopic eye. Most of this visual impairment /blindness ( $64 \%$ ) would occur after age 65.
- The effectiveness of interventions in improving amblyopia is fairly contentious. The USPSTF noted an average improvement of approximately one line on the Snellen eye chart. ${ }^{10}$ Others suggest a clinically significant improvement resulting from treatment of between $26 \%$ and $75 \% .^{11,12}$ We have used the mid-point of this range ( $51 \%$ ) in our base model and the range in sensitivity analysis (Table 1, row $m$ ).
- Other assumptions used in assessing the clinically preventable burden are detailed in the Reference Document.

Based on these assumptions, the CPB associated with screening for amblyopia in children ages 3 to 5 is 23.0 QALYs (Table 1 , row $n$ ).

[^2]Table 1: CPB of Screening for Amblyopia in 3-5 Year-Olds in a Birth Cohort of 40,000 (B.C.)

| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | \% survival at age 4 | 0.9957 | $\checkmark$ |
| b | 4 Year olds in cohort | 39,828 | $=a * 40,000$ |
| C | Prevalence of amblyopia | 3.85\% | $\checkmark$ |
| d | 4 year-olds with amblyopia in birth cohort | 1,533 | $=\mathrm{b}^{*} \mathrm{c}$ |
| e | \% of amblyopia that are undetected (asymptomatic) | 70\% | $\checkmark$ |
| f | 4 year-olds with amblyopia in birth cohort detected through screening | 1,073 | $=d^{*} \mathrm{e}$ |
| g | Average life expectancy of a 4 year old | 78.6 | Ref Doc |
| h | Incidence of permanent visual impairment or blindness -5-15 yrs | 0.00004 | $\checkmark$ |
| i | Incidence of permanent visual impairment or blindness - 16-64 yrs | 0.00005 | $\checkmark$ |
| j | Incidence of permanent visual impairment or blindness - 65+ yrs | 0.00046 | $\checkmark$ |
| k | Change in QoL associated with permanent visual impairment or blindness | 0.187 | Ref Doc |
| 1 | Estimated QALYs lost | 45.6 | Calculated |
| m | Effectiveness of intervention | 51\% | $\checkmark$ |
| n | QALYs gained, CPB | 23.0 | $=1 * \mathrm{~m}$ |

$\checkmark=$ Estimates from the literature
We also modified several major assumptions and recalculated the CPB as follows:

- Assume the prevalence of amblyopia is reduced from $3.85 \%$ to $2.9 \%$ (Table 1, row c): $\mathrm{CPB}=17.5$
- Assume the prevalence of amblyopia is increased from $3.85 \%$ to $4.8 \%$ (Table 1, row c): $\mathrm{CPB}=29.0$
- Assume the effectiveness of interventions in improving amblyopia is reduced from $51 \%$ to $26 \%$ (Table 1, row m): $\mathrm{CPB}=11.9$
- Assume the effectiveness of interventions in improving amblyopia is increased from $51 \%$ to $75 \%$ (Table 1, row m): $\mathrm{CPB}=34.2$
- Assume the incidence of permanent visual impairment or blindness is at the low end of the range (Table 1 , rows $\mathrm{h}, \mathrm{i}, \mathrm{j}$ ): $\mathrm{CPB}=17.0$
- Assume the incidence of permanent visual impairment or blindness is at the high end of the range (Table 1 , rows $\mathrm{h}, \mathrm{i}, \mathrm{j}$ ): $\mathrm{CPB}=30.2$
- Assume the disutility associated with permanent visual impairment or blindness is reduced from -0.187 to -0.124 (Table 1, row k): $\mathrm{CPB}=15.3$
- Assume the disutility associated with permanent visual impairment or blindness is increased from -0.187 to -0.260 (Table 1, row k): $\mathrm{CPB}=32.0$


## Modelling Cost-Effectiveness

In this section, we will calculate the CE associated with screening all children at least once between the ages of 3 and 5 years, to detect the presence of amblyopia or its risk factors.

In modelling $C E$, we made the following assumptions:

- The estimated cost of screening (Table 2, row $d$ ) and interventions (Table 2, row $g$ ) are based on information in the economic evaluation by Carlton et al. ${ }^{13}$ The cost of screening is estimated at $12.90(95 \%$ CI of 8.38 to 18.38$)$ in 2006 British Pounds Sterling ( $£$ ) or $\$ 27.56$ ( $95 \%$ CI of $\$ 17.90$ to $\$ 39.26$ ) in 2017 CAD. The cost of an intervention is estimated at $1,015(95 \%$ CI of 907 to 1,122$)$ in 2006 British Pounds Sterling (£) or $\$ 2,168(95 \%$ CI of $\$ 1,938$ to $\$ 2,397)$ in 2017 CAD.
- For patient time and travel costs, we estimated two hours of patient time required per physician visit.
- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the CE associated with screening for amblyopia in children ages 3 to 5 is $\$ 546,597$ per QALY (Table 2, row $n$ ).

| Table 2: CE of Screening for Amblyopia in 3-5 Year-Olds in a Birth Cohort of |  |  |  |
| :---: | :---: | :---: | :---: |
| 40,000 (B.C.) |  |  |  |
| Row Label | Variable | Base Case | Data Source |
| a | 4 Year olds in cohort | 39,828 | Table 1 row b |
| b | Screening rate | 93\% | Ref Doc |
| c | \# of screens | 37,040 | $=\mathrm{a}$ * b |
|  | Costs of screening |  |  |
| d | Estimated screening cost | \$27.56 | $\checkmark$ |
| e | Value of patient time and travel for office visit | \$59.38 | Ref Doc |
| f | Cost of screening over lifetime of birth cohort | \$3,220,261 | $=c^{*}(\mathrm{~d}+\mathrm{e})$ |
|  | Costs of interventions |  |  |
| g | Estimated intervention cost | \$2,168 | $\checkmark$ |
| h | \# of interventions | 1,073 | Table 1 row f |
| i | Total cost over lifetime of birth cohort | \$2,327,506 | = $\mathrm{g}^{*} \mathrm{~h}$ |
|  | CE calculation |  |  |
| j | Cost of screening over lifetime of birth cohort | \$3,220,261 | = f |
| k | Costs of intervention | \$2,327,506 | = i |
| I | QALYs saved (0\% discount rate) | 23.0 | Table 1 row n |
| m | QALYs saved (1.5\% discount rate) | 10.1 | Calculated |
| n | CE (\$/QALY saved) | \$546,597 | $=(j+k) / m$ |

$\checkmark=$ Estimates from the literature
We also modified several major assumptions and recalculated the cost per QALY as follows:

- Assume the prevalence of amblyopia is reduced from $3.85 \%$ to $2.9 \%$ (Table 1, row c): $\mathrm{CE}=\$ 650,532$

[^3]- Assume the prevalence of amblyopia is increased from 3.85\% to $4.8 \%$ (Table 1, row c): $\mathrm{CE}=\$ 483,802$
- Assume the effectiveness of interventions in improving amblyopia is reduced from $51 \%$ to $26 \%$ (Table 1, row m): $\mathrm{CE}=\$ 1,061,659$
- Assume the effectiveness of interventions in improving amblyopia is increased from $51 \%$ to $75 \%$ (Table 1, row m): $\mathrm{CE}=\$ 368,042$
- Assume the incidence of permanent visual impairment or blindness is at the low end of the range (Table 1, rows $\mathrm{h}, \mathrm{i}, \mathrm{j}$ ): $\mathrm{CE}=\$ 766,266$
- Assume the incidence of permanent visual impairment or blindness is at the high end of the range (Table 1, rows $\mathrm{h}, \mathrm{i}, \mathrm{j}$ ): $\mathrm{CE}=409,817$
- Assume the disutility associated with permanent visual impairment or blindness is reduced from -0.187 to -0.124 (Table 1, row k ): $\mathrm{CE}=\$ 824,303$
- Assume the disutility associated with permanent visual impairment or blindness is increased from -0.187 to -0.260 (Table 1, row k): $\mathrm{CE}=\$ 393,129$
- Assume the screening cost is reduced from $\$ 27.56$ per screen to $\$ 17.90$ (Table 2, row b): $\mathrm{CE}=\$ 511,355$
- Assume the screening cost is increased from $\$ 27.56$ per screen to $\$ 39.26$ (Table 2, row $b$ ): $\mathrm{CE}=\$ 589,300$
- Assume the cost per intervention is reduced from $\$ 2,168$ to $\$ 1,938$ (Table 2, row $f$ ): $\mathrm{CE}=\$ 522,196$
- Assume the cost per intervention is increased from $\$ 2,168$ to $\$ 2,397$ (Table 2, row $f$ ): $\mathrm{CE}=\$ 570,771$


## Summary

| Table 3: Screening for Amblyopia in 3-5 Year-Olds in a Birth Cohort of 40,000 |  |  |  |
| :---: | :---: | :---: | :---: |
| Summary |  |  |  |
| $\begin{aligned} & \text { Base } \\ & \text { Case } \\ & \hline \end{aligned}$ |  | Range |  |
| CPB (Potential QALYs Gained) |  |  |  |
| Assume No Current Service |  |  |  |
| 1.5\% Discount Rate | 10.1 | 5.2 | 15.1 |
| 3\% Discount Rate | 4.6 | 2.4 | 6.8 |
| 0\% Discount Rate | 23.0 | 11.9 | 34.2 |
| CE (\$/QALY) including patient time costs |  |  |  |
| 1.5\% Discount Rate | \$546,597 | \$368,042 | \$1,061,659 |
| 3\% Discount Rate | \$1,213,089 | \$816,814 | \$2,356,193 |
| 0\% Discount Rate | \$240,992 | \$162,268 | \$468,081 |
| CE (\$/QALY) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | \$329,896 | \$222,130 | \$640,579 |
| 3\% Discount Rate | \$732,155 | \$492,984 | \$1,422,069 |
| 0\% Discount Rate | \$145,450 | \$97,936 | \$282,508 |

## Screening for Major Depressive Disorder in Youth

## United States Preventive Services Task Force Recommendations ${ }^{14}$

This recommendation applies to children and adolescents aged 18 years or younger who do not have a diagnosis of MDD [major depressive disorder].

The USPSTF recommends screening for MDD in adolescents aged 12 to 18 years. Screening should be implemented with adequate systems in place to ensure accurate diagnosis, effective treatment, and appropriate follow-up. (B recommendation)

The USPSTF concludes that the current evidence is insufficient to assess the balance of benefits and harms of screening for MDD in children aged 11 years or younger. (I statement)

## Canadian Task Force on Preventive Health Care Recommendations

The CTFPHC does not have a specific recommendation on depression screening for children or adolescents. ${ }^{15}$

## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with screening for MDD in adolescents ages 12 to 18 .

In modelling CPB, we made the following assumptions:

- The USPSTF "found no evidence on appropriate or recommended screening intervals, and the optimal interval is unknown...opportunistic screening may be appropriate for adolescents, who may have infrequent health care visits. ${ }^{16}$ For adolescents with risk factors for MDD, "repeated screening may be most productive. ${ }^{17}$
- Rand and colleagues evaluated primary care visits by US adolescents and found that many did not have any primary care visits during a 12-month period. ${ }^{18}$ Averaging the data presented for the relevant $12-18$ year old group, $56.9 \%$ had a primary care visit during the last 12 -month period.
- Skehar and colleagues found that adolescents $12-14$ years old who were continuously enrolled in private insurance in the US made an average of 0.58 wellcare visits per year. ${ }^{19}$

[^4]- Using data provided by the BC Ministry of Health, Health Sector Information, Analysis and Reporting Division ${ }^{20}$ we were able to generate BC-specific rates of primary care visits and average visits per year for the fiscal years ending in 2012/13 to 2016/17, in total and by sex, as shown in Table 1 below.
- For the five years considered, the average proportion of adolescents ages 10-19 visiting a GP is $70 \%$, and the average number of GP visits per adolescent is 2.07 per year. The proportion of males visiting a GP was $65.4 \%$ and for females it was $75.0 \%$. The average number of visits per male in the population was 1.75 and for females was 2.42 .

Table 1: General Practitioner Visits by Adolescents
British Columbia, 2012/13 to 2016/17

| Age Group | Population in Each Age Group |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | 2012/13 | 2013/14 | 2014/15 | 2015/16 | 2016/17 | Total |
| 10-14 | 234,780 | 231,544 | 230,178 | 230,177 | 232,010 | 1,158,689 |
| 15-19 | 284,482 | 282,214 | 279,997 | 276,909 | 272,677 | 1,396,279 |
| Total | 519,262 | 513,758 | 510,175 | 507,086 | 504,687 | 2,554,968 |
|  | Number of Unique Individuals with GP Visit |  |  |  |  |  |
| 10-14 | 163,332 | 160,912 | 158,653 | 160,260 | 159,826 | 802,983 |
| 15-19 | 205,821 | 200,410 | 196,629 | 192,566 | 189,547 | 984,973 |
| Total | 369,153 | 361,322 | 355,282 | 352,826 | 349,373 | 1,787,956 |
|  | Proportion of Individuals with a GP Visit |  |  |  |  |  |
| 10-14 | 69.6\% | 69.5\% | 68.9\% | 69.6\% | 68.9\% | 69.3\% |
| 15-19 | 72.3\% | 71.0\% | 70.2\% | 69.5\% | 69.5\% | 70.5\% |
| Total | 71.1\% | 70.3\% | 69.6\% | 69.6\% | 69.2\% | 70.0\% |
|  | Number of GP Visits |  |  |  |  |  |
| 10-14 | 429,881 | 422,188 | 412,182 | 413,411 | 407,442 | 2,085,104 |
| 15-19 | 681,806 | 659,038 | 641,316 | 619,790 | 601,925 | 3,203,875 |
| Total | 1,111,687 | 1,081,226 | 1,053,498 | 1,033,201 | 1,009,367 | 5,288,979 |
|  | GP Visits per Individual in Total Population |  |  |  |  |  |
| 10-14 | 1.83 | 1.82 | 1.79 | 1.80 | 1.76 | 1.80 |
| 15-19 | 2.40 | 2.34 | 2.29 | 2.24 | 2.21 | 2.29 |
| Total | 2.14 | 2.10 | 2.06 | 2.04 | 2.00 | 2.07 |

[^5]Table 1: General Practitioner Visits by Adolescents British Columbia, 2012/13 to 2016/17

Males

| Age Group | Population in Each Age Group |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | 2012/13 | 2013/14 | 2014/15 | 2015/16 | 2016/17 | Total |
| 10-14 | 121,031 | 119,378 | 118,720 | 118,572 | 119,586 | 597,287 |
| 15-19 | 149,279 | 147,563 | 145,417 | 143,117 | 140,451 | 725,827 |
| Total | 270,310 | 266,941 | 264,137 | 261,689 | 260,037 | 1,323,114 |
|  | Number of Unique Males with GP Visit |  |  |  |  |  |
| 10-14 | 82,970 | 81,960 | 80,756 | 81,067 | 80,862 | 407,615 |
| 15-19 | 95,992 | 93,224 | 91,170 | 89,118 | 87,596 | 457,100 |
| Total | 178,962 | 175,184 | 171,926 | 170,185 | 168,458 | 864,715 |
|  | Proportion of Males with a GP Visit |  |  |  |  |  |
| 10-14 | 68.6\% | 68.7\% | 68.0\% | 68.4\% | 67.6\% | 68.2\% |
| 15-19 | 64.3\% | 63.2\% | 62.7\% | 62.3\% | 62.4\% | 63.0\% |
| Total | 66.2\% | 65.6\% | 65.1\% | 65.0\% | 64.8\% | 65.4\% |
|  | Number of GP Visits |  |  |  |  |  |
| 10-14 | 215,841 | 211,444 | 206,909 | 206,013 | 202,386 | 1,042,593 |
| 15-19 | 270,303 | 259,637 | 253,874 | 244,381 | 238,257 | 1,266,452 |
| Total | 486,144 | 471,081 | 460,783 | 450,394 | 440,643 | 2,309,045 |
|  | GP Visits per Male in Total Population |  |  |  |  |  |
| 10-14 | 1.78 | 1.77 | 1.74 | 1.74 | 1.69 | 1.75 |
| 15-19 | 1.81 | 1.76 | 1.75 | 1.71 | 1.70 | 1.74 |
| Total | 1.80 | 1.76 | 1.74 | 1.72 | 1.69 | 1.75 |

Table 1: General Practitioner Visits by Adolescents British Columbia, 2012/13 to 2016/17

Females

| Age Group | Population in Each Age Group |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | 2012/13 | 2013/14 | 2014/15 | 2015/16 | 2016/17 | Total |
| 10-14 | 113,749 | 112,166 | 111,458 | 111,605 | 112,424 | 561,402 |
| 15-19 | 135,203 | 134,651 | 134,580 | 133,792 | 132,226 | 670,452 |
| Total | 248,952 | 246,817 | 246,038 | 245,397 | 244,650 | 1,231,854 |
|  | Number of Unique Females with GP Visit |  |  |  |  |  |
| 10-14 | 80,381 | 78,955 | 77,909 | 79,202 | 78,985 | 395,432 |
| 15-19 | 109,865 | 107,210 | 105,496 | 103,488 | 101,995 | 528,054 |
| Total | 190,246 | 186,165 | 183,405 | 182,690 | 180,980 | 923,486 |
|  | Proportion of Females with a GP Visit |  |  |  |  |  |
| 10-14 | 70.7\% | 70.4\% | 69.9\% | 71.0\% | 70.3\% | 70.4\% |
| 15-19 | 81.3\% | 79.6\% | 78.4\% | 77.3\% | 77.1\% | 78.8\% |
| Total | 76.4\% | 75.4\% | 74.5\% | 74.4\% | 74.0\% | 75.0\% |
|  | Number of GP Visits |  |  |  |  |  |
| 10-14 | 214,033 | 210,738 | 205,270 | 207,393 | 205,052 | 1,042,486 |
| 15-19 | 411,487 | 399,386 | 387,411 | 375,393 | 363,660 | 1,937,337 |
| Total | 625,520 | 610,124 | 592,681 | 582,786 | 568,712 | 2,979,823 |
|  | GP Visits per Female in Total Population |  |  |  |  |  |
| 10-14 | 1.88 | 1.88 | 1.84 | 1.86 | 1.82 | 1.86 |
| 15-19 | 3.04 | 2.97 | 2.88 | 2.81 | 2.75 | 2.89 |
| Total | 2.51 | 2.47 | 2.41 | 2.37 | 2.32 | 2.42 |

Source: BC Ministry of Health, Health Sector Information, Analysis and Reporting Division
Calculations by H. Krueger \& Associates, Inc.

- In our model, we assume a maximum (best in the world) adolescent depression screening rate of $7.4 \%\left(10.6 \%^{21}\right.$ times $\left.70.0 \%\right)$ and that screening for this $7.4 \%$ of adolescents (Table 6, row $a h$ ) is completed at each well-care visit, or 2.07 times per year (Table 6, row $a g$ ), ${ }^{22}$ during the seven years of an adolescent's life between 12 and 18 years of age.
- In our model for males, we assume a maximum (best in the world) depression screening rate of $6.9 \%\left(10.6 \%{ }^{23}\right.$ times $\left.65.4 \%\right)$ and that screening for this $6.9 \%$ of male adolescents (Table 6a, row $a h$ ) is completed at each well-care visit, or 1.75 times per year (Table 6a, row $a g$ ), ${ }^{24}$ during the seven years of an adolescent's life between 12 and 18 years of age.
- In our model for females, we assume a maximum (best in the world) depression screening rate of $8.0 \%\left(10.6 \%{ }^{25}\right.$ times $\left.75.0 \%\right)$ and that screening for this $8.0 \%$ of female adolescents (Table 6b, row ah) is completed at each well-care visit, or 2.42 times per year (Table 6 b, row $a g$ ), ${ }^{26}$ during the seven years of an adolescent's life between 12 and 18 years of age.
- Patten et al. estimate that, for the Canadian population aged 15-25, the annual prevalence of MDD was $5.0 \%$ ( $95 \%$ CI $4.2 \%-5.7 \%$ ) and the lifetime prevalence was $8.8 \%$ ( $95 \%$ CI $7.9 \%-9.7 \%$ ). ${ }^{27}$
- Avenevoli et al. report that the annual and lifetime prevalence of MDD in $13-18$ year olds in the US is $7.5 \%$ and $11.0 \%$ respectively. ${ }^{28}$
- Using data from the US National Survey on Drug Use and Health (NSDUH), Mojtabai and colleagues found that the annual prevalence of MDD in the US has increased from $5.6 \%$ in 2005 to $7.2 \%$ in 2014 for 12-13 year olds, $9.1 \%$ to $11.8 \%$ in $14-15$ year olds and $11.2 \%$ to $14.7 \%$ in 16-17 year olds. ${ }^{29}$
- Vasiliadis and colleagues found that there was no significant difference between Canadian and US rates of depression and subsequent use of mental health services. ${ }^{30}$

[^6]- Using the detailed data tables publicly available from the US NSDUH, we calculated the aggregate rates of 12-month major depressive episodes for the years 2014 (the end of Mojtabai and colleague's data) through 2017, using the tables from $2015^{31}$ (containing data for 2014 and 2015) and $2017^{32}$ (containing data for 2016 and 2017), splitting the results by age and sex. The results, shown in Table 2, indicate a substantial difference in major depressive episodes between the sexes, with the annual prevalence of MDE being consistently lower in males than females.
- Similar overall data to the US NSDUH has been reported in the McCreary Centre's Balance and Connection in BC report summarizing the results of the 2018 BC Adolescent Health Survey. Adolescents in grades 7 through 12 were surveyed and $10 \%$ of males reported "mental health conditions", while $20 \%$ of females reported the same. ${ }^{33}$

[^7]Table 2: (US) National Survey on Drug Use and Health 12-Month MDE Events, By Age and Sex

2014-2017 Results

| 12 Year Olds |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Male |  |  | Female |  |  | Calculated Total |  |  |
| Year | Sample Size | MDE \% | MDE ( n ) | Sample Size | MDE \% | MDE ( n ) | Sample Size | MDE \% | MDE ( n ) |
| 2014 | 1,347 | 2.8\% | 38 | 1,293 | 8.9\% | 115 | 2,640 | 5.8\% | 153 |
| 2015 | 1,346 | 2.2\% | 30 | 1,307 | 8.7\% | 114 | 2,653 | 5.4\% | 143 |
| 2016 | 1,323 | 3.1\% | 41 | 1,291 | 6.9\% | 89 | 2,614 | 5.0\% | 130 |
| 2017 | 1,329 | 2.7\% | 36 | 1,269 | 7.0\% | 89 | 2,598 | 4.8\% | 125 |
| Total | 5,345 | 2.7\% | 144 | 5,160 | 7.9\% | 407 | 10,505 | 5.2\% | 551 |
| 13 Year Olds |  |  |  |  |  |  |  |  |  |
|  |  | Male |  |  | Female |  |  | ulated To |  |
|  | Sample Size | MDE \% | MDE ( n ) | Sample Size | MDE \% | MDE ( n ) | Sample Size | MDE \% | MDE ( n ) |
| 2014 | 1,433 | 3.9\% | 56 | 1,388 | 13.8\% | 192 | 2,821 | 8.8\% | 247 |
| 2015 | 1,428 | 3.9\% | 56 | 1,394 | 16.8\% | 234 | 2,822 | 10.3\% | 290 |
| 2016 | 1,479 | 3.8\% | 56 | 1,414 | 15.3\% | 216 | 2,893 | 9.4\% | 273 |
| 2017 | 1,507 | 3.6\% | 54 | 1,423 | 14.5\% | 206 | 2,930 | 8.9\% | 261 |
| Total | 5,847 | 3.8\% | 222 | 5,619 | 15.1\% | 848 | 11,466 | 9.3\% | 1,070 |
| 14 Year Olds |  |  |  |  |  |  |  |  |  |
|  |  | Male |  |  | Female |  |  | ulated To |  |
|  | Sample Size | MDE \% | MDE ( n ) | Sample Size | MDE \% | MDE ( n ) | Sample Size | MDE \% | MDE ( n ) |
| 2014 | 1,491 | 4.6\% | 69 | 1,443 | 17.1\% | 247 | 2,934 | 10.7\% | 315 |
| 2015 | 1,491 | 4.1\% | 61 | 1,411 | 19.0\% | 268 | 2,902 | 11.3\% | 329 |
| 2016 | 1,484 | 5.2\% | 77 | 1,432 | 20.5\% | 294 | 2,916 | 12.7\% | 371 |
| 2017 | 1,492 | 5.2\% | 78 | 1,385 | 19.0\% | 263 | 2,877 | 11.8\% | 341 |
| Total | 5,958 | 4.8\% | 284 | 5,671 | 18.9\% | 1,072 | 11,629 | 11.7\% | 1,356 |
| 15 Year Olds |  |  |  |  |  |  |  |  |  |
|  |  | Male |  |  | Female |  |  | ulated To |  |
| Year | Sample Size | MDE \% | MDE ( n ) | Sample Size | MDE \% | MDE ( n ) | Sample Size | MDE \% | MDE ( n ) |
| 2014 | 1,483 | 5.5\% | 82 | 1,451 | 20.7\% | 300 | 2,934 | 13.0\% | 382 |
| 2015 | 1,438 | 5.3\% | 76 | 1,486 | 26.7\% | 397 | 2,924 | 16.2\% | 473 |
| 2016 | 1,512 | 6.5\% | 98 | 1,498 | 21.0\% | 315 | 3,010 | 13.7\% | 413 |
| 2017 | 1,460 | 7.4\% | 108 | 1,427 | 27.2\% | 388 | 2,887 | 17.2\% | 496 |
| Total | 5,893 | 6.2\% | 364 | 5,862 | 23.9\% | 1,400 | 11,755 | 15.0\% | 1,764 |
| 16 Year Olds |  |  |  |  |  |  |  |  |  |
|  |  | Male |  |  | Female |  |  | ulated To |  |
|  | Sample Size | MDE \% | MDE ( n ) | Sample Size | MDE \% | MDE ( n ) | Sample Size | MDE \% | MDE ( n ) |
| 2014 | 1,467 | 7.5\% | 110 | 1,469 | 20.7\% | 304 | 2,936 | 14.1\% | 414 |
| 2015 | 1,459 | 9.9\% | 144 | 1,384 | 22.3\% | 309 | 2,843 | 15.9\% | 453 |
| 2016 | 1,487 | 9.4\% | 140 | 1,409 | 25.8\% | 364 | 2,896 | 17.4\% | 503 |
| 2017 | 1,508 | 9.8\% | 148 | 1,389 | 24.1\% | 335 | 2,897 | 16.7\% | 483 |
| Total | 5,921 | 9.2\% | 542 | 5,651 | 23.2\% | 1,311 | 11,572 | 16.0\% | 1,853 |
| 17 Year Olds |  |  |  |  |  |  |  |  |  |
|  | Male |  |  | Female |  |  | Calculated Total |  |  |
|  | Sample Size | MDE \% | $\operatorname{MDE}(\mathrm{n})$ | Sample Size | MDE \% | MDE ( n ) | Sample Size | MDE \% | $\operatorname{MDE}(\mathrm{n})$ |
| 2014 | 1,392 | 9.7\% | 135 | 1,350 | 21.0\% | 284 | 2,742 | 15.3\% | 419 |
| 2015 | 1,434 | 9.1\% | 130 | 1,333 | 21.5\% | 287 | 2,767 | 15.1\% | 417 |
| 2016 | 1,415 | 9.7\% | 137 | 1,337 | 24.7\% | 330 | 2,752 | 17.0\% | 467 |
| 2017 | 1,419 | 11.6\% | 165 | 1,418 | 25.5\% | 362 | 2,837 | 18.5\% | 526 |
| Total | 5,660 | 10.0\% | 567 | 5,438 | 23.2\% | 1,262 | 11,098 | 16.5\% | 1,829 |

Source for Sample Size and MDE \%: National Survey on Drug Use and Health, 2014-2017
Calculations by H. Krueger \& Associates, Inc.

- Based on the data in Table 2, we assume an annual prevalence of MDD of $5.2 \%$ in 12 year olds (Table 6, row b), $7.9 \%$ in 12 year old females (Table 6b, row $b$ ) and $2.7 \%$ in 12 year old males (Table 6a, row $b$ ).
- We assume an annual prevalence of MDD of $9.3 \%$ in 13 year olds (Table 6 , row $f$ ), $15.1 \%$ in 13 year old females (Table 6b, row $f$ ) and $3.8 \%$ in 13 year old males (Table 6 a , row $f$ ).
- We assume an annual prevalence of MDD of $11.7 \%$ in 14 year olds (Table 6, row $j$ ), $18.9 \%$ in 14 year old females (Table 6b, row $j$ ) and $4.8 \%$ in 14 year old males (Table 6 a , row $j$ ).
- We assume an annual prevalence of MDD of $15.0 \%$ in 15 year olds (Table 6, row $n$ ), $23.9 \%$ in 15 year old females (Table 6 b row $n$ ) and $6.2 \%$ in 15 year old males (Table 6a, row $n$ ).
- We assume an annual prevalence of MDD of $16.0 \%$ in 16 year olds (Table 6, row $r$ ), $23.2 \%$ in 16 year old females (Table 6 b row $r$ ) and $9.2 \%$ in 16 year old males (Table 6 a, row $r$ ).
- We assume an annual prevalence of MDD of $16.5 \%$ in 17 and 18 year olds (Table 6, row $v$ ), $23.2 \%$ in 17 and 18 year old females (Table 6 b row $v$ ) and $10.0 \%$ in 17 and 18 year old males (Table 6a, row $v$ ).
- In 2017, $17.2 \%$ of US high school students had seriously considered attempting suicide during the previous 12 months, $13.6 \%$ had made a plan about how they would attempt suicide, $7.4 \%$ had actually attempted suicide and $2.4 \%$ had made a suicide attempt resulting in an injury, poisoning or overdose that had to be treated by a doctor or nurse. ${ }^{34}$
- In BC in 2013, 12.2\% of students in grades 7-12 had seriously considered attempting suicide during the previous 12 months and $6.2 \%$ had actually attempted suicide. ${ }^{35}$
- Suicide mortality among youth ages $15-19$ in BC between 2011 and 2013 is 4.7 / 100,000 population. ${ }^{36}$
- The ratio of attempted suicides to completed suicides among adolescents is estimated to be $50: 1$ to $100: 1 .{ }^{37}$
- Rohde and colleagues report that $19 \%$ ( $95 \%$ CI of $14.4 \%-22.9 \%$ ) of adolescents with MDD had at least one suicide attempt by age 30, compared with $3 \%$ ( $95 \%$ CI of $1.6 \%$ and $5.1 \%$ ) of adolescents without MDD. ${ }^{38}$

[^8]- A 2018 systematic review by Johnson et al. found that adolescent depression increased the risk of adult depression by 2.78 times (OR of 2.78; 95\% CI of 1.97 3.93). ${ }^{39}$
- Based on the evidence from Rohde et al. ${ }^{40}$ and Johnson et al. ${ }^{41}$ noted above, we have assumed that the effect of adolescent depression on suicide would continue until age 34.
- Based on data from the $2013^{42}, 2014^{43}$ and $2015^{44}$ BC Vital Statistics annual reports, $24.3 \%$ of deaths in males and $15.5 \%$ of deaths in females ages 15-19 are due to intentional self-harm (see Table 3).

|  | Table 3: Total Deaths and Deaths Attributable to Intentional Self-Harm (ISH) British Columbia, 2013 to 2015 |  |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Males |  |  |  |  |  |  |  |  |  |  |  |
|  | 2013 |  |  | 2014 |  |  | 2015 |  |  | 2013-2015 Combined |  |  |
|  |  | Deaths | \% of Deaths |  | Deaths | \% of Deaths |  | Deaths | \% of Deaths |  | Deaths | \% of Deaths |
| Age Group | All <br> Deaths | Attributable to ISH | Attributable to ISH | All <br> Deaths | Attributable to ISH | Attributable to ISH | All <br> Deaths | Attributable $\qquad$ to ISH | Attributable to ISH | All <br> Deaths | Attributable $\qquad$ to ISH | Attributable $\qquad$ to ISH |
| 10-14 | 10 | 1 | 10.0\% | 12 | 2 | 16.7\% | 12 | 1 | 8.3\% | 34 | 4 | 11.8\% |
| 15-19 | 58 | 5 | 8.6\% | 64 | 24 | 37.5\% | 59 | 15 | 25.4\% | 181 | 44 | 24.3\% |
| 20-24 | 119 | 16 | 13.4\% | 99 | 22 | 22.2\% | 110 | 22 | 20.0\% | 328 | 60 | 18.3\% |
| 25-44 | 650 | 107 | 16.5\% | 669 | 119 | 17.8\% | 757 | 89 | 11.8\% | 2,076 | 315 | 15.2\% |
|  | 837 | 129 | 15.4\% | 844 | 167 | 19.8\% | 938 | 127 | 13.5\% | 2,619 | 423 | 16.2\% |
|  | Females |  |  |  |  |  |  |  |  |  |  |  |
|  | 2013 |  |  | 2014 |  |  | 2015 |  |  | 2013-2015 Combined |  |  |
|  |  | Deaths | \% of Deaths | Deaths \% of Deaths |  |  | Deaths \% of Deaths |  |  | Deaths |  | \% of Deaths |
| Age Group | All <br> Deaths | Attributable to ISH | Attributable to ISH | All <br> Deaths | Attributable to ISH | Attributable to ISH | All <br> Deaths | Attributable | Attributable to ISH | All <br> Deaths | Attributable | Attributable to ISH |
| 10-14 | 11 | 0 | 0.0\% | 3 | 0 | 0.0\% | 5 | 0 | 0.0\% | 19 | 0 | 0.0\% |
| 15-19 | 29 | 6 | 20.7\% | 26 | 3 | 11.5\% | 29 | 4 | 13.8\% | 84 | 13 | 15.5\% |
| 20-24 | 55 | 15 | 27.3\% | 37 | 9 | 24.3\% | 43 | 9 | 20.9\% | 135 | 33 | 24.4\% |
| 25-44 | 368 | 42 | 11.4\% | 392 | 44 | 11.2\% | 337 | 25 | 7.4\% | 1,097 | 111 | 10.1\% |
|  | 463 | 63 | 13.6\% | 458 | 56 | 12.2\% | 414 | 38 | 9.2\% | 1,335 | 157 | 11.8\% |

- Tables 4 and 5 provide data on the expected number of deaths in a BC birth cohort of 20,000 males (see Table 4) and 20,000 females (see Table 5) and how many of those deaths would be attributable to intentional self-harm (see Table 3). Total deaths and deaths attributable to intentional self-harm (ISH) from age 12 to 34 are considered.

[^9]- In the birth cohort of 20,000 males, 45 of the 267 ( $17.0 \%$ ) deaths between the ages of 12 and 34 are due to ISH, resulting in 2,159 life-years lost due to ISH (see Table 4). In the birth cohort of 20,000 females, 17 of $131(13.2 \%)$ deaths between the ages of 12 and 34 are due to ISH, resulting in 1,030 life-years lost due to ISH (see Table 5).

| Self-Harm |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| in a British Columbia Male Birth Cohort of 20,000 |  |  |  |  |  |  |
| Age Group | Individuals in Birth Cohort | Deaths | \% of Deaths due to Intentional Self-Harm | \# of Deaths due to Intentional Self-Harm | Average Life Years Lived | Life Years Lost due to Intentional Self-Harm |
| 11 | 19,898 |  |  |  |  |  |
| 12 | 19,896 | 2 | 11.8\% | 0.2 | 68.6 | 13 |
| 13 | 19,894 | 2 | 11.8\% | 0.3 | 67.6 | 17 |
| 14 | 19,892 | 3 | 11.8\% | 0.3 | 66.6 | 20 |
| 15 | 19,888 | 3 | 24.3\% | 0.8 | 65.7 | 54 |
| 16 | 19,884 | 4 | 24.3\% | 1.0 | 64.7 | 66 |
| 17 | 19,878 | 6 | 24.3\% | 1.4 | 63.7 | 87 |
| 18 | 19,871 | 7 | 24.3\% | 1.8 | 62.7 | 110 |
| 19 | 19,862 | 9 | 24.3\% | 2.2 | 61.7 | 138 |
| 20 | 19,850 | 12 | 18.3\% | 2.1 | 60.8 | 129 |
| 21 | 19,837 | 14 | 18.3\% | 2.5 | 59.8 | 149 |
| 22 | 19,821 | 16 | 18.3\% | 2.9 | 58.9 | 168 |
| 23 | 19,805 | 17 | 18.3\% | 3.0 | 57.9 | 176 |
| 24 | 19,788 | 17 | 18.3\% | 3.1 | 57.0 | 175 |
| 25 | 19,772 | 16 | 15.2\% | 2.5 | 56.0 | 138 |
| 26 | 19,756 | 15 | 15.2\% | 2.3 | 55.1 | 127 |
| 27 | 19,742 | 15 | 15.2\% | 2.2 | 54.1 | 120 |
| 28 | 19,727 | 15 | 15.2\% | 2.2 | 53.1 | 118 |
| 29 | 19,713 | 14 | 15.2\% | 2.2 | 52.2 | 114 |
| 30 | 19,698 | 15 | 15.2\% | 2.2 | 51.2 | 115 |
| 31 | 19,683 | 15 | 15.2\% | 2.3 | 50.2 | 117 |
| 32 | 19,666 | 16 | 15.2\% | 2.5 | 49.3 | 121 |
| 33 | 19,649 | 17 | 15.2\% | 2.6 | 48.3 | 125 |
| 34 | 19,631 | 18 | 15.2\% | 2.7 | 47.4 | 129 |
| Total |  | 267 | 17.0\% | 45 |  | 2,159 |



- Depression has an important influence on a person's QoL. Studies have also shown that individuals with current or treated depression report lower preference scores for depression health states than the general population. ${ }^{45,46}$ Pyne and colleagues suggest that "public stigma may result in the general population being less sympathetic to the suffering of individuals with depression and less willing to validate the impact of depression symptoms. ${ }^{3{ }^{47}}$ Revicki and Wood, based on input from patients with depression who had completed at least eight weeks of anti-depressant (AD) medication, identified the following health state utilities: severe depression $=0.30$, moderate depression $=0.55$ to 0.63 , mild depression $=0.64$ to 0.73 and

[^10]antidepressant maintenance therapy $=0.72$ to $0.83 .{ }^{48}$ Whiteford and colleagues ${ }^{49}$ suggest the following health utilities:

- Severe depression
0.35 ( $95 \%$ CI of $0.18-0.53$ )
- Moderate depression
0.59 ( $95 \%$ CI of $0.45-0.72$ )
- Mild depression
0.84 ( $95 \%$ CI of 0.78-0.89)
- For modelling purposes, we assumed an equal proportion of individuals with mild, moderate and severe depression and used the average quality of life provided by Whiteford and colleagues of 0.59 ( $95 \%$ CI of 0.47 to 0.72 ). Based on a general population QoL of 0.85 (see Reference Document), depression results in a reduction in QoL of $31 \%(0.85-0.59 / 0.85)(95 \%$ CI of $15 \%$ to $45 \%$ ) (see Table 6, row $z$ ).
- When a longitudinal perspective is taken, $30 \%$ of adult patients with depression remain undetected at 1 year and only $14 \%$ at the end of 3 years, or approximately one out of seven patients with treatable depression. ${ }^{50,51,52}$
- Applying the adult rate of undiagnosed treatable depression to adolescents may result in understating the number of adolescents with undetected depression in BC as adolescents are more likely than adults to seek advice from peers rather than seek professional help. ${ }^{53}$
- For modelling purposes, we assumed that $25 \%$ of adolescent major depressive disorder is undiagnosed treatable depression and varied this between $15 \%$ and $35 \%$ in the sensitivity analysis (Table 6, row ae).
- The USPSTF only found two screening methods that it deemed adequate for use with adolescents, the Patient Health Questionnaire for Adolescents (PHQ-A) and the Beck Depression Inventory (BDI). The sensitivity of a screening instrument refers to the number of people with the illness, in this case, depression correctly identified by the test. The specificity of the test is the number of people without the illness that are correctly identified by the test.
- For the PHQ-A, Johnson et al. found a sensitivity of $73 \%$ and a specificity of $94 \%$. $^{54}$ They report a positive predictive value (probability that the disease is present when the test is positive) of $56 \%$ for MDD and a negative predictive value of $97 \%$. The PHQ-A has been validated compared to a structured clinical interview.

[^11]- In their analysis of the BDI, Canals et al. found for a cut-off score of 11 (i.e. 11 and higher $=$ depressed) the sensitivity of BDI was $90 \%$, the specificity was $86 \%$ and the positive predictive value was $20 \%$. ${ }^{55}$
- Roberts et al. found sensitivity of BDI at $83.7 \%$, specificity at $80.9 \%$ and positive predictive value at $10.2 \%$ when referenced against DSM III clinical diagnosis. ${ }^{56}$
- The USPSTF considers the PHQ-A to be the best test to use in assessing adolescent depression. We will therefore assume use of the PHQ-A in our base model (with a sensitivity of $73 \%$ and a specificity of $94 \%$ ) (Table 6 , rows ai \& aj). We will assume use of the BDI in our sensitivity analysis, taking the average of the Canals and Roberts studies for sensitivity ( $86.9 \%$ ) and specificity ( $83.5 \%$ ) of the BDI. Because of the potential harms of misdiagnosis, it is useful to apply a second test if individuals test positive with the PHQ-A. When this is modelled, we begin with the PHQ-A and then apply the BDI. In the base model, the second test sensitivity is set to $100 \%$ and the specificity to $0 \%$ in order to correctly carry through the all first tests results to the rest of the model (Table 6, rows $a m$ \& $a n$ ).
- Merikangas and colleagues found that $40.9 \%$ of female and $36.5 \%$ of male adolescents in the US aged 13-17 years with major depressive disorder received mental health services for their illness. ${ }^{57}$
- Mojtabai and colleagues found a similar overall rate in 2005 , reporting that $36.4 \%$ of adolescents 12-17 sought treatment. This rate increased modestly to $42.0 \%$ in 2014 in US adolescents aged 12-17. ${ }^{58}$
- On the other hand, research by Ghandour et al. based on 2016 survey results in the US found that $79.0 \%$ ( $95 \%$ CI of $74.4 \%$ to $83.0 \%$ ) of adolescents aged 12-17 with diagnosed depression received mental health treatment or counselling. ${ }^{59}$ In females 3 -17 years old (the only sex breakdown available), the number was $80.7 \%$ ( $95 \% \mathrm{CI}$ of 76.2 to $84.5 \%$ ) and in males $3-17$ years old it was $75.2 \%$ ( $95 \%$ CI of 67.9 to $81.3 \%$ ). Unfortunately, the study by Ghandour et al. does not provide information on the extent of that treatment or the type of treatment.
- Updating Mojtabai and colleague's numbers using the 2016 and 2017 data from the NSDUH shows that a total of $40.3 \%$ of individuals with a 12 -month major depressive episode either saw or talked to a health professional or used prescription medication. Averaging the rates for the two years, the number is $31.8 \%$ for males and $43.3 \%$ for females. ${ }^{60}$

[^12]- Mojtabai and colleagues found that of those US adolescents aged 12-17 seeking treatment for their MDD, $20.0 \%$ reported use of prescription medication while $50.7 \%$ reported receiving counselling or therapy. ${ }^{61}$ No sex breakdown of counselling or therapy rates was available. NSDUH data for 2016 and 2017 show medication rates of $17.3 \%$ for males and $21.7 \%$ for females. ${ }^{62}$
- The Mental Health Parity and Addiction Equity Act in the US "generally prevents group health plans and health insurance issuers that provide mental health or substance use disorder (MH/SUD) benefits from imposing less favorable benefit limitations on those benefits than on medical/surgical benefits." ${ }^{63}$ The lack of similar legislation in BC may result in treatment seeking rates being lower in BC than are reflected in the US data, especially for non-pharmacological interventions (e.g. counselling). ${ }^{64}$
- In our model, we reduce the US treatment rate(s) by an absolute value of $10 \%$ to account for possibly lower treatment rates in BC.
- Data provided by the BC Ministry of Health indicate that for fiscal years 2011/12 through 2015/16 (5 years), $15.7 \%$ of BC adolescents (12-18) diagnosed with major depression had a prescription for fluoxetine filled within one month of diagnosis, $19.7 \%$ within three months of diagnosis (i.e. an additional $4 \%$ ) and $22.2 \%$ within six months of diagnosis (i.e. an additional $2.5 \%$ since the three-month point). These rates are $14.1 \%, 17.5 \%$ and $19.5 \%$, respectively, for males and $16.6 \%, 20.9 \%$ and $23.6 \%$, respectively, for females. ${ }^{65}$
- It is not uncommon to see wait times of $2-6$ months for non-pharmacological depression interventions (e.g. cognitive behavioural therapy or individual counselling) in BC. ${ }^{66}$
- We consider four distinct groups in our model, that branch from the group of individuals who received a positive screen for major depressive disorder as follows:

[^13]

- We model each group over different time horizons:
- False Positives (no MDD) are modeled as being treated for six months after which time we assume that it becomes clear that this group has been incorrectly screened positive and treatments cease for this group.
- The group with correctly diagnosed MDD that ends up being single event MDD, is also modeled as receiving treatment for six months after which time we assume that no further treatments are undertaken or necessary.
- The group with correctly diagnosed MDD that ends up being recurrent is modeled as receiving treatment for one year after the index event. We model that this group receives treatment for seven subsequent events during their lifetime, each lasting one year.
- The group with correctly diagnosed MDD that ends up being persistent is modeled as receiving treatment for twenty years after the index event. We model that this group continues to use anti-depressants throughout this time.
- For modelling purposes, we assume that $50.5 \%(60.5 \%-10 \%)$ of adolescents with MDD seek treatment ( $60.5 \%$ is the mid-point of $42 \%^{67}$ and $79 \% \%^{68}$ ) and vary this from $32 \%$ to $69 \%$ in our sensitivity analysis (Table 6 , rows $b e, b u$ \& co).
- Of those seeking treatment, $50.7 \%$ receive counselling or therapy (Table 6 , rows $b f$, $b v \& c p$ ).
- In modelling for males, we assume that $43.5 \%$ ( $53.5 \%-10 \%$ ) of male adolescents with MDD seek treatment ( $53.5 \%$ is the mid-point of $31.8 \%{ }^{69}$ and $75.2 \%^{70}$ ) and vary this from $21.8 \%$ to $65.2 \%$ in our sensitivity analysis (Table 6 a, rows $b e, b u \& c o$ ).

[^14]- In modelling for females, we assume that $52.0 \%$ (62.0\%-10\%) of female adolescents with MDD seek treatment ( $62.0 \%$ is the mid-point of $43.3 \%^{71}$ and $80.7 \%^{72}$ ) and vary this from $33.3 \%$ to $70.7 \%$ in our sensitivity analysis (Table 6b, rows $b e, b u \& c o$ ).
- In our model, we assume that $19.7 \%$ (Table 6, row ap) of all individuals screened positive for depression will fill anti-depressant prescriptions during the first three months of treatment and that this increases to $22.2 \%$ during months $4-6$ after a positive screen (Table 6, row ar).
- In our model for males, we assume that $17.5 \%$ (Table 6 a , row $a p$ ) of all males screened positive for depression will fill anti-depressant prescriptions during the first three months of treatment and that this increases to $19.5 \%$ during months $4-6$ after a positive screen (Table 6a, row $a r$ ).
- In our model for females, we assume that $20.9 \%$ (Table 6 b , row $a p$ ) of all females screened positive for depression will fill anti-depressant prescriptions during the first three months of treatment and that this increases to $23.6 \%$ during months $4-6$ after a positive screen (Table 6b, row $a r$ ).
- We model anti-depressant use among recurrent MDD cases and the first year of persistent MDD at $22.2 \%$ (Table 6, row bo) and assume that after the first year, all of the persistent MDD cases are taking anti-depressant medication (Table 6, row $c j$ )
- In males, we model anti-depressant use among recurrent MDD cases and the first year of persistent MDD at $19.5 \%$ (Table 6a, row bo) and assume that after the first year, all of the persistent MDD cases are taking anti-depressant medication (Table 6, row $c j$ )
- In females, we model anti-depressant use among recurrent MDD cases and the first year of persistent MDD at $23.6 \%$ (Table 6 b, row bo) and assume that after the first year, all of the persistent MDD cases are taking anti-depressant medication (Table 6, row $c j$ )
- Cognitive behavioural therapy (CBT) is considered to be a "well-established intervention" for depression in adolescents. ${ }^{73}$
- The systematic review prepared by Forman-Hoffman and colleagues for the USPSTF estimated that CBT leads to a clinical improvement in MDD for $12.1 \%$ (Table 6, row $a u)$ of adolescents receiving this therapy compared to a placebo. ${ }^{74}$

[^15]- Cipriani and colleagues conducted a meta-analysis on efficacy and tolerability of antidepressants in adolescents with major depressive disorder and concluded that "only fluoxetine was statistically significantly more effective than placebo."75
- In the clinical guideline for the USPSTF, Siu only identifies one type of selective serotonin reuptake inhibitor (SSRI) with a "good" quality study supporting its use in treating MDD in adolescents: fluoxetine. ${ }^{76}$
- The systematic review prepared by Forman-Hoffman and colleagues for the USPSTF estimated that fluoxetine alone leads to a clinical improvement in MDD for $25.7 \%$ ( $95 \% \mathrm{CI}$ of $16.2 \%$ to $35.2 \%$ ) of adolescents taking it.
- The systematic review prepared by Forman-Hoffman and colleagues for the USPSTF estimated that when fluoxetine is combined with CBT, the clinical improvement in MDD increases to $36.2 \%$ ( $95 \%$ CI of $27.2 \%$ to $45.2 \%$ ) (Table 6, row $a v$ ).
- The Canadian Network for Mood and Anxiety Treatments (CANMAT) 2016 Clinical Guidelines recommend two treatment phases for depression: ${ }^{77}$
- an acute phase, lasting 8 to 12 weeks, targeting symptom remission and restoration of functioning
- a maintenance phase, lasting 6 to 24 months, targeting prevention of recurrence and return to full functioning and quality of life
- Depression is a highly recurrent disorder. ${ }^{78}$ On average, half of individuals experiencing at least one MDE during their lifetime will experience between 5-9 recurrent episodes during their lifetime. ${ }^{79,80,81}$
- In a follow-up of individuals using anti-depressants, Colman and colleagues reported that $24 \%$ of patients were still using anti-depressants 10 -years later. ${ }^{82}$
- In our model, we assume that $50 \%$ of the MDD cases are single events and the remainder will be recurrent or persistent MDD (Table 6, row $a x$ ).
- We model that $5.3 \%$ of the MDD cases are persistent ( $22.2 \%$ 6-month anti-depressant use in BC adolescents x $24 \%$ still using anti-depressants 10 years later $=5.3 \%$ of MDD) (Table 6, row $c c$ ), which leaves $44.7 \%$ of the initial MDD cases that recur multiple times in an individual's lifetime ( $100 \%-50 \%-5.3 \%=44.7 \%$ ) (Table 6, row $b m$ ).

[^16]- For males, we model that $4.7 \%$ of the MDD cases are persistent ( $19.5 \%$ 6-month anti-depressant use in BC adolescents x $24 \%$ still using anti-depressants 10 years later $=4.7 \%$ of MDD) (Table 6, row $c c$ ), which leaves $45.3 \%$ of the initial MDD cases that recur multiple times in an individual's lifetime ( $100 \%-50 \%-4.7 \%=$ 45.3\%) (Table 6, row bm).
- For females, we model that $5.7 \%$ of the MDD cases are persistent ( $23.6 \%$ 6-month anti-depressant use in BC adolescents x $24 \%$ still using anti-depressants 10 years later $=5.7 \%$ of MDD $)($ Table 6 , row $c c)$, which leaves $44.3 \%$ of the initial MDD cases that recur multiple times in an individual's lifetime ( $100 \%-50 \%-5.7 \%=$ $44.3 \%$ ) (Table 6, row bm).
- We have modeled an additional 7 episodes after the index MDD episode for a total of eight (8) MDD events for recurrent MDD (Table 6, row bs). For discounting purposes, we model these as occurring eight years apart throughout the lifetime of the affected individuals.
- Approximately $60 \%$ of patients stay on anti-depressant medication for at least 3 months and $45 \%$ for at least 6 months. ${ }^{83,84}$ For those diagnosed with depression and taking medication, an average of $71 \%$ of days in a 180-day period had anti-depressant use and $62 \%$ of days in a 365 -day period had anti-depressant use. ${ }^{85}$ On average, antidepressants are taken on 226 days each year. ${ }^{86}$
- The average length of an adolescent depressive episode has been reported to range between 24.4 and 27 weeks. ${ }^{87,88}$
- Van der Voort and colleagues report that single episodes of MDD recover within six months of onset and that individuals with syndromal (recurrent) MDD take up to twelve months to recover fully. ${ }^{89}$
- Following van der Voort and colleagues, we model single episodes of MDD as recovering within 6 months (Table 6, row $b c$ ) and recurrent episodes as recovering within one year (Table 6 , row $b r$ ). We model persistent MDD as requiring treatment throughout the lifetime (Table 6, row $c t$ ). We model persistent treatment for the 20 years from 15 years old (mid-point of the 12 - 18 year old cohort) to 34 years of age, consistent with Tables 4 \& 5 .

[^17]- Several recent meta-analyses suggest that internet-based cognitive behavioural therapy may be effective in treating general depression in adults. ${ }^{90,91}$ The evidence that is currently available is insufficient to justify modeling this approach for adolescents with MDD.
- We model treatment for those with a positive MDD screen by time period as follows:
- $0-3$ months after screening: $19.7 \%$ of positive screened adolescents ( $17.5 \%$ males, $20.9 \%$ females) are taking anti-depressants.
- 4-6 months after screening: $22.2 \%$ of positive screen adolescents are taking anti-depressants and $25.6 \%$ are in counselling or therapy (Table 6 rows $b g$, $b w \& c q$ ), with half of the therapy group in individual sessions and half in group sessions. The $25.6 \%$ is based on $50.5 \%$ seeking treatment multiplied by $50.7 \%$ of those seeking treatment attending therapy / counselling.
- For males the counselling rate is $22.1 \%$ ( $43.5 \%$ treatment seeking x $50.7 \%$ counselling rate among treatment seekers) (Table 6a rows bg, $b w \& c q)$.
- For females the counselling rate is $26.4 \%$ ( $52.0 \%$ treatment seeking x $50.7 \%$ counselling rate among treatment seekers) (Table 6a rows bg, $b w \& c q)$.
- 7-12 months after screening: $22.2 \%$ of correctly diagnosed adolescents with recurrent or persistent MDD are on anti-depressants and $25.6 \%$ are in counselling or therapy, with half of the therapy group in individual sessions and half in group sessions.
- 13+ months after screening: all of the correctly diagnosed adolescents with persistent MDD are on anti-depressants. We assume that the $25.6 \%$ in counselling or therapy receive four (4) individual sessions annually.
- Recurrent MDD: for each year of recurrent MDD, $22.2 \%$ of individuals with recurrent MDD take anti-depressants and $25.6 \%$ receive therapy ( 5 sessions).

[^18]Treatment Modeling for Positive MDD Screens

|  |  |  | e Positive Scree |  | False Positive |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Single Event | Recurrent | Persistent | Screens |
| 0-3 Months | Pharmacological | 19.7\% anti-depressant rate |  |  |  |
|  | Therapeutic | None |  |  |  |
| 4-6 Months | Pharmacological | 22.2\% anti-depressant rate |  |  |  |
|  | Therapeutic | 25.6\% receiving therapy |  |  |  |
| 7-12 Months | Pharmacological | No treatment | 22.2\% anti-depressant rate |  | No treatment |
|  | Therapeutic |  | 25.6\% receiving therapy |  |  |
| 13+ Months | Pharmacological |  | No Treatment | 100\% antidepressant rate |  |
|  | Therapeutic |  |  | $25.6 \%$ receiving therapy |  |

- Revicki and Wood found that antidepressant maintenance therapy resulted in a weighted average QoL of 0.78 ( $95 \% \mathrm{CI}$ of 0.63 to 0.93 ). ${ }^{92}$ Based on a general population QoL of 0.85 (see Reference Document), antidepressant maintenance therapy results in a reduction in QoL of $8 \%(0.85-0.78 / 0.85)(95 \% \mathrm{CI}$ of $26 \%$ to no reduction) (Table 6, row $b g$ ).

[^19] depression severity and antidepressant medications. Journal of Affective Disorders. 1998; 48(1): 25-36.

## CPB for Both Sexes

Based on these assumptions, the CPB associated with screening for major depressive disorder in adolescents (both sexes) ages 12 to 18 is 222 QALYs (see Table 6, row da).

| Table 6: CPB of Screening for MDD in Adolescents Ages 12-18 <br> In a BC Birth Cohort of 40,000 |  |  |  |
| :---: | :---: | :---: | :---: |
| Row <br> Label | Variable | Base case | Data Source |
| a | Number of life years, 12 year olds | 39,804 | BC Life Table |
| b | Annual rate of MDD, 12 year olds | 5.2\% | $\checkmark$ |
| c | Life years with MDD, 12 year olds | 2,070 | = a*b |
| d | Life years without MDD, 12 year olds | 37,735 | =a-c |
| e | Number of life years, 13 year olds | 39,801 | BC Life Table |
| $f$ | Annual rate of MDD, 13 year olds | 9.3\% | $\checkmark$ |
| g | Life years with MDD, 13 year olds | 3,702 | $=e^{*} \mathrm{f}$ |
| h | Life years without MDD, 13 year olds | 36,100 | $=\mathrm{e}-\mathrm{g}$ |
| i | Number of life years, 14 year olds | 39,797 | BC Life Table |
| j | Annual rate of MDD, 14 year olds | 11.7\% | $\checkmark$ |
| k | Life years with MDD, 14 year olds | 4,656 | $=\mathrm{i}^{*} \mathrm{j}$ |
| 1 | Life years without MDD, 14 year olds | 35,141 | = i-k |
| m | Number of life years, 15 year olds | 39,792 | BC Life Table |
| n | Annual rate of MDD, 15 year olds | 15.0\% | $\checkmark$ |
| 0 | Life years with MDD, 15 year olds | 5,969 | $=\mathrm{m}$ * n |
| p | Life years without MDD, 15 year olds | 33,823 | = m - 0 |
| q | Number of life years, 16 year olds | 39,784 | BC Life Table |
| r | Annual rate of MDD, 16 year olds | 16.0\% | $\checkmark$ |
| S | Life years with MDD, 16 year olds | 6,365 | $=q^{*} r$ |
| t | Life years without MDD, 16 year olds | 33,419 | =q-s |
| u | Number of life years, 17 and 18 year olds | 79,534 | BC Life Table |
| v | Annual rate of MDD, 17 and 18 year olds | 16.5\% | $\checkmark$ |
| w | Life years with MDD, 17 and 18 year olds | 13,123 | $=u^{*} \mathrm{v}$ |
| x | Life years without MDD, 17 and 18 year olds | 66,411 | = $\mathrm{u}-\mathrm{w}$ |
| y | Life years with MDD between 12 and 18 | 35,885 | $=c+g+k+o+s+w$ |
| z | QoL decrement due to depression | 0.31 | $\checkmark$ |
| aa | QALYs lost during adolescence due to depression | 11,124 | $=y^{*} \mathrm{z}$ |
| ab | Deaths attributable to ISH between the ages of 12 and 34 | 65 | Tables 4\&5 |
| ac | QALYS lost due to deaths attributable to ISH between the ages of 12 and 34 | 3,189 | Tables 4 \& 5 |
| ad | Total QALYs lost due to depression in adolescence | 14,313 | = aa + ac |
| ae | \% MDD undetected in lifetime | 25.0\% | $\checkmark$ |
| af | Life years with undetected MDD in cohort between 12-18 years of age | 8,971 | $=y^{*} \mathrm{ae}$ |
| ag | Number of well care visits per year | 2.07 | $\checkmark$ |
| ah | Depression screening rate | 7.4\% | $\checkmark$ |
| ai | Sensitivity (rate of true positives), initial test | 73.0\% | V |
| aj | Specificity (rate of true negatives), initial test | 94.0\% | $\checkmark$ |
| ak | Number of MDD cases correctly identified, initial test | 1,003 | = af * ag * $\mathrm{hh}^{*}$ ai |
| al | Number of MDD cases diagnosed incorrectly, initial test | 2,230 | $=(\mathrm{d}+\mathrm{h}+\mathrm{l}+\mathrm{p}+\mathrm{t}+\mathrm{x})^{*} \mathrm{ag} * \mathrm{ah} *(1-\mathrm{aj})$ |
| am | Sensitivity (rate of true positives), 2nd test | 100.0\% | No second test in base model |
| an | Specificity (rate of true negatives), 2nd test | 0.0\% | No second test in base model |
|  | Incorrectly Diagnosed MDD Cases |  |  |
| ao | Number of MDD cases diagnosed incorrectly, overall | 2,230 | = al * (1-an) |
| ap | Rate of anti-depressants, months 0-3 | 19.7\% | V |
| aq | Number taking anti-depressants months 0-3 | 439 | = ao * ap |
| ar | Rate of anti-depressants, months 4-6 | 22.2\% | $\checkmark$ |
| as | Number taking anti-depressants months 4-6 | 495 | = ao * ar |
| at | Life years on anti-depressants | 234 | $=\left(\mathrm{aq}^{*} 0.25\right)+\left(\mathrm{as}^{*} 0.25\right)$ |
| au | QoL decrement due to antidepressant therapy | 0.08 | $\checkmark$ |
| av | QALYs Gained (or Lost), Incorrectly Diagnosed MDD | -18.7 | $=-$ (at * au) |

Table 6: CPB of Screening for MDD in Adolescents Ages 12-18
In a BC Birth Cohort of 40,000

|  | Correctly Diagnosed MDD Cases |  |  |
| :---: | :---: | :---: | :---: |
|  | Single Event MDD |  |  |
| aw | Number of MDD cases correctly identified, overall | 1,003 | = ak * am |
| ax | Rate of single event MDD in correct diagnoses | 50.0\% | $\checkmark$ |
| ay | Number of single event MDD cases | 502 | = aw * $\mathrm{ax}^{\text {a }}$ |
| az | Rate of 6-month antidepressant use | 22.2\% | $\checkmark$ |
| ba | Number on anti-depressants | 111 | = ay * az |
| bb | Clinical improvement rate due to anti-depressants | 25.7\% | V |
| bc | Length of single event MDD, years | 0.5 | $\checkmark$ |
| bd | Depression-free life years gained due to anti-depressants | 14.3 | $=a b^{*} \mathrm{bb}{ }^{*} \mathrm{bc}$ |
| be | Treatment seeking rate | 50.5\% | $\checkmark$ |
| bf | Rate counselling among treatment seekers | 50.7\% | $\checkmark$ |
| bg | Overall counselling rate | 25.6\% | $=\mathrm{be}$ * bf |
| bh | Number in counselling | 128 | = ay * bg |
| bi | Clinical improvement rate due to counselling | 12.1\% | $\checkmark$ |
| bj | Length of single event MDD counselling, years | 0.25 | $\checkmark$ |
| bk | Depression-free life years gained due to counselling | 3.9 | $=\mathrm{bh}{ }^{*} \mathrm{bi}^{*} \mathrm{bj}$ |
| Recurrent MDD |  |  |  |
| bl | Number of MDD cases correctly identified, overall | 1,003 | = ak * am |
| bm | Rate of recurrent MDD in correct diagnoses | 44.7\% | $\checkmark$ |
| bn | Number of recurrent MDD cases | 448 | $=\mathrm{bl}$ * bm |
| bo | Rate of 12-month antidepressant use | 22.2\% | $\checkmark$ |
| bp | Number on anti-depressants | 99 | $=\mathrm{bn}$ * bo |
| bq | Clinical improvement rate due to anti-depressants | 25.7\% | $\checkmark$ |
| br | Length of recurrent MDD event, years | 1.0 | $\checkmark$ |
| bs | Number of recurrent episodes, lifetime | 8.0 | $\checkmark$ |
| bt | Depression-free life years gained due to anti-depressants | 205 | $=b p$ * ${ }^{\text {d }}$ * br*bs |
| bu | Treatment seeking rate | 50.5\% | $\checkmark$ |
| bv | Rate counselling among treatment seekers | 50.7\% | $\checkmark$ |
| bw | Overall counselling rate | 25.6\% | $=\mathrm{bu}$ * bv |
| bx | Number in counselling | 115 | = bn * bw |
| by | Clinical improvement rate due to counselling | 12.1\% | $\checkmark$ |
| bz | Length of recurrent MDD counselling, years | 0.75 | $\checkmark$ |
| ca | Depression-free life years gained due to counselling | 83 | = bx * by * bz *bs |
| Persistent MDD |  |  |  |
| cb | Number of MDD cases correctly identified, overall | 1,003 | = ak * am |
| cc | Rate of persistent MDD in correct diagnoses | 5.3\% | $\checkmark$ |
| cd | Number of persistent MDD cases | 53 | $=\mathrm{cb}$ * cc |
| ce | Rate of first year antidepressant use | 22.2\% | $\checkmark$ |
| cf | Number on anti-depressants | 12 | $=\mathrm{cd}^{*} \mathrm{ce}$ |
| cg | Clinical improvement rate due to anti-depressants | 25.7\% | $\checkmark$ |
| ch | Length of treatment | 1.0 | $\checkmark$ |
| ci | Depression-free life years gained due to anti-depressants, year 1 | 3.0 | $=\mathrm{cf} * \mathrm{cg}^{*} \mathrm{ch}$ |
| cj | Rate of antidepressant use years 2-20 | 100.0\% | $\checkmark$ |
| ck | Number on anti-depressants | 53 | = $\mathrm{cd}^{*} \mathrm{cj}$ |
| cl | Clinical improvement rate due to anti-depressants | 25.7\% | $\checkmark$ |
| cm | Length of treatment | 19.0 | $\checkmark$ |
| cn | Depression-free life years gained due to anti-depressants, years 2-20 | 261 | $=\mathrm{ck} * \mathrm{cl}^{*} \mathrm{~cm}$ |
| co | Treatment seeking rate | 50.5\% | $\checkmark$ |
| cp | Rate counselling among treatment seekers | 50.7\% | $\checkmark$ |
| cq | Overall counselling rate | 25.6\% | = $\mathrm{co}^{*} \mathrm{cp}$ |
| cr | Number in counselling | 14 | $=\mathrm{cd}^{*} \mathrm{cq}$ |
| cs | Clinical improvement rate due to counselling | 12.1\% | $\checkmark$ |
| ct | Length of effect persistent event MDD counselling, years | 20.0 | $\checkmark$ |
| cu | Depression-free life years gained due to counselling | 33 | $=\mathrm{cr}$ * cs * ct |
| Summary of QALYs Gained with Screening |  |  |  |
| cv | Individuals with MDD helped by treatment | 88 | = aw * ( az * bb) + (bg * bi)) |
| CW | Depression free life years due to screening, correctly diagnosed MDD | 603 | $=(b d+b k)+(b t+c a)+(c i+c n+c u)$ |
| CX | Reduction in \% of total life years with MDD due to screening | 1.68\% | = cw $/ \mathrm{y}$ |
| cy | QALYs gained due to screening, correctly diagnosed MDD | 241 | = cx* ad |
| cz | QALYs due to treating incorrectly diagnosed MDD | -19 | = av |
| da | Net QALYs as a result of screening (CPB) | 222 | = cy +cz |

For the sensitivity analysis of the base model (both sexes), we modified a number of major assumptions and recalculated the CPB as follows:

- Assume the rate of undetected MDD decreases from $25 \%$ to $15 \%$ (Table 6, row $a e$ ): $\mathrm{CPB}=126$
- Assume the rate of undetected MDD increases from $25 \%$ to $35 \%$ (Table 6, row $a e$ ): $\mathrm{CPB}=318$
- Assume a second round of screening (with BDI) is introduced, with a sensitivity of $86.9 \%$ and a specificity of $83.5 \%$ (Table 6, rows $a m$ \& an): $\mathrm{CPB}=206$
- Assume the rate of treatment seeking increases from $50.5 \%$ to $69 \%$ (Table 6, row aq): $\mathrm{CPB}=239$
- Assume the rate of treatment seeking decreases from $50.5 \%$ to $32 \%$ (Table 6 , row $a q): \mathrm{CPB}=204$
- Assume the QoL decrement for depression is reduced from 31\% to $15 \%$ (Table 6, row $z$ ) and the QoL decrement for anti-depressant maintenance therapy is reduced from $8 \%$ to $0 \%$ (i.e. no decrement) (Table 6, row $b g$ ): $\mathrm{CPB}=144$
- Assume the QoL decrement for depression is increased from 31\% to 45\% (Table 6, row $z$ ) and the QoL decrement for anti-depressant maintenance therapy is increased from $8 \%$ to $26 \%$ (Table 6, row $b g$ ): $\mathrm{CPB}=264$
- Assume that the screening rate is only applied to one visit per year per patient, rather than 2.07 (Table 6, row $a g$ ): $\mathrm{CPB}=107$


## CPB for Males

Based on the above assumptions for males, the CPB associated with screening for major depressive disorder in male adolescents' ages 12 to 18 is 83 QALYs (see Table 6 , row $d a$ ).

Table 6a: CPB of Screening for MDD in Male Adolescents Ages 12-18 In a BC Birth Cohort of 40,000

| Row <br> Label | Variable | Base case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Number of life years, 12 year olds | 19,896 | BC Life Table |
| b | Annual rate of MDD, 12 year olds | 5.2\% | V |
| c | Life years with MDD, 12 year olds | 1,035 | = ${ }^{*} \mathrm{~b}$ |
| d | Life years without MDD, 12 year olds | 18,862 | $=a-c$ |
| e | Number of life years, 13 year olds | 19,894 | BC Life Table |
| f | Annual rate of MDD, 13 year olds | 9.3\% | $\checkmark$ |
| g | Life years with MDD, 13 year olds | 1,850 | = ${ }^{*} \mathrm{f}$ |
| h | Life years without MDD, 13 year olds | 18,044 | =e-g |
| i | Number of life years, 14 year olds | 19,892 | BC Life Table |
| j | Annual rate of MDD, 14 year olds | 11.7\% | $\checkmark$ |
| k | Life years with MDD, 14 year olds | 2,327 | $=i^{*} \mathrm{j}$ |
| I | Life years without MDD, 14 year olds | 17,564 | = i-k |
| m | Number of life years, 15 year olds | 19,888 | BC Life Table |
| n | Annual rate of MDD, 15 year olds | 15.0\% | V |
| 0 | Life years with MDD, 15 year olds | 2,983 | $=\mathrm{m}^{*} \mathrm{n}$ |
| p | Life years without MDD, 15 year olds | 16,905 | = m - 0 |
| q | Number of life years, 16 year olds | 19,884 | BC Life Table |
| r | Annual rate of MDD, 16 year olds | 16.0\% | $\checkmark$ |
| s | Life years with MDD, 16 year olds | 3,181 | $=q^{*} r$ |
| t | Life years without MDD, 16 year olds | 16,703 | $=q-s$ |
| u | Number of life years, 17 and 18 year olds | 39,750 | BC Life Table |
| v | Annual rate of MDD, 17 and 18 year olds | 16.5\% | $\checkmark$ |
| w | Life years with MDD, 17 and 18 year olds | 6,559 | $=u^{*} \mathrm{v}$ |
| x | Life years without MDD, 17 and 18 year olds | 33,191 | = u-w |
| y | Life years with MDD between 12 and 18 | 17,935 | $=c+g+k+o+s+w$ |
| z | QoL decrement due to depression | 0.31 | $\checkmark$ |
| aa | QALYs lost during adolescence due to depression | 5,560 | = ${ }^{*} \mathrm{z}$ |
| ab | Deaths attributable to ISH between the ages of 12 and 34 | 45 | Table 4 |
| ac | QALYS lost due to deaths attributable to ISH between the ages of 12 and 34 | 2,159 | Table 4 |
| ad | Total QALYs lost due to depression in adolescence | 7,719 | = aa + ac |
| ae | \% MDD undetected in lifetime | 25.0\% | $\checkmark$ |
| af | Life years with undetected MDD in cohort between 12-18 years of age | 4,484 | = ${ }^{*}$ * ${ }^{\text {e }}$ |
| ag | Number of well care visits per year | 1.75 | V |
| ah | Depression screening rate | 6.9\% | $\checkmark$ |
| ai | Sensitivity (rate of true positives), initial test | 73.0\% | $\checkmark$ |
| aj | Specificity (rate of true negatives), initial test | 94.0\% | $\checkmark$ |
| ak | Number of MDD cases correctly identified, initial test | 395 | = af *ag *ah *ai |
| al | Number of MDD cases diagnosed incorrectly, initial test | 879 | $=(\mathrm{d}+\mathrm{h}+\mathrm{l}+\mathrm{p}+\mathrm{t}+\mathrm{x}) * \mathrm{ag} * \mathrm{ah} *(1-\mathrm{aj})$ |
| am | Sensitivity (rate of true positives), 2nd test | 100.0\% | No second test in base model |
| an | Specificity (rate of true negatives), 2nd test | 0.0\% | No second test in base model |
|  | Incorrectly Diagnosed MDD cases |  |  |
| ao | Number of MDD cases diagnosed incorrectly, overall | 879 | = al * (1-an) |
| ap | Rate of anti-depressants, months 0-3 | 17.5\% | V |
| aq | Number taking anti-depressants months 0-3 | 154 | = ao * ap |
| ar | Rate of anti-depressants, months 4-6 | 19.5\% | V |
| as | Number taking anti-depressants months 4-6 | 171 | = ao * ar |
| at | Life years on anti-depressants | 81 | $=(\mathrm{aq} * 0.25)+(\mathrm{as} * 0.25)$ |
| au | QoL decrement due to antidepressant therapy | 0.08 | $\checkmark$ |
| av | QALYs Gained (or Lost), Incorrectly Diagnosed MDD | -6.5 | =- (at * au) |

Table 6a: CPB of Screening for MDD in Male Adolescents Ages 12-18
In a BC Birth Cohort of 40,000

|  | Correctly Diagnosed MDD cases |  |  |
| :---: | :---: | :---: | :---: |
|  | Single Event MDD |  |  |
| aw | Number of MDD cases correctly identified, overall | 395 | = ak * am |
| ax | Rate of single event MDD in correct diagnoses | 50.0\% | $\checkmark$ |
| ay | Number of single event MDD cases | 198 | = aw * $\mathrm{ax}^{\text {d }}$ |
| az | Rate of 6-month antidepressant use | 19.5\% | $\checkmark$ |
| ba | Number on anti-depressants | 39 | = ay * az |
| bb | Clinical improvement rate due to anti-depressants | 25.7\% | $\checkmark$ |
| bc | Length of single event MDD, years | 0.5 | $\checkmark$ |
| bd | Depression-free life years gained due to anti-depressants | 5.0 | $=a b * b b^{*} \mathrm{bc}$ |
| be | Treatment seeking rate | 43.5\% | $\checkmark$ |
| bf | Rate counselling among treatment seekers | 50.7\% | $\checkmark$ |
| bg | Overall counselling rate | 22.1\% | $=\mathrm{be}$ * bf |
| bh | Number in counselling | 44 | = ay * bg |
| bi | Clinical improvement rate due to counselling | 12.1\% | V |
| bj | Length of single event MDD counselling, years | 0.25 | $\checkmark$ |
| bk | Depression-free life years gained due to counselling | 1.3 | $=\mathrm{bh}{ }^{*} \mathrm{bi}^{*} \mathrm{bj}$ |
|  | Recurrent MDD |  |  |
| bl | Number of MDD cases correctly identified, overall | 395 | = ak * am |
| bm | Rate of recurrent MDD in correct diagnoses | 45.3\% | $\checkmark$ |
| bn | Number of recurrent MDD cases | 179 | $=\mathrm{bl}$ * bm |
| bo | Rate of 12-month antidepressant use | 19.5\% | $\checkmark$ |
| bp | Number on anti-depressants | 35 | = bn * bo |
| bq | Clinical improvement rate due to anti-depressants | 25.7\% | $\checkmark$ |
| br | Length of recurrent MDD event, years | 1.0 | $\checkmark$ |
| bs | Number of recurrent episodes, lifetime | 8.0 | $\checkmark$ |
| bt | Depression-free life years gained due to anti-depressants | 72 | $=b p^{*} b q^{*} b r^{*}$ bs |
| bu | Treatment seeking rate | 43.5\% | $\checkmark$ |
| bv | Rate counselling among treatment seekers | 50.7\% | $\checkmark$ |
| bw | Overall counselling rate | 22.1\% | $=\mathrm{bu}$ * bv |
| bx | Number in counselling | 39 | = bn * bw |
| by | Clinical improvement rate due to counselling | 12.1\% | $\checkmark$ |
| bz | Length of recurrent MDD counselling, years | 0.75 | $\checkmark$ |
| ca | Depression-free life years gained due to counselling | 29 | = bx * by * bz *bs |
|  | Persistent MDD |  |  |
| cb | Number of MDD cases correctly identified, overall | 395 | = $\mathrm{ak}^{*} \mathrm{am}$ |
| cc | Rate of persistent MDD in correct diagnoses | 4.7\% | $\checkmark$ |
| cd | Number of persistent MDD cases | 19 | $=\mathrm{cb}{ }^{*} \mathrm{cc}$ |
| ce | Rate of first year antidepressant use | 19.5\% | $\checkmark$ |
| cf | Number on anti-depressants | 4 | = cd * ce |
| cg | Clinical improvement rate due to anti-depressants | 25.7\% | $\checkmark$ |
| ch | Length of treatment | 1.0 | $\checkmark$ |
| ci | Depression-free life years gained due to anti-depressants, year 1 | 0.9 | = cf * $\mathrm{cg}^{*} \mathrm{ch}$ |
| cj | Rate of antidepressant use years 2-20 | 100.0\% | $\checkmark$ |
| ck | Number on anti-depressants | 19 | = $\mathrm{cd}^{*} \mathrm{cj}$ |
| cl | Clinical improvement rate due to anti-depressants | 25.7\% | $\checkmark$ |
| cm | Length of treatment | 19.0 | $\checkmark$ |
| cn | Depression-free life years gained due to anti-depressants, years 2-20 | 91 | $=\mathrm{ck} * \mathrm{cl}^{*} \mathrm{~cm}$ |
| co | Treatment seeking rate | 43.5\% | $\checkmark$ |
| cp | Rate counselling among treatment seekers | 50.7\% | $\checkmark$ |
| cq | Overall counselling rate | 22.1\% | = $\mathrm{co}^{*} \mathrm{cp}$ |
| cr | Number in counselling | 4 | $=\mathrm{cd} * \mathrm{cq}$ |
| cs | Clinical improvement rate due to counselling | 12.1\% | $\checkmark$ |
| ct | Length of effect persistent event MDD counselling, years | 20.0 | $\checkmark$ |
| cu | Depression-free life years gained due to counselling | 10 | $=\mathrm{cr}^{*} \mathrm{cs} * \mathrm{ct}$ |
|  | Summary of QALYs Gained with Screening |  |  |
| CV | Individuals with MDD helped by treatment | 30 | $=\mathrm{aw}$ * ((az * bb) + (bg * bi)) |
| cw | Depression free life years due to screening, correctly diagnosed MDD | 208 | $=(b d+b k)+(b t+c a)+(c i+c n+c u)$ |
| cx | Reduction in \% of total life years with MDD due to screening | 1.16\% | = cw $/ \mathrm{y}$ |
| cy | QALYs gained due to screening, correctly diagnosed MDD | 90 | = cx* ad |
| cz | QALYs due to treating incorrectly diagnosed MDD | -7 | = av |
| da | Net QALYs as a result of screening (CPB) | 83 | = cy + cz |

[^20]For the sensitivity analysis of the base model for males, we modified a number of major assumptions and recalculated the CPB as follows:

- Assume the rate of undetected MDD decreases from 25\% to $15 \%$ (Table 6a, row $a e$ ): $\mathrm{CPB}=47$
- Assume the rate of undetected MDD increases from $25 \%$ to $35 \%$ (Table 6a, row $a e$ ): $\mathrm{CPB}=119$
- Assume a second round of screening (with BDI) is introduced, with a sensitivity of $86.9 \%$ and a specificity of $83.5 \%$ (Table 6a, rows $a m$ \& an): CPB $=77$
- Assume the rate of treatment seeking increases from $43.5 \%$ to $65.2 \%$ (Table 6a, row $a q): \mathrm{CPB}=92$
- Assume the rate of treatment seeking decreases from $43.5 \%$ to $21.8 \%$ (Table 6a, row aq): $\mathrm{CPB}=75$
- Assume the QoL decrement for depression is reduced from 31\% to $15 \%$ (Table 6a, row $z$ ) and the QoL decrement for anti-depressant maintenance therapy is reduced from $8 \%$ to $0 \%$ (i.e. no decrement) (Table 6 , row $b g$ ): $\mathrm{CPB}=56$
- Assume the QoL decrement for depression is increased from 31\% to $45 \%$ (Table 6a, row $z$ ) and the QoL decrement for anti-depressant maintenance therapy is increased from $8 \%$ to $26 \%$ (Table 6a, row $b g$ ): CPB $=98$
- Assume that the screening rate is only applied to one visit per year per patient, rather than 1.75 (Table 6a, row $a g$ ): CPB $=48$


## CPB for Females

Based on the above assumptions for females, the CPB associated with screening for major depressive disorder in female adolescents' ages 12 to 18 is 135 QALYs (see Table 6 b , row $d a$ ).

Table 6b: CPB of Screening for MDD in Female Adolescents Ages 12-18 In a BC Birth Cohort of 40,000

| Row <br> Label | Variable | Base case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Number of life years, 12 year olds | 19,911 | BC Life Table |
| b | Annual rate of MDD, 12 year olds | 5.2\% | V |
| c | Life years with MDD, 12 year olds | 1,035 | = ${ }^{*} \mathrm{~b}$ |
| d | Life years without MDD, 12 year olds | 18,876 | $=a-c$ |
| e | Number of life years, 13 year olds | 19,910 | BC Life Table |
| f | Annual rate of MDD, 13 year olds | 9.3\% | $\checkmark$ |
| g | Life years with MDD, 13 year olds | 1,852 | $=e^{*} \mathrm{f}$ |
| h | Life years without MDD, 13 year olds | 18,059 | =e-g |
| i | Number of life years, 14 year olds | 19,909 | BC Life Table |
| j | Annual rate of MDD, 14 year olds | 11.7\% | $\checkmark$ |
| k | Life years with MDD, 14 year olds | 2,329 | $={ }^{*}{ }^{\text {j }}$ |
| I | Life years without MDD, 14 year olds | 17,580 | $=\mathrm{i}-\mathrm{k}$ |
| m | Number of life years, 15 year olds | 19,907 | BC Life Table |
| n | Annual rate of MDD, 15 year olds | 15.0\% | $\checkmark$ |
| $\bigcirc$ | Life years with MDD, 15 year olds | 2,986 | $=\mathrm{m}^{*} \mathrm{n}$ |
| p | Life years without MDD, 15 year olds | 16,921 | = m - 0 |
| q | Number of life years, 16 year olds | 19,904 | BC Life Table |
| r | Annual rate of MDD, 16 year olds | 16.0\% | $\checkmark$ |
| s | Life years with MDD, 16 year olds | 3,185 | $=q^{*} \mathrm{r}$ |
| t | Life years without MDD, 16 year olds | 16,720 | $=q-s$ |
| $u$ | Number of life years, 17 and 18 year olds | 39,794 | BC Life Table |
| v | Annual rate of MDD, 17 and 18 year olds | 16.5\% | $\checkmark$ |
| w | Life years with MDD, 17 and 18 year olds | 6,566 | = ${ }^{*}$ v |
| x | Life years without MDD, 17 and 18 year olds | 33,228 | = $u$ - w |
| y | Life years with MDD between 12 and 18 | 17,953 | $=c+g+k+o+s+w$ |
| z | Qol decrement due to depression | 0.31 | $\checkmark$ |
| aa | QALYs lost during adolescence due to depression | 5,565 | = ${ }^{*} \mathrm{z}$ |
| ab | Deaths attributable to ISH between the ages of 12 and 34 | 20 | Table 5 |
| ac | QALYS lost due to deaths attributable to ISH between the ages of 12 and 34 | 1,030 | Table 5 |
| ad | Total QALYs lost due to depression in adolescence | 6,596 | $=a \mathrm{a}+\mathrm{ac}$ |
| ae | \% MDD undetected in lifetime | 25.0\% | $\checkmark$ |
| af | Life years with undetected MDD in cohort between 12-18 years of age | 4,488 | = ${ }^{*}$ ae |
| ag | Number of well care visits per year | 2.42 | V |
| ah | Depression screening rate | 8.0\% | V |
| ai | Sensitivity (rate of true positives), initial test | 73.0\% | $\checkmark$ |
| aj | Specificity (rate of true negatives), initial test | 94.0\% | $\checkmark$ |
| ak | Number of MDD cases correctly identified, initial test | 630 | = af * ag *ah * ai |
| al | Number of MDD cases diagnosed incorrectly, initial test | 1,401 | $=(\mathrm{d}+\mathrm{h}+1+\mathrm{p}+\mathrm{t}+\mathrm{x}) * \mathrm{ag} * \mathrm{ah} *(1-\mathrm{aj})$ |
| am | Sensitivity (rate of true positives), 2nd test | 100.0\% | No second test in base model |
| an | Specificity (rate of true negatives), 2nd test | 0.0\% | No second test in base model |
|  | Incorrectly Diagnosed MDD cases |  |  |
| ao | Number of MDD cases diagnosed incorrectly, overall | 1,401 | = al * (1-an) |
| ap | Rate of anti-depressants, months 0-3 | 20.9\% | V |
| aq | Number taking anti-depressants months 0-3 | 293 | = ao * ap |
| ar | Rate of anti-depressants, months 4-6 | 23.6\% | $\checkmark$ |
| as | Number taking anti-depressants months 4-6 | 331 | = ao * $\mathrm{ar}^{\text {r }}$ |
| at | Life years on anti-depressants | 156 | $=(\mathrm{aq} * 0.25)+(\mathrm{as} * 0.25)$ |
| au | Qol decrement due to antidepressant therapy | 0.08 | $\checkmark$ |
| av | QALYs Gained (or Lost), Incorrectly Diagnosed MDD | -12.5 | =- (at * au) |

Table 6b: CPB of Screening for MDD in Female Adolescents Ages 12-18
In a BC Birth Cohort of 40,000

|  | Correctly Diagnosed MDD cases |  |  |
| :---: | :---: | :---: | :---: |
|  | Single Event MDD |  |  |
| aw | Number of MDD cases correctly identified, overall | 630 | = ak * am |
| ax | Rate of single event MDD in correct diagnoses | 50.0\% | $\checkmark$ |
| ay | Number of single event MDD cases | 315 | = aw * $\mathrm{ax}^{\text {a }}$ |
| az | Rate of 6-month antidepressant use | 23.6\% | $\checkmark$ |
| ba | Number on anti-depressants | 74 | = ay * az |
| bb | Clinical improvement rate due to anti-depressants | 25.7\% | V |
| bc | Length of single event MDD, years | 0.5 | $\checkmark$ |
| bd | Depression-free life years gained due to anti-depressants | 9.6 | $=a b * b b^{*} \mathrm{bc}$ |
| be | Treatment seeking rate | 52.0\% | $\checkmark$ |
| bf | Rate counselling among treatment seekers | 50.7\% | $\checkmark$ |
| bg | Overall counselling rate | 26.4\% | $=\mathrm{be}$ * bf |
| bh | Number in counselling | 83 | = ay * bg |
| bi | Clinical improvement rate due to counselling | 12.1\% | $\checkmark$ |
| bj | Length of single event MDD counselling, years | 0.25 | $\checkmark$ |
| bk | Depression-free life years gained due to counselling | 2.5 | $=\mathrm{bh}{ }^{*} \mathrm{bi}^{*} \mathrm{bj}$ |
|  | Recurrent MDD |  |  |
| bl | Number of MDD cases correctly identified, overall | 630 | = ak * am |
| bm | Rate of recurrent MDD in correct diagnoses | 44.3\% | $\checkmark$ |
| bn | Number of recurrent MDD cases | 279 | $=\mathrm{bl}$ * bm |
| bo | Rate of 12-month antidepressant use | 23.6\% | $\checkmark$ |
| bp | Number on anti-depressants | 66 | $=\mathrm{bn}$ * bo |
| bq | Clinical improvement rate due to anti-depressants | 25.7\% | $\checkmark$ |
| br | Length of recurrent MDD event, years | 1.0 | $\checkmark$ |
| bs | Number of recurrent episodes, lifetime | 8.0 | $\checkmark$ |
| bt | Depression-free life years gained due to anti-depressants | 135 | $=\mathrm{bp}$ * bq * br * bs |
| bu | Treatment seeking rate | 52.0\% | $\checkmark$ |
| bv | Rate counselling among treatment seekers | 50.7\% | $\checkmark$ |
| bw | Overall counselling rate | 26.4\% | = bu* bv |
| bx | Number in counselling | 74 | = bn * bw |
| by | Clinical improvement rate due to counselling | 12.1\% | $\checkmark$ |
| bz | Length of recurrent MDD counselling, years | 0.75 | $\checkmark$ |
| ca | Depression-free life years gained due to counselling | 53 | = bx*by *bz*bs |
|  | Persistent MDD |  |  |
| cb | Number of MDD cases correctly identified, overall | 630 | = ak * am |
| cc | Rate of persistent MDD in correct diagnoses | 5.7\% | $\checkmark$ |
| cd | Number of persistent MDD cases | 36 | = cb * cc |
| ce | Rate of first year antidepressant use | 23.6\% | $\checkmark$ |
| cf | Number on anti-depressants | 8 | = cd * ce |
| cg | Clinical improvement rate due to anti-depressants | 25.7\% | $\checkmark$ |
| ch | Length of treatment | 1.0 | $\checkmark$ |
| ci | Depression-free life years gained due to anti-depressants, year 1 | 2.2 | $=\mathrm{cf} * \mathrm{cg}^{*} \mathrm{ch}$ |
| cj | Rate of antidepressant use years 2-20 | 100.0\% | $\checkmark$ |
| ck | Number on anti-depressants | 36 | = $\mathrm{cd}^{*} \mathrm{cj}$ |
| cl | Clinical improvement rate due to anti-depressants | 25.7\% | $\checkmark$ |
| cm | Length of treatment | 19.0 | $\checkmark$ |
| cn | Depression-free life years gained due to anti-depressants, years 2-20 | 175 | $=\mathrm{ck} * \mathrm{cl}^{*} \mathrm{~cm}$ |
| co | Treatment seeking rate | 52.0\% | $\checkmark$ |
| cp | Rate counselling among treatment seekers | 50.7\% | $\checkmark$ |
| cq | Overall counselling rate | 26.4\% | = $\mathrm{co} *{ }^{*} \mathrm{cp}$ |
| cr | Number in counselling | 9 | $=\mathrm{cd} * \mathrm{cq}$ |
| cs | Clinical improvement rate due to counselling | 12.1\% | $\checkmark$ |
| ct | Length of effect persistent event MDD counselling, years | 20.0 | $\checkmark$ |
| cu | Depression-free life years gained due to counselling | 23 | $=\mathrm{cr}$ * cs * ct |
|  | Summary of QALYs Gained with Screening |  |  |
| cv | Individuals with MDD helped by treatment | 58 | $=\mathrm{aw}$ * ( az * bb) + (bg * bi)) |
| cw | Depression free life years due to screening, correctly diagnosed MDD | 402 | $=(\mathrm{bd}+\mathrm{bk})+(\mathrm{bt}+\mathrm{ca})+(\mathrm{ci}+\mathrm{cn}+\mathrm{cu})$ |
| CX | Reduction in \% of total life years with MDD due to screening | 2.24\% | = cw $/ \mathrm{y}$ |
| cy | QALYs gained due to screening, correctly diagnosed MDD | 148 | = cx* ad |
| cz | QALYs due to treating incorrectly diagnosed MDD | -12 | = av |
| da | Net QALYs as a result of screening (CPB) | 135 | = cy +cz |

For the sensitivity analysis of the base model for females, we modified a number of major assumptions and recalculated the CPB as follows:

- Assume the rate of undetected MDD decreases from $25 \%$ to $15 \%$ (Table 6 b , row $a e$ ): $\mathrm{CPB}=76$
- Assume the rate of undetected MDD increases from $25 \%$ to $35 \%$ (Table 6 b, row $a e$ ): CPB $=194$
- Assume a second round of screening (with BDI) is introduced, with a sensitivity of $86.9 \%$ and a specificity of $83.5 \%$ (Table 6 b , rows am \& an ): $\mathrm{CPB}=126$
- Assume the rate of treatment seeking increases from $52.0 \%$ to $70.7 \%$ (Table 6 b, row $a q): \mathrm{CPB}=145$
- Assume the rate of treatment seeking decreases from $52.0 \%$ to $33.3 \%$ (Table 6 b, row $a q): \mathrm{CPB}=125$
- Assume the QoL decrement for depression is reduced from $31 \%$ to $15 \%$ (Table 6b, row $z$ ) and the QoL decrement for anti-depressant maintenance therapy is reduced from $8 \%$ to $0 \%$ (i.e. no decrement) (Table 6 b, row $b g$ ): $\mathrm{CPB}=83$
- Assume the QoL decrement for depression is increased from 31\% to $45 \%$ (Table 6b, row $z$ ) and the QoL decrement for anti-depressant maintenance therapy is increased from $8 \%$ to $26 \%$ (Table 6 , row $b g$ ): $\mathrm{CPB}=163$
- Assume that the screening rate is only applied to one visit per year per patient, rather than 2.42 (Table 6 , row $a g$ ): $\mathrm{CPB}=56$


## Modelling Cost-Effectiveness

In this section, we will calculate the CE associated with screening for major depressive disorder in adolescents.

In modelling CE, we made the following assumptions:

- An adolescent depression screening rate of 7.4\% (Table 7, row $c$ ), completed at each well-care visit, or 2.07 times per year (Table 7 , row $b$ ), ${ }^{93}$ during the seven years of an adolescent's life between 12 and 18 years of age. We model the number available for screening as the sum of adolescents of each age in the cohort (Table 7, row $a$ ).
- The cost of each 10 -minute primary care provider office visit is $\$ 34.85$ (see Reference Document) (Table 7, row $e$ ).
- The value of patient time for each visit to a primary care office is $\$ 59.38$ (see Reference Document) (Table 7, row $f$ ).
- The proportion of each office visit attributable to screening is $50 \%$ (see Reference Document) (Table 7, row $g$ ).
- If a second screening is applied (Table 7, row $k$ ), then all individuals with a positive screen on the first test make another visit to their primary care provider for the second screen. $50 \%$ of the office visit time is assumed to be used for the second screen (Table 7, row $g$ ).
- Both the PHQ-A ${ }^{94}$ and BDI are available online. The PHQ-A is free, but the BDI is copyright (though unlicensed copies exist online) and therefore each use of the BDI is considered to occur through properly licensed channels and cost $\$ 4.40$ per use (Table 7, row $n$ ). ${ }^{95}$
- We have assumed that each positive depression diagnosis results in one (1) follow-up visit to the primary care provider. It is assumed that the entire visit is devoted to the depression diagnosis ( $100 \%$ of office visit cost and patient cost) (Table 7, row $r$ ).
- We have assumed that each depression diagnosis resulting in a course of antidepressant medication results in two (2) additional visits to a primary care provider to monitor prescription effectiveness (Table 7, row $a b$ ).
- We model treatment for those with a positive MDD screen by time period as follows:
- 0-3 months after screening: $19.7 \%$ of positive screened adolescents are taking anti-depressants (Table 7, row $t$ ).
- For males this rate is $17.5 \%$ (Table 7a, row $t$ )
- For females this rate is $20.9 \%$ (Table 7b, row $t$ )
- 4-6 months after screening: $22.2 \%$ of positive screen adolescents are taking anti-depressants and $25.6 \%$ are in counselling or therapy (Table 7 row $a d$ ), with half of the therapy group in individual sessions and half in group sessions.

[^21]- For males the counselling rate is $22.1 \%$ (Table 7a row $a d$ ).
- For females the counselling rate is $26.4 \%$ (Table 7 b row $a d$ ).
- 7-12 months after screening: $22.2 \%$ of correctly diagnosed adolescents with recurrent or persistent MDD are on anti-depressants and $25.6 \%$ are in counselling or therapy, with half of the therapy group in individual sessions and half in group sessions (To avoid double-counting, counselling for these individuals is modelled in the $4-6$ month time period).
- 13+ months after screening: all of the correctly diagnosed adolescents with persistent MDD are on anti-depressants. We assume that the $25.6 \%$ in counselling or therapy receive four (4) individual sessions annually (Table 7 row $b k$ ).
- For males the counselling rate is $22.1 \%$ (Table 7a row $b k$ ).
- For females the counselling rate is $26.4 \%$ (Table 7 b row $b k$ ).
- Recurrent MDD: for each year of recurrent MDD, $22.2 \%$ of individuals with recurrent MDD take anti-depressants and $25.6 \%$ receive therapy (Table 7 row $c c$ ).
- For males the counselling rate is $22.1 \%$ (Table 7a row $c c$ ).
- For females the counselling rate is $26.4 \%$ (Table 7 b row $c c$ ).

|  |  | True Positive Screens |  |  | False Positive Screens |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Single Event | Recurrent | Persistent |  |
| 0-3 Months | Pharmacological | 19.7\% anti-depressant rate |  |  |  |
|  | Therapeutic | None |  |  |  |
| 4-6 Months | Pharmacological | 22.2\% anti-depressant rate |  |  |  |
|  | Therapeutic | 25.6\% receiving therapy |  |  |  |
| 7-12 Months | Pharmacological | No treatment | 22.2\% anti-depressant rate |  | No treatment |
|  | Therapeutic |  | 25.6\% receiving therapy |  |  |
| 13+ Months | Pharmacological |  | No Treatment | 100\% antidepressant rate |  |
|  | Therapeutic |  |  | $25.6 \%$ receiving therapy |  |

- $50 \%$ of the MDD cases are single events and $50 \%$ will be recurrent (Table 7, row $a x$ ), split into $5.3 \%$ (Table 7 , row $b f$ ) of the total that are persistent (i.e. requiring continuing treatment) and $44.7 \%$ of the total that occur on a recurrent basis (Table 7, row $b u$ ).
- For males, $50 \%$ of MDD cases will be recurrent (Table 7a, row $a x$ ), split into $4.7 \%$ (Table 7a, row $b f$ ) of the total that are persistent (i.e. requiring continuing treatment) and $45.3 \%$ of the total that occur on a recurrent basis (Table 7a, row $b u$ ).
- For females, $50 \%$ of MDD cases will be recurrent (Table 7, row $a x$ ), split into 5.7\% (Table 7, row $b f$ ) of the total that are persistent (i.e. requiring continuing treatment) and $44.3 \%$ of the total that occur on a recurrent basis (Table 7, row $b u$ ).
- Each patient with persistent MDD visits their primary care provider an additional 2 times each year for mental health related matters. ${ }^{96,97}$ (Table 7, row bs)
- Treatment length for persistent MDD is modelled at 20 years, in keeping with Tables 4 \& 5 .
- For recurrent cases, there are an additional 7 episodes after the index MDD episode (Table 7, row $b w$ ). For discounting purposes, we model these as occurring eight years apart throughout the lifetime of the affected individuals.
- When group CBT is given, it is typically provided in a group setting of 10 individuals and lasts between $10-15$ sessions. Each session is approximately 1.5 hours long (Table 7, row an). ${ }^{98}$
- We assume one hour of total travel time per patient to attend each CBT session (Table 7, row ao).
- We assume that each session is provided by a grade VI clinical social worker, Level 16 with 6 years of experience. We assume $25 \%$ benefits and $40 \%$ non-worked hours and a wage rate of $\$ 48.65 / \mathrm{hr}^{99}$ for a total cost per worked hour of $\$ 80.27$ ( $\$ 48.65+$ $(\$ 48.65 * 0.25)+(\$ 48.65 * 0.40)$ ).
- We assume that each of 12 group CBT sessions lasts 1.5 hours and that the preparation time is also 1.5 hours, for a total cost of $\$ 240.82$ ( 3 hours * $\$ 80.27$ ) per session for the clinical social worker (Table 7, row ai, bm \& ch).
- We model that half ( $50 \%$ ) of adolescents receiving counselling interventions receive 12 group CBT sessions (Table 7, rows aq) lasting 1.5 hours in groups of 10 (Table 7, rows $a r$ ) for their initial sessions. Subsequent CBT requirements as a result of recurring MDD are reduced to 5 sessions each time (Table 7, row $c p$ ).
- We model that the other half ( $50 \%$ ) of adolescents receiving counselling interventions receive 12 individual counselling sessions with a clinical social worker (Table 7, rows $a h$ ). These sessions also last 1.5 hours.
- Individuals with persistent MDD receive four sessions of individual counselling each year (Table 7, row $b l$ ).
- March and colleagues' report, upon which the USPSTF recommendation was based, started the treatment at 10 mg of fluoxetine daily, increased to $20 \mathrm{mg} /$ day after one week and, if necessary, up to a maximum of $40 \mathrm{mg} /$ day by week 8 of the twelve week trial. ${ }^{100}$

[^22]- Fluoxetine is available in 10 mg and 20 mg doses. ${ }^{101}$ We model daily treatment with 20mg fluoxetine (or generic equivalent). The cost ranges between $\$ 0.35-0.88$ per 20 mg pill for the "BC, Canada" and "Vancouver, BC" geographies. The dispensing fee ranges from $\$ 10-13.99 .{ }^{102}$ Using the mid-point of the above ranges and assuming a 30 -day dose is dispensed each time, the modelled annual cost of treatment is $\$ 368.48((\$ 0.615 * 365)+(12 * \$ 12.00))$ (Table 7, row $a j)$. Using the high and low numbers of the ranges above, we use a high of $\$ 489$ and low of $\$ 248$ / year in our sensitivity analysis.
- Clayton and Barcelo estimated the direct costs associated with a completed suicide in the province of New Brunswick to be $\$ 5,693$ (in 1996 CAD) or $\$ 8,129$ in 2017 CAD, including ambulance, hospital, physician, autopsy, and funeral services plus the cost of police investigations. ${ }^{103}$
- Kinchin and Doran estimated the direct costs per youth suicide in Australia to be \$9,721 (in 2014 AUD) or \$8,336 in 2017 CAD. ${ }^{104}$
- Shepard et al. estimated that the direct costs per nonfatal suicide attempt are $10 \%$ higher than the direct costs per completed suicide in the US. ${ }^{105}$
- For modelling purposes, we have assumed the direct costs per completed suicide in BC to be $\$ 8,233(\$ 8,129+\$ 8,336 / 2)$ (Table 7, row $d b$ ) and the direct cost per suicide attempt to be $\$ 9,056$ ( $\$ 8,233$ * 1.1) (Table 7, row $d c$ ).
- The ratio of attempted suicides to completed suicides among adolescents is estimated to be $50: 1$ to $100: 1 .{ }^{106}$ One-third ( $33 \%$ ) of suicide attempts in adolescents require medical attention. ${ }^{107}$ For modelling purposes, we assumed that there would be 25 attempted suicides requiring medical attention per completed suicide (Table 7, row $d f$ ) (based on the midpoint between 50 and 100 times $33 \%$ ) and varied this from 17 to 33 in the sensitivity analysis.
- In a US study by Wright and colleagues, adolescents ages 13-17 who screened negative for depression utilized $\$ 2,357$ (in 2013 USD) in health care services in the 12 -month period following the screening. By comparison, adolescents who screened positive for moderate to severe depression utilized $\$ 8,173$ in health care services in the 12 -month period following the screening. ${ }^{108} \mathrm{We}$ assumed that the difference of $\$ 5,816(\$ 8,173-\$ 2,357)$ would be avoided in those adolescents for whom treatment for MDD was effective. This comes to $\$ 5,251$ (2017) CAD (Table 7, row di).

[^23]
## CE for Both Sexes

Based on these assumptions, the CE associated with screening for major depressive disorder in adolescents ages 12 to 18 is $\$ 28,215 /$ QALY (Table 7, row $d p$ ).

Table 7: CE of Screening for MDD in Adolescents Ages 12-18 In a BC Birth Cohort of 40,000

| Row <br> Label | Variable | Base case | Source |
| :---: | :---: | :---: | :---: |
| a | Life years, 12 to 18 year olds | 278,512 | Table 6, rows a $+\mathrm{e}+\mathrm{l}+\mathrm{m}+\mathrm{q}+\mathrm{u}$ |
| b | Number of well care visits per year | 2.07 | $v$ |
| c | Depression screening rate | 7.4\% | $\checkmark$ |
| d | Number of screens conducted, cohort total | 42,662 | =a*b*c |
| e | Cost of 10 minute office visit | \$34.85 | Ref Doc |
| f | Value of patient time and travel for office visit | \$59.38 | Ref Doc |
| g | Portion of 10-minute visit for screening | 50\% | Ref Doc |
| h | Initial screening cost | \$2,010,042 | $=d^{*}(e+f) * g$ |
| i | Number of MDD cases correctly identified, initial test | 1,003 | Table 6, row ak |
| j | Number of MDD cases diagnosed incorrectly, initial test | 2,230 | Table 6, row al |
| k | Second screen applied | NO | Table 6, row am |
| I | Number to be re-screened | 0 | = i +j (if applicable) |
| m | Cost of second screening test, each | \$4.40 | $\checkmark$ |
| n | Cost of second screening | \$0 | $=1 *((e+f) * g)+\mathrm{m})$ |
| 0 | Number of MDD cases correctly identified, overall | 1,003 | Table 6, row ao |
| $p$ | Number of MDD cases diagnosed incorrectly, overall | 2,230 | Table 6, row ap |
| q | Total number of MDD cases diagnosed | 3,233 | = $0+p$ |
| r | Follow up visits, each diagnosed depression | 1 | Assumed |
| $s$ | Follow up visit cost | \$304,656 | $=q^{*}(e+f) * r$ |
|  | Treatment 0-3 months post diagnosis (All positive screens) |  |  |
| t | Anti-depressant rate, 0-3 months | 19.7\% | $\checkmark$ |
| $u$ | Number on anti-depressants | 637 | $=q^{*} \mathrm{t}$ |
| v | Cost of medication, per year | \$368 | $\checkmark$ |
| w | Cost of medication, 0-3 months | \$58,673 | $=u *{ }^{*} 0.25$ |
|  | Treatment 4-6 months post diagnosis (All positive screens) |  |  |
| x | Anti-depressant rate, 4-6 months | 22.2\% | $\checkmark$ |
| y | Number on anti-depressants | 718 | $=q^{*} \mathrm{x}$ |
| z | Cost of medication, per year | \$368 | $\checkmark$ |
| aa | Cost of medication, 4-6 months | \$66,118 | $=y^{*}{ }^{*} 0.25$ |
| ab | Follow up visits for medication review, per patient | 1 | $\checkmark$ |
| ac | Cost of medication follow-up | \$67,634 | $=y^{*} \mathrm{~b}^{*}(\mathrm{e}+\mathrm{f})$ |
| ad | Counselling rate | 25.6\% | Table 6 |
| ae | Number receiving counselling | 828 | = $\mathrm{q}^{*}$ ad |
| af | Rate of individual counselling | 50.0\% | V |
| ag | Number receiving individual counselling | 414 | = ae * $\mathrm{af}^{\text {f }}$ |
| ah | Number of CBT sessions | 12 | $\checkmark$ |
| ai | Cost of clinical social worker per session | \$240.82 | $\checkmark$ |
| aj | Cost of offering individual CBT (social worker) | \$1,196,090 | = ag * ah * ai |
| ak | Session length, in hours | 1.5 | V |
| al | Travel time, in hours | 1.0 | $\checkmark$ |
| am | Patient time, cost per hour | \$29.69 | Ref Doc |
| an | Patient time cost, individual CBT treatment sessions | \$368,656 | = ag * $\mathrm{ah} *(\mathrm{ak}+\mathrm{al})$ * am |
| ao | Rate of group counselling | 50.0\% | $\checkmark$ |
| ap | Number receiving individual counselling | 414 | = ae * ao |
| aq | Number of CBT sessions | 12 | $\checkmark$ |
| ar | Number of individuals in each session | 10 | $\checkmark$ |
| as | Cost of offering group CBT (social worker) | \$119,609 | $=(\mathrm{ap} / \mathrm{ar}) * \mathrm{aq}$ * ai |
| at | Session length, in hours | 1.5 | $\checkmark$ |
| au | Travel time, in hours | 1.0 | $\checkmark$ |
| av | Patient time cost per hour | \$29.69 | Ref Doc |
| aw | Patient time cost, group CBT treatment sessions | \$368,656 | = ap *aq * $\mathrm{at}+\mathrm{au}$ ) * av |
|  | Treatment 7-12 months post diagnosis (recurrent and persistent MDD only) |  |  |
| ax | Rate of recurrent and persistent MDD, correctly diagnosed | 50.0\% | $\checkmark$ |
| ay | Anti-depressant rate, 7-12 months | 22.2\% | $\checkmark$ |
| az | Number on anti-depressants | 111 | =o*ax*ay |
| ba | Cost of medication, per year | \$368 | $\checkmark$ |
| bb | Cost of medication, 7-12 months | \$20,515 | = az * ba * 0.5 |
| bc | Counselling costs | \$0 | Included in 4-6 month counselling costs |

Table 7: CE of Screening for MDD in Adolescents Ages 12-18
In a BC Birth Cohort of 40,000

|  | Treatment 13+ months post diagnosis (persistent MDD only) |  |  |
| :---: | :---: | :---: | :---: |
| be | Anti-depressant rate, 13+ months | 100.0\% | $\checkmark$ |
| bf | Rate of persistent MDD, correctly diagnosed | 5.3\% | $\checkmark$ |
| bg | Number on anti-depressants | 53 | = ${ }^{*}$ be * bf |
| bh | Cost of medication, per year | \$368 | $\checkmark$ |
| bi | Additional years of medication | 19 | $\checkmark$ |
| bj | Cost of medication, 2-20 years | \$374,198 | $=\mathrm{bg}$ * bh * bi |
| bk | Counselling rate, for persistent MDD | 25.6\% | V |
| bl | Number of CBT sessions, per year | 4 | $\checkmark$ |
| bm | Cost of clinical social worker per session | \$240.82 | $\checkmark$ |
| bn | Cost of offering individual CBT (social worker), years 2-20 | \$250,464 | $=\mathrm{bg} * \mathrm{bi} * \mathrm{bl}$ * bk *bm |
| bo | Session length, in hours | 1.5 | $\checkmark$ |
| bp | Travel time, in hours | 1.0 | $\checkmark$ |
| bq | Patient time cost per hour | \$29.69 | Ref Doc |
| br | Patient time cost, first CBT treatment sessions | \$301,512 | $=\mathrm{bg} * \mathrm{bi} * \mathrm{bl}{ }^{*}(\mathrm{bo}+\mathrm{bp}) * \mathrm{bq}$ |
| bs | Additional physician visits due to anti-depressant medication, each year | 2 | $V$ |
| bt | Cost of additional physician visits, persistent MDD | \$191,387 | $=\mathrm{bg} * \mathrm{bi}^{*} \mathrm{bs} *(\mathrm{e}+\mathrm{f})$ |
|  | Treatment for Recurrent MDD (after index event) |  |  |
| bu | Rate of recurrent MDD, correctly diagnosed | 44.7\% | $\checkmark$ |
| bv | Number of individuals with recurrent MDD | 448 | =o * bu |
| bw | Number of additional recurrent MDD events after index event | 7 | $\checkmark$ |
| bx | Length of each recurrent MDD event, years | 1 | $\checkmark$ |
| by | Anti-depressant rate, recurrent MDD | 22.2\% | $\checkmark$ |
| bz | Number on anti-depressants | 99 | $=\mathrm{bv}^{*} \mathrm{by}$ |
| ca | Cost of medication, per year | \$368 | $\checkmark$ |
| cb | Cost of medication, recurrent MDD | \$256,608 | = bz* ca* bw * bx |
| cc | Counselling rate, for recurrent MDD | 25.6\% | $\checkmark$ |
| cd | Number individuals in therapy, per recurrent MDD event | 115 | $=\mathrm{bv} * \mathrm{cc}$ |
| ce | Rate of individual counselling | 50.0\% | $\checkmark$ |
| cf | Number receiving individual counselling | 57 | = cd * ce |
| cg | Number of CBT sessions | 5 | V |
| ch | Cost of clinical social worker per session | \$240.82 | $\checkmark$ |
| ci | Cost of offering individual CBT (social worker) | \$483,550 | =cf*cg*ch*bw |
| cj | Session length, in hours | 1.5 | $\checkmark$ |
| ck | Travel time, in hours | 1.0 | $\checkmark$ |
| cl | Patient time cost per hour | \$29.69 | Ref Doc |
| cm | Patient time cost, individual CBT sessions, recurrent MDD | \$149,039 | = cf * cg * (cj +ck ) * cl ${ }^{\text {b }} \mathrm{bw}$ |
| cn | Rate of group counselling | 50.0\% | $\checkmark$ |
| co | Number receiving group counselling | 57 | = $\mathrm{cd}^{*} \mathrm{cn}$ |
| cp | Number of CBT sessions | 5 | $\checkmark$ |
| cq | Number of individuals in each session | 10 | $\checkmark$ |
| cr | Cost of offering group CBT (social worker) | \$48,355 | $=(\mathrm{co} / \mathrm{cq}) * \mathrm{cp} * \mathrm{ch} * \mathrm{bw}$ |
| cs | Session length, in hours | 1.5 | V |
| ct | Travel time, in hours | 1.0 | $\checkmark$ |
| cu | Patient time cost per hour | \$29.69 | Ref Doc |
| cv | Patient time cost, group CBT, recurrent MDD | \$149,039 | $=\mathrm{co} \mathrm{*} \mathrm{cp} \mathrm{*} \mathrm{cs}+\mathrm{ct}$ ) ${ }^{\text {cu }}$ * bw |
| cw | Sub-total, Screening \& Screening Follow-up Cost | \$2,314,698 | $=h+n+s$ |
| CX | Sub-total, Medication and Medication Follow-up Cost | \$1,035,133 | $=w+a a+a c+b b+b j+b t+c b$ |
| cy | Sub-total, Individual Counselling Cost | \$2,749,310 | $=a j+a n+b n+b r+c i+c m$ |
| cz | Sub-total, Group Counselling Cost | \$685,659 | $=a s+a w+c r+c v$ |
| da | Total Cost of Intervention | \$6,784,800 | = cw $+\mathrm{cx}+\mathrm{cy}+\mathrm{cz}$ |
|  | Potential Costs Avoided |  |  |
| db | Direct costs per completed suicide | \$8,233 | $\checkmark$ |
| dc | Direct cost per attempted suicide | \$9,056 | $\checkmark$ |
| dd | Completed suicides avoided due to screening | 1.09 | Table 6, row ab * Table 6, row cx |
| de | Costs avoided due to suicides avoided | \$8,988 | $=\mathrm{db}$ * dd |
| df | Attempted suicides requiring medical attention per completed suicide | 25 | $\checkmark$ |
| dg | Costs avoided due to suicide attempts avoided | \$247,171 | $=\mathrm{dc}^{*} \mathrm{dd}^{*} \mathrm{df}$ |
| dh | Number of people for whom treatment is effective | 88.3 | Table 6, row cv |
| di | Health care cost avoided in first 12 months after screening due to effective treatment | \$5,251 | $\checkmark$ |
| dj | Health care cost avoided, total | \$463,735 | $=\mathrm{dh} * \mathrm{di}$ |
| dk | Net Costs of Intervention | \$6,064,907 | $=\mathrm{da}-\mathrm{de}-\mathrm{dg}-\mathrm{dj}$ |
| dl | Net QALYs Gained | 221.9 | Table 6, row da |
| dm | Cost Effectiveness (CE) of Intervention, \$/QALY | \$27,331 | = $\mathrm{dk} / \mathrm{dl}$ |
| dn | Net Cost of Intervention (1.5\% Discount) | \$5,375,723 | Calculated |
| do | Net QALYs Gained (1.5\% Discount) | 190.5 | Calculated |
| dp | Cost Effectiveness (CE) of Intervention, \$/QALY (1.5\% Discount) | \$28,215 | = dn / do |

$V=$ Estimates from the literature

For the sensitivity analysis, we modified a number of major assumptions and recalculated the CE as follows:

- Assume the rate of undetected MDD decreases from $25 \%$ to $15 \%$ (Table 6, row $a e$ ): $\mathrm{CE}=\$ 43,932$
- Assume the rate of undetected MDD increases from $25 \%$ to $35 \%$ (Table 6, row $a e$ ): CE $=\$ 22,091$
- Assume a second round of screening (with BDI) is introduced, with a sensitivity of $86.9 \%$ and a specificity of $83.5 \%$ (Table 6 , rows $a m$ \& $a m$ ): $\mathrm{CE}=\$ 21,555$
- Assume the rate of treatment seeking increases from $50.5 \%$ to $69 \%$ (Table 6, row aq): $\mathrm{CE}=\$ 30,645$
- Assume the rate of treatment seeking decreases from $50.5 \%$ to $32 \%$ (Table 6 , row aq): $\mathrm{CE}=\$ 25,361$
- Assume the QoL decrement for depression is reduced from 31\% to $15 \%$ (Table 6, row $z$ ) and the QoL decrement for anti-depressant maintenance therapy is reduced from $8 \%$ to $0 \%$ (i.e. no decrement) (Table 6, row $b g$ ): $\mathrm{CE}=\$ 45,994$
- Assume QoL decrement for depression is increased from $31 \%$ to $45 \%$ (Table 6, row $z$ ) and the QoL decrement for anti-depressant maintenance therapy is increased from $8 \%$ to $26 \%$ (Table 6, row $b g$ ): $\mathrm{CE}=\$ 23,446$
- Assume number of visits after depression diagnosis increases from 1 to 2 (Table 7, row $r$ ): $\mathrm{CE}=\$ 29,745$
- Assume the cost of medication increases from $\$ 368 /$ year to $\$ 489 /$ year (Table 7, rows $v, z, b a, b h \& c a)$ : $\mathrm{CE}=\$ 29,251$
- Assume the cost of medication decreases from $\$ 368 /$ year to $\$ 248 /$ year (Table 7 , rows $v, z, b a, b h \& c a): \mathrm{CE}=\$ 27,177$
- Assume the number of suicide attempts per completed suicide is increased from 25 to 33 (Table 7, row $d f$ ): $\mathrm{CE}=\$ 27,869$
- Assume the number of suicide attempts per completed suicide is reduced from 25 to 17 (Table 7, row $d f$ ): $\mathrm{CE}=\$ 28,561$
- Assume the direct cost of completed suicide doubles from $\$ 8,233$ to $\$ 16,466$ (Table 7, row $d b$ ) and the direct cost of attempted suicide doubles from $\$ 9,056$ to $\$ 18,112$ (Table 7, row $d c$ ): $\mathrm{CE}=\$ 27,094$
- Assume that the screening rate is only applied to one visit per year per patient, rather than 2.07 (Table 6, row $a g$ ): $\mathrm{CE}=\$ 28,215$ (i.e. no change)


## CE for Males

Based on the above assumptions for males, the CE associated with screening for major depressive disorder in male adolescents' ages 12 to 18 is $\$ 27,595$ (see Table 7a, row $d p$ ).

| Table 7a: CE of Screening for MDD in Male Adolescents Ages 12-18 In a BC Birth Cohort of 40,000 |  |  |  |
| :---: | :---: | :---: | :---: |
| Row <br> Label | Variable | Base case | Source |
| a | Life years, 12 to 18 year olds | 139,204 | Table 6, rows a + e $+1+m+q+u$ |
| b | Number of well care visits per year | 1.75 | $\checkmark$ |
| c | Depression screening rate | 6.9\% | $\checkmark$ |
| d | Number of screens conducted, cohort total | 16,809 | $=a^{*}{ }^{*} \mathrm{c}$ |
| e | Cost of 10 minute office visit | \$34.85 | Ref Doc |
| f | Value of patient time and travel for office visit | \$59.38 | Ref Doc |
| g | Portion of 10-minute visit for screening | 50\% | Ref Doc |
| h | Initial screening cost | \$791,951 | $=d^{*}(e+f) * g$ |
| i | Number of MDD cases correctly identified, initial test | 395 | Table 6, row ak |
| j | Number of MDD cases diagnosed incorrectly, initial test | 879 | Table 6, row al |
| k | Second screen applied | NO | Table 6, row am |
| I | Number to be re-screened | 0 | = i +j (if applicable) |
| m | Cost of second screening test, each | \$4.40 | $\checkmark$ |
| n | Cost of second screening | \$0 | $\left.\left.=1^{*}(()+\mathrm{f}) * \mathrm{~g}\right)+\mathrm{m}\right)$ |
| o | Number of MDD cases correctly identified, overall | 395 | Table 6, row ao |
| p | Number of MDD cases diagnosed incorrectly, overall | 879 | Table 6, row ap |
| q | Total number of MDD cases diagnosed | 1,274 | = $0+p$ |
| r | Follow up visits, each diagnosed depression | 1 | Assumed |
| s | Follow up visit cost | \$120,033 | $=q^{*}(e+f) * r$ |
|  | Treatment 0-3 months post diagnosis (All positive screens) |  |  |
| t | Anti-depressant rate, 0-3 months | 17.5\% | $\checkmark$ |
| $u$ | Number on anti-depressants | 223 | $=q^{*}$ t |
| v | Cost of medication, per year | \$368 | $\checkmark$ |
| w | Cost of medication, 0-3 months | \$20,535 | $=u *{ }^{*} 0.25$ |
|  | Treatment 4-6 months post diagnosis (All positive screens) |  |  |
| x | Anti-depressant rate, 4-6 months | 19.5\% | $\checkmark$ |
| y | Number on anti-depressants | 248 | $=q^{*} \mathrm{x}$ |
| z | Cost of medication, per year | \$368 | $\checkmark$ |
| aa | Cost of medication, 4-6 months | \$22,882 | $=y^{*}{ }^{*} 0.25$ |
| ab | Follow up visits for medication review, per patient | 1 | $\checkmark$ |
| ac | Cost of medication follow-up | \$23,406 | $=y^{*} \mathrm{ab}^{*}(\mathrm{e}+\mathrm{f})$ |
| ad | Counselling rate | 22.1\% | Table 6 |
| ae | Number receiving counselling | 281 | = $\mathrm{q}^{*}$ ad |
| af | Rate of individual counselling | 50.0\% | $\checkmark$ |
| ag | Number receiving individual counselling | 140 | = ae * af |
| ah | Number of CBT sessions | 12 | $\checkmark$ |
| ai | Cost of clinical social worker per session | \$240.82 | $\checkmark$ |
| aj | Cost of offering individual CBT (social worker) | \$405,932 | = ag * $\mathrm{ah}^{*}$ ai |
| ak | Session length, in hours | 1.5 | V |
| al | Travel time, in hours | 1.0 | $\checkmark$ |
| am | Patient time, cost per hour | \$29.69 | Ref Doc |
| an | Patient time cost, individual CBT treatment sessions | \$125,115 | = ag * ah * ( $\mathrm{ak}+\mathrm{al}$ ) * am |
| ao | Rate of group counselling | 50.0\% | $\checkmark$ |
| ap | Number receiving individual counselling | 140 | = ae * ao |
| aq | Number of CBT sessions | 12 | $\checkmark$ |
| ar | Number of individuals in each session | 10 | $\checkmark$ |
| as | Cost of offering group CBT (social worker) | \$40,593 | $=(\mathrm{ap} / \mathrm{ar}) *$ aq * ai |
| at | Session length, in hours | 1.5 | $\checkmark$ |
| au | Travel time, in hours | 1.0 | $\checkmark$ |
| av | Patient time cost per hour | \$29.69 | Ref Doc |
| aw | Patient time cost, group CBT treatment sessions | \$125,115 | = ap *aq * $\mathrm{at}+\mathrm{au}$ ) * av |
|  | Treatment 7-12 months post diagnosis (recurrent and persistent MDD only) |  |  |
| ax | Rate of recurrent and persistent MDD, correctly diagnosed | 50.0\% | $\checkmark$ |
| ay | Anti-depressant rate, 7-12 months | 19.5\% | $\checkmark$ |
| az | Number on anti-depressants | 39 | =o*ax*ay |
| ba | Cost of medication, per year | \$368 | V |
| bb | Cost of medication, 7-12 months | \$7,100 | = az * ba * 0.5 |
| bc | Counselling costs | \$0 | Included in 4-6 month counselling costs |

Table 7a: CE of Screening for MDD in Male Adolescents Ages 12-18
In a BC Birth Cohort of 40,000

|  | Treatment 13+ months post diagnosis (persistent MDD only) |  |  |
| :---: | :---: | :---: | :---: |
| be | Anti-depressant rate, 13+ months | 100.0\% | $\checkmark$ |
| bf | Rate of persistent MDD, correctly diagnosed | 4.7\% | $\checkmark$ |
| bg | Number on anti-depressants | 19 | = o * be * bf |
| bh | Cost of medication, per year | \$368 | $\checkmark$ |
| bi | Additional years of medication | 19 | $\checkmark$ |
| bj | Cost of medication, 2-20 years | \$130,053 | $=\mathrm{bg}$ * $\mathrm{bh} * \mathrm{bi}$ |
| bk | Counselling rate, for persistent MDD | 22.1\% | $\checkmark$ |
| bl | Number of CBT sessions, per year | 4 | $\checkmark$ |
| bm | Cost of clinical social worker per session | \$240.82 | $\checkmark$ |
| bn | Cost of offering individual CBT (social worker), years 2-20 | \$74,983 | $=\mathrm{bg} * \mathrm{bi}^{*} \mathrm{bl}{ }^{*} \mathrm{bk} * \mathrm{bm}$ |
| bo | Session length, in hours | 1.5 | $\checkmark$ |
| bp | Travel time, in hours | 1.0 | $\checkmark$ |
| bq | Patient time cost per hour | \$29.69 | Ref Doc |
| br | Patient time cost, first CBT treatment sessions | \$104,791 | $=\mathrm{bg} * \mathrm{bi}^{*} \mathrm{bl}{ }^{*}(\mathrm{bo}+\mathrm{bp}) * \mathrm{bq}$ |
| bs | Additional physician visits due to anti-depressant medication, each year | 2 | $\frac{1}{}$ |
| bt | Cost of additional physician visits, persistent MDD | \$66,517 | $=\mathrm{bg} * \mathrm{bi}{ }^{*} \mathrm{bs} *(\mathrm{e}+\mathrm{f})$ |
|  | Treatment for Recurrent MDD (after index event) |  |  |
| bu | Rate of recurrent MDD, correctly diagnosed | 45.3\% | $\checkmark$ |
| bv | Number of individuals with recurrent MDD | 179 | = ${ }^{*}$ bu |
| bw | Number of additional recurrent MDD events after index event | 7 | $\checkmark$ |
| bx | Length of each recurrent MDD event, years | 1 | $\checkmark$ |
| by | Anti-depressant rate, recurrent MDD | 19.5\% | $\checkmark$ |
| bz | Number on anti-depressants | 35 | $=\mathrm{bv}$ * by |
| ca | Cost of medication, per year | \$368 | $\checkmark$ |
| cb | Cost of medication, recurrent MDD | \$90,054 | $=\mathrm{bz} * \mathrm{ca} * \mathrm{bw} * \mathrm{bx}$ |
| cc | Counselling rate, for recurrent MDD | 22.1\% | $\checkmark$ |
| cd | Number individuals in therapy, per recurrent MDD event | 39 | $=\mathrm{bv}$ * cc |
| ce | Rate of individual counselling | 50.0\% | $\checkmark$ |
| cf | Number receiving individual counselling | 20 | $=\mathrm{cd}^{*} \mathrm{ce}$ |
| cg | Number of CBT sessions | 5 | $\checkmark$ |
| ch | Cost of clinical social worker per session | \$240.82 | $\checkmark$ |
| ci | Cost of offering individual CBT (social worker) | \$166,413 | $=\mathrm{cf} \mathrm{*} \mathrm{cg}^{*} \mathrm{ch} *$ bw |
| cj | Session length, in hours | 1.5 | $\checkmark$ |
| ck | Travel time, in hours | 1.0 | $\checkmark$ |
| cl | Patient time cost per hour | \$29.69 | Ref Doc |
| cm | Patient time cost, individual CBT sessions, recurrent MDD | \$51,292 | = cf * cg * (cj + ck) * cl * bw |
| cn | Rate of group counselling | 50.0\% | $\checkmark$ |
| co | Number receiving group counselling | 20 | $=\mathrm{cd}^{*} \mathrm{cn}$ |
| cp | Number of CBT sessions | 5 | $\checkmark$ |
| cq | Number of individuals in each session | 10 | $\checkmark$ |
| cr | Cost of offering group CBT (social worker) | \$16,641 | $=(\mathrm{co} / \mathrm{cq}) * \mathrm{cp}^{*} \mathrm{ch} * \mathrm{bw}$ |
| CS | Session length, in hours | 1.5 | $\checkmark$ |
| ct | Travel time, in hours | 1.0 | $\checkmark$ |
| cu | Patient time cost per hour | \$29.69 | Ref Doc |
| cv | Patient time cost, group CBT, recurrent MDD | \$51,292 | $=\mathrm{co}{ }^{*} \mathrm{cp} *(\mathrm{cs}+\mathrm{ct}) * \mathrm{cu} * \mathrm{bw}$ |
| CW | Sub-total, Screening \& Screening Follow-up Cost | \$911,984 | $=h+n+s$ |
| cx | Sub-total, Medication and Medication Follow-up Cost | \$360,547 | $=w+a a+a c+b b+b j+b t+c b$ |
| cy | Sub-total, Individual Counselling Cost | \$928,526 | $=a j+a n+b n+b r+c i+c m$ |
| cz | Sub-total, Group Counselling Cost | \$233,641 | $=a s+a w+c r+c v$ |
| da | Total Cost of Intervention | \$2,434,699 | = cw $+\mathrm{cx}+\mathrm{cy}+\mathrm{cz}$ |
|  | Potential Costs Avoided |  |  |
| db | Direct costs per completed suicide | \$8,233 | $\checkmark$ |
| dc | Direct cost per attempted suicide | \$9,056 | $\checkmark$ |
| dd | Completed suicides avoided due to screening | 0.53 | Table 6, row ab * Table 6, row cx |
| de | Costs avoided due to suicides avoided | \$4,326 | $=\mathrm{db}$ * dd |
| df | Attempted suicides requiring medical attention per completed suicide | 25 | $\checkmark$ |
| dg | Costs avoided due to suicide attempts avoided | \$118,972 | $=\mathrm{dc}^{*} \mathrm{dd}^{*} \mathrm{df}$ |
| dh | Number of people for whom treatment is effective | 30.4 | Table 6, row cv |
| di | Health care cost avoided in first 12 months after screening due to effective treatment | \$5,251 | $\checkmark$ |
| dj | Health care cost avoided, total | \$159,394 | $=\mathrm{dh}^{*} \mathrm{di}$ |
| dk | Net Costs of Intervention | \$2,152,006 | $=\mathrm{da}-\mathrm{de}-\mathrm{dg}-\mathrm{dj}$ |
| d | Net QALYs Gained | 83.1 | Table 6, row da |
| dm | Cost Effectiveness (CE) of Intervention, \$/QALY | \$25,887 | $=\mathrm{dk} / \mathrm{dl}$ |
| dn | Net Cost of Intervention (1.5\% Discount) | \$1,916,383 | Calculated |
| do | Net QALYs Gained (1.5\% Discount) | 69.4 | Calculated |
| dp | Cost Effectiveness (CE) of Intervention, \$/QALY (1.5\% Discount) | \$27,595 | = dn /do |

$V=$ Estimates from the literature

For the sensitivity analysis of the base model for males, we modified a number of major assumptions and recalculated the CE as follows:

- Assume the rate of undetected MDD decreases from $25 \%$ to $15 \%$ (Table 6a, row ae): $C E=\$ 43,386$
- Assume the rate of undetected MDD increases from $25 \%$ to $35 \%$ (Table 6a, row $a e$ ): CE $=\$ 21,415$
- Assume a second round of screening (with BDI) is introduced, with a sensitivity of $86.9 \%$ and a specificity of $83.5 \%$ (Table 6a, rows $a m$ \& $a m$ ): $C E=\$ 21,583$
- Assume the rate of treatment seeking increases from $43.5 \%$ to $65.2 \%$ (Table 6 a, row aq): $\mathrm{CE}=\$ 30,523$
- Assume the rate of treatment seeking decreases from $43.5 \%$ to $21.8 \%$ (Table 6 a , row $a q): \mathrm{CE}=\$ 23,984$
- Assume the QoL decrement for depression is reduced from $31 \%$ to $15 \%$ (Table 6 a , row $z$ ) and the QoL decrement for anti-depressant maintenance therapy is reduced from $8 \%$ to $0 \%$ (i.e. no decrement) (Table 6a, row $b g$ ): $\mathrm{CE}=\$ 43,489$
- Assume QoL decrement for depression is increased from $31 \%$ to $45 \%$ (Table 6a, row $z$ ) and the QoL decrement for anti-depressant maintenance therapy is increased from $8 \%$ to $26 \%$ (Table 6 , row $b g$ ): $\mathrm{CE}=\$ 23,168$
- Assume number of visits after depression diagnosis increases from 1 to 2 (Table 7a, row $r$ ): $\mathrm{CE}=\$ 29,249$
- Assume the cost of medication increases from \$368/year to \$489/year (Table 7a, rows $v, z, b a, b h \& c a): \mathrm{CE}=\$ 28,586$
- Assume the cost of medication decreases from \$368/year to \$248/year (Table 7a, rows $v, z, b a, b h \& c a): \mathrm{CE}=\$ 26,603$
- Assume the number of suicide attempts per completed suicide is increased from 25 to 33 (Table 7a, row $d f$ ): $\mathrm{CE}=\$ 27,138$
- Assume the number of suicide attempts per completed suicide is reduced from 25 to 17 (Table 7a, row $d f$ ): $\mathrm{CE}=\$ 28,052$
- Assume the direct cost of completed suicide doubles from $\$ 8,233$ to $\$ 16,466$ (Table 7 a , row $d b$ ) and the direct cost of attempted suicide doubles from $\$ 9,056$ to $\$ 18,112$ (Table 7a, row $d c$ ): $\mathrm{CE}=\$ 26,116$
- Assume that the screening rate is only applied to one visit per year per patient, rather than 1.75 (Table 6a, row $a g$ ): $\mathrm{CE}=\$ 27,595$ (i.e. no change)


## CE for Females

Based on the above assumptions for males, the CE associated with screening for major depressive disorder in male adolescents' ages 12 to 18 is $\$ 29,368$ (see Table 7 b , row $d p$ ).


Table 7b: CE of Screening for MDD in Female Adolescents Ages 12-18 In a BC Birth Cohort of 40,000

|  | Treatment 13+ months post diagnosis (persistent MDD only) |  |  |
| :---: | :---: | :---: | :---: |
| be | Anti-depressant rate, 13+ months | 100.0\% | $\checkmark$ |
| bf | Rate of persistent MDD, correctly diagnosed | 5.7\% | $\checkmark$ |
| bg | Number on anti-depressants | 36 | = ${ }^{*}$ be * bf |
| bh | Cost of medication, per year | \$368 | $\checkmark$ |
| bi | Additional years of medication | 19 | $\checkmark$ |
| bj | Cost of medication, $2-20$ years | \$251,548 | $=\mathrm{bg}$ * $\mathrm{bh}^{*} \mathrm{bi}$ |
| bk | Counselling rate, for persistent MDD | 26.4\% | $\checkmark$ |
| bl | Number of CBT sessions, per year | 4 | V |
| bm | Cost of clinical social worker per session | \$240.82 | $\checkmark$ |
| bn | Cost of offering individual CBT (social worker), years 2-20 | \$173,371 | $=\mathrm{bg} * \mathrm{bi} * \mathrm{bl}{ }^{\text {b }} \mathrm{bk} * \mathrm{bm}$ |
| bo | Session length, in hours | 1.5 | $\checkmark$ |
| bp | Travel time, in hours | 1.0 | $\checkmark$ |
| bq | Patient time cost per hour | \$29.69 | Ref Doc |
| br | Patient time cost, first CBT treatment sessions | \$202,685 | $=\mathrm{bg} * \mathrm{bi} * \mathrm{bl} *(\mathrm{bo}+\mathrm{bp}) * \mathrm{bq}$ |
| bs | Additional physician visits due to anti-depressant medication, each year | 2 | $V$ |
| bt | Cost of additional physician visits, persistent MDD | \$128,656 | $=\mathrm{bg}$ * $\mathrm{bi}^{*} \mathrm{bs} *(\mathrm{e}+\mathrm{f})$ |
|  | Treatment for Recurrent MDD (after index event) |  |  |
| bu | Rate of recurrent MDD, correctly diagnosed | 44.3\% | $\checkmark$ |
| bv | Number of individuals with recurrent MDD | 279 | =o*bu |
| bw | Number of additional recurrent MDD events after index event | 7 | v |
| bx | Length of each recurrent MDD event, years | 1 | $\checkmark$ |
| by | Anti-depressant rate, recurrent MDD | 23.6\% | $\checkmark$ |
| bz | Number on anti-depressants | 66 | $=\mathrm{bv}^{*}$ by |
| ca | Cost of medication, per year | \$368 | $\checkmark$ |
| cb | Cost of medication, recurrent MDD | \$169,983 | $=\mathrm{bz} * \mathrm{ca}$ * bw * bx |
| cc | Counselling rate, for recurrent MDD | 26.4\% | $\checkmark$ |
| cd | Number individuals in therapy, per recurrent MDD event | 74 | $=\mathrm{bv}^{*} \mathrm{cc}$ |
| ce | Rate of individual counselling | 50.0\% | $\checkmark$ |
| cf | Number receiving individual counselling | 37 | = cd * ce |
| cg | Number of CBT sessions | 5 | $\checkmark$ |
| ch | Cost of clinical social worker per session | \$240.82 | $\checkmark$ |
| ci | Cost of offering individual CBT (social worker) | \$310,262 | $=\mathrm{cf} * \mathrm{cg} * \mathrm{ch} * \mathrm{bw}$ |
| cj | Session length, in hours | 1.5 | $\checkmark$ |
| ck | Travel time, in hours | 1.0 | $\checkmark$ |
| cl | Patient time cost per hour | \$29.69 | Ref Doc |
| cm | Patient time cost, individual CBT sessions, recurrent MDD | \$95,628 | = cf * cg * (cj +ck ) * cl ${ }^{*}$ bw |
| cn | Rate of group counselling | 50.0\% | V |
| co | Number receiving group counselling | 37 | = $\mathrm{cd}^{*} \mathrm{cn}$ |
| cp | Number of CBT sessions | 5 | $\checkmark$ |
| cq | Number of individuals in each session | 10 | $\checkmark$ |
| cr | Cost of offering group CBT (social worker) | \$31,026 | $=(\mathrm{co} / \mathrm{cq}) * \mathrm{cp}{ }^{\text {* ch }}$ * bw |
| cs | Session length, in hours | 1.5 | V |
| ct | Travel time, in hours | 1.0 | $\checkmark$ |
| cu | Patient time cost per hour | \$29.69 | Ref Doc |
| cV | Patient time cost, group CBT, recurrent MDD | \$95,628 | $=\mathrm{co} \mathrm{*} \mathrm{cp} \mathrm{*} \mathrm{(cs}+\mathrm{ct}$ ) * cu* bw |
| cw | Sub-total, Screening \& Screening Follow-up Cost | \$1,454,427 | $=\mathrm{h}+\mathrm{n}+\mathrm{s}$ |
| cx | Sub-total, Medication and Medication Follow-up Cost | \$692,346 | $=w+a a+a c+b b+b j+b t+c b$ |
| cy | Sub-total, Individual Counselling Cost | \$1,794,354 | = aj $+\mathrm{an}+\mathrm{bn}+\mathrm{br}+\mathrm{ci}+\mathrm{cm}$ |
| Cz | Sub-total, Group Counselling Cost | \$442,567 | = as +aw + cr + cv |
| da | Total Cost of Intervention | \$4,383,695 | = cw $+\mathrm{cx}+\mathrm{cy}+\mathrm{cz}$ |
|  | Potential Costs Avoided |  |  |
| db | Direct costs per completed suicide | \$8,233 | V |
| dc | Direct cost per attempted suicide | \$9,056 | $\checkmark$ |
| dd | Completed suicides avoided due to screening | 0.44 | Table 6, row ab * Table 6, row cx |
| de | Costs avoided due to suicides avoided | \$3,627 | $=\mathrm{db}$ * dd |
| df | Attempted suicides requiring medical attention per completed suicide | 25 | $\checkmark$ |
| dg | Costs avoided due to suicide attempts avoided | \$99,741 | $=\mathrm{dc}^{*} \mathrm{dd}$ * df |
| dh | Number of people for whom treatment is effective | 58.3 | Table 6, row cv |
| di | Health care cost avoided in first 12 months after screening due to effective treatment | \$5,251 | V |
| dj | Health care cost avoided, total | \$306,347 | $=\mathrm{dh}^{*} \mathrm{di}$ |
| dk | Net Costs of Intervention | \$3,973,980 | = da - de - dg - dj |
| dl | Net QALYs Gained | 135.1 | Table 6, row da |
| dm | Cost Effectiveness (CE) of Intervention, \$/QALY | \$29,425 | = $\mathrm{dk} / \mathrm{dl}$ |
| dn | Net Cost of Intervention (1.5\% Discount) | \$3,514,247 | Calculated |
| do | Net QALYs Gained (1.5\% Discount) | 119.7 | Calculated |
| dp | Cost Effectiveness (CE) of Intervention, \$/QALY (1.5\% Discount) | \$29,368 | = dn /do |

V = Estimates from the literature

For the sensitivity analysis of the base model for females, we modified a number of major assumptions and recalculated the CE as follows:

- Assume the rate of undetected MDD decreases from $25 \%$ to $15 \%$ (Table 6 b , row $a e$ ): CE $=\$ 45,560$
- Assume the rate of undetected MDD increases from $25 \%$ to $35 \%$ (Table 6b, row $a e$ ): CE $=\$ 23,098$
- Assume a second round of screening (with BDI) is introduced, with a sensitivity of $86.9 \%$ and a specificity of $83.5 \%$ (Table 6 b , rows $\mathrm{am} \& \mathrm{am}$ ): $\mathrm{CE}=\$ 22,321$
- Assume the rate of treatment seeking increases from $52.0 \%$ to $70.7 \%$ (Table 6 b , row $a q): \mathrm{CE}=\$ 31,878$
- Assume the rate of treatment seeking decreases from $52.0 \%$ to $33.3 \%$ (Table 6 b, row $a q): \mathrm{CE}=\$ 26,434$
- Assume the QoL decrement for depression is reduced from $31 \%$ to $15 \%$ (Table 6b, row $z$ ) and the QoL decrement for anti-depressant maintenance therapy is reduced from $8 \%$ to $0 \%$ (i.e. no decrement) (Table 6b, row $b g$ ): $\mathrm{CE}=\$ 49,734$
- Assume QoL decrement for depression is increased from 31\% to 45\% (Table 6b, row $z$ ) and the QoL decrement for anti-depressant maintenance therapy is increased from $8 \%$ to $26 \%$ (Table 6 , row $b g$ ): $\mathrm{CE}=\$ 24,171$
- Assume number of visits after depression diagnosis increases from 1 to 2 (Table 7b, row $r$ ): $\mathrm{CE}=\$ 30,899$
- Assume the cost of medication increases from $\$ 368 /$ year to $\$ 489 /$ year (Table 7 b , row aj): $\mathrm{CE}=\$ 30,472$
- Assume the cost of medication decreases from $\$ 368 /$ year to $\$ 248 /$ year (Table 7 b , row aj): $\mathrm{CE}=\$ 28,264$
- Assume the number of suicide attempts per completed suicide is increased from 25 to 33 (Table 7b, row $d f$ ): $\mathrm{CE}=\$ 29,146$
- Assume the number of suicide attempts per completed suicide is reduced from 25 to 17 (Table 7b, row $d f$ ): $\mathrm{CE}=\$ 29,591$
- Assume the direct cost of completed suicide doubles from $\$ 8,233$ to $\$ 16,466$ (Table 7 b , row $d b$ ) and the direct cost of attempted suicide doubles from $\$ 9,056$ to $\$ 18,112$ (Table 7b, row $d c$ ): $\mathrm{CE}=\$ 28,649$
- Assume that the screening rate is only applied to one visit per year per patient, rather than 2.42 (Table 6b, row $a g$ ): $\mathrm{CE}=\$ 29,368$ (i.e. no change)


## Summary

The clinically preventable burden (CPB) associated with screening for, and treatment of, major depressive disorder (MDD) in adolescents ages 12 to 18 is estimated to be 191 qualityadjusted life years (QALYs) while the cost-effectiveness (CE) is estimated at $\$ 28,215$ per QALY (see Table 8). In male adolescents ages 12-18, the CPB with screening for, and treatment of, MDD is estimated to be 69 QALYs while the CE is estimated at $\$ 27,595$ per QALY (see Table 8a). In female adolescents ages 12-18, the CPB with screening for, and treatment of, MDD is estimated to be 120 QALYs while the CE is estimated at $\$ 29,368$ per QALY (see Table 8 b ).

Table 8: Screening for MDD in Adolescents
Ages 12-18 in a BC Birth Cohort of 40,000
Summary

|  | Base Case | Range |  |
| :---: | :---: | :---: | :---: |
| CPB (Potential QALYs Gained) |  |  |  |
| Assume No Current Service |  |  |  |
| 1.5\% Discount Rate | 191 | 92 | 274 |
| 3\% Discount Rate | 171 | 83 | 247 |
| 0\% Discount Rate | 222 | 107 | 318 |
| CE (\$/QALY) including patient time costs |  |  |  |
| 1.5\% Discount Rate | \$28,215 | \$21,555 | \$45,994 |
| 3\% Discount Rate | \$28,892 | \$21,422 | \$48,789 |
| 0\% Discount Rate | \$27,331 | \$21,661 | \$42,094 |
| CE (\$/QALY) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | \$14,063 | \$9,656 | \$22,925 |
| 3\% Discount Rate | \$14,201 | \$9,298 | \$23,981 |
| 0\% Discount Rate | \$13,998 | \$10,199 | \$21,558 |

Table 8a: Screening for MDD in Male Adolescents Ages 12-18 in a BC Birth Cohort of 40,000

Summary

|  | $\begin{aligned} & \text { Base } \\ & \text { Case } \end{aligned}$ | Range |  |
| :---: | :---: | :---: | :---: |
| CPB (Potential QALYs Gained) |  |  |  |
| Assume No Current Service |  |  |  |
| 1.5\% Discount Rate | 69 | 39 | 100 |
| 3\% Discount Rate | 61 | 34 | 88 |
| 0\% Discount Rate | 83 | 47 | 119 |
| CE (\$/QALY) including patient time costs |  |  |  |
| 1.5\% Discount Rate | \$27,595 | \$21,415 | \$43,489 |
| 3\% Discount Rate | \$28,858 | \$22,004 | \$47,491 |
| 0\% Discount Rate | \$25,887 | \$2,061 | \$38,218 |
| CE (\$/QALY) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | \$13,264 | \$10,301 | \$20,904 |
| 3\% Discount Rate | \$13,693 | \$10,395 | \$22,535 |
| 0\% Discount Rate | \$12,788 | \$10,264 | \$18,879 |

Table 8b: Screening for MDD in Female Adolescents Ages 12-18 in a BC Birth Cohort of $\mathbf{4 0 , 0 0 0}$ Summary Base Case Range CPB (Potential QALYs Gained)

Assume No Current Service

| 1.5\% Discount Rate | 120 | 49 | 173 |
| :---: | :---: | :---: | :---: |
| 3\% Discount Rate | 110 | 45 | 158 |
| 0\% Discount Rate | 135 | 56 | 194 |
| CE (\$/QALY) including patient time costs |  |  |  |
| 1.5\% Discount Rate | \$29,368 | \$22,321 | \$49,734 |
| 3\% Discount Rate | \$29,432 | \$21,724 | \$51,078 |
| 0\% Discount Rate | \$29,425 | \$23,174 | \$47,720 |
| CE (\$/QALY) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | \$14,934 | \$10,282 | \$25,291 |
| 3\% Discount Rate | \$14,742 | \$9,689 | \$25,585 |
| 0\% Discount Rate | \$15,378 | \$11,210 | \$24,940 |

## Behavioural Counselling Interventions

Promotion of Breastfeeding
Canadian Task Force on Preventive Health Care (2004)
Breast-feeding has been shown in both developing and developed countries to improve the health of infants and their mothers, making it the optimal method of infant nutrition.

The CTFPHC concludes that there is good evidence to recommend providing structured antepartum educational programs and postpartum support to promote breastfeeding initiation and duration. (A recommendation)

Unfortunately, advice from a woman's primary clinician (such as family physician, obstetrician or midwife) has not been sufficiently evaluated, and a research gap remains in this area.

The CTFPHC concludes that there is insufficient evidence to make a recommendation regarding advice by primary caregivers to promote breastfeeding. (I Recommendation) ${ }^{109}$

## United States Preventive Services Task Force Recommendations (2008)

The USPSTF recommends interventions during pregnancy and after birth to promote and support breastfeeding. This is a grade B recommendation.

There is convincing evidence that breastfeeding provides substantial health benefits for children and adequate evidence that breastfeeding provides moderate health benefits for women.

Adequate evidence indicates that interventions to promote and support breastfeeding increase the rates of initiation, duration, and exclusivity of breastfeeding.

The USPSTF concludes that there is moderate certainty that interventions to promote and support breastfeeding have a moderate net benefit.

Interventions may include multiple strategies, such as formal breastfeeding education for mothers and families, direct support of mothers during breastfeeding observations, and the training of health professional staff about breastfeeding and techniques for breastfeeding support.

Although the activities of individual clinicians to promote and support breastfeeding are likely to be positive, additional benefit may result from efforts that are integrated into systems of care. ${ }^{110}$

[^24]
## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with interventions aimed at improving longer term ( 6 months) exclusive breastfeeding rates in a British Columbia birth cohort of 40,000.

Breastfeeding promotion interventions in developed countries are associated with a $28 \%$ increase (odds ratio or $\mathrm{OR}=1.28,95 \% \mathrm{CI}$ of $1.11-1.48$ ) in short-term ( $1-3$ months) exclusive breastfeeding and a $44 \%$ increase ( $\mathrm{OR}=1.44,95 \% \mathrm{CI}$ of $1.13-1.84$ ) in long-term (6-8 months) exclusive breastfeeding. ${ }^{111}$

Research evidence does not clearly identify which types or components of breastfeeding promotion interventions are effective. In their review for the USPSTF, Chung and colleagues "did not find that formal or structured breastfeeding education or individual-level professional support significantly affected the breastfeeding outcomes. [They] did find that lay support significantly increased the rate of any and exclusive breastfeeding in the shortterm." They also noted that interventions including both pre- and post-natal components are important. Finally, "the BFHI (Baby Friendly Hospital Initiative) is effective in increasing exclusive breastfeeding rates, at least up to 6 months after delivery." ${ }^{112}$

From the perspective of a CPS, then, it may be most important for the clinician to refer their pregnant patient or new mother to an intervention including lay support.

Breastfeeding is associated with the following health benefits for the infant:

- Any breastfeeding is associated with a $40 \%$ reduction ( $\mathrm{OR}=0.60,95 \% \mathrm{CI}$ of $0.46-$ 0.78 ) in the risk of otitis media (OM) compared to no breastfeeding (Table 2, row $k$ ). ${ }^{113}$ The overall incidence of OM is 1.9 episodes in the first year of life (Table 2, row j). ${ }^{114}$
- Exclusive breastfeeding for 3 months or longer is associated with a $42 \%$ reduction ( $\mathrm{OR}=0.58,95 \% \mathrm{CI}$ of $0.41-0.92$ ) in the risk of atopic dermatitis (AD) compared to exclusive breastfeeding for less than 3 months (Table 2, row $n$ ). ${ }^{115} \mathrm{AD}$ has a cumulative incidence of 0.165 in the first two years of life (Table 2 , row $m$ ). ${ }^{116}$
- Any breastfeeding is associated with a $64 \%$ reduction ( $\mathrm{OR}=0.36,95 \% \mathrm{CI}$ of 0.32 0.41 ) in the risk of gastrointestinal infection (GI) compared to no breastfeeding (Table 2, row $q$ ). ${ }^{117} \mathrm{GI}$ is associated with 0.222 ambulatory visits (Table 2, row $p$ ) and 0.00298 hospitalizations per infant < 1 year old. ${ }^{118}$

[^25]- Exclusive breastfeeding for 4 months or longer is associated with a $72 \%$ reduction ( $\mathrm{OR}=0.28,95 \% \mathrm{CI}$ of $0.14-0.54$ ) in the risk of lower respiratory tract infection (LRTI) compared to formula feeding (Table 2, row $t$ ). ${ }^{119}$ The overall incidence of LRTI in infants is 0.0409 cases (Table 2, row $s$ ) with a death rate of 0.0000732 (Table 2, row $v$ ). ${ }^{120}$
- Breastfeeding for 3 months or longer is associated with a $27 \%$ reduction ( $\mathrm{OR}=0.73$, $95 \%$ CI of $0.59-0.92$ ) in the risk of asthma compared to no breastfeeding in families without a history of asthma (Table 2, row $a a$ ). ${ }^{121}$ The cumulative incidence of asthma during childhood is 0.127 (Table 2, row $z$ ) with a death rate of 0.00000273 (Table 2, row $c c$ ). ${ }^{122}$
- Any breastfeeding is associated with a $24 \%$ reduction ( $\mathrm{OR}=0.76,95 \% \mathrm{CI}$ of 0.67 0.86 ) in the risk of overweight or obesity compared to no breastfeeding (Table 2, row $h h \& m m$ ). Each month of breastfeeding is associated with a $4 \%$ reduced risk of overweight or obesity. ${ }^{123}$ The 2010 rate of overweight and obesity by age group in BC is detailed in Figure $1 .{ }^{124}$ Based on this rate and mean survival rates by age group, a birth cohort of 40,000 in BC would be expected to include 878,446 years in a 'state' of overweight and 348,584 years in a 'state' of obesity (see Table 1).
Overweight/obesity is associated with a reduced life expectancy of approximately 0.6 and 2.6 years, respectively (see Reference Document). Given the average life expectancy in BC of 82.2 years, this represents a reduction in life expectancy of $0.73 \%(0.6$ / 82.2) associated with overweight (Table 2, row $j j$ ) and 3.16\% (2.6 / 82.2) for obesity (Table 2, row oo).

[^26]

Table 1: Years of Life as Overweight or Obese in a Birth Cohort of 40,000

| Age Group | Mean Survival Rate | Years of Life in Birth Cohort | \% Overweight | Years of Life Overweight | \% Obese | Years of Life Obese |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 0-4 | 99.6\% | 199,198 | 11.3\% | 22,572 | 2.9\% | 5,711 |
| 5-9 | 99.5\% | 199,088 | 11.3\% | 22,560 | 2.9\% | 5,708 |
| 10-14 | 99.5\% | 199,022 | 11.3\% | 22,552 | 2.9\% | 5,706 |
| 15-19 | 99.4\% | 198,868 | 14.1\% | 28,034 | 4.0\% | 7,856 |
| 20-24 | 99.2\% | 198,408 | 19.5\% | 38,776 | 10.1\% | 19,990 |
| 25-29 | 98.9\% | 197,850 | 23.2\% | 45,921 | 9.1\% | 18,075 |
| 30-34 | 98.6\% | 197,290 | 25.5\% | 50,330 | 11.1\% | 21,927 |
| 35-39 | 98.3\% | 196,550 | 33.3\% | 65,453 | 10.1\% | 19,818 |
| 40-44 | 97.8\% | 195,526 | 29.6\% | 57,851 | 11.5\% | 22,580 |
| 45-49 | 97.0\% | 194,070 | 32.0\% | 62,018 | 15.0\% | 29,161 |
| 50-54 | 96.0\% | 191,948 | 35.2\% | 67,489 | 18.3\% | 35,177 |
| 55-59 | 94.4\% | 188,786 | 36.6\% | 69,177 | 18.0\% | 34,041 |
| 60-64 | 92.0\% | 183,998 | 44.5\% | 81,961 | 17.4\% | 31,970 |
| 65-69 | 88.3\% | 176,658 | 34.5\% | 60,915 | 15.0\% | 26,517 |
| 70-74 | 82.7\% | 165,362 | 38.2\% | 63,193 | 15.4\% | 25,408 |
| 75-79 | 74.1\% | 148,142 | 36.0\% | 53,308 | 14.3\% | 21,158 |
| 80+ | 59.5\% | 214,284 | 31.0\% | 66,334 | 8.3\% | 17,784 |
| Total |  | 3,245,048 | 27.1\% | 878,446 | 10.7\% | 348,584 |

- Breastfeeding for 3 months or longer is associated with a $19 \%$ reduction ( $\mathrm{OR}=0.81$, $95 \%$ CI of $0.74-0.89$ ) in the risk of type 1 diabetes compared to breastfeeding for less than 3 months (Table 2, row $r$ r). ${ }^{125}$ The overall incidence of type 1 diabetes is 0.000186 (Table 2, row $q q$ ) with a death rate of 0.00000121 (Table 1-2, row $t t$ ). ${ }^{126}$
- Breastfeeding for less than 6 months is associated with a $12 \%$ reduction ( $\mathrm{OR}=0.88$, $95 \%$ CI of $0.80-0.96$ ) in the risk of childhood leukemia while breastfeeding for more than 6 months is associated with a $24 \%$ reduction ( $\mathrm{OR}=0.76,95 \%$ CI of $0.68-$ 0.84 ) in the risk of childhood leukemia compared to no breastfeeding (Table 2, row $y y$ ). ${ }^{127112}$ The overall incidence of childhood leukemia is 0.0000321 (Table 2, row $x x$ ) with a five-year death rate $39.8 \%$ (Table 2, row $a a a$ ) for children younger than $15 .{ }^{128}$
- Any breastfeeding is associated with a $36 \%$ reduction ( $\mathrm{OR}=0.64,95 \% \mathrm{CI}$ of $0.51-$ 0.81 ) in the risk of sudden infant death syndrome (SIDS) compared to no breastfeeding (Table 2, row fff). ${ }^{129}$ The overall incidence of SIDS is 0.00054 (Table 2 , row eee). ${ }^{130}$

Breastfeeding is associated with the following health benefits for the mother:

- The risk of breast cancer is reduced by $4.3 \%$ for each year of breastfeeding. ${ }^{131} \mathrm{We}$ have assumed a reduced risk of $2.15 \%$ for each 6 months of breastfeeding (Table 2, row $j j j$ ). The lifetime probability of developing (female) breast cancer is $11.5 \%$ (Table 2, row iii). ${ }^{132}$ Breast cancer is associated with a reduced life expectancy of 12.9 years (see Reference Document, Table 2, row mmm).
- Any breastfeeding is associated with a $21 \%$ reduction ( $\mathrm{OR}=0.79,95 \% \mathrm{CI}$ of $0.68-$ 0.91 ) in the risk of ovarian cancer compared to no breastfeeding (Table 1-2, row $p p p)$. Cumulative breastfeeding of at least 12 months is associated with a $28 \%$ reduction ( $\mathrm{OR}=0.72,95 \% \mathrm{CI}$ of $0.54-0.97$ ) in the risk of ovarian cancer compared to no breastfeeding. ${ }^{133112}$ Ovarian cancer is associated with a reduced life expectancy of 16.5 years (see reference Document, Table 2, row sss).
- Other assumptions used in assessing the clinically preventable burden are detailed in the Reference Document.

[^27]Based on these assumptions, the CPB associated with interventions aimed at improving rates of exclusive breastfeeding at 6 months from $0 \%$ to $60 \%$ is 5,002 QALYs (Table 2, row $v v v$ ).

## Table 2: CPB of Promotion of Breastfeeding in a Birth Cohort of 40,000

| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Infants in birth cohort | 40,000 |  |
| b | Current proportion exclusively breastfed for 6 months | 41\% | $\checkmark$ |
| c | Number exclusively breastfed for 6 months | 16,400 | $=(\mathrm{a} * \mathrm{c}$ ) |
| d | Effectiveness of breastfeeding promotion interventions in increasing adherence to breastfeeding for 6 months | 44\% | $\checkmark$ |
| e | Increase in exclusive 6-month breastfeeding with 100\% adherence | 10,384 | $=(\mathrm{a}-\mathrm{c}) * \mathrm{~d}$ |
| f | Estimated adherence with intervention | 75\% | Assumed |
| g | Increase in exclusive 6-month breastfeeding with intervention | 7,788 | $=(e * f)$ |
| h | Total proportion exclusively breastfed for 6 months with intervention | 60\% | $=(\mathrm{c}+\mathrm{g}) / \mathrm{a}$ |
|  | Health Benefits for the Infant |  |  |
| i | Average life expectancy of an infant in BC | 82.2 | $\checkmark$ |
| j | Average cases of otitis media (OM) in first year | 1.90 | $\checkmark$ |
| k | Effectiveness of breastfeeding in reducing risk of OM | 40.0\% | $\checkmark$ |
| I | Reduced cases of OM with intervention | 5,919 | $=(\mathrm{g}$ * j$) * \mathrm{k}$ |
| m | Average cases of atopic dermatitis (AD) in first 2 years | 0.165 | $\checkmark$ |
| n | Effectiveness of breastfeeding in reducing risk of AD | 42.0\% | $\checkmark$ |
| 0 | Reduced cases of AD with intervention | 540 | $=(\mathrm{g} * \mathrm{~m}) * \mathrm{n}$ |
| p | Average cases of gastrointestinal infection (GI) in first year | 0.222 | $\checkmark$ |
| q | Effectiveness of breastfeeding in reducing risk of GI | 64.0\% | $\checkmark$ |
| r | Reduced cases of Gl with intervention | 1,107 | $=(\mathrm{g} * \mathrm{p}) * \mathrm{q}$ |
| s | Average cases of lower respiratory tract infection (LTRI) in first year | 0.041 | $\checkmark$ |
| t | Effectiveness of breastfeeding in reducing risk of LTRI | 72.0\% | $\checkmark$ |
| $u$ | Reduced cases of LTRI with intervention | 229 | $=(\mathrm{g} * \mathrm{~s}) * \mathrm{t}$ |
| v | Average rate of death due to LTRI | 0.0000732 | $\checkmark$ |
| w | Effectiveness of breastfeeding in reducing risk of LTRI | 72.0\% | $\checkmark$ |
| X | Reduced deaths due to LTRI with intervention | 0.41 | $=(\mathrm{g} * \mathrm{v}){ }^{*} \mathrm{w}$ |
| y | Life years gained with intervention | 33.7 | = ${ }^{*}$ i |
| z | Average cases of childhood asthma | 0.127 | $\checkmark$ |
| aa | Effectiveness of breastfeeding in reducing risk of asthma | 27.0\% | $\checkmark$ |
| bb | Reduced cases of asthma with intervention | 267 | $=(\mathrm{g} * \mathrm{z}) *$ aa |
| cc | Average rate of death due to asthma | 0.0000027 | $\checkmark$ |
| dd | Effectiveness of breastfeeding in reducing risk of asthma | 27.0\% | $\checkmark$ |
| ee | Reduced deaths due to asthma with intervention | 0.01 | $=(\mathrm{g} * \mathrm{cc}) * \mathrm{dd}$ |
| ff | Life years gained with intervention | 0.5 | $=\mathrm{ee}$ * i |
| gg | Average \% of years as overweight | 27.1\% | Table 1-1 |
| hh | Effectiveness of breastfeeding in reducing risk of overweight | 24\% | $\checkmark$ |
| ii | Reduced years as overweight with intervention | 41,591 | $=\mathrm{g}^{*} \mathrm{i}^{*} \mathrm{gg} * \mathrm{hh}$ |
| jj | \% of life years lost with overweight | 0.73\% | $\checkmark$ |
| kk | Life years gained with intervention | 304 | = ii * jj |
| 11 | Average \% of years as obese | 10.7\% | Table 1 |
| mm | Effectiveness of breastfeeding in reducing risk of obesity | 24\% | $\checkmark$ |
| nn | Reduced years as obese with intervention | 16,504 | $=\mathrm{g} * \mathrm{i}$ * $\mathrm{ll}^{*} \mathrm{~mm}$ |
| о0 | \% of life years lost with obesity | 3.16\% | $\checkmark$ |
| pp | Life years gained with intervention | 522 | = nn * 00 |
| qq | Average cases of type 1 diabetes in children | 0.0001860 | $\checkmark$ |
| rr | Effectiveness of breastfeeding in reducing risk of type 1 diabetes | 19.0\% | $\checkmark$ |
| ss | Reduced cases of type 1 diabetes with intervention | 0.28 | $=(\mathrm{g} * \mathrm{qq}) * \mathrm{rr}$ |
| tt | Average rate of death due to type 1 diabetes | 0.0000012 | $\checkmark$ |
| uu | Effectiveness of breastfeeding in reducing risk of type 1 diabetes | 19.0\% | $\checkmark$ |
| vv | Reduced deaths due to type 1 diabetes with intervention | 0.002 | $=(\mathrm{g} * \mathrm{tt}) * \mathrm{uu}$ |
| ww | Life years gained with intervention | 0.15 | $=\mathrm{vv} * \mathrm{i}$ |


| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| xx | Average cases of childhood leukemia | 0.0000321 | $\checkmark$ |
| yy | Effectiveness of breastfeeding in reducing risk of childhood leukemia | 24.0\% | $\checkmark$ |
| zz | Reduced cases of childhood leukemia with intervention | 0.06 | $=(\mathrm{g} * \mathrm{xx}){ }^{*} \mathrm{yy}$ |
| aaa | 5 year death rate due to childhood leukemia | 39.8\% | $\checkmark$ |
| bbb | Effectiveness of breastfeeding in reducing risk of childhood leukemia | 24.0\% | $\checkmark$ |
| ccc | Reduced deaths due to childhood leukemia with intervention | 0.006 | = zz * aaa * bbb |
| ddd | Life years gained with intervention | 0.47 | = ccc* ${ }^{\text {i }}$ |
| eee | Average rate of death due to Sudden Infant Death Syndrome (SIDS) | 0.00054 | $\checkmark$ |
| fff | Effectiveness of breastfeeding in reducing risk of SIDS | 36.0\% | $\checkmark$ |
| ggg | Reduced deaths due to SIDS with intervention | 1.514 | $=\left(\mathrm{g}\right.$ * eee) ${ }^{*} \mathrm{fff}$ |
| hhh | Life years gained with intervention | 124.4 | = ggg *i |
|  | Health Benefits for the Mother |  |  |
| iii | Lifetime probability of developing breast cancer | 11.5\% | $\checkmark$ |
| jjj | Effectiveness of breastfeeding in reducing risk of breast cancer | 2.15\% | $\checkmark$ |
| kkk | Reduced breast cancer cases due to intervention | 19.3 | $=(\mathrm{g} * \mathrm{iii}) * \mathrm{jjj}$ |
| III |  |  |  |
| mmm | Life years lost per breast cancer | 12.9 | Ref Doc |
| nnn | Life years gained with intervention | 248.4 | = kkk * mmm |
| ooo | Lifetime probability of developing ovarian cancer | 1.4\% | $\checkmark$ |
| ppp | Effectiveness of breastfeeding in reducing risk of ovarian cancer | 21\% | $\checkmark$ |
| qqq | Reduced ovarian cancer cases due to intervention | 22.9 | $=\left(\mathrm{g} *\right.$ 000) ${ }^{*} \mathrm{ppp}$ |
| rrr |  |  |  |
| sss | Life years lost per ovarian cancer | 16.5 | Ref Doc |
| ttt | Life years gained with intervention | 377.8 | = qqq*sss |
| uuu | Potential QALYs gained, Intervention increasing from 41\% to 60\% | 1,611 | $\begin{gathered} =\mathrm{y}+\mathrm{ff}+\mathrm{kk}+\mathrm{pp}+\mathrm{ww}+ \\ \mathrm{ddd}+\mathrm{hhh}+\mathrm{nnn}+\mathrm{ttt} \end{gathered}$ |
| vvv | Potential QALYs gained, Intervention increasing from 0\% to 60\% | 5,002 | =(uuu/g) * $\mathrm{c}+\mathrm{g}$ ) |

$v=$ Estimates from the literature
We also modified a number of major assumptions and recalculated the CPB as follows:

- Assume the effectiveness of interventions aimed at improving rates of exclusive breastfeeding at 6 months is reduced from $44 \%$ to $13 \%$ (Table 2, row $d$ ): CPB = 3,868 QALYs
- Assume the effectiveness of interventions aimed at improving rates of exclusive breastfeeding at 6 months is increased from $44 \%$ to $84 \%$ (Table 2, row $d$ ): $\mathrm{CPB}=$ 6,466 QALYs
- Assume the effectiveness of breastfeeding in reducing overweight and obesity is reduced from $24 \%$ to $14 \%$ (Table 2 , row $h h \& m m$ ): $\mathrm{CPB}=3,934$ QALYs
- Assume the effectiveness of breastfeeding in reducing overweight and obesity is increased from $24 \%$ to $33 \%$ (Table 2, row $h h \& m m$ ): CPB = 5,963 QALYs


## Modelling Cost-Effectiveness

In this section, we will calculate the CPB associated with interventions aimed at improving longer term ( 6 months) exclusive breastfeeding rates in a British Columbia birth cohort of 40,000.

In modelling CE, we made the following assumptions:

- Patient time costs for office visit - We assumed that two hours of patient time would be required, including travel to and from the appointment.
- Patient time costs for breastfeeding support groups - We assumed that a new mother would attend a breastfeeding support group once per month (lasting two hours) for six months. We assumed an additional hour for travel time for a total patient time commitment of 18 hours.
- Otitis media - Two estimates from the US suggest a direct cost (ambulatory care and antibiotics) per case of $\$ 156$ (2007 USD) ${ }^{134}$ and $\$ 106$ (2004 USD). ${ }^{135}$ A Canadian study suggested additional hospital costs over and above physician and drug costs of $15.6 \%$. ${ }^{136}$ We have converted the $\$ 156$ to 2017 Canadian dollars and then added $15.6 \%$ to this cost per case to reflect hospital costs for a total cost per case of \$251 (Table 3, row $p$ ).
- Atopic dermatitis - The mean duration of atopic dermatitis is 10 years with $45 \%$ of cases being mild in severity, $45 \%$ moderate and $10 \%$ severe. ${ }^{137}$ The direct annual costs per mild, moderate and severe case are $\$ 175, \$ 300$, and $\$ 405$, respectively. The average weighted cost totalled $\$ 254$ CAD in $2001^{138}$ or $\$ 342$ (in 2017 CAD) per case per year. Lifetime costs were estimated at $\$ 3,420$ (Table 3, row $s$ ).
- Gastrointestinal infection - A US study suggests the direct costs for gastrointestinal infections and lower respiratory tract infections are $\$ 331$ per case (in 1995 USD) ${ }^{139}$ or $\$ 462$ in 2017 CAD (Table 3, rows $v$ ).
- Lower respiratory tract infection - See above (Table 3, rows y).
- Asthma - A BC study estimated the annual direct costs attributable to asthma at $\$ 444$ per person year (in 2006 CAD ) ${ }^{140}$ or $\$ 523$ in 2017 CAD. Based on an average treatment duration of 10 years, ${ }^{141}$ the total costs attributable to childhood asthma would be $\$ 5,230$ per case (Table 3 , row $b b$ ).
- Type 1 diabetes - The lifetime cost per case in the US has been estimated at $\$ 77,463$ (in 2007 USD) ${ }^{142}$ or $\$ 76,598$ in 2017 CAD (Table 3, row $k k$ ).
- Childhood leukemia - The lifetime cost per case in the US has been estimated at $\$ 136,444$ (in 2007 USD) ${ }^{143}$ or $\$ 134,920$ in 2017 CAD (Table 3, row $n n$ ).
- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the CE associated with interventions aimed at improving rates of exclusive breastfeeding at 6 months is $-\$ 9,021$ per QALY (Table 3, row $b b b$ ).

[^28]| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Women eligible for screening/referral in primary care | 40,000 |  |
| b | Proportion already exclusively breastfeeding for 6 months | 41\% | Table 2, row b |
| c | Number exclusively breastfeeding for 6 months | 16,400 | = a* ${ }^{\text {a }}$ |
| d | Women eligible for intervention (support group) | 23,600 | = a-c |
| e | Estimated adherence with intervention | 75\% | Assumed |
| f | Women attending intervention (support group) | 17,700 | $=d^{*} \mathrm{f}$ |
| g | Effectiveness of breastfeeding promotion interventions in increasing adherence to breastfeeding for 6 months | 44\% | Table 2, row d |
| h | \# of women attending intervention (support group) who exclusively breastfeed for 6 months | 7,788 | $=\mathrm{f}$ * g |
|  | Costs of intervention |  |  |
| i | Cost of 10-minute office visit | \$34.85 | Ref Doc |
| j | Value of patient time and travel for office visit | \$59.38 | =2 * 29.69 |
| k | Portion of 10-minute office visit for screen/referral | 50\% | Ref Doc |
| I | Estimated cost of screening | \$1,884,600 | $=a *(1+j) * k$ |
| m | Value of patient time and travel for intervention | \$534 | = 18 * \$29.69 |
| n | Estimated cost of intervention over lifetime of birth cohort | \$9,451,800 | = ${ }^{*} \mathrm{~m}$ |
|  | Cost avoided |  |  |
| 0 | Cases of otitis media avoided | 5,919 | Table 2, row 1 |
| p | Cost per case | \$251 | $\checkmark$ |
| q | Costs avoided | \$1,485,639 | = ${ }^{*} \mathrm{p}$ |
| r | Cases of atopic dermatitis avoided | 540 | Table 2, row o |
| s | Cost per person with atopic dermatitis | \$3,420 | $\checkmark$ |
| t | Costs avoided | \$1,845,803 | = ${ }^{*}$ s |
| u | Cases of gastrointestinal infection avoided | 1,107 | Table 2, row r |
| v | Cost per case | \$462 | $\checkmark$ |
| w | Costs avoided | \$511,212 | $=u^{*} \mathrm{v}$ |
| x | Cases of lower respiratory tract infection avoided | 229 | Table 2, row u |
| y | Cost per case | \$462 | $\checkmark$ |
| z | Costs avoided | \$105,956 | = ${ }^{*} \mathrm{y}$ |
| aa | Cases of asthma avoided | 267 | Table 2, row bb |
| bb | Cost per case | \$5,230 | $\checkmark$ |
| cc | Costs avoided | \$1,396,674 | = aa * bb |
| dd | Years of overweight avoided | 41,591 | Table 2, row ii |
| ee | Cost per year | \$227 | Ref Doc |
| ff | Costs avoided | \$9,441,234 | = dd * ee |
| gg | Years of obesity avoided | 16,504 | Table 2, row nn |
| hh | Cost per year | \$805 | Ref Doc |
| ii | Costs avoided | \$13,285,924 | = gg* hh |
| jj | Cases of type 1 diabetes avoided | 0.3 | Table 2, row ss |
| kk | Cost per case | \$76,598 | $\checkmark$ |
| 11 | Costs avoided | \$21,082 | $=\mathrm{jj}$ * kk |
| mm | Cases of childhood leukemia avoided | 0.06 | Table 2, row zz |
| nn | Cost per case | \$134,920 | $\checkmark$ |
| оо | Costs avoided | \$8,095 | $=\mathrm{mm}$ * nn |
| pp | Cases of breast cancer avoided | 19.3 | Table 2, row kkk |
| qq | Cost per case | \$29,707 | Ref Doc |
| rr | Costs avoided | \$572,033 | = pp * qq |
| ss | Cases of ovarian cancer avoided | 22.9 | Table 2, row qqq |
| tt | Cost per case | \$84,534 | Ref Doc |
| uu | Costs avoided | \$1,935,551 | = ss * tt |
|  | CE calculation |  |  |
| vv | Cost of intervention over lifetime of birth cohort | \$11,336,400 | $=1+n$ |
| ww | Costs avoided | \$30,609,203 | $\begin{gathered} =\mathrm{q}+\mathrm{t}+\mathrm{w}+\mathrm{z}+\mathrm{cc}+\mathrm{ff}+\mathrm{ii} \\ \\ +\mathrm{ll}+\mathrm{oo}+\mathrm{rr}+\mathrm{uu} \end{gathered}$ |
| xx | QALYs saved | 1,611 | Table 2, row uuu |
| yy | Cost of intervention over lifetime of birth cohort (1.5\% discount) | \$11,336,400 | Calculated |
| zz | Costs avoided (1.5\% discount) | \$19,827,768 | Calculated |
| aaa | QALYs saved (1.5\% discount) | 941 | Calculated |
| bbb | CE (\$/QALY saved) | -\$9,021 | $=(\mathrm{yy}$-zz)/aaa |

V = Estimates from the literature

We also modified several major assumptions and recalculated the cost per QALY as follows:

- Assume the effectiveness of interventions aimed at improving rates of exclusive breastfeeding at 6 months is reduced from $44 \%$ to $13 \%$ (Table 2, row $d$ ): $\mathrm{CE}=$ \$19,699 per QALY
- Assume the effectiveness of interventions aimed at improving rates of exclusive breastfeeding at 6 months is increased from $44 \%$ to $84 \%$ (Table 2, row $d$ ): $\mathrm{CE}=$ -\$14,757 per QALY
- Assume the effectiveness of breastfeeding in reducing overweight and obesity is reduced from $24 \%$ to $14 \%$ (Table 2, rows $h h \& m m$ ): $\mathrm{CE}=-\$ 3,995$ per QALY
- Assume the effectiveness of breastfeeding in reducing overweight and obesity is increased from $24 \%$ to $33 \%$ (Table 2, rows $h h \& m m$ ): $\mathrm{CE}=-\$ 12,006$ per QALY
- Assume the proportion of an office visit required for screening/referral is reduced from $50 \%$ to $33 \%$ (Table 3, row $k$ ): $\mathrm{CE}=-\$ 9,702$ per QALY
- Assume the proportion of an office visit required for screening/referral is increased from $50 \%$ to $67 \%$ (Table 3, row $k$ ): CE $=-\$ 8,341$ per QALY


## Summary

| Table 4: Promotion of Breastfeeding in a Birth Cohort of 40,000 Summary |  |  |  |
| :---: | :---: | :---: | :---: |
|  | $\begin{aligned} & \text { Base } \\ & \text { Case } \\ & \hline \end{aligned}$ | Range |  |
| CPB (Potential QALYs Gained) |  |  |  |
| Assume No Current Service |  |  |  |
| 1.5\% Discount Rate | 2,923 | 2,260 | 3,779 |
| 3\% Discount Rate | 1,853 | 1,433 | 2,396 |
| 0\% Discount Rate | 5,002 | 3,868 | 6,466 |
| Gap between B.C. Current and Best in the World |  |  |  |
| 1.5\% Discount Rate | 941 | 278 | 1,797 |
| 3\% Discount Rate | 597 | 176 | 1,139 |
| 0\% Discount Rate | 1,611 | 476 | 3,075 |
| $\overline{\mathrm{CE}(\$ / \mathrm{QALY}) \text { including patient time costs }}$ |  |  |  |
| 1.5\% Discount Rate | -\$9,021 | -\$14,757 | \$19,699 |
| 3\% Discount Rate | -\$4,745 | -\$13,791 | \$40,557 |
| 0\% Discount Rate | -\$11,966 | -\$15,318 | \$4,818 |
| CE ( $\$ /$ /AALY) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | -\$20,325 | -\$20,678 | -\$18,599 |
| 3\% Discount Rate | -\$22,574 | -\$23,130 | -\$19,789 |
| 0\% Discount Rate | -\$18,572 | -\$18,778 | -\$17,540 |

## Growth Monitoring and Healthy Weight Management in Children and Youth

## Canadian Task Force on Preventive Health Care (2015) ${ }^{144}$

We recommend growth monitoring ${ }^{145}$ at all appropriate ${ }^{146}$ primary care visits using the 2014 WHO Growth Charts for Canada. (Strong recommendation; very low quality evidence)

This growth monitoring recommendation applies to all children and youth 0-17 years of age who present to primary care.

We recommend that primary care practitioners not routinely offer structured interventions ${ }^{147}$ aimed at preventing overweight and obesity in healthy weight children and youth. (Weak recommendation; very low quality evidence)

This prevention recommendation applies to all children and youth $0-17$ years of age who have a healthy weight. They do not apply to children and youth with eating disorders, or who are underweight, overweight, or obese.

For children and youth aged 2 to 17 years who are overweight or obese, we recommend that primary care practitioners offer or refer to structured behavioural interventions aimed at healthy weight management. (Weak recommendation; moderate quality evidence)

For children and youth aged 2 to 11 years who are overweight or obese, we recommend that primary care practitioners not offer Orlistat ${ }^{148}$ aimed at healthy weight management. (Strong recommendation; very low quality evidence)

For children and youth aged 12 to 17 years who are overweight or obese, we recommend that primary care practitioners not routinely offer Orlistat aimed at healthy weight management. (Weak recommendation; moderate quality evidence)

For children and youth aged 2 to 17 years who are overweight or obese, we recommend that primary care practitioners not routinely refer for surgical interventions. (Strong recommendation; very low quality evidence)

[^29]These management recommendations apply to children and youth 2-17 years of age who are overweight or obese. Children and youth with health conditions where weight management is inappropriate are excluded.
The CTFPHC concludes that "the most effective behavioural interventions were those that were delivered by a specialized interdisciplinary team, involved group sessions, and incorporated family and parent involvement". Furthermore, "where structured behavioural interventions for weight management in children and youth are not yet available in Canada, primary care practitioners and policy makers should consider their development a priority." ${ }^{149}$

## United States Preventive Services Task Force Recommendations (2017)

The USPSTF recommends that clinicians screen for obesity in children and adolescents 6 years and older and offer or refer them to comprehensive, intensive behavioral interventions to promote improvements in weight status. (Grade $B$ recommendation) ${ }^{150}$

## Modelling the Clinically Preventable Burden

In this section, we model CPB associated with growth monitoring in children and youth ages 0-17 and the offer of, or referral to, structured behavioural interventions aimed at healthy weight management for children and youth aged 2 to 17 years who are overweight or obese.

In modelling CPB , we made the following assumptions:

- There were 865,080 children and youth ages $0-17$ living in BC in 2017. The majority of these children and youth would be eligible for growth monitoring. Based on measured height and weight as calculated for the 2004 Canadian Community Health Survey (CCHS), $26.5 \%$ of BC children and youth ages 1-17 are either overweight or obese. ${ }^{151}$ An estimated $19.9 \%$ are overweight (or 172,583 individuals) while a further $6.6 \%$ are obese (or 56,749 individuals) (see Table 1). The 56,749 children and youth with obesity are most likely to be offered structured behavioural interventions aimed at healthy weight management.

[^30]| Table 1: Estimated Number of Overweight and Obese Children and Youth In British Columbia |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Male |  | Female |  | Total |  |
| Population |  |  |  |  |  |  |
| <1 | 23,780 |  | 22,580 |  | 46,360 |  |
| 1 to 3 | 71,340 |  | 67,740 |  | 139,080 |  |
| 4 to 8 | 121,140 |  | 113,540 |  | 234,680 |  |
| 9 to 13 | 122,260 |  | 114,340 |  | 236,600 |  |
| 14 to 17 | 107,640 |  | 100,720 |  | 208,360 |  |
| Total | 446,160 |  | 418,920 |  | 865,080 |  |
|  | Overweig | Obese | Overwei | Obese | Overweight | Obese |
| Prevalence |  |  |  |  |  |  |
| <1 | - | - | - | - | - | - |
| 1 to 3 | 11.5\% | 8.5\% | 13.9\% | 2.1\% | 12.9\% | 4.7\% |
| 4 to 8 | 17.3\% | 2.2\% | 11.4\% | 13.6\% | 14.2\% | 8.2\% |
| 9 to 13 | 32.8\% | 6.1\% | 22.2\% | 4.7\% | 27.6\% | 5.4\% |
| 14 to 17 | 20.0\% | 10.1\% | 18.5\% | 3.8\% | 19.2\% | 6.8\% |
| Total | 23.1\% | 6.3\% | 17.1\% | 6.8\% | 19.9\% | 6.6\% |
|  | Overweig | Obese | Overweig | Obese | Overweight | Obese |
| \# of Individuals |  |  |  |  |  |  |
| <1 | - | - | - | - | - | - |
| 1 to 3 | 8,177 | 6,042 | 9,447 | 1,454 | 18,003 | 6,532 |
| 4 to 8 | 21,016 | 2,704 | 12,930 | 15,463 | 33,336 | 19,332 |
| 9 to 13 | 40,084 | 7,515 | 25,427 | 5,323 | 65,281 | 12,806 |
| 14 to 17 | 21,502 | 10,884 | 18,643 | 3,851 | 40,010 | 14,155 |
| Total | 102,881 | 28,249 | 71,665 | 28,356 | 172,583 | 56,749 |

- Evidence suggests that excess weight in children/youth often persists into adulthood. ${ }^{152,153,154}$ We assumed that, without any intervention, the $20.0 \%$ of $14-17$ year old males and $18.5 \%$ of 14-17 year old females who are overweight would remain so for the rest of their lives (see Table 1). A similar assumption was made for the $10.1 \%$ of 14-17 year old males and $3.8 \%$ of 14-17 year old females who are obese. Based on this assumption, of the total 1.5 million life years in the male birth cohort (see Table 3, row $a$ ), 310,760 would be lived as overweight (see Table 3, row $b$ ) and 143,044 as obese (see Table 3, row $c$ ). Similarly, of the total 1.6 million life years in the female birth cohort (see Table 3, row $d$ ), 287,637 would be lived as overweight (see Table 3, row $e$ ) and 69,962 as obese (see Table 3 , row $f$ ).

[^31]| Age | Mean Survival Rate |  | Individuals in Birth Cohort |  | Years of Life in Birth Cohort |  | \% Overweight |  | Years of Life Overweight |  | \% Obese |  | Years of Life Obese |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Group | Male | Female | Male | Female | Male | Female | Male | Female | Male | Female | Male | Female | Male | Female |
| 0-4 | 99.55\% | 99.63\% | 19,910 | 19,926 | 99,551 | 99,629 | 11.5\% | 13.9\% | 11,411 | 13,894 | 8.5\% | 2.1\% | 8,432 | 2,138 |
| 5-9 | 99.51\% | 99.58\% | 19,903 | 19,915 | 99,513 | 99,577 | 17.3\% | 11.4\% | 17,264 | 11,340 | 2.2\% | 13.6\% | 2,221 | 13,561 |
| 10-14 | 99.48\% | 99.55\% | 19,895 | 19,911 | 99,476 | 99,553 | 32.8\% | 22.2\% | 32,614 | 22,139 | 6.1\% | 4.7\% | 6,115 | 4,635 |
| 15-19 | 99.37\% | 99.48\% | 19,875 | 19,897 | 99,374 | 99,484 | 20.0\% | 18.5\% | 19,851 | 18,415 | 10.1\% | 3.8\% | 10,048 | 3,804 |
| 20-24 | 99.07\% | 99.32\% | 19,813 | 19,865 | 99,065 | 99,323 | 20.0\% | 18.5\% | 19,789 | 18,385 | 10.1\% | 3.8\% | 10,017 | 3,797 |
| 25-29 | 98.67\% | 99.16\% | 19,734 | 19,833 | 98,672 | 99,163 | 20.0\% | 18.5\% | 19,711 | 18,355 | 10.1\% | 3.8\% | 9,977 | 3,791 |
| 30-34 | 98.29\% | 98.98\% | 19,658 | 19,795 | 98,289 | 98,975 | 20.0\% | 18.5\% | 19,634 | 18,320 | 10.1\% | 3.8\% | 9,938 | 3,784 |
| 35-39 | 97.80\% | 98.71\% | 19,560 | 19,741 | 97,798 | 98,706 | 20.0\% | 18.5\% | 19,536 | 18,271 | 10.1\% | 3.8\% | 9,889 | 3,774 |
| 40-44 | 97.13\% | 98.31\% | 19,427 | 19,662 | 97,134 | 98,311 | 20.0\% | 18.5\% | 19,403 | 18,197 | 10.1\% | 3.8\% | 9,822 | 3,759 |
| 45-49 | 96.20\% | 97.73\% | 19,241 | 19,546 | 96,203 | 97,730 | 20.0\% | 18.5\% | 19,217 | 18,090 | 10.1\% | 3.8\% | 9,727 | 3,737 |
| 50-54 | 94.86\% | 96.87\% | 18,971 | 19,375 | 94,855 | 96,873 | 20.0\% | 18.5\% | 18,948 | 17,931 | 10.1\% | 3.8\% | 9,591 | 3,704 |
| 55-59 | 92.85\% | 95.59\% | 18,570 | 19,118 | 92,852 | 95,591 | 20.0\% | 18.5\% | 18,548 | 17,694 | 10.1\% | 3.8\% | 9,389 | 3,655 |
| 60-64 | 89.84\% | 93.63\% | 17,967 | 18,726 | 89,835 | 93,630 | 20.0\% | 18.5\% | 17,945 | 17,331 | 10.1\% | 3.8\% | 9,083 | 3,580 |
| 65-69 | 85.26\% | 90.57\% | 17,052 | 18,113 | 85,261 | 90,567 | 20.0\% | 18.5\% | 17,032 | 16,764 | 10.1\% | 3.8\% | 8,621 | 3,463 |
| 70-74 | 78.34\% | 85.72\% | 15,668 | 17,144 | 78,342 | 85,720 | 20.0\% | 18.5\% | 15,650 | 15,867 | 10.1\% | 3.8\% | 7,921 | 3,277 |
| 75-79 | 68.08\% | 78.04\% | 13,616 | 15,608 | 68,078 | 78,041 | 20.0\% | 18.5\% | 13,599 | 14,445 | 10.1\% | 3.8\% | 6,884 | 2,984 |
| 80+ | 53.10\% | 65.90\% | 10,620 | 13,180 | 53,100 | 65,900 | 20.0\% | 18.5\% | 10,607 | 12,198 | 10.1\% | 3.8\% | 5,369 | 2,520 |
| Total |  |  |  |  | 1,547,398 | 1,596,773 | 20.1\% | 18.0\% | 310,760 | 287,637 | 9.2\% | 4.4\% | 143,044 | 69,962 |

- The systematic review and meta-analysis for the CTFPHC found that the overall effectiveness of interventions resulted in a -0.53 drop in BMI ( $95 \%$ CI from -0.69 to 0.36 ). This decrease, however, was not maintained 6-12 months after the intervention ( 0.08 change in $\mathrm{BMI}, 95 \% \mathrm{CI}$ from -0.07 to 0.23 ). The most effective interventions included a focus on both diet and exercise ( -1.09 drop in BMI, $95 \%$ CI from -1.84 to -0.34). The review also found a statistically significant improvement in QoL. ${ }^{155}$
- Interventions reduced the prevalence of overweight from $40 \%$ to $35 \%$ and obesity from $33 \%$ to $31 \%$ over a duration of up to 36 months. ${ }^{156}$
- Improvements in QoL appear to be positively correlated with weight loss. ${ }^{157}$ One small study found a clinically important improvement in $22 \%$ (4 of 18) of the children/youth who successfully completed a multidisciplinary lifestyle program. ${ }^{158}$
- For modelling purposes, we assumed that a weight management program would reduce overweight by $12.5 \%$ (Table 3, row $a k$ ) and obesity by $6.1 \%$ (Table 3, row al) (based on the reduction in the prevalence of overweight from $40 \%$ to $35 \%$ and obesity from $33 \%$ to $31 \%$ noted above ${ }^{159}$ ). We also assumed the increase in QoL associated with the successful completion of a weight management program would be maintained long-term for $22 \%$ of participants (Table 3, rows an \& ao). This

[^32]assumption was varied in the sensitivity analysis from $12.5 \%$ for overweight and $6.1 \%$ for obese to $30 \%$ for both overweight and obese.

- Children in families that do not have a regular primary care provider (PCP) are unlikely to enter a weight monitoring/management process. Based on 2012 CCHS data, $89 \%$ of families in BC have a regular PCP (Table 3, row $a d$ ). ${ }^{160}$
- We noted earlier that the regular assessment of BMI by primary care providers is relatively poor. For modelling purposes, we assumed that $13 \%$ of PCPs would regularly monitor BMI (Table 3, row $a e$ ) and that $70 \%$ of these PCPs would refer overweight and obese children youth to a weight management program (Table 3, row af ). Furthermore, we assumed that $39 \%$ of families referred to a weight management program would successfully complete the program (Table 3, row $a g$ with a range from $29 \%$ to $49 \%$ ). Between January 2013 and June 2015, 1,071 children and their parent(s) were referred to Shapedown BC. ${ }^{161}$ Between January and June of 2015, $39 \%$ of those referred to the program ultimately completed it.
- The USPSTF review grouped interventions by intensity as follows: very low (<10 hours), low (10-25 hours), moderate ( $26-75$ hours) or high (>75 hours). The comprehensiveness of the interventions was determined by a focus on both diet and physical activity as well as instruction in and support for the use of behavioural management techniques. Only comprehensive interventions of moderate to high intensity were effective (a reduction of between 1.9 to $3.3 \mathrm{~kg} / \mathrm{m}^{2}$ at 12 months). ${ }^{162,163}$
- Other assumptions used in assessing the clinically preventable burden are detailed in the Reference Document.
Based on these assumptions, the CPB associated with growth monitoring in children and youth ages 0-17 along with the offer of, or referral to, structured behavioural interventions aimed at healthy weight management for children and youth aged 2 to 17 years who are overweight or obese is 80 QALYs (see Table 3, row $a r$ ). The CPB of 80 represents the gap between no coverage and the 'best in the world' growth monitoring coverage, which was estimated at $13 \%$.

[^33]Table 3: CPB of Growth Monitoring and Healthy Weight Management in Children / Youth in a Birth Cohort of 40,000

| Row Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
|  | Current State |  |  |
| a | Years of life lived in the birth cohort - males | 1,547,398 | Table 2 |
| b | Years of life lived with overweight in the birth cohort - males | 310,760 | Table 2 |
| c | Years of life lived with obesity in the birth cohort - males | 143,044 | Table 2 |
| d | Years of life lived in the birth cohort - females | 1,596,773 | Table 2 |
| e | Years of life lived with overweight in the birth cohort - females | 287,637 | Table 2 |
| f | Years of life lived with obesity in the birth cohort - females | 69,962 | Table 2 |
| g | Disutility associated with overweight | 0.0\% | Ref Doc |
| h | Disutility associated with obesity | 5.9\% | Ref Doc |
| i | QALYs lost due to overweight - males | 0 | = ${ }^{*} \mathrm{~g}$ |
| j | QALYs lost due to obesity - males | 8,440 | = ${ }^{*}$ h |
| k | QALYs lost due to overweight - females | 0 | = ${ }^{*} \mathrm{~g}$ |
| I | QALYs lost due to obesity - females | 4,128 | = ${ }^{*}$ h |
| m | Overweight males at age 18 | 3,970 | Table 2 |
| n | Obese males at age 18 | 2,010 | Table 2 |
| 0 | Overweight females at age 18 | 3,683 | Table 2 |
| p | Obese females at age 18 | 761 | Table 2 |
| q | Life years lost due to overweight per individual | 0.6 | Ref Doc |
| r | Life years lost due to obesity per individual | 2.6 | Ref Doc |
| s | Life years lost due to overweight - males | 2,382 | $=\mathrm{m}^{*} \mathrm{q}$ |
| t | Life years lost due to obesity - males | 5,225 | $=\mathrm{n}^{*} \mathrm{r}$ |
| $u$ | Life years lost due to overweight - females | 2,210 | = ${ }^{*} \mathrm{q}$ |
| v | Life years lost due to obesity - females | 1,978 | = ${ }^{*}$ r |
| w | Total QALYs lost due to overweight - males | 2,382 | = i + s |
| x | Total QALYs lost due to obesity - males | 13,665 | = $\mathrm{j}+\mathrm{t}$ |
| y | Total QALYs lost due to excess weight in males | 16,047 | $=w+x$ |
| $z$ | Total QALYs lost due to overweight - females | 2,210 | = $\mathrm{k}+\mathrm{u}$ |
| aa | Total QALYs lost due to obesity - females | 6,106 | = $1+\mathrm{v}$ |
| ab | Total QALYs lost due to excess weight in females | 8,315 | $=z+a \mathrm{a}$ |
| ac | Total QALYs lost due to excess weight in birth cohort | 24,362 | $=y+a b$ |
|  | Effect of Intervention |  |  |
| ad | BC families with a regular primary care provider (PCP) | 89\% | $\checkmark$ |
| ae | Proportion of PCPs who regularly assess BMI | 13\% | Ref Doc |
| af | Proportion of PCPs who regularly assess BMI who would refer children/youth with excess weight to a weight management program | 70\% | Assumed |
| ag | Proportion of children/youth who would successfully complete a weight management program | 39\% | $\checkmark$ |
| ah | Number of overweight individuals who would successfully complete a weight management program | 125 | = m * ad * ae * af * ag |
| ai | Number of obese individuals who would successfully complete a weight management program | 63 | = n * ad * ae * af * ag |
| aj | Years of life lived by an 8 -year old in this subgroup | 74 | $\checkmark$ |
| ak | Decrease in prevalence of overweight associated with intervention | 12.5\% | V |
| al | Decrease in prevalence of obesity associated with intervention | 6.1\% | $\checkmark$ |
| am | Life-years gained with intervention | 19 | $=(a h * q * a k)+(a i * r *$ <br> al) |
| an | Proportion of individuals with overweight benefitting from an improvement in QoL | 22.0\% | v |
| ao | Proportion of individuals with obesity benefitting from an improvement in QoL | 22.0\% | $\checkmark$ |
| ap | QALYs gained due to intervention | 61 | $\begin{gathered} =(\text { ah } * a j * g * a n)+(a i * \\ a j * h * a o) \end{gathered}$ |
| ar | Potential QALYs gained, Intervention increasing from 0\% to 13\% | 80 | = am +ap |

We also modified a number of major assumptions and recalculated the CPB as follows:

- Assume that the proportion of children/youth who successfully complete a weight management program after being referred is reduced from $39 \%$ to $29 \%$ (Table 3, row $a g): ~ \mathrm{CPB}=60$.
- Assume that the proportion of children/youth who would successfully complete a weight management program after being referred is increased from $39 \%$ to $49 \%$ (Table 3, row $a g$ ): $\mathrm{CPB}=101$.
- Assume that the proportion of children/youth who maintain improvement in QoL after successfully completing a weight management program is reduced from $22 \%$ to $12.5 \%$ and $6.1 \%$ for children / youth who are overweight/obese (Table 3, row an \& $a o): ~ \mathrm{CPB}=36$.
- Assume that the proportion of children/youth who maintain improvement in QoL after successfully completing a weight management program is increased from $22 \%$ to $30 \%$ (Table 3, row an \& ao): $\mathrm{CPB}=103$.


## Modeling Cost-Effectiveness

In modelling CE, we made the following assumptions:

- Frequency of screening - The CTFPHC recommends growth monitoring at all appropriate primary care visits. Appropriate primary care visits are defined as "scheduled health supervision visits, visits for immunizations or medication renewal, episodic care or acute illness, and other visits where the primary care practitioner deems it appropriate. Primary care visits are completed at primary health care settings, including those outside of a physician's office (e.g. public health nurses carrying out a well-child visit at a community setting)." ${ }^{164}$ The Canadian Paediatric Association recommends that well-child visits take place at 1 week, at $2,4,6$ and 12 months, annually from ages 2-5 and then every year or two until the child is 18 years of age. ${ }^{165}$ For modelling purposes, we have assumed that growth monitoring would occur annually between the ages of $0-17$ at a well-child visit (Table 4, row $d$ ).
- Program costs - Holingworth and colleagues estimated a range of program costs between $£ 108$ and $£ 662$ (in 2009 British pounds) per child based on a review of ten lifestyle interventions to treat overweight and obesity in children. ${ }^{166} \mathrm{We}$ converted these costs to equivalent Canadian health care costs in 2017, for a cost of $\$ 214$ to $\$ 1,310$ per child. For modelling purposes we used the mid-point for the base case scenario (\$762) and the range in the sensitivity analysis (Table 4, row $l \& m$ ).
- We assumed that the excess costs associated with overweight and obesity would be avoided during the remaining lifetime of the individual after a successful weight management program. We also modified this assumption so that costs would only be avoided for a five year period after a successful weight management program.
- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.

[^34]- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the CE associated with growth monitoring in children and youth ages 0-17 and the offer of, or referral to, structured behavioural interventions aimed at healthy weight management for children and youth ages 2 to 17 years who are overweight or obese is \$77,441 / QALY (Table 4, row $a c$ ).

Table 4: CE of Growth Monitoring and Healthy Weight Management in Children / Youth in a Birth Cohort of $\mathbf{4 0 , 0 0 0}$

| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Years of life lived in birth cohort from 0-17 | 716,614 | Table 2 |
| b | BC families with a regular primary care provider (PCP) | 89\% | Table 3, row ad |
| c | Proportion of PCPs who regularly assess BMI | 13\% | Table 3, row ae |
| d | Number of assessments per year | 1 | Assumed |
| e | Total number of screens | 82,912 | $=a * b * c * d$ |
|  | Costs of Screening |  |  |
| f | Cost of 10-minute office visit | \$34.85 | Ref Doc |
| g | Value of patient time and travel for office visit | \$59.38 | Ref Doc |
| h | Portion of 10-minute office visit for screen/referral | 50\% | Assumed |
| i | Estimated cost of screening | \$3,906,409 | $=\left(e{ }^{*}{ }^{*} \mathrm{~h}\right)+\left(\mathrm{e}^{*} \mathrm{~g}\right.$ * h$)$ |
|  | Costs of Intervention |  |  |
| j | Number of obese individuals successfully completing a weight management program | 63 | Table 3, row ai |
| k | Number of overweight individuals successfully completing a weight management program | 125 | Table 3, row ah |
| I | Cost of intervention per obese individual | \$762 | $\checkmark$ |
| m | Cost of intervention per overweight individual | \$762 | $\checkmark$ |
| n | Cost of intervention | \$143,925 | $=(\mathrm{j} * \mathrm{I})+(\mathrm{k} * \mathrm{~m})$ |
| 0 | Value of patient time and travel per intervention | \$891 | $\checkmark$ |
| p | Total value of patient time and travel for interventions | \$168,290 | $=(\mathrm{j}+\mathrm{k}) * \mathrm{O}$ |
|  | Cost avoided |  |  |
| q | Years of overweight avoided | 1,160 | Table 3, row ah * Table 3, row aj * Table 3, row ak |
| r | Medical care costs per year associated with overweight | \$227 | Ref Doc |
| S | Costs avoided | \$263,314 | $=q^{*} r$ |
| t | Years of obesity avoided | 287 | Table 3, row ai * Table 3, row aj * Table 3, row al |
| u | Medical care costs per year associated with obesity | \$805 | Ref Doc |
| v | Costs avoided | \$230,655 | = ${ }^{*}$ u |
|  | CE calculation |  |  |
| w | Cost of intervention over lifetime of birth cohort | \$4,218,624 | = $\mathrm{i}+\mathrm{n}+\mathrm{p}$ |
| x | Costs avoided | \$493,969 | = $\mathrm{s}+\mathrm{v}$ |
| y | QALYs saved | 80 | Table 3, row ar |
| z | Cost of intervention over lifetime of birth cohort (1.5\% discount) | \$3,704,213 | Calculated |
| aa | Costs avoided (1.5\% discount) | \$272,147 | Calculated |
| ab | QALYs saved (1.5\% discount) | 44 | Calculated |
| ac | CE (\$/QALY saved) | \$77,441 | = (z - aa) / ab |

V = Estimates from the literature

We also modified a number of major assumptions and recalculated the CE as follows:

- Assume that the proportion of children/youth who successfully complete a weight management program after being referred is reduced from $39 \%$ to $29 \%$ (Table 3, row ag): $\mathrm{CE}=\$ 104,129$.
- Assume that the proportion of children/youth who would successfully complete a weight management program after being referred is increased from $39 \%$ to $49 \%$ (Table 3, row $a g$ ): $\mathrm{CE}=\$ 61,646$.
- Assume that the proportion of children/youth who maintain improvement in QoL after successfully completing a weight management program is reduced from $22 \%$ to $12.5 \%$ and $6.1 \%$ for children/youth who are overweight/obese (Table 3, rows an \& ao): $\mathrm{CE}=\$ 171,245$.
- Assume that the proportion of children/youth who maintain improvement in QoL after successfully completing a weight management program is increased from $22 \%$ to $30 \%$ (Table 3, rows an \& ao): $\mathrm{CE}=\$ 60,709$.
- Assume that the proportion of an office visit for weight measurement is decreased from $50 \%$ to $33 \%$ (Table 4, row $h$ ): $\mathrm{CE}=\$ 51,126$.
- Assume that the proportion of an office visit for weight measurement is increased from $50 \%$ to $67 \%$ (Table 4, row $h$ ): $\mathrm{CE}=\$ 103,755$.
- Assume that the cost of the weight management program per individual is reduced from $\$ 762$ to $\$ 214$ (Table 4, row $l \& m$ ): CE $=\$ 75,390$.
- Assume that the cost of the weight management program per individual is increased from $\$ 762$ to $\$ 1,310$ (Table 4, row $l \& m$ ): $\mathrm{CE}=\$ 79,491$.
- Assume that costs avoided would only last for five years, rather than a lifetime, after a successful weight management program (Table 3, rows $a j$ ): $\mathrm{CE}=\$ 283,574$.


## Summary

Table 5: Growth Monitoring and Healthy Weight Management in Children / Youth in a Birth Cohort of 40,000 Summary Base


## Preventing Tobacco Use

## Canadian Task Force on Preventive Health Care Recommendations (2017)

We recommend asking children and youth (age 5-18 yr) or their parents about tobacco use by the child or youth and offering brief information and advice, as appropriate, during primary care visits to prevent tobacco smoking among children and youth (weak recommendation, low-quality evidence).

We recommend asking children and youth (age 5-18 yr) or their parents about tobacco use by the child or youth and offering brief information and advice, as appropriate, during primary care visits to treat tobacco smoking among children and youth (weak recommendation, low-quality evidence). ${ }^{167}$

## United States Preventive Services Task Force Recommendations (2013)

The USPSTF recommends that primary care clinicians provide interventions, including education or brief counseling, to prevent initiation of tobacco use in school-aged children and adolescents. (B Recommendation) ${ }^{168}$

In their review of the evidence, ${ }^{169}$ the USPSTF noted that the 2012 Surgeon General's Report concluded that there is a "large, robust, and consistent" evidence base that documents known effective strategies for reducing tobacco use among youths and young adults. ${ }^{170}$ These strategies include coordinated, multi-component campaigns that combine media campaigns, price increases, school-based policies and programs and community-wide changes in policies and norms. The purpose of the USPSTF review was not to reconsider the evidence covered by the Surgeon General's Report, but rather "to review the evidence for the efficacy and harms of primary-care relevant interventions that aim to reduce tobacco use among children and adolescents." ${ }^{171}$

## Modelling the Clinically Preventable Burden

In this section, we model CPB associated with asking children and youth or their parents about tobacco use by the child or youth and offering brief information and advice, as appropriate, during primary care visits to prevent and / or treat tobacco smoking among children and youth.

[^35]In modelling CPB, we made the following assumptions:

- Interventions aimed at reducing smoking initiation among non-smoking children and adolescents have an effectiveness of $18 \%$ (RR $0.82,95 \% \mathrm{CI}$ of 0.72 to 0.94 ). ${ }^{172}$
- Interventions aimed at smoking cessation among children and adolescents have an effectiveness of $34 \%$ (RR $1.34,95 \%$ CI of 1.05 to 1.69 ). ${ }^{173}$
- An estimated $12.34 \%$ of 19 year-olds were daily or occasional smokers in BC in 2010 (see Table 1). ${ }^{174}$

| Table 1: Smokers in British Columbia in 2010 <br> Based on 2010 CCHS Data <br> Ages 12 to 19 |  |  |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Total Population |  |  | Daily Smokers |  |  | Occasional Smokers |  |  | Current Smokers as \% of Pop. |  |  |
| Age Group | Males | Females | Total | Males | Females | Total | Males | Females | Total | Males | Females | Total |
| 12-14 | 73,171 | 68,779 | 141,950 | 459 | - | 459 | 97 | - | 97 | 0.76\% | 0.00\% | 0.39\% |
| 15-17 | 81,088 | 74,831 | 155,919 | 4,383 | 2,994 | 7,377 | 1,274 | 208 | 1,482 | 6.98\% | 4.28\% | 5.68\% |
| 18-19 | 57,055 | 55,256 | 112,311 | 4,661 | 4,479 | 9,140 | 3,541 | 1,175 | 4,716 | 14.38\% | 10.23\% | 12.34\% |
| Total | 211,314 | 198,866 | 410,180 | 9,503 | 7,473 | 16,976 | 4,912 | 1,383 | 6,295 | 6.82\% | 4.45\% | 5.67\% |

- On average, $57.3 \%$ of smokers would quit (become former smokers) by the age of $25-34$ (Table 3, row $e$ ), $60.4 \%$ by age $35-44$ (Table 3 , row $h$ ) and $68.9 \%$ by age $45-54$ (Table 3, row $k$ ) (see Table 2). ${ }^{175}$

| Table 2: Smoking Occurrence British Columbia, 2010 |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| SMOKING CATEGORY | AGE GROUP |  |  |  |  |  |
|  | 18-24 | 25-34 | 35-44 | 45-54 | 55-64 | 65+ |
| DAILY SMOKER | 50,238 | 91,696 | 94,232 | 114,679 | 70,612 | 47,346 |
| OCCASIONAL SMOKER (FORMER DAILY SMOKER) | 17,203 | 27,935 | 21,481 | 18,486 | 9,914 | 12,950 |
| ALWAYS AN OCCASIONAL SMOKER | 31,786 | 18,272 | 15,056 | 7,787 | 6,320 | 296 |
| FORMER DAILY SMOKER | 27,365 | 77,671 | 110,446 | 203,967 | 183,720 | 256,094 |
| FORMER OCCASIONAL SMOKER | 53,224 | 107,195 | 89,353 | 108,870 | 83,717 | 92,489 |
| NEVER SMOKED | 225,389 | 267,255 | 288,143 | 265,911 | 209,738 | 223,185 |
| SMOKERS | 179,816 | 322,769 | 330,568 | 453,789 | 354,283 | 409,175 |
| \% of FORMER SMOKERS | 44.8\% | 57.3\% | 60.4\% | 68.9\% | 75.5\% | 85.2\% |

- An average of 11.5 life years lost per smoker (Table 3, row $c$ ). An average of 10.5 of those life-years can be regained by stopping smoking at age 30 (Table 3 , row $g$ ), 9.5 by stopping smoking at age 40 (Table 3 , row $j$ ) and 6.5 by stopping smoking at age 50 (Table 3, row $l$ ). ${ }^{176}$

[^36]- Other assumptions used in assessing the clinically preventable burden are detailed in the Reference Document.

Based on these assumptions, the CPB associated with interventions aimed at preventing and / or treating tobacco smoking among children and youth is 4,123 QALYs (Table 3, row $g g$ ). The CPB of 4,123 represents the gap between no coverage and the 'best in the world' coverage, which was estimated at $53 \%$.

Table 3: CPB of Interventions for Tobacco Use Prevention and Cessation in Children and Youth for Birth Cohort of $\mathbf{4 0 , 0 0 0}$ Individuals (B.C.)

| Estimate of Life Years Lost without Intervention | Base Case | Data Source |
| :---: | :---: | :---: |
| a \% of 19 year-olds who smoke in B.C. | 12.34\% | Table 1 |
| b Estimated \# in birth cohort initiating smoking by age 19 | 4,935 | = a* 40,000 |
| c Life-years lost per smoker | 11.5 | $\checkmark$ |
| d Potential life-years lost | 56,751 | $=c^{*} \mathrm{~b}$ |
| e Proportion former smokers at age 30 | 57.3\% | Table 2 |
| f Former smokers at age 30 | 2,828 | $=e^{*} \mathrm{~b}$ |
| g Life-years gained by stopping smoking at age 30 | 10.5 | $\checkmark$ |
| h Proportion former smokers at age 40 | 60.4\% | Table 2 |
| i Former smokers at age 40 | 2,981 | $=h^{*} \mathrm{~b}$ |
| j Life-years gained by stopping smoking at age 40 | 9.5 | $\checkmark$ |
| k Proportion former smokers at age 50 | 68.9\% | Table 2 |
| I Life-years gained by stopping smoking at age 50 | 6.5 | $\checkmark$ |
| m Former smokers at age 50 | 3,400 | = ${ }^{*}$ b |
| $n \quad$ Life-years gained by stopping smoking | 33,871 | $\begin{gathered} =\left(f^{*} \mathrm{~g}\right)+(\mathrm{i}- \\ \mathrm{f}) * \mathrm{j}+(\mathrm{m}-\mathrm{i})^{*} \end{gathered}$ |
| o Estimated Life Years Lost without Intervention | 22,881 | $=\mathrm{d}-\mathrm{n}$ |
| Estimate of Life Years Lost with Intervention |  |  |
| $p$ Effectiveness of intervention | 34.0\% | $\checkmark$ |
| q Estimated \# in birth cohort initiating smoking by age 19 | 3,257 | $\begin{gathered} =a *(1-p) \\ * 40,000 \end{gathered}$ |
| r Life-years lost per smoker | 11.5 | $\checkmark$ |
| s Potential life-years lost | 37,456 | $=r^{*} \mathrm{q}$ |
| t Proportion former smokers at age 30 | 57.3\% | Table 2 |
| u Former smokers at age 30 | 1,866 | = t ${ }^{\text {q }}$ |
| $v$ Life-years gained by stopping smoking at age 30 | 10.5 | $\checkmark$ |
| w Proportion former smokers at age 40 | 60.4\% | Table 2 |
| $x$ Former smokers at age 40 | 1,967 | = ${ }^{*}$ q |
| y Life-years gained by stopping smoking at age 40 | 9.5 | $\checkmark$ |
| z Proportion former smokers at age 50 | 68.9\% | Table 2 |
| aa Life-years gained by stopping smoking at age 50 | 6.5 | $\checkmark$ |
| bb Former smokers at age 50 | 2,244 | $=z^{*} \mathrm{q}$ |
| cc Life-years gained by stopping smoking | 22,355 | $\begin{gathered} =\left(u^{*} v\right)+(x- \\ u)^{*} y+(b b-x) * a a \end{gathered}$ |
| dd Estimated Life Years Lost with Intervention | 15,101 | =s-cc |
| Calculation of CPB |  |  |
| ee Life-years gained with 100\% adherence | 7,779 | = o-dd |
| ff Potential coverage of this service | 53\% | Ref Doc |
| gg Potential CPB in BC | 4,123 | $=\mathrm{ee}$ *ff |

$\checkmark=$ Estimates from the literature
We also modified a major assumption and recalculated the CPB as follows:

- Assume the effectiveness of interventions aimed at smoking cessation among children and adolescents is reduced from $34 \%$ to $5 \%$ (Table 3, row $p$ ): $\mathrm{CPB}=606$.
- Assume the effectiveness of interventions aimed at smoking cessation among children and adolescents is increased from $34 \%$ to $69 \%$ (Table 3, row $p$ ): CPB $=$ 8,367.


## Modelling Cost-Effectiveness

In this section, we model CE associated with asking children and youth or their parents about tobacco use by the child or youth and offering brief information and advice, as appropriate, during primary care visits to prevent and/or treat tobacco smoking among children and youth.

In estimating CE, we made the following assumptions:

- The USPSTF evidence review suggests that the effectiveness of the intervention lasts for at least two years. ${ }^{177}$ We have assumed that an intervention would be required seven times between the ages of 5 and 19 for maximum effect (Table 4, row $d$ ).
- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the CE associated with interventions to prevent and/or treat tobacco smoking among children and youth.is $-\$ 7,349$ per QALY (Table 4, row $p$ ).

[^37]
## Table 4: Cost Effectiveness of Interventions for Tobacco Use Prevention in Children

 and Youth for Birth Cohort of $\mathbf{4 0 , 0 0 0}$ Individuals (B.C.)|  |  | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| Cost of counseling |  |  |  |
| a | Cost of 10-minute office visit | \$34.85 | Ref Doc |
| b | Cost of patient time and travel for office visit | \$59.38 | Ref Doc |
| c | Portion of office visit needed for counseling | 50\% | Ref Doc |
| d | \# of interventions | 7.0 | $\checkmark$ |
| e | Total cost of counseling per individual | \$329.81 | $=(a+b) *{ }^{*}{ }^{*}$ |
| f | Estimated Cost of Counselling | \$13,192,200 | = e * 40,000 |
| Estimated Cost Avoidance |  |  |  |
| g | Annual medical costs avoided per additional year as never smoker | \$1,195 | Ref Doc |
| h | Years of smoking avoided due to intervention | 43,950 | Calculated |
| i | Costs avoided | \$52,520,012 | = g * h |
| CE calculation |  |  |  |
| j | Estimated Cost of Counselling | \$13,192,200 | = f |
| k | Costs avoided | \$52,520,012 | = i |
| 1 | Potential QALYs saved | 4,123 | = Table 3, row gg |
| m | Estimated Cost of Counselling (1.5\% discount rate) | \$11,830,577 | Calculated |
| n | Costs avoided (1.5\% discount rate) | \$27,965,774 | Calculated |
| 0 | Potential QALYs saved (1.5\% discount rate) | 2,195 | Calculated |
| p | Cost per QALY (CE) | -\$7,349 | $=(\mathrm{m}-\mathrm{n}) / \mathrm{o}$ |

Notes: $\quad V=$ Estimates from the literature

We also modified several major assumptions and recalculated the cost per QALY as follows:

- Assume the effectiveness of interventions aimed at smoking cessation among children and adolescents is reduced from 34\% to 5\% (Table 3, row $p$ ): \$/QALY = \$23,905.
- Assume the effectiveness of interventions aimed at smoking cessation among children and adolescents is increased from $34 \%$ to $69 \%$ (Table 3, row $p$ ): $\$ / \mathrm{QALY}=$ - $\$ 10,083$.
- Assume the portion of an office visit needed for counseling is reduced from $50 \%$ to $33 \%$ (Table 4, row $c$ ): $\$ / \mathrm{QALY}=-\$ 9,182$.
- Assume the portion of an office visit needed for counseling is increased from $50 \%$ to $67 \%$ (Table 4, row $c$ ): \$/QALY = -\$5,517.


## Summary

Table 5: Interventions for Tobacco Use Prevention and Cessation in Children and Youth for Birth Cohort of 40,000

Summary
Base
Case
Range
CPB (Potential QALYs Gained)

| Assume No Current Service |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: |
| 1.5\% Discount Rate | $\mathbf{2 , 1 9 5}$ |  | 323 | 4,455 |
| 3\% Discount Rate | 1,206 |  | 177 | 2,447 |
| 0\% Discount Rate | 4,123 |  | 606 | 8,367 |
| CE (\$/QALY) including patient time costs |  |  |  |  |
| 1.5\% Discount Rate | $-\$ 7,349$ |  | $-\$ 10,083$ | $\$ 23,905$ |
| 3\% Discount Rate | $-\$ 3,909$ |  | $-\$ 8,388$ | $\$ 47,299$ |
| 0\% Discount Rate | $-\$ 9,538$ |  | $-\$ 11,161$ | $\$ 9,019$ |
| CE (\$/QALY) excluding patient time costs |  |  |  |  |
| 1.5\% Discount Rate | $-\$ 10,745$ |  | $-\$ 11,756$ | $\$ 814$ |
| 3\% Discount Rate | $-\$ 9,473$ |  | $-\$ 11,129$ | $\$ 9,466$ |
| 0\% Discount Rate | $-\$ 11,555$ |  | $-\$ 12,155$ | $-\$ 4,691$ |

## Preventive Medication / Devices

Fluoride Varnish and Fissure Sealants for Dental Health in Children
United States Preventive Service Task Force Recommendations (2014)
Dental caries is the most common chronic disease in children in the United States. According to the 1999-2004 National Health and Nutrition Examination Survey (NHANES), $\sim 42 \%$ of children ages 2 to 11 years have dental caries in their primary teeth. After decreasing from the early 1970s to the mid-1990s, the prevalence of dental caries in children has been increasing, particularly in young children ages 2 to 5 years.

The U.S. Preventive Services Task Force recommends that primary care clinicians prescribe oral fluoride supplementation starting at age 6 months for children whose water supply is deficient in fluoride. (B recommendation)

The U.S. Preventive Services Task Force recommends that primary care clinicians apply fluoride varnish to the primary teeth of all infants and children starting at the age of primary tooth eruption. (B recommendation) ${ }^{178}$

## Canadian Task Force on Preventive Health Care Recommendations (1994)

Lower dental caries prevalence and the need for efficiency in the provision of preventive and therapeutic dental services require selective use of dental caries preventives and targeting of services toward persons at greatest risk. The following recommendations are based on a review of the available evidence.
There is good evidence of effectiveness of the following measures in preventing dental caries (A Recommendation):

1. Water fluoridation for preventing coronal and root caries;
2. Fluoride supplements in low fluoride areas with careful adherence to low dosage schedules;
3. Professional topical fluoride applications and self-administered fluoride mouth rinses for those with very active decay or at high future risk for dental caries;
4. Fluoride dentifrices, with special supervision and the use of small amounts for young children;
5. Professionally-applied fissure sealants for selective use on permanent molar teeth soon after their eruption. ${ }^{179}$

## The Cochrane Oral Health Group (2017)

Resin-based sealants applied on occlusal surfaces of permanent molars are effective for preventing caries in children and adolescents. Our review found moderate-quality evidence that resin-based sealants reduced caries by between $11 \%$ and $51 \%$ compared to no sealant, when measured at 24 months. ${ }^{180}$

## Fluoride Varnish - Modelling the Clinically Preventable Burden

In this section, we model the CPB associated with applying fluoride varnish every six months between the ages of one and five for the prevention of dental caries in children.

In modelling CPB, we made the following assumptions:

- In 2012/13, 91.8\% of BC kindergarten children were screened for dental health. Of these, $67.3 \%$ were caries free, $18.1 \%$ had treated caries and $14.6 \%$ had visible decay (Table 1, row $a$ ). ${ }^{181}$

[^38]- The effectiveness of fluoride varnish in reducing decayed, missing and filled teeth is $37 \%$ with a $95 \%$ CI of $24 \%$ to $51 \%$ (Table 1, row b). ${ }^{182}$
- Other assumptions used in assessing the clinically preventable burden are detailed in the Reference Document.

Based on these assumptions, the CPB associated with applying fluoride varnish every six months between the ages of one and five for the prevention of dental caries in children is 150 (Table 1, row $i$ ).

| Years of Age in a Birth Cohort of 40,000 (B.C.) |  |  |  |
| :---: | :---: | :---: | :---: |
| Row <br> Label | Variable | Base Case | Data Source |
| a | Proportion of B.C. kindergarten children caries free | 67.3\% | $\checkmark$ |
| b | Effectiveness of fluoride varnish in reducing decayed, missing and filled tooth surfaces | 37.0\% | $\checkmark$ |
| c | Adherence with intervention | 62\% | Ref Doc |
| d | Children with treated caries or visible decay | 13,080 | $=(1-a) * 40,000$ |
| e | Children benefitting from intervention | 3,001 | $=(\mathrm{d} * \mathrm{c}) * \mathrm{~b}$ |
| f | Years of benefits (from ages 1 to 5) per child | 5.0 | $\checkmark$ |
| g | Life-years lived with poor oral health | 15,003 | $=e^{*} \mathrm{f}$ |
| h | Change in QoL associated with improved oral health | 0.01 | Ref Doc |
| i | Potential QALYs gained, CPB | 150 | = $\mathrm{F}^{*} \mathrm{~h}$ |

$V=$ Estimates from the literature
We also modified several major assumptions and recalculated the CPB as follows:

- Assume the effectiveness of fluoride varnish in reducing decayed, missing and filled teeth is reduced from $37 \%$ to $24 \%$ (Table 1, row b): CPB $=97$
- Assume the effectiveness of fluoride varnish in reducing decayed, missing and filled teeth is increased from $37 \%$ to $51 \%$ (Table 1, row $b$ ): $\mathrm{CPB}=207$
- Assume the change in QoL associated with improved oral health is reduced from 0.01 to 0.005 (Table 1, row $h$ ): $\mathrm{CPB}=75$
- Assume the change in QoL associated with improved oral health is increased from 0.01 to 0.019 (Table 1, row $h$ ): $\mathrm{CPB}=285$


## Fluoride Varnish - Modelling Cost-Effectiveness

In this section, we model the CE associated with applying fluoride varnish every six months between the ages of one and five for the prevention of dental caries in children.

In modelling CE, we made the following assumptions:

- Fluoride varnish would be available for application to all children in BC with a $62 \%$ adherence rate (Table 2, row $b$ ).
- Assume fluoride varnish would need to be applied once every six months from age 1 to age 5 for a total of 9 applications (Table 2, row $f$ ). ${ }^{183}$

[^39]- For patient time and travel costs, we assumed an hour of patient time required per dental visit and three hours of patient time for dental day surgery. Dental day surgery in BC lasts an average of 83 minutes. ${ }^{184}$
- Assume 2.9 new carious surfaces per untreated 5 year-old (Table 2, row $g$ ). ${ }^{185}$
- The prevalence for day surgery for dental cavities in BC is estimated to be $1.38 \%$ of children (Table 2, row $l$ ). ${ }^{186}$
- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the CE associated with applying fluoride varnish every six months between the ages of one and five for the prevention of dental caries in children is $\$ 43,048$ per QALY (Table 2, row $y$ ).

[^40]Table 2: CE of Flouride Varnish for the Prevention of Dental Caries in Children < 5 Years of Age in a Birth Cohort of 40,000 (B.C.)

| Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Children eligible for intervention | 40,000 | $\checkmark$ |
| b | Adherence with intervention | 62\% | = Table 1 row c |
| C | Children with treated caries or visible decay | 13,080 | = Table 1 row d |
|  | Costs of intervention |  |  |
| d | Cost of flouride varnish application | \$10.61 | Ref Doc |
| e | Value of patient time and travel for office visit | \$29.69 | Ref Doc |
| f | \# of times flouride varnish applied from age 1 to 5 | 9 | $\checkmark$ |
| g | Estimated cost of intervention over lifetime of birth cohort | \$8,994,960 | $=(\mathrm{d}+\mathrm{e}) * \mathrm{f} * \mathrm{a}$ * b |
|  | Cost avoided |  |  |
| h | New carious surfaces per untreated 5 year-old | 2.9 | $\checkmark$ |
| i | Dental caries avoided | 14,035 | = g* ${ }^{*}$ Table 1 row b |
| j | Cost per filling | \$92.75 | Ref Doc |
| k | Value of patient time and travel for office visit | \$59.38 | Ref Doc |
| I | Filling costs avoided | -\$2,135,120 | $=(\mathrm{i}+\mathrm{j}) * \mathrm{~h}$ |
| m | Prevalence of day surgery for caries | 1.38\% | $\checkmark$ |
| n | Day surgeries without intervention in birth cohort | 552 | = ${ }^{*} \mathrm{~m}$ |
| 0 | Day surgeries avoided with intervention in birth cohort | 204 | $=\mathrm{m}^{*}$ Table 1 row b |
| p | Cost of day surgery | \$1,884 | Ref Doc |
| q | Value of patient time and travel for day surgery | \$89.07 | Ref Doc |
| $r$ | Day surgery costs avoided | -\$402,980 | $=(\mathrm{p}+\mathrm{q}) *$ o |
|  | CE calculation |  |  |
| S | Cost of intervention over lifetime of birth cohort | \$8,994,960 | = g |
| t | Costs avoided | -\$2,538,100 | $=1+r$ |
| $u$ | QALYs saved | 150 | Table 8-1 row i |
| v | Cost of intervention over lifetime of birth cohort (1.5\% discount) | \$8,605,388 | Calculated |
| w | Costs avoided (1.5\% discount) | -\$2,428,175 | Calculated |
| x | QALYs saved (1.5\% discount) | 144 | Calculated |
| y | CE (\$/QALY saved) | \$43,038 | $=(v+w) / x$ |

$v=$ Estimates from the literature
We also modified several major assumptions and recalculated the cost per QALY as follows:

- Assume the effectiveness of fluoride varnish in reducing decayed, missing and filled teeth is reduced from $37 \%$ to $24 \%$ (Table 1 , row $b$ ): $\mathrm{CE}=\$ 75,514$
- Assume the effectiveness of fluoride varnish in reducing decayed, missing and filled teeth is increased from $37 \%$ to $51 \%$ (Table 1, row b): $\mathrm{CE}=\$ 26,579$
- Assume the change in QoL associated with improved oral health is reduced from 0.01 to 0.005 (Table 1, row $h$ ): $\mathrm{CE}=\$ 86,076$
- Assume the change in QoL associated with improved oral health is increased from 0.01 to 0.019 (Table 1, row $h$ ): $\mathrm{CE}=\$ 22,651$
- Assume that the application of fluoride varnish is equally effective if applied annually (versus every six months) (Table 2, row $f$ ). The evidence on frequency of applications is inconclusive ${ }^{187}$ : $\mathrm{CE}=\$ 16,391$
- Assume that the cost per filling is reduced from $\$ 92.75$ to $\$ 83.10$ (Table 2, row $j$ ): $C E=\$ 43,941$

[^41]- Assume that the cost per filling is increased from $\$ 92.75$ to $\$ 102.40$ (Table 2, row $j$ ): $C E=\$ 42,135$


## Fluoride Varnish - Summary

| Table 3: Application of Years of Age in | de Var <br> Coho mary |  | $\begin{aligned} & \text { ildren } \\ & 10 \end{aligned}$ |
| :---: | :---: | :---: | :---: |
|  | $\begin{aligned} & \text { Base } \\ & \text { Case } \\ & \hline \end{aligned}$ |  |  |
| CPB (Potential QALYs Gained) |  |  |  |
| Assume No Current Service |  |  |  |
| 1.5\% Discount Rate | 144 | 72 | 273 |
| 3\% Discount Rate | 137 | 69 | 261 |
| 0\% Discount Rate | 150 | 75 | 285 |
| $\overline{\mathrm{CE}}$ (\$/QALY) including patient time costs |  |  |  |
| 1.5\% Discount Rate | \$43,038 | \$16,391 | \$86,076 |
| 3\% Discount Rate | \$43,038 | \$16,391 | \$86,076 |
| 0\% Discount Rate | \$43,038 | \$16,391 | \$86,076 |
| CE (\$/QALY) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | \$4,543 | -\$2,472 | \$9,087 |
| 3\% Discount Rate | \$4,543 | -\$2,472 | \$9,087 |
| 0\% Discount Rate | \$4,543 | - $\$ 2,472$ | \$9,087 |

## Dental Sealants - Modelling the Clinically Preventable Burden

While the focus of the USPSTF is on improving dental health in preschool children, there is also a body of evidence indicating that the use of dental sealants is effective in preventing decayed, missing and filled teeth in children six years of age and older with permanent teeth. ${ }^{188}$

In this section, we model the CPB associated with applying dental sealants for the prevention of dental caries in children and youth with permanent teeth.

In modelling CPB, we made the following assumptions:

- Dental sealants would be placed on the $1^{\text {st }}$ molars at age six, the $1^{\text {st }}$ and $2^{\text {nd }}$ bicuspids at age 10 and the $2^{\text {nd }}$ molars at age 12 .
- The effectiveness of dental sealants in reducing decayed, missing and filled teeth is $84 \%$ at year 1, decreasing to $55 \%$ at year 9 . Effectiveness beyond nine years is unknown. ${ }^{189}$
- An estimated $12.2 \%$ of Canadians avoid certain foods because of problems with their teeth or mouth, and $11.6 \%$ of Canadians sometimes or always have pain in their mouth. ${ }^{190}$ Based on this information, we assumed that $12 \%$ of children/youth with

[^42]caries would have significant enough pain to reduce their quality of life (Table 4, row j).

- Other assumptions used in assessing the clinically preventable burden are detailed in the Reference Document.

Based on these assumptions, the CPB associated with preventing decayed, missing and filled teeth in children with permanent teeth is 157 (Table 4, row $m$ ). The CPB of 157 represents the gap between no coverage and improving coverage to $59 \%$.

Table 4: CPB of Dental Sealants in Children/Youth with Permanent Teeth in a Birth Cohort of 40,000 (B.C.)

| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | \# of 6-year olds in a birth cohort of 40,000 | 39,818 | Ref Doc |
| b | Adherence with intervention | 59\% | Ref Doc |
| c | Children 'accepting' intervention | 23,492 | = a * b |
| d | Estimated new caries between ages 6-20 per child - untreated | 7.69 | Calculated |
| e | Estimated new caries between ages 6-20 per child - treated | 2.46 | Calculated |
| $f$ | Estimated new caries without intervention | 180,615 | = ${ }^{*} \mathrm{~d}$ |
| g | Estimated new caries with intervention | 57,718 | = ${ }^{*}$ e |
| h | New caries avoided with intervention | 122,898 | = f-g |
| 1 | Life-years lived without caries due to intervention | 130,643 | Calculated |
| j | Proportion of children living with caries with significant pain | 12.0\% | $\checkmark$ |
| k | Life-years lived without caries or pain due to intervention | 15,677 | $=1^{*} \mathrm{j}$ |
| 1 | Change in QoL associated with improved oral health | 0.01 | Ref Doc |
| m | Potential QALYs gained, Intervention increasing from 0\% to 59\% | 157 | = ${ }^{*}$ ] |

$V=$ Estimates from the literature
We also modified a major assumption and recalculated the CPB as follows:

- Assume the change in QoL associated with improved oral health is reduced from 0.01 to 0.005 (Table 4, row $l$ ): $\mathrm{CPB}=78$
- Assume the change in QoL associated with improved oral health is increased from 0.01 to 0.019 (Table 4, row $l$ ): $\mathrm{CPB}=298$


## Dental Sealants - Modelling Cost-Effectiveness

In this section, we model the CE associated with applying dental sealants for the prevention of dental caries in children and youth with permanent teeth.

In modelling CE, we made the following assumptions:

- The cost of applying sealants is estimated at $\$ 19.74$ for the first tooth in a quadrant and $\$ 10.83$ for each additional tooth in the quadrant (see Reference Document). The costs of applying dental sealants on the $1^{\text {st }}$ molars at age six would therefore be $\$ 78.96$, the $1^{\text {st }}$ and $2^{\text {nd }}$ bicuspids at age 10 would be $\$ 122.32$ and the $2^{\text {nd }}$ molars at age 12 would be $\$ 78.96$ for a total cost of $\$ 280.24$ (Table 5, row $d$ ).
- For patient time and travel costs, we estimated two hours of patient time per dental visit.
- An average of 1.84 fillings would be treated each time fillings are required (Table 5, row $l$ ). ${ }^{191}$
- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the CE associated with preventing dental caries in children with permanent teeth is $-\$ 24,690$ per QALY (Table 5, row $v$ ).

Table 5: CE of Dental Sealants in Children/Youth with Permanent Teeth in a Birth Cohort of $\mathbf{4 0 , 0 0 0}$ (B.C.)

| Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Children eligible for intervention | 39,818 | = Table 4, row a |
| b | Adherence with intervention | 59\% | = Table 4, row b |
| C | Children 'accepting' intervention | 23,492 | = Table 4, row c |
|  | Costs of intervention |  |  |
| d | Cost of dental sealant applications | \$280.24 | $\checkmark$ |
| e | Value of patient time and travel for office visit | \$59.38 | Ref Doc |
| f | \# of sealant applications (at age 6, 10 and 12) | 3 | $\checkmark$ |
| g | Estimated cost of intervention over lifetime of birth cohort | \$6,583,506 | $=c^{*} \mathrm{~d}$ |
| h | Estimated cost of patient time over lifetime of birth cohort | \$4,184,933 | $=c^{*} e^{*} \mathrm{f}$ |
|  | Cost avoided |  |  |
| i | Dental caries avoided with intervention | 122,898 | Calculated |
| j | Cost per filling | \$92.75 | Ref Doc |
| k | Value of patient time and travel for office visit | \$59.38 | Ref Doc |
| 1 | \# of fillings per visit | 1.84 | $\checkmark$ |
| m | \# of dental visits avoided | 66,792 | = i/l |
| n | Filling costs avoided | -\$11,398,770 | $=i^{*} \mathrm{j}$ |
| 0 | Patient costs avoided | -\$3,966,125 | $=\mathrm{m}^{*} \mathrm{k}$ |
|  | CE calculation |  |  |
| p | Cost of intervention over lifetime of birth cohort | \$10,768,439 | $=\mathrm{g}+\mathrm{h}$ |
| q | Costs avoided | -\$15,364,896 | $=\mathrm{n}+\mathrm{o}$ |
| $r$ | QALYs saved | 157 | Table 4, row k |
| S | Cost of intervention over lifetime of birth cohort (1.5\% discount) | \$10,096,096 | Calculated |
| t | Costs avoided (1.5\% discount) | -\$13,499,918 | Calculated |
| u | QALYs saved (1.5\% discount) | 138 | Calculated |
| v | CE (\$/QALY saved) | -\$24,690 | $=(\mathrm{s}+\mathrm{t}) / \mathrm{u}$ |

V = Estimates from the literature
We also modified several major assumptions and recalculated the cost per QALY as follows:

- Assume the change in QoL associated with improved oral health is reduced from 0.01 to 0.005 (Table 4, row $l$ ): $\mathrm{CE}=-\$ 24,359$
- Assume the change in QoL associated with improved oral health is increased from 0.01 to 0.019 (Table 4, row $l$ ): $\mathrm{CE}=-\$ 24,851$
- Assume that the cost per filling is reduced from $\$ 92.75$ to $\$ 83.10$ (Table 5, row $j$ ): $C E=-\$ 17,132$

[^43] Center for Health Statistics. 2007; 11(248): 1-104.

- Assume that the cost per filling is increased from $\$ 92.75$ to $\$ 102.40$ (Table 5, row $j$ ): CE $=-\$ 32,248$


## Dental Sealants - Summary



## Clinical Prevention in Adults

## Screening for Asymptomatic Disease or Risk Factors

Screening for Breast Cancer

## Canadian Task Force on Preventive Health Care Recommendations (2011)

For women aged 40-49 we recommend not routinely screening with mammography. (Weak recommendation; moderate quality evidence)

For women aged 50-69 years we recommend routinely screening with mammography every 2 to 3 years. (Weak recommendation; moderate quality evidence)

For women aged 70-74 we recommend routinely screening with mammography every 2 to 3 years. (Weak recommendation; low quality evidence) ${ }^{192}$

## United States Preventive Services Task Force Recommendations (2016)

The USPSTF recommends biennial screening mammography for women aged 50 to 74 years. (B recommendation) ${ }^{193}$

## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with screening women ages 50 to 74 years of age for breast cancer every 2 to 3 years.

In modelling CPB, we made the following assumptions:

- Based on BC life tables for 2010 to 2012, a total of 3,938 deaths would be expected in females between the ages of $50-79$ in a BC birth cohort of 40,000 (see Table 1). While routine screening occurs to age 74 , we have assumed the protective effect of that routine screening would continue to age 79 .
- Based on BC vital statistics data, there were 1,990 deaths in females between the ages of 45 and 64 in BC in 2012, with 215 ( $10.8 \%$ ) of these deaths due to breast cancer (ICD-10 codes C50). There were also 3,566 deaths between the ages of 65 and 79 that year, with $230(6.4 \%)$ of these deaths due to breast cancer. ${ }^{194}$ This suggests that 288 of the $3,938(7.3 \%)$ of the female deaths in the BC birth cohort between the ages of 50 and 79 would be due to breast cancer (see Table 1).

[^44]

- Screening mammography in women ages 50-74 leads to a reduction in breast cancer mortality of $21 \%$ (RR $0.79,95 \%$ CI of $0.68-0.90$ ). This is based on 10 trials in which the attendance rates at first screening were approximately $85 \%$. ${ }^{195}$
- For every death avoided, 204 women will have false positive results. ${ }^{196} \mathrm{We}$ have assumed a one-time QALY loss of 0.013 (4.7 days) after a false-positive mammography result. ${ }^{197}$
- For every death avoided, 26 women will have an unnecessary biopsy. ${ }^{198}$
- For every death avoided, 3 women will have an unnecessary lumpectomy or mastectomy (with a 3:1 ratio for lumpectomy vs. mastectomy). ${ }^{199}$
- Other assumptions used in assessing the clinically preventable burden are detailed in the Reference Document.

Based on these assumptions, the CPB associated with screening women ages 50 to 74 years of age for breast cancer every 2 to 3 years is 1,189 QALYs saved (Table 2, row o). The CPB of 1,189 represents the gap between no coverage and the 'best in the world' coverage estimated at $88 \%$. The CPB of 486 QALYs saved (see Table 2, row $p$ ) represents the gap between the current coverage of $52 \%$ and the 'best in the world' coverage estimated at $88 \%$.

[^45]Table 2. Calculation of Clinically Preventable Burden of Breast Cancer Screening Being Offered to a Birth Cohort of 40,000 Between the Ages of 50 to 74

| Row | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
|  | Estimated Current Status |  |  |
| a | Estimated deaths due to breast cancer in birth cohort between ages 50-79 | 288 | Table 1 |
| b | Effectiveness of mammography screening in preventing mortality (based on $85 \%$ adherence in clinical trials) | 21.0\% | $\checkmark$ |
| C | Effectiveness of mammography screening in preventing mortality (assuming 100\% adherence in clinical trials) | 24.7\% | $={ }^{\text {b* }} 1.1764$ |
| d | Frequency of screening in last 30 months | 52\% | Ref Doc |
| e | Potential adherence | 88\% | Ref Doc |
| f | Predicted deaths in the absence of screening | 331 | $=a /\left(1-d^{*} \mathrm{c}\right)$ |
|  | Benefits of Screening |  |  |
| g | Deaths avoided - 100\% adherence | 82 | $=\mathrm{f}^{*} \mathrm{c}$ |
| h | Deaths avoided - 88\% adherence | 72 | $=\mathrm{g}^{*} \mathrm{e}$ |
| i | Deaths avoided - 52\% adherence | 42 | $=\mathrm{g}^{*} \mathrm{~d}$ |
| j | Life expectancy at average age of breast cancer death | 19.2 | Table 1 |
| k | QALYs saved with 88\% adherence to screening | 1,379 | = ${ }^{*}{ }^{\text {j }}$ |
|  | Harms Associated with Screening |  |  |
| 1 | False positive results per death avoided | 204 | $\checkmark$ |
| m | Reduced QALYs per false positive | 0.013 | $\checkmark$ |
| n | Reduced QALYs associated with false positives | -191 | $=\mathrm{h}^{*} \\|^{*} \mathrm{~m}$ |
|  | Summary of Benefits and Harms |  |  |
| 0 | Potential QALYs saved - Utilization increasing from 0\% to 88\% | 1,189 | $=\mathrm{k}+\mathrm{n}$ |
| p | Potential QALYs saved - Utilization increasing from 52\% to 88\% | 486 | $=0^{*}(e-d) / e$ |

We modified the following major assumptions and recalculated the CPB as follows:

- Assume the effectiveness of screening mammography in reducing deaths from breast cancer is reduced from $21 \%$ to $10 \%$ (Table 2, row $b$ ): $\mathrm{CPB}=526$.
- Assume the effectiveness of screening mammography in reducing deaths from breast cancer is increased from $21 \%$ to $32 \%$ (Table 2, row $b$ ): $\mathrm{CPB}=1,963$.


## Modelling Cost-Effectiveness

In this section, we will calculate the CE associated with screening women ages 50 to 74 years of age for breast cancer every 2 to 3 years.

In estimating the CE of screening mammography, we made the following assumptions:

- Costs of screening - Information from the BC Cancer Agency Screening

Mammography Program indicates a cost of $\$ 79.35$ per screen in 2015/16. ${ }^{200}$ There are a total of 462,381 life years lived in females ages $50-74$ in a BC birth cohort of 40,000 (see Table 1). We assumed that, on average, women would participate in screening once every 30 months (i.e., every 2.5 years), resulting in 184,952 screens for the birth cohort assuming $100 \%$ adherence. At $88 \%$ adherence, the number of screens would be reduced to 162,758 (Table 3, row $a \& b$ ).

[^46]- Costs associated with overtreatment - For every death avoided, 3 women will have an unnecessary lumpectomy or mastectomy (with a 75:25 ratio for lumpectomy vs. mastectomy) with a cost per lumpectomy of $\$ 5,152$ and a mastectomy of $\$ 7,260$ (see reference document) for a weighted cost of $\$ 5,679$ (Table 2 , row $k$ ).
- Patient time and travel costs - For patient time and travel costs, we assumed an estimated two hours of patient time required per screening visit of $\$ 57.56,7.5$ for a biopsy and 37.5 hours for a lumpectomy or mastectomy.
- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the CE associated with screening women ages 50 to 74 years of age for breast cancer every 2 to 3 years would be $\$ 19,720$ / QALY (Table 3, row $u$ ).

Table 3. Summary of CE Estimate for Breast Cancer Screening
B.C. Birth Cohort of $\mathbf{4 0 , 0 0 0}$

| Row | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Screening visits with 100\% Adherence | 184,952 | $\checkmark$ |
| b | Screening visits with 88\% Adherence | 162,758 | = a * Table 2, row e |
| C | Cost per screen | \$79.35 | Ref Doc |
| d | Value of patient time (per hour) | \$29.69 | Ref Doc |
| e | Screening costs | \$12,914,856 | $=b^{*} \mathrm{c}$ |
| f | Patient time costs | \$9,664,577 | $=(\mathrm{b} * \mathrm{~d}) * 2$ |
| g | Deaths avoided | 72 | Table 2, row h |
| h | Costs avoided per death prevented | -\$47,230 | Ref Doc |
| i | Costs avoided due to deaths prevented | -\$3,394,150 | $=\mathrm{g}$ * h |
| j | Unnecessary lumpectomies / mastectomies for every death avoided | 3 | $\checkmark$ |
| k | Costs per lumpectomy / mastectomy | \$5,679 | Ref Doc |
| I | Costs associated with unnecessary lumpectomies / mastectomies | \$1,224,352 | $=\mathrm{g}^{*} \mathrm{j}^{*} \mathrm{k}$ |
| m | Unnecessary biopsies per death avoided | 26 | $\checkmark$ |
| n | Cost per unnecessary biopsy | \$386 | Ref Doc |
| 0 | Costs for unnecessary biopsies | \$721,230 | $=\mathrm{n} * \mathrm{f} * \mathrm{o}$ |
| p | Patient time and travel costs associated with unnecessary procedures | \$656,098 | $\begin{gathered} =((\mathrm{g} * \mathrm{j} * 7.5)+(\mathrm{g} * \mathrm{~m} * \\ 37.5)) * \mathrm{~d} \end{gathered}$ |
| q | Net costs undiscounted | \$21,786,962 | $=e+f+i+l+o+p$ |
| r | CPB undiscounted | 1,189 | Table 2, row o |
| S | Net costs 1.5\% discount | \$18,103,440 | Calculated |
| t | CPB 1.5\% discount | 918 | Calculated |
| u | CE (\$/QALY saved)-1.5\% discount | \$19,720 | = s/t |

$V=$ Estimates from the literature
We also modified the major assumption and recalculated the cost per QALY as follows:

- Assume the effectiveness of screening mammography in reducing deaths from breast cancer is reduced from $21 \%$ to $10 \%$ (Table 2 , row $b$ ): $\$ /$ QALY $=\$ 45,514$.
- Assume the effectiveness of screening mammography in reducing deaths from breast cancer is increased from $21 \%$ to $32 \%$ (Table 2, row $b$ ): $\$ /$ QALY $=\$ 11,659$.


## Summary

## Table 4: Breast Cancer Screening Being Offered to a

Birth Cohort of 40,000 Between the Ages of 50 to 74

## Summary

Base
Case Range
CPB (Potential QALYs Gained)
Assume No Current Service

| 1.5\% Discount Rate | 918 | 406 | 1,516 |
| :--- | :---: | :---: | :---: |
| 3\% Discount Rate | 721 | 319 | 1,191 |
| O\% Discount Rate | 1,189 | 526 | 1,963 |

Gap between B.C. Current (52\%) and 'Best in the World' (88\%)

| 1.5\% Discount Rate | 376 |  | 166 | 620 |
| :---: | :---: | :---: | :---: | :---: |
| 3\% Discount Rate | 295 |  | 131 | 487 |
| 0\% Discount Rate | 486 |  | 215 | 803 |
| CE (\$/QALY) including patient time costs |  |  |  |  |
| 1.5\% Discount Rate | $\$ 19,720$ |  | $\$ 11,659$ | $\$ 45,514$ |
| 3\% Discount Rate | $\$ 21,048$ |  | $\$ 12,444$ | $\$ 48,580$ |
| 0\% Discount Rate | $\$ 18,326$ |  | $\$ 10,835$ | $\$ 42,298$ |

CE (\$/QALY) excluding patient time costs

| 1.5\% Discount Rate | $\$ 10,378$ | $\$ 5,769$ | $\$ 25,132$ |
| :--- | :--- | :--- | :--- |
| $3 \%$ Discount Rate | $\$ 11,077$ | $\$ 6,156$ | $\$ 26,825$ |
| $0 \%$ Discount Rate | $\$ 9,645$ | $\$ 5,360$ | $\$ 23,356$ |

## Screening (Cytology-Based) for Cervical Cancer

## Canadian Task Force on Preventive Health Care Recommendations (2013)

The following recommendations refer to cytologic screening, using either conventional or liquid-based methods, whether manual or computer-assisted.
For women aged 20-24 years, we recommend not routinely screening for cervical cancer. (Weak recommendation; moderate-quality evidence)
For women aged 25-29 years, we recommend routine screening for cervical cancer every 3 years. (Weak recommendation; moderate-quality evidence)
For women aged 30-69 years, we recommend routine screening for cervical cancer every 3 years. (Strong recommendation; high-quality evidence)
For women aged 70 years and older who have undergone adequate screening (i.e., 3 successive negative Pap test results in the previous 10 years), we recommend that routine screening may end. For women aged 70 years and older who have not undergone adequate screening, we recommend continued screening until 3 negative test results have been obtained. (Weak recommendation; low-quality evidence) ${ }^{201}$

## United States Preventive Services Task Force Recommendations (2017)

The USPSTF recommends screening for cervical cancer in women age 21 to 65 years with cytology (Pap smear) every 3 years or, for women age 30 to 65 years who want to lengthen the screening interval, screening with a combination of cytology and human papillomavirus (HPV) testing every 5 years. ${ }^{202}$

## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with screening women ages 25 to 69 years of age for cervical cancer, using cytology screening, every 3 years.

In modelling CPB, we made the following assumptions:

- Based on BC life tables for 2010 to 2012, a total of 2,721 deaths would be expected in females between the ages of $25-74$ in a BC birth cohort of 40,000 (see Table 1). While routine screening occurs to age 69 , we have assumed the protective effect of that routine screening would continue to age 74 .
- Based on BC vital statistics data, there were 357 deaths in females between the ages of 25 and 44 in BC in 2012, with 8 ( $2.2 \%$ ) of these deaths due to cervical cancer (ICD-10 codes C53). There were also 1,990 deaths between the ages of 45 and 64 that year, with $20(1.0 \%)$ of these deaths due to cervical cancer. Finally, there were 3,566 deaths between the ages of 65 and 79 that year, with $10(1.0 \%)$ of these deaths due to cervical cancer. ${ }^{203}$ This suggests that 18 of the $2,721(0.7 \%)$ of the female deaths in the BC birth cohort between the ages of 25 and 74 would be due to cervical cancer (see Table 1).

[^47]| Table 1: Mortality Due to Cervical Cancer <br> Between the Ages of $\mathbf{2 5}$ and 74 <br> in a British Columbia Birth Cohort of 40,000 |  |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Age Group | Mean Survival Rate | Individuals in Birth |  |  |  | Deaths in Birth Cohort |  | Cervical Cancer |  | Life Years Lost Per |  |
|  | Males Females | Males | Females | Total | Life Years Lived | \% | \# | \% | \# | Death | Total |
| 20-24 | 0.993 |  | 19,865 |  |  |  |  |  |  |  |  |
| 25-29 | 0.992 |  | 19,833 |  | 99,163 | 0.2\% | 32 | 2.2\% | 0.7 | 57.8 | 41 |
| 30-34 | 0.990 |  | 19,795 |  | 98,975 | 0.2\% | 38 | 2.2\% | 0.8 | 52.9 | 45 |
| 35-39 | 0.987 |  | 19,741 |  | 98,706 | 0.3\% | 54 | 2.2\% | 1.2 | 48.1 | 58 |
| 40-44 | 0.983 |  | 19,662 |  | 98,311 | 0.4\% | 79 | 2.2\% | 1.8 | 43.2 | 76 |
| 45-49 | 0.977 |  | 19,546 |  | 97,730 | 0.6\% | 116 | 1.0\% | 1.2 | 38.5 | 45 |
| 50-54 | 0.969 |  | 19,375 |  | 96,873 | 0.9\% | 171 | 1.0\% | 1.7 | 33.8 | 58 |
| 55-59 | 0.956 |  | 19,118 |  | 95,591 | 1.3\% | 256 | 1.0\% | 2.6 | 29.2 | 75 |
| 60-64 | 0.936 |  | 18,726 |  | 93,630 | 2.1\% | 392 | 1.0\% | 3.9 | 24.7 | 97 |
| 65-69 | 0.906 |  | 18,113 |  | 90,567 | 3.4\% | 613 | 0.3\% | 1.6 | 20.4 | 32 |
| 70-74 | 0.857 |  | 17,144 |  | 85,720 | 5.7\% | 969 | 0.3\% | 2.5 | 16.3 | 40 |
|  |  |  |  |  |  |  | 2,721 | 0.7\% | 18.0 | 31.6 | 568 |

- Cervical cancer screening in women ages 25-69 leads to a reduction in cervical cancer mortality of $35 \%$ (RR $0.65,95 \%$ CI of 0.47 to 0.90 ). ${ }^{204}$
- Cervical cancer screening in women ages 25-69 leads to a reduction in cervical cancer incidence of $44 \%$ (RR $0.56,95 \%$ CI of 0.42 to 0.75 ). ${ }^{205}$
- Potential harms associated with cervical cancer screening include anxiety caused by false positive screening results and pain, bleeding or discharge after an unnecessary biopsy or loop electrosurgical excision and an increase in preterm births caused by excisional treatment of CIN. ${ }^{206}$
- The false positive rate associated with cytology screening ranges from $3.2 \%$ to $6.5 \%{ }^{207}$ We have used the midpoint for our base case ( $4.9 \%$ ) and the range in our sensitivity analysis. A false-positive Pap smear result is associated with a disutility of 0.046 for a period of approximately 10 months (or a one-time QALY loss of $0.038) .{ }^{208}$
- Other assumptions used in assessing the clinically preventable burden are detailed in the Reference Document.

Based on these assumptions, the CPB associated with screening women ages 25 to 69 years of age for cervical cancer every 3 years is 1,471 QALYs saved (Table 2, row $v$ ). The CPB of 1,471 represents the gap between no coverage and the 'best in the world' coverage estimated at $88 \%$. The CPB of 317 QALYs saved (see Table 2, row $w$ ) represents the gap between the current coverage of $69 \%$ and the 'best in the world' coverage estimated at $88 \%$.

[^48]Table 2. Calculation of Clinically Preventable Burden for Cervical Cancer in Average Risk Women in a BC Birth Cohort of $\mathbf{4 0 , 0 0 0}$

| Row | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
|  | Estimated Current Status |  |  |
| a | Total cervical cancer mortality in a birth cohort of 40,000 between the ages of 25 and 74 | 18.0 | Table 1 |
| b | Ratio of nonfatal cervical cancers per fatal cervical cancer | 10.1 | Ref Doc |
| C | Estimated nonfatal cervical cancers | 181.4 | $=a^{*} \mathrm{~b}$ |
| d | Effectiveness of screening in reducing mortality | 35\% | $\checkmark$ |
| e | Effectiveness of screening in reducing incidence | 44\% | $\checkmark$ |
| $f$ | Current screening rate in BC | 69\% | Ref Doc |
| g | Potential screening rate | 88\% | Ref Doc |
| h | Predicted deaths in the absence of screening | 23.7 | $=a /\left(1-f^{*} d\right)$ |
| i | Predicted nonfatal cervical cancers in absence of screening | 260.5 | $=c /\left(1-f^{*} e\right)$ |
|  | Benefits of Screening |  |  |
| j | Deaths avoided - 100\% adherence | 8.3 | $=h^{*} \mathrm{~d}$ |
| k | Deaths avoided - 88\% adherence | 7.3 | $=j^{*} \mathrm{~g}$ |
| 1 | Deaths avoided - 69\% adherence | 5.7 | $=j^{*} \mathrm{f}$ |
| m | Nonfatal cancers avoided - 100\% adherence | 114.6 | $=i^{*} \mathrm{e}$ |
| n | Nonfatal cancers avoided - 88\% adherence | 100.9 | $=m^{*} \mathrm{~g}$ |
| 0 | Nonfatal cancers avoided - 69\% adherence | 79.1 | $=m^{*} \mathrm{f}$ |
| p | LE at average age of cervical cancer death | 31.6 | Table 1 |
| q | Life years lost per nonfatal cervical cancer | 17 | Ref Doc |
| $r$ | QALYs saved with 88\% adherence to screening | 1,945 | $=\left(k^{*} \mathrm{p}\right)+\left(\mathrm{n}^{*} \mathrm{q}\right)$ |
|  | Harms Associated with Screening |  |  |
| S | False-positive screening rate | 4.9\% | $\checkmark$ |
| t | Reduced QALYs per false positive | 0.038 | $\checkmark$ |
| u | Reduced QALYs associated with false positives | -475 | $\begin{gathered} =-(\mathrm{s} * \text { Table } 3, \text { row } \\ c) * t \end{gathered}$ |
|  | Summary of Benefits and Harms |  |  |
| V | Potential QALY saved - Utilization increasing from 0\% to 88\% | 1,471 | $=r+u$ |
| w | Potential QALY saved - Utilization increasing from 69\% to 88\% | 317 | $=v^{*}(g-f) / g$ |

V = Estimates from the literature
We also modified several major assumptions and recalculated the CPB as follows:

- Assume the effectiveness of screening in reducing cervical cancer deaths is reduced from $35 \%$ to $10 \%$ and the effectiveness of reducing cervical cancer incidence is reduced from $44 \%$ to $25 \%$ (Table 2, rows $d \& e$ ): CPB $=399$.
- Assume the effectiveness of screening in reducing cervical cancer deaths is increased from $35 \%$ to $53 \%$ and the effectiveness of reducing cervical cancer incidence is increased from $44 \%$ to $58 \%$ (Table 2, rows $d \& e$ ): $\mathrm{CPB}=2,567$.
- Assume that the false-positive screening rate is reduced from $4.9 \%$ to $3.2 \%$ (Table 2, row $s$ ): $\mathrm{CPB}=1,635$.
- Assume that the false-positive screening rate is increased from $4.9 \%$ to $6.5 \%$ (Table 2 , row $s$ ): $\mathrm{CPB}=1,315$.


## Modelling Cost-Effectiveness

In this section, we will calculate the CE associated with screening women ages 25 to 69 years of age for cervical cancer every 3 years.
In estimating the CE of screening for cervical cancer, we made the following assumptions:

- We assumed a screening rate of once every 3 years starting at age 25 . There are an estimated 869,546 life years lived by women between the ages of 25 and 69 in a BC birth cohort of 40,000 , resulting in an estimated 255,067 screens (with $88 \%$ adherence) between the ages of 25 and 69 in this birth cohort. We have also assumed that $5 \%$ of screens would have a mildly abnormal Pap resulting in a rescreen. ${ }^{209}$ Total screens in this cohort are therefore estimated at 267,820 (Table 3, row $d$ ).
- Based on the BC HPV FOCAL study, the colposcopy referral rate is $3.1 \%$ (with a $95 \%$ CI of $2.8 \%$ to $3.5 \%$ ). The participation rate for these referrals is approximately $85 \%{ }^{210}$ Women are typically recalled for multiple follow-ups if something is identified on the initial colposcopy. We have assumed an average of two colposcopies per accepted referral, ${ }^{211}$ yielding a colposcopy rate of $5.3 \%$ ( 0.031 * $0.85 * 2$ ).
- In 2007, the rate of detection of CIN2/3 lesions in BC was 5.9 per 1,000 screens (Table 3, row $o$ ). ${ }^{212}$ These would typically be treated by a loop electrosurgical excision procedure (LEEP) as an ambulatory procedure in a colposcopy suite. Three Canadian studies estimated the cost per treatment for a precancerous lesion to be $\$ 965^{213}, \$ 1,032^{214}$ and $\$ 1,071^{215}$ in 2005 or 2006 CAD. We updated these estimates to 2017 CAD and then used the average for the base case estimate and the extremes in the sensitivity analysis ( $\$ 1,216$ with a range from $\$ 1,137$ to $\$ 1,295$, in 2017 CAD).
- For patient time and travel costs, we estimated two hours of patient time would be required per screening visit and 7.5 hours per colposcopy or treatment for a precancerous lesion.
- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the CE associated with screening women ages 25 to 69 years of age for cervical cancer every 3 years would be $\$ 25,542$ / QALY (Table 3, row $a f$ ).

[^49]Table 3. Summary of CE Estimate for Cervical Cancer Screening
B.C. Birth Cohort of 40,000

| Row | Variable | Base Case <br> Ages 25-69 | Data Source |
| :---: | :---: | :---: | :---: |
|  | Costs of Screening and Treatment |  |  |
| a | Life years lived between age 25 and 69 in birth cohort | 869,546 | Table 1 |
| b | Screening visits at 100\% adherence | 289,849 | = a/3 |
| c | Screening visits at 88\% adherence | 255,067 | = ${ }^{\text {* }}$ Table 2, row g |
| d | Screening visits with 5\% rescreen rate | 267,820 | = c * 1.05 |
| e | Cost per screening visit | \$70 | Ref Doc |
| f | Screening costs | \$18,747,412 | $=e^{*} \mathrm{~d}$ |
| g | Value of patient time (per hour) | \$29.69 | Ref Doc |
| h | Patient time per screening visit (in hours) | 2 | Ref Doc |
| i | Value of patient time - screening | \$15,903,162 | $=d^{*} h^{*} \mathrm{~g}$ |
| j | Rate of colposcopies per screen | 5.3\% | $\checkmark$ |
| k | Cost per colposcopy | \$251 | Ref Doc |
| I | Colposcopy costs | \$3,562,812 | $=j^{*} \mathrm{~d}^{*} \mathrm{k}$ |
| m | Patient time per colposcopy (in hours) | 7.5 | $\checkmark$ |
| n | Value of patient time - colposcopy | \$3,160,753 | $=d^{*}{ }^{*} \mathrm{~m}^{*} \mathrm{~g}$ |
| 0 | Proportion of screens resulting in treatment for CIN2 or 3 | 0.59\% | $\checkmark$ |
| p | Treatment costs per CIN2/3 | \$1,216 | Ref Doc |
| q | Treatment costs for CIN2/3 | \$1,921,449 | = ${ }^{*}{ }^{\text {o }}$ * p |
| r | Patient time per treatment for CIN2/3 (in hours) | 7.5 | $\checkmark$ |
| s | Value of patient time - treatment of CIN2/3 | \$351,857 | $=\mathrm{d}^{*} \mathrm{o}^{*} \mathrm{r}$ * |
| t | Costs of screening and treatment | \$43,647,445 | $=f+i+l+n+q+s$ |
|  | Costs Avoided |  |  |
| u | Deaths prevented | 7.3 | Table 2, row k |
| v | Costs avoided per death prevented | -\$46,603 | Ref Doc |
| w | Costs avoided due to deaths prevented | -\$339,908 | = $\mathrm{u}^{*} \mathrm{v}$ |
| x | \# of cervical cancers prevented | 100.9 | Table 2, row n |
| y | Costs avoided per cervical cancer prevented | -\$36,021 | Ref Doc |
| z | Costs avoided due to cervical cancers prevented | -\$3,633,357 | = ${ }^{*} \mathrm{y}$ |
| aa | Costs avoided | -\$3,973,265 | = w +z |
| ab | Net costs | \$39,674,180 | = $\mathrm{t}+\mathrm{aa}$ |
| ac | CPB undiscounted | 1,471 | Table 2, row v |
| ad | Net costs (1.5\% discount) | \$24,509,536 | Calculated |
| ae | CPB (1.5\% discount) | 960 | Calculated |
| af | CE (\$/QALY saved) | \$25,542 | = ad / ae |

$\checkmark$ = Estimates from the literature

We also modified several major assumptions and recalculated the cost per QALY as follows:

- Assume the effectiveness of screening in reducing cervical cancer deaths is reduced from $35 \%$ to $10 \%$ and the effectiveness of reducing cervical cancer incidence is reduced from $44 \%$ to $25 \%$ (Table 2, rows $d \& e$ ): $\mathrm{CE}=\$ 99,328$.
- Assume the effectiveness of screening in reducing cervical cancer deaths is increased from $35 \%$ to $53 \%$ and the effectiveness of reducing cervical cancer incidence is increased from $44 \%$ to $58 \%$ (Table 2, rows $d \& e$ ): $\mathrm{CE}=\$ 13,818$.
- Assume that the false-positive screening rate is reduced from $4.9 \%$ to $3.2 \%$ (Table 2, row $s$ ): $\mathrm{CE}=\$ 22,968$.
- Assume that the false-positive screening rate is increased from $4.9 \%$ to $6.5 \%$ (Table 2 , row $s$ ): $\mathrm{CE}=\$ 28,553$.
- Assume the cost per screening visit is reduced from $\$ 70$ to $\$ 33$ (Table 3, row e): CE = \$19, 162 .
- Assume the cost per screening visit is increased from $\$ 70$ to $\$ 108$ (Table 3, row e): CE $=\$ 32,094$.
- Assume the cost per colposcopy is reduced from $\$ 251$ to $\$ 176$ (Table 3, row $k$ ): CE $=\$ 24,857$.
- Assume the cost per colposcopy is increased from $\$ 251$ to $\$ 392$ (Table 3, row $k$ ): CE $=\$ 26,831$.


## Summary

| Table 4: Cervical Cancer Screening Being Offered to a Birth Cohort of 40,000 Women Between the Ages of 25 to 69 Summary |  |  |  |
| :---: | :---: | :---: | :---: |
|  | $\begin{aligned} & \text { Base } \\ & \text { Case } \end{aligned}$ | Range |  |
| CPB (Potential QALYs Gained) |  |  |  |
| Assume No Current Service |  |  |  |
| 1.5\% Discount Rate | 960 | 260 | 1,675 |
| 3\% Discount Rate | 657 | 178 | 1,147 |
| 0\% Discount Rate | 1,471 | 399 | 2,567 |
| Gap between B.C. Current (69\%) and 'Best in the World' (88\%) |  |  |  |
| 1.5\% Discount Rate | 207 | 56 | 362 |
| 3\% Discount Rate | 142 | 38 | 248 |
| 0\% Discount Rate | 318 | 86 | 554 |
| CE (\$/OALY) including patient time costs |  |  |  |
| 1.5\% Discount Rate | \$25,542 | \$13,818 | \$99,328 |
| 3\% Discount Rate | \$28,928 | \$15,524 | \$113,289 |
| 0\% Discount Rate | \$26,980 | \$14,596 | \$104,919 |
| CE ( $\$ / \mathrm{QALY}$ ) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | \$13,042 | \$6,658 | \$53,225 |
| 3\% Discount Rate | \$14,594 | \$7,314 | \$60,424 |
| 0\% Discount Rate | \$13,776 | \$7,033 | \$56,221 |

## Screening (HPV-Based) for Cervical Cancer

United States Preventive Services Task Force Recommendations (2017)
The USPSTF recommends screening for cervical cancer in women age 21 to 65 years with cytology (Pap smear) every 3 years or, for women age 30 to 65 years who want to lengthen the screening interval, screening with a combination of cytology and human papillomavirus (HPV) testing every 5 years. ${ }^{216}$

## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with incorporating HPV-based screening in females ages 30-65 in a BC birth cohort of 40,000 .

In modelling CPB, we made the following assumptions:

- Based on BC life tables for 2010 to 2012, a total of 1,719 deaths would be expected in females between the ages of $30-69$ in a BC birth cohort of 40,000 (see Table 1). While routine HPV-based screening occurs to age 65 , we have assumed the protective effect of routine screening would continue to age 69 .
- Based on BC vital statistics data, there were 357 deaths in females between the ages of 25 and 44 in BC in 2012, with 8 ( $2.2 \%$ ) of these deaths due to cervical cancer (ICD-10 codes C53). There were also 1,990 deaths between the ages of 45 and 64 that year, with $20(1.0 \%)$ of these deaths due to cervical cancer. Finally, there were 3,566 deaths between the ages of 65 and 79 that year, with $10(1.0 \%)$ of these deaths due to cervical cancer. ${ }^{217}$ This suggests that 14.8 of the $1,719(0.9 \%)$ of the female deaths in the BC birth cohort between the ages of 30 and 69 would be due to cervical cancer (see Table 1).

- HPV-based screening is associated with a $55 \%$ reduction in the incidence of cervical cancers (RR of $0.45,95 \% \mathrm{CI}$ of 0.25 to 0.81 ) in females ages $30-64$, when

[^50]compared to cytology-based screening. ${ }^{218}$ The effectiveness of HPV-based screening is observed primarily in the reduction in adenocarcinomas. We assumed that the effectiveness of HPV-based screening in reducing mortality from cervical cancers would be the same as the observed effectiveness in reducing the incidence of cervical cancers.

- The cumulative incidence of cervical cancer is lower at 5.5 years after a negative HPV test than 3.5 years after a negative cytology test, indicating that 5 year screening intervals with HPV testing are safer than 3 year screening intervals with cytology testing. ${ }^{219}$

In estimating the effect of the additional CPB associated with incorporating HPV-based we first re-ran the model for cytology-based screening above but modified the age range to 30-69 (from 25-74). The result is a modest reduction in QALYs saved, from 1,471 (based on ages $25-74$ ) to 1,188 (based on ages $30-69$ ) (see Table 2).

Table 2. Calculation of Clinically Preventable Burden for Cervical Cancer in
Average Risk Women in a BC Birth Cohort of 40,000

| Row | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
|  | Estimated Current Status |  |  |
| a | Total cervical cancer mortality in a birth cohort of 40,000 between the ages of 30 and 69 | 14.8 | Table 1 |
| b | Ratio of nonfatal cervical cancers per fatal cervical cancer | 10.1 | Ref Doc |
| C | Estimated nonfatal cervical cancers | 149.3 | = a*b |
| d | Effectiveness of screening in reducing mortality | 35\% | $\checkmark$ |
| e | Effectiveness of screening in reducing incidence | 44\% | $\checkmark$ |
| $f$ | Current screening rate in BC | 69\% | Ref Doc |
| g | Potential screening rate | 88\% | Ref Doc |
| h | Predicted deaths in the absence of screening | 19.5 | $=a /(1-f * d)$ |
| i | Predicted nonfatal cervical cancers in absence of screening | 214.4 | $=c /(1-f * e)$ |
|  | Benefits of Screening |  |  |
| j | Deaths avoided - 100\% adherence | 6.8 | $=\mathrm{h} * \mathrm{~d}$ |
| k | Deaths avoided - 88\% adherence | 6.0 | $=j^{*} \mathrm{~g}$ |
| 1 | Deaths avoided - 69\% adherence | 4.7 | $=j^{*} \mathrm{f}$ |
| m | Nonfatal cancers avoided - 100\% adherence | 94.3 | $=i^{*} \mathrm{e}$ |
| n | Nonfatal cancers avoided - 88\% adherence | 83.0 | $=\mathrm{m}^{*} \mathrm{~g}$ |
| 0 | Nonfatal cancers avoided - 69\% adherence | 65.1 | $=m$ * |
| p | LE at average age of cervical cancer death | 32.9 | Table 1 |
| q | Life years lost per nonfatal cervical cancer | 17 | Ref Doc |
| $r$ | QALYs saved with 88\% adherence to screening | 1,609 | $=(\mathrm{k} * \mathrm{p})+\left(\mathrm{n}^{*} \mathrm{q}\right)$ |
|  | Harms Associated with Screening |  |  |
| S | False-positive screening rate | 4.9\% | $\checkmark$ |
| t | Reduced QALYs per false positive | 0.038 | $\checkmark$ |
| u | Reduced QALYs associated with false positives | -421 | $=-(\mathrm{s} * \text { Table 4, row }$ $\text { c) } * \mathrm{t}$ |
|  | Summary of Benefits and Harms |  |  |
| V | Potential QALY saved - Utilization increasing from 0\% to 88\% | 1,188 | $=r+u$ |
| w | Potential QALY saved - Utilization increasing from 69\% to 88\% | 257 | $=v^{*}(g-f) / g$ |

V = Estimates from the literature

[^51]We then adjusted the assumptions in this table to reflect HPV-based screening. This meant that the effectiveness of HPV-based screening improved by $55 \%$ compared to cytology-based screening (Table 3, row $j$ ) while the false-positive screening rate increased from $4.9 \%$ to $7.28 \%$ (Table 3, row $p$ ). ${ }^{220}$

The result is a gain of 975 QALYs saved, from 1,188 (see Table 2, row $v$ ) to 2,163 (Table 3, row $s$ ) associated with incorporating HPV-based screening in females ages $30-65$ in a BC birth cohort of 40,000 .

Table 3. Calculation of CPB for HPV-Based Cervical Cancer Screening in Average
Risk Women in a BC Birth Cohort of $\mathbf{4 0 , 0 0 0}$

| Row | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
|  | Estimated Current Status - Cytology-based Screening |  |  |
| a | Total cervical cancer mortality in a birth cohort of 40,000 between the ages of $\mathbf{3 0}$ and 69 | 14.8 | Table 1 |
| b | Ratio of nonfatal cervical cancers per fatal cervical cancer | 10.1 | Ref Doc |
| C | Estimated nonfatal cervical cancers | 149.3 | $=\mathrm{a}$ * b |
| d | Effectiveness of screening in reducing mortality | 35\% | Table 2, row d |
| e | Effectiveness of screening in reducing incidence | 44\% | Table 2, row e |
| f | Current screening rate in BC | 69\% | Ref Doc |
| g | Potential screening rate | 88\% | Ref Doc |
| h | Predicted deaths in the absence of screening | 19.5 | Table 2, row h |
| i | Predicted nonfatal cervical cancers in absence of screening | 214.4 | Table 2, row i |
|  | Benefits of HPV-based Screening |  |  |
| j | Rate ratio comparing HPV- to cytology-based screening | 55\% | $\checkmark$ |
| k | Deaths avoided - 88\% adherence | 9.3 | $\begin{aligned} & \text { = Table } 2 \text {, row } \mathrm{k}+ \\ & \text { (Table 2, row k }{ }^{\mathrm{j}} \text { ) } \end{aligned}$ |
| 1 | Nonfatal cancers avoided - 88\% adherence | 128.7 | = Table 2 , row $n+$ <br> (Table 2, row $n * j$ ) |
| m | LE at average age of cervical cancer death | 32.9 | Table 1 |
| n | Life years lost per nonfatal cervical cancer | 17 | Ref Doc |
| 0 | QALYs saved with 88\% adherence to screening | 2,494 | $=\left(k^{*} \mathrm{I}\right)+\left(\mathrm{I}^{*} \mathrm{n}\right)$ |
|  | Harms Associated with Screening |  |  |
| p | False-positive screening rate | 7.28\% | $\checkmark$ |
| q | Reduced QALYs per false positive | 0.038 | $\checkmark$ |
| $r$ | Reduced QALYs associated with false positives | -331 | $=-(p *$ Table 5, row <br> e) $* q$ |
|  | Summary of Benefits and Harms |  |  |
| S | Potential QALY saved - Utilization increasing from 0\% to 88\% | 2,163 | $=0+r$ |
| $V=$ Estimates from the literature |  |  |  |

We also modified a major assumption and recalculated the CE as follows:

- Assume that the effectiveness of HPV-based screening compared to cytology-based screening is reduced from $55 \%$ to $19 \%$ (Table 3, rows $j$ ): CPB $=395$.
- Assume that the effectiveness of HPV-based screening compared to cytology-based screening is reduced from $55 \%$ to $75 \%$ (Table 3, rows $j$ ): $\mathrm{CPB}=1,296$.

[^52]
## Modelling Cost-effectiveness

Note that in modelling cost-effectiveness we are trying to tease out the additional benefits and costs associated with HPV-based screening to generate a cost/QALY associated with moving from cytology-based screening every three years in women ages 30-69 to HPV-based screening every five years in women ages 30-65 in a BC birth cohort of 40,000.
In estimating the effect on CE associated with incorporating HPV-based screening, we first re-ran the model for cytology-based screening used in the previous section but modified the age range to $30-69$ (from 25-74). The result is a reduction in net costs from $\$ 39,674,180$ (based on ages 25-74) to $\$ 35,399,781$ (based on ages 30-69) (see Table 4, row $a b$ ).

## Table 4. Summary of Net Costs for Cervical Cancer Screening

## B.C. Birth Cohort of 40,000

| Row | Variable | Base Case <br> Ages 30-69 | Data Source |
| :---: | :---: | :---: | :---: |
|  | Costs of Screening and Treatment |  |  |
| a | Life years lived between age 30 and 69 in birth cohort | 770,383 | Table 1 |
| b | Screening visits at 100\% adherence | 256,794 | = a/3 |
| c | Screening visits at 88\% adherence | 225,979 | = b * Table 2, row g |
| d | Screening visits with 5\% rescreen rate | 237,278 | = c * 1.05 |
| e | Cost per screening visit | \$70 | Ref Doc |
| f | Screening costs | \$16,609,457 | $=e$ * d |
| g | Value of patient time (per hour) | \$29.69 | Ref Doc |
| h | Patient time per screening visit (in hours) | 2 | Ref Doc |
| i | Value of patient time - screening | \$14,089,566 | $=d^{*} h^{*} \mathrm{~g}$ |
| J | Rate of colposcopies per screen | 5.3\% | $\checkmark$ |
| k | Cost per colposcopy | \$251 | Ref Doc |
| I | Colposcopy costs | \$3,156,509 | $=j^{*} \mathrm{~d}^{*} \mathrm{k}$ |
| m | Patient time per colposcopy (in hours) | 7.5 | $\checkmark$ |
| n | Value of patient time - colposcopy | \$2,800,301 | $=d^{*}{ }^{*} \mathrm{~m}^{*} \mathrm{~g}$ |
| 0 | Proportion of screens resulting in treatment for CIN2 or 3 | 0.59\% | $\checkmark$ |
| p | Treatment costs per CIN2/3 | \$1,216 | Ref Doc |
| q | Treatment costs for CIN2/3 | \$1,702,327 | = ${ }^{*} \mathrm{o}^{*} \mathrm{p}$ |
| r | Patient time per treatment for CIN2/3 (in hours) | 7.5 | $\checkmark$ |
| s | Value of patient time - treatment of CIN2/3 | \$311,732 | $=d^{*} o^{*}{ }^{*} \mathrm{~g}$ |
| t | Costs of screening and treatment | \$38,669,892 | $=f+i+l+n+q+s$ |
|  | Costs Avoided |  |  |
| u | Deaths prevented | 6.0 | Table 2, row k |
| v | Costs avoided per death prevented | -\$46,603 | Ref Doc |
| w | Costs avoided due to deaths prevented | -\$279,754 | = ${ }^{*}$ v |
| x | \# of cervical cancers prevented | 83.0 | Table 2, row n |
| y | Costs avoided per cervical cancer prevented | -\$36,021 | Ref Doc |
| z | Costs avoided due to cervical cancers prevented | -\$2,990,356 | = ${ }^{*} \mathrm{y}$ |
| aa | Costs avoided | -\$3,270,110 | = w +z |
| ab | Net costs | \$35,399,781 | = $\mathrm{t}+\mathrm{aa}$ |

$V=$ Estimates from the literature
We then estimated the net costs of incorporating HPV-based screening in females ages 30-65 in a BC birth cohort of 40,000 . In doing so, we made the following assumptions:

- Number of HPV-based screens - We assumed a screening rate of once every five years starting at age 30. Based on the initial results of the HPV FOCAL trial, $91.9 \%$ of tests are negative and the woman is recalled at 5 years. The $8.1 \%$ of women with
hr-HPV positive tests (Table 5, row $f$ ) are reflexed to cytology (Table 5, row $g$ ). Cytology results are negative for $64 \%$ of these women (Table 5, row $h$ ). Women with positive results are referred to colposcopy. Women who are hr-HPV positive but cytology negative are retested with HPV and cytology after 6-12 months. $43 \%$ of these women are both HPV and cytology negative and move into routine HPV-based screening at 5-year intervals. The $57 \%$ of women who are HPV and/or cytology positive are referred to colposcopy. ${ }^{221}$ This approach results in 125,850 HPV-based screens (Table 5, row $l$ ) and 15,894 cytology-based screens (Table 5 , row $m$ ) in females between the ages of 30 and 65 in a BC birth cohort of 40,000 .
- Based on the BC HPV FOCAL study, the colposcopy referral rate associated with cytology-based screening is $3.1 \%$ (with a $95 \%$ CI of $2.8 \%$ to $3.5 \%$ ) while the colposcopy referral rate associated with HPV-based screening is $5.9 \%$ (with a $95 \%$ CI of $5.5 \%$ to $6.3 \%$ ). ${ }^{222}$ The participation rate for these referrals is approximately $85 \%{ }^{223}$ Women are typically recalled for multiple follow-ups if something is identified on the initial colposcopy. We have assumed an average of two colposcopies per accepted referral, ${ }^{224}$ yielding a HPV-based colposcopy rate of $10.0 \%$ ( $0.059 * 0.85 * 2$ ).
- In 2007, the rate of detection of CIN2/3 lesions in BC was 5.9 per 1,000 screens. ${ }^{225}$ Based on the BC HPV FOCAL study, the detection rate of CIN2/3 lesions is increased by $50 \%$ with HPV-based screening, to 8.85 per 1,000 screens. ${ }^{226}$ These lesions would typically be treated by a loop electrosurgical excision procedure (LEEP) as an ambulatory procedure in a colposcopy suite.
- For patient time and travel costs, we estimated two hours of patient time would be required per screening visit and 7.5 hours per colposcopy or treatment for a precancerous lesion.
- Other costs and assumptions used in assessing net costs are detailed in the Reference Document.

Based on these assumptions, the estimated net costs of incorporating HPV-based screening in females ages $30-65$ in a BC birth cohort of 40,000 is $\$ 22,776,189$ (see Table 5, row $a k$ ). This is $\$ 12,623,593$ less than the estimated net costs associated with the current cytology-based screening (ref. Table 4, row $a b$ ) for females ages 30-69 in a BC birth cohort of 40,000.

[^53]Table 5. Summary of Net Cost for HPV-Based Cervical Cancer Screening

| Row Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
|  | Costs of Screening and Treatment |  |  |
| a | Life years lived between age 30 and 65 in birth cohort | 679,816 | Table 1 |
| b | Annual frequency of HPV-based screening | 20\% | $\checkmark$ |
| c | Number of HPV-based screens - 100\% adherence | 135,963 | $=\mathrm{a}$ * b |
| d | Adherence with HPV-based screening | 88\% | Table 3, row g |
| e | Number of HPV-based screens - 88\% adherence | 119,648 | $=c^{*} \mathrm{~d}$ |
| f | Proportion of screens hrHPV-positive | 8.1\% | $\checkmark$ |
| g | Number of reflex cytology screens | 9,691 | $=e^{*} \mathrm{f}$ |
| h | Proportion of reflex cytology screens negative | 64\% | $\checkmark$ |
| i | Number of reflex cytology screens negative | 6,203 | $=\mathrm{g}^{*} \mathrm{~h}$ |
| j | Number of follow-up cytology screens | 6,203 | = i |
| k | Number of follow-up HPV screens | 6,203 | = i |
| I | HPV-based screening - number of HPV-based screens | 125,850 | $=\mathrm{e}+\mathrm{k}$ |
| m | HPV-based screening - number of cytology-based screens | 15,894 | $=\mathrm{g}+\mathrm{j}$ |
| n | Cost per HPV-based screen | \$96 | Ref Doc |
| 0 | Cost for HPV-based screening | \$12,081,614 | = ${ }^{*} \mathrm{n}$ |
| p | Value of patient time (per hour) | \$29.69 | Ref Doc |
| q | Patient time per screening visit (in hours) | 2 | $\checkmark$ |
| $r$ | Value of patient time - screening | \$8,416,767 | $=(\mathrm{l}+\mathrm{m}) * \mathrm{q} * \mathrm{p}$ |
| s | Rate of colposcopies per screen | 10.0\% | $\checkmark$ |
| t | Cost per colposcopy | \$251 | Ref Doc |
| u | Colposcopy costs | \$3,158,839 | $=1 * s * t$ |
| v | Patient time per colposcopy (in hours) | 7.5 | $\checkmark$ |
| w | Value of patient time - colposcopy | \$2,664,253 | =e*s*v*p |
| x | Proportion of screens resulting in treatment for CIN2 or 3 | 0.885\% | $\checkmark$ |
| y | Treatment costs per CIN2/3 | \$1,216 | Ref Doc |
| z | Treatment costs for CIN2/3 | \$1,287,600 | = ${ }^{*} \mathrm{x}^{*} \mathrm{y}$ |
| aa | Patient time per treatment for CIN2/3 (in hours) | 7.5 | $\checkmark$ |
| ab | Value of patient time - treatment of CIN2/3 | \$235,786 | $=e^{*}{ }^{*}$ aa *p |
| ac | Costs of screening and treatment | \$27,844,859 | $=0+r+u+w+z+a b$ |
|  | Costs Avoided |  |  |
| ad | Deaths prevented | 9.3 | Table 3, row k |
| ae | Costs avoided per death prevented | -\$46,603 | Ref Doc |
| af | Costs avoided due to deaths prevented | -\$433,618 | = ad * ae |
| ag | \# of cervical cancers prevented | 128.7 | Table 3, row I |
| ah | Costs avoided per cervical cancer prevented | -\$36,021 | Ref Doc |
| ai | Costs avoided due to cervical cancers prevented | -\$4,635,053 | = ag * ah |
| aj | Costs avoided | -\$5,068,671 | = af + ai |
| ak | Net costs | \$22,776,189 | = af +aj |

$V=$ Estimates from the literature

After discounting costs and QALYs by $1.5 \%$, the cost per QALY associated with cytologybased cervical cancer screening is $\$ 33,340$ (see Table 6 , row $i$ ) compared to the cost per QALY associated with HPV-based cervical cancer screening of $\$ 11,784$ (see Table 6, row $l$ ). Implementing HPV-based cervical cancer screening in females ages 30-65 in a BC birth cohort of 40,000 is estimated to cost $\$ 21,556$ less per QALY than the current cytology-based screening in this cohort (see Table 6 , row $m$ ).

Table 6. Summary of CE Estimate for HPV-Based Cervical Cancer Screening B.C. Birth Cohort of $\mathbf{4 0 , 0 0 0}$

| Row | Variable | Base Case Ages 30-65 | Data Source |
| :---: | :---: | :---: | :---: |
|  | Undiscounted Cost / QALY |  |  |
| a | Net costs for cytology-based cervical cancer screening | \$35,399,781 | Table 4, row ab |
| b | QALYs gained with cytology-based cervical cancer screening | 1,188 | Table 2, row v |
| c | Undiscounted cost / QALY | \$29,796 | $=a / c$ |
| d | Net costs for HPV-based cervical cancer screening | \$22,776,189 | Table 5, row ak |
| e | QALYs gained with HPV-based cervical cancer screening | 2,163 | Table 3, row s |
| f | Undiscounted cost / QALY | \$10,531 | = d/e |
|  | Discounted Cost / QALY - 1.5\% |  |  |
| g | Net costs for cytology-based cervical cancer screening | \$26,636,256 | Calculated |
| h | QALYs gained with cytology-based cervical cancer screening | 799 | Calculated |
| i | Discounted cost / QALY | \$33,340 | = g/h |
| j | Net costs for HPV-based cervical cancer screening | \$17,137,744 | Calculated |
| k | QALYs gained with HPV-based cervical cancer screening | 1,454 | Calculated |
| I | Discounted cost / QALY | \$11,784 | = j/k |
| m | Cost / QALY saved with incorporating HPV-based cervical cancer screening | -\$21,556 | $=\mathrm{l}-\mathrm{i}$ |

We also modified a major assumption and recalculated the CE as follows:

- Assume that the effectiveness of HPV-based screening compared to cytology-based screening is reduced from $55 \%$ to $19 \%$ (Table 3, rows $j$ ): $\mathrm{CE}=-\$ 16,414$.
- Assume that the effectiveness of HPV-based screening compared to cytology-based screening is reduced from $55 \%$ to $75 \%$ (Table 3, rows $j$ ): $\mathrm{CE}=-\$ 23,377$.


## Summary

Table 7: HPV-based Cervical Cancer Screening Being Offered to a Birth Cohort of 40,000 Between the Ages of 30 and 65

| Summary |  |  |  |
| :---: | :---: | :---: | :---: |
| Base |  |  |  |
| CPB (Potential QALYs gained in moving from cytology- to HPB-based screening) Gap between B.C. Current (0\%) and 'Best in the World' (88\%) |  |  |  |
|  |  |  |  |
| 1.5\% Discount Rate | 655 | 266 | 872 |
| 3\% Discount Rate | 459 | 186 | 611 |
| 0\% Discount Rate | 975 | 395 | 1,296 |
| CE (\$/QALY) including patient time costs |  |  |  |
| 1.5\% Discount Rate | -\$21,556 | -\$16,414 | -\$23,377 |
| 3\% Discount Rate | -\$23,624 | -\$17,989 | -\$25,620 |
| 0\% Discount Rate | -\$19,264 | -\$14,669 | -\$20,892 |
| CE (\$/QALY) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | -\$11,210 | -\$8,210 | -\$12,273 |
| 3\% Discount Rate | -\$12,286 | -\$8,998 | -\$13,450 |
| 0\% Discount Rate | -\$10,019 | -\$7,337 | -\$10,968 |

## Screening for Colorectal Cancer

## Canadian Task Force on Preventive Health Care Recommendations (2016)

We recommend screening adults aged 50 to 59 years for colorectal cancer with FOBT (gFOBT or FIT) every two years or flexible sigmoidoscopy every 10 years. (Weak recommendation; moderate-quality evidence)
We recommend screening adults aged 60 to 74 years for colorectal cancer with FOBT (gFOBT or FIT) every two years or flexible sigmoidoscopy every 10 years. (Strong recommendation; moderate-quality evidence) ${ }^{227}$

## United States Preventive Services Task Force Recommendations (2016)

The USPSTF recommends screening for colorectal cancer starting at age 50 years and continuing until age 75 years. (A recommendation $)^{228}$

## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with screening adults aged 50 to 74 years of age for colorectal cancer with a fecal occult blood test (with either a guaiac fecal occult blood test [gFOBT] or a fecal immunochemical test [FIT]) every two years or flexible sigmoidoscopy / colonoscopy every 10 years.

In modelling CPB, we made the following assumptions:

- Based on BC life tables for 2010 to 2012, a total of 9,340 deaths would be expected between the ages of 50-79 in a BC birth cohort of 40,000 (see Table 1). Routine screening occurs to age 74 , and we have assumed the protective effect of routine screening continues to age 79 .
- Based on BC vital statistics data, there were 5,117 deaths between the ages of 45 and 64 in BC in 2012, with $257(5.0 \%)$ of these deaths due to CRC (ICD-10 codes C1820). There were also 8,674 deaths between the ages of 65 and 79 that year, with 379 $(4.4 \%)$ of these deaths due to CRC. ${ }^{229}$ This suggests that 423 of the $9,340(4.5 \%)$ of the deaths in the BC birth cohort between the ages of 50 and 79 would be due to CRC (see Table 1).

[^54]| Table 1: Mortality Due to Colorectal Cancer Between the Ages of 50 and 79 in a British Columbia Birth Cohort of 40,000 |  |  |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Age <br> Group | Mean Survival Rate |  | Individuals in Birth Cohort |  |  | Life Years Lived | Deaths in Birth Cohort |  | Deaths due to Colorectal Cancer |  | Life Years Lost Per |  |
| 45-49 | 0.963 | 0.977 | 19,263 | 19,546 | 38,809 |  |  |  |  |  |  |  |
| 50-54 | 0.950 | 0.969 | 19,003 | 19,375 | 38,378 | 191,890 | 1.1\% | 431 | 5.0\% | 22 | 32.2 | 694 |
| 55-59 | 0.931 | 0.956 | 18,619 | 19,118 | 37,737 | 188,686 | 1.7\% | 641 | 5.0\% | 32 | 27.7 | 888 |
| 60-64 | 0.902 | 0.936 | 18,041 | 18,726 | 36,767 | 183,834 | 2.6\% | 970 | 5.0\% | 49 | 23.4 | 1,135 |
| 65-69 | 0.858 | 0.906 | 17,164 | 18,113 | 35,277 | 176,387 | 4.2\% | 1,489 | 4.4\% | 66 | 19.2 | 1,258 |
| 70-74 | 0.792 | 0.857 | 15,837 | 17,144 | 32,981 | 164,903 | 7.0\% | 2,297 | 4.4\% | 101 | 15.3 | 1,546 |
| 75-79 | 0.693 | 0.780 | 13,861 | 15,608 | 29,469 | 147,346 | 11.9\% | 3,511 | 4.4\% | 155 | 11.8 | 1,823 |
|  |  |  |  |  |  |  |  | 9,340 | 4.5\% | 423 | 17.4 | 7,344 |

- The overall screening delivery rate for BC in 2012 is $49.6 \%$, with an equal mix of fecal immunochemical testing (FIT) at $31.3 \%$ of the population ages 50-74 and sigmoidoscopy/colonoscopy at $31.1 \%$. Across Canada, approximately $40 \%$ of those who have a FIT also have a sigmoidoscopy or colonoscopy. ${ }^{230}$
- Screening with gFOBT reduces the risk of mortality from CRC by $18 \%$ (RR of 0.82 with a $95 \%$ CI of 0.73 to 0.92 ) and the incidence of late stage CRC by $8 \%$ (RR of 0.92 with a $95 \%$ CI of 0.85 to 0.99 ). Screening with flexible sigmoidoscopy reduces the risk of mortality from CRC by $26 \%$ (RR of 0.74 with a $95 \%$ CI of 0.67 to 0.82 ) and the incidence of late stage CRC by $27 \%$ (RR of 0.73 with a $95 \% \mathrm{CI}$ of 0.66 to 0.82 ). ${ }^{231}$
- Approximately $25 \%$ of CRCs are diagnosed as late stage cancers (stage III or IV). The life expectancy for an individual diagnosed with a late-stage CRC is approximately 30 months ( 2.5 years). ${ }^{232}$ The average individual with CRC survives for 6.6 years (see Reference Document) so early detection is estimated to save 4.1 years ( 6.6 minus 2.5).
- Harms associated with screening for CRC include a false positive rate of $1.22 \%$ for gFOBT and between $5.55 \%$ and $12.89 \%$ for FIT. Harms following flexible sigmoidoscopy are rare but include intestinal perforation ( $0.001 \%$ of patients), minor bleeding ( $0.05 \%$ of patients), major bleeding ( $0.009 \%$ of patients) and death $(0.015 \%$ of patients). ${ }^{233}$
- Other assumptions used in assessing the clinically preventable burden are detailed in the Reference Document.

Based on these assumptions, the CPB associated with screening adults aged 50 to 74 years of age for CRC with FOBT every two years or flexible sigmoidoscopy / colonoscopy every 10 years is 1,734 QALYs saved (Table 2, row $a h$ ). The CPB of 1,734 QALYs saved represents the gap between no coverage and the 'best in the world' coverage estimated at $76 \%$. The CPB of 593 QALYs saved (see Table 2, row ai) represents the gap between the current coverage of $50 \%$ and the 'best in the world' coverage estimated at $76 \%$.

[^55]Table 2. Calculation of Clinically Preventable Burden (CPB) Estimate for Colorectal
Cancer Screening in a BC Birth Cohort of 40,000

| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
|  | Estimated Current Status |  |  |
| a | Colorectal cancer deaths ages 55-79 | 423 | Table 1 |
| b | Predicted CRC deaths ages 55-79 in the absence of screening | 475 | =a / (1-w ${ }^{*}$ ) |
| c | Weighted life expectancy at death | 17.4 | Table 1 |
| d | Life years lost due to CRC deaths | 8,252 | $=b^{*} \mathrm{c}$ |
| e | Ratio of nonfatal CRC per fatal CRC | 4.32 | Ref Doc |
| $f$ | Nonfatal CRCs | 1,828 | = ${ }^{*}$ e |
| g | Average age of CRC incidence | 70.4 | Ref Doc |
| h | Life years lost per CRC case | 9.9 | Ref Doc |
| i | Life years lost due to CRC incidence | 18,099 | = f * h |
| j | Years lived with CRC per case | 6.6 | Ref Doc |
| k | Total years lived with CRC | 12,066 | = * $^{\text {j }}$ |
| 1 | QoL disutility for CRC survivors | 0.061 | Ref Doc |
| m | QALYs lost for cancer survivors | 740 | = ${ }^{*}$ \\| |
| n | Total QALYs lost due to CRC | 27,091 | $=d+i+m$ |
|  | Benefits if 100\% Adherence with Screening |  |  |
| 0 | Effectiveness in reducing the risk of mortality from CRC - gFOBT | 18.0\% | V |
| p | Effectiveness in reducing the risk of mortality from CRC - flexible sigmoidoscopy | 26.0\% | V |
| q | Weighted effectiveness | 22.0\% | $=(0 * u)+\left(p^{*} v\right)$ |
| $r$ | Effectiveness in reducing the incidence of late-stage CRC - gFOBT | 8.0\% | $\checkmark$ |
| S | Effectiveness in reducing the incidence of late-stage CRC - flexible sigmoidoscopy | 27.0\% | V |
| t | Proportion of CRCs detected as late-stage (III or IV) | 25.0\% | $\checkmark$ |
| u | Proportion of screening via gFOBT / FIT | 50.0\% | $\checkmark$ |
| V | Proportion of screening via flexible sigmoidoscopy / colonoscopy | 50.0\% | $\checkmark$ |
| W | Weighted proportion screened | 50.0\% | $=(u+v) / 2$ |
| X | CRC deaths avoided via gFOBT / FIT | 42.8 | $=(b * u) * o$ |
| y | CRC deaths avoided via flexible sigmoidoscopy / colonoscopy | 61.8 | $=\left(b^{*} v\right)^{*} p$ |
| z | Proportion of CRC deaths avoided via screening | 22.0\% | $=(x+y) / b$ |
| aa | Life years lost due to CRC deaths avoided | 1,815 | $=d^{*} z$ |
| ab | Late stage CRCs avoided via gFOBT / FIT | 59.4 | $=(f * t) * u * p$ |
| ac | Late stage CRCs avoided via flexible sigmoidoscopy / colonoscopy | 61.7 | $=(f * t) * v * s$ |
| ad | Life years saved per CRC due to earlier detection of CRC | 4.1 | $\checkmark$ |
| ae | Life years saved due to earlier detection of CRC | 497 | $=(a b+a c) * a d$ |
| af | QALYs lost for cancer survivors | -30 | $=-a e^{*}$ |
| ag | Potential QALYs saved with 100\% Utilization of Screening | 2,282 | $=\mathrm{aa}+\mathrm{ae}+\mathrm{af}$ |
| ah | Potential QALYs saved (CPB) - Utilization increasing from 0\% to 76\% | 1,734 | = ag * 0.76 |
| ai | Potential QALYs saved (CPB) - Utilization increasing from 50\% to 76\% | 593 | $=\mathrm{ah}-(\mathrm{ag} * 0.50)$ |

$V=$ Estimates from the literature
We modified several major assumptions and recalculated the CPB as follows:

- Assume the QoL disutility for CRC survivors is reduced from 0.061 to 0.039 (Table 2 , row $l$ ): $\mathrm{CPB}=1,742$.
- Assume the QoL disutility for CRC survivors is increased from 0.061 to 0.090 (Table 2, row $l$ ): $\mathrm{CPB}=1,723$.
- Assume the effectiveness of gFOBT in reducing the risk of mortality from CRC is reduced from $18 \%$ to $8 \%$ (Table 2, row $o$ ), the effectiveness of flexible sigmoidoscopy / colonoscopy in reducing the risk of mortality from CRC is reduced from $26 \%$ to $18 \%$ (Table 2, row $p$ ), the effectiveness of gFOBT in reducing the
incidence of late-stage CRC is reduced from $8 \%$ to $1 \%$ (Table 2, row $r$ ) and the effectiveness of flexible sigmoidoscopy / colonoscopy in reducing the incidence of late-stage CRC is reduced from $27 \%$ to $18 \%$ (Table 2, row $s$ ): CPB $=1,017$.
- Assume the effectiveness of gFOBT in reducing the risk of mortality from CRC is increased from $18 \%$ to $27 \%$ (Table 2, row $o$ ), the effectiveness of flexible sigmoidoscopy / colonoscopy in reducing the risk of mortality from CRC is increased from $26 \%$ to $33 \%$ (Table 2, row $p$ ), the effectiveness of gFOBT in reducing the incidence of late-stage CRC is increased from $8 \%$ to $15 \%$ (Table 2, row $r$ ) and the effectiveness of flexible sigmoidoscopy / colonoscopy in reducing the incidence of late-stage CRC is increased $27 \%$ to $34 \%$ (Table 2, row $s$ ): $\mathrm{CPB}=2,418$.


## Modelling Cost-Effectiveness

In this section, we will calculate the CE associated with screening adults aged 50 to 74 years of age for colorectal cancer with FOBT (gFOBT or FIT) every two years or flexible sigmoidoscopy every 10 years.

In modelling the estimated CE of colorectal cancer screening, we made the following assumptions:

- Costs of screening - We assumed a biennial FIT test would cost $\$ 14.74$. This is based on a $\$ 5.36$ fee for sample collection (MSP Fee 92007 Fecal immunochemical test - For sample collection only) and a $\$ 9.38$ fee for analysis (MSP Fee 92006 Fecal immunochemical test - For analysis only). A colonoscopy every 10 years would cost $\$ 593.40$. This is based on the assumption that $16 \%$ of colonoscopies would involve the removal of polyps. Colonoscopy with polyp removal could cost $\$ 850.39$ ( $\$ 250$ for facility fee, $\$ 347.55$ for physician fee [MSP fee \#S33374], $\$ 65.48$ for anesthesia fee [MSP fee \#01172] and $\$ 187.36$ for laboratory fees). Colonoscopy without polyp removal could cost \$544.45 (\$250 for facility fee, \$228.97 [MSP fee \#S10731] for physician fee and $\$ 65.48$ for anesthesia fee).
- Patient time and travel costs - For patient time and travel costs, we assumed that two hours of patient time would be required per FIT screening visit and that 7.5 hours of patient time would be required for a colonoscopy.
- Costs of follow-up colonoscopies - An average of $9.8 \%$ of FIT tests are positive, ranging from $5.3 \%$ to $14.2 \% .{ }^{234}$ Each positive FIT test would be followed by a colonoscopy. Approximately $40 \%$ of these colonoscopies would be positive for polyps. Individuals in whom a colonoscopy is positive for polyps would require a further follow-up colonoscopy. ${ }^{235}$
- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the estimated cost per QALY would be $\$ 47,265$ (see Table 3, row $a h$ ).

[^56]Table 3. Summary of Cost Effectiveness (CE) Estimate for Colorectal Cancer Screening in BC

| Row Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Life years lived between age 50-74 in the birth cohort | 905,700 | Table 1 |
| b | Estimated total screens with $76 \%$ screening adherence | 260,570 | = $\mathrm{e}+\mathrm{f}$ |
| c | Proportion receiving a biennial FIT screen | 48.0\% | $\checkmark$ |
| d | Proportion receiving a colonoscopy every 10 years | 47.7\% | , |
| e | Number receiving a FIT screen | 217,368 | $=\left(a^{*} \mathrm{c}\right) / 2$ |
| f | Number receiving a colonoscopy screen | 43,202 | $=(\mathrm{a}$ * f$) / 10$ |
| g | Cost per screen - FIT | \$14.74 | V |
| h | Cost per screen - Colonoscopy (no polyps - 84\%) | \$544.45 | V |
| i | Cost per screen - Colonoscopy (polyps - 16\%) | \$850.39 | $\checkmark$ |
| j | Weighted cost per screen - Colonoscopy | \$593.40 | $=(\mathrm{h} * 0.84)+(\mathrm{i} * 0.16)$ |
| k | Cost of screening | \$28,840,023 | $=\left(\mathrm{e}^{*} \mathrm{~g}\right)+\left(\mathrm{f}^{*} \mathrm{j}\right)$ |
| I | Cost of 10-minute office visit | \$34.85 | Ref Doc |
| m | Value of patient time (per hour) | \$29.69 | Ref Doc |
| 0 | Proportion of office visit for screening | 50.0\% | Ref Doc |
| p | Value of patient time | \$22,527,293 | $\begin{gathered} =((\mathrm{e} * 2)+(\mathrm{f} * 7.5)) \\ \mathrm{m} \end{gathered}$ |
| q | Total cost of office visits | \$4,540,430 | = ${ }^{*} 1^{*}$ o |
| $r$ | Proportion of FIT tests positive | 9.8\% | $\checkmark$ |
| s | \% of Follow-up colonoscopies with polyps | 40.0\% | $\checkmark$ |
| t | Follow-up colonoscopies | 21,302 | $=e^{*} \mathrm{r}$ |
| $u$ | Further follow-up colonoscopies | 8,521 | = ${ }^{*} \mathrm{t}$ |
| v | Weighted cost per follow-up colonoscopy | \$666.83 | $=\left(h^{*} 0.6\right)+\left({ }^{*} 0.4\right)$ |
| w | Cost of follow-up colonoscopies | \$14,204,770 | = ${ }^{*}$ v |
| x | Cost of further follow-up colonoscopies | \$5,056,261 | = $\mathrm{u}^{*} \mathrm{j}$ |
| y | Patient time costs associated with follow-up colonoscopies | \$6,640,812 | $=((\mathrm{t}+\mathrm{u}) * 7.5))^{*} \mathrm{~m}$ |
| z | Total Costs of Screening and Follow-up | \$81,809,590 | $=\mathrm{k}+\mathrm{p}+\mathrm{q}+\mathrm{w}+\mathrm{x}+\mathrm{y}$ |
| aa | Deaths prevented | 105 | Table 2, row $x+y$ |
| ab | Costs avoided per death prevented | -\$49,197 | Ref Doc |
| ac | Costs avoided due to deaths prevented | -\$5,146,491 | = aa * ab |
| ad | Net screening and patient costs (undiscounted) | \$76,663,099 | $=\mathrm{ff}+\mathrm{dd}+\mathrm{aa}$ |
| ae | QALYs saved (undiscounted) | 1,734 | Table 2, row ah |
| af | Net screening and patient costs (1.5\% discount) | \$63,701,669 | Calculated |
| ag | QALYs saved (1.5\% discount) | 1,348 | Calculated |
| ah | CE (\$/QALY saved) | \$47,265 | = af/ag |

$\checkmark=$ Estimates from the literature
We also modified several major assumptions and recalculated the cost per QALY as follows:

- Assume the effectiveness of gFOBT in reducing the risk of mortality from CRC is reduced from $18 \%$ to $8 \%$ (Table 2, row $o$ ), the effectiveness of flexible sigmoidoscopy / colonoscopy in reducing the risk of mortality from CRC is reduced from $26 \%$ to $18 \%$ (Table 2, row $p$ ), the effectiveness of gFOBT in reducing the incidence of late-stage CRC is reduced from $8 \%$ to $1 \%$ (Table 2, row $r$ ) and the effectiveness of flexible sigmoidoscopy / colonoscopy in reducing the incidence of late-stage CRC is reduced from $27 \%$ to $18 \%$ (Table 2, row $s$ ): $\mathrm{CE}=\$ 82,979$.
- Assume the effectiveness of gFOBT in reducing the risk of mortality from CRC is increased from $18 \%$ to $27 \%$ (Table 2, row $o$ ), the effectiveness of flexible sigmoidoscopy / colonoscopy in reducing the risk of mortality from CRC is increased from $26 \%$ to $33 \%$ (Table 2, row $p$ ), the effectiveness of gFOBT in reducing the
incidence of late-stage CRC is increased from $8 \%$ to $15 \%$ (Table 2, row $r$ ) and the effectiveness of flexible sigmoidoscopy / colonoscopy in reducing the incidence of late-stage CRC is increased $27 \%$ to $34 \%$ (Table 2, row $s$ ): $\mathrm{CE}=\$ 32,923$.
- Assume that the proportion of FIT tests that are positive is decreased from $9.8 \%$ to $5.3 \%$ (Table 3, row $r$ ): $\mathrm{CE}=\$ 39,932$.
- Assume that the proportion of FIT tests that are positive is increased from $9.8 \%$ to $14.2 \%$ (Table 3, row $r$ ): $\mathrm{CE}=\$ 54,434$.


## Summary

| Table 4: Colorectal Cancer Screening Being Offered to a Birth Cohort of 40,000 Between the Ages of 50 and 74 Summary |  |  |  |
| :---: | :---: | :---: | :---: |
|  | Base <br> Case | Range |  |
| CPB (Potential QALYs Gained) |  |  |  |
| Assume No Current Service |  |  |  |
| 1.5\% Discount Rate | 1,348 | 790 | 1,879 |
| 3\% Discount Rate | 1,065 | 624 | 1,484 |
| 0\% Discount Rate | 1,734 | 1,017 | 2,418 |
| Gap between B.C. Current (50\%) and 'Best in the World' (76\%) |  |  |  |
| 1.5\% Discount Rate | 461 | 270 | 643 |
| 3\% Discount Rate | 364 | 213 | 508 |
| 0\% Discount Rate | 593 | 348 | 827 |
| $\overline{\mathbf{C E}(\$ / \mathrm{QLLY}}$ ) including patient time costs |  |  |  |
| 1.5\% Discount Rate | \$47,265 | \$32,923 | \$82,979 |
| 3\% Discount Rate | \$50,162 | \$34,942 | \$88,066 |
| 0\% Discount Rate | \$44,213 | \$30,798 | \$77,622 |
| CE (\$/QALY) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | \$29,282 | \$20,027 | \$52,309 |
| 3\% Discount Rate | \$31,077 | \$21,254 | \$55,515 |
| 0\% Discount Rate | \$27,391 | \$18,734 | \$48,931 |

## Screening for Lung Cancer

## Canadian Task Force on Preventive Health Care (2016)

We recommend screening for lung cancer among adults 55 to 74 years of age with at least a 30 pack-year smoking history, who smoke or quit smoking less than 15 years ago, with low-dose computed tomography (CT) every year up to three consecutive years. Screening should only be done in health care settings with access to expertise in early diagnosis and treatment of lung cancer. (Weak recommendation, low-quality evidence.)

We recommend not screening all other adults, regardless of age, smoking history or other risk factors, for lung cancer with low-dose CT. (Strong recommendation, very low quality evidence.)

We recommend that chest radiography, with or without sputum cytology, not be used to screen for lung cancer. (Strong recommendation, low-quality evidence.) ${ }^{236}$

## United States Preventive Services Task Force Recommendations (2014)

The USPSTF recommends annual screening for lung cancer with low-dose computed tomography in adults aged 55 to 80 years who have a 30 pack-year smoking history and currently smoke or have quit within the past 15 years. Screening should be discontinued once a person has not smoked for 15 years or develops a health problem that substantially limits life expectancy or the ability or willingness to have curative lung surgery. (Grade B recommendation) ${ }^{237}$

The relevant BC population includes all adults aged 55 to 74 years who have a 30 pack-year smoking history and currently smoke or have quit within the past 15 years. To estimate the relevant BC population, we used data from the 2012 Canadian Community Health Survey (CCHS) to determine the proportion of the population by age group who were current daily smokers, former daily (now occasional) smokers and former daily (now non-) smokers (variable SMKDSTY, type of smoker). ${ }^{238}$ This information was combined with data on the number of years smoked (variable SMKDYCS), years since stopped smoking daily (variable SMK_G09C), number of cigarettes smoked/day for daily smokers (variable SMK_204) and number of cigarettes smoked/day for former daily smokers (variable SMK_208) to calculate the proportion of smokers or former smokers who meet the criteria of a 30 pack-year smoking history and currently smoke or have quit within the past 15 years.

The data suggest that approximately 90,900 individuals between the ages of 55 to 74 meet the criteria for lung cancer screening in BC, or $8.7 \%$ of this population (see Table 1).

[^57]Table 1: Proportion of Population Eligible for Lung Cancer (LC) Screening British Columbia, 2013 by Age Group, Based on CCHS Data 2012

|  | Age Group (years) |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  | 55 to 59 | 60 to 64 | 65 to 69 | 70 to 74 | 55 to 74 |
| BC Population 2013 | 335,332 | 293,907 | 244,139 | 175,627 | 1,049,005 |
| Current Daily Smokers |  |  |  |  |  |
| Proportion of the Population in BC who are CD Smokers | 14.44\% | 10.04\% | 6.84\% | 5.78\% |  |
| Proportion of CD Smokers who Meet Criteria | 48.64\% | 48.96\% | 54.80\% | 48.34\% |  |
| Number of CD Smokers Eligible for LC Screening | 23,560 | 14,452 | 9,154 | 4,910 | 52,076 |
| Former Daily (Now Occasional) Smokers |  |  |  |  |  |
| Proportion of the Population in BC who are FD(NO) Smokers | 0.43\% | 0.33\% | 0.38\% | 0.00\% |  |
| Proportion of FD(NO) Smokers who Meet Criteria | 53.10\% | 89.86\% | 18.40\% | 0.00\% |  |
| Number of FD(NO) Smokers Eligible for LC Screening | 760 | 859 | 172 | 0 | 1,791 |
| Former Daily (Now Non-) Smokers |  |  |  |  |  |
| Proportion of the Population in BC who are FD(NN) Smokers | 6.44\% | 5.00\% | 6.00\% | 3.57\% |  |
| Proportion of FD(NN) Smokers who Meet Criteria | 50.9\% | 67.7\% | 81.5\% | 66.0\% |  |
| Number of FD(NN) Smokers Eligible for LC Screening | 11,002 | 9,957 | 11,939 | 4,140 | 37,038 |
| BC Population Eligible for LC Screening, by Age Group | 35,323 | 25,268 | 21,264 | 9,050 | 90,905 |
| Proportion of the BC Population Eligible for LC Screening, by Age Group | 10.5\% | 8.6\% | 8.7\% | 5.2\% | 8.7\% |
| CD=current daily; $\mathrm{FD}(\mathrm{NO}$ ) = former (now occasional); $\mathrm{FD}(\mathrm{NN})$ = former daily (now non-) |  |  |  |  |  |

Note that this estimate is lower than the Canadian average based on the Cancer Risk Management Model (CRMM). In a cost-effectiveness analysis using the CRMM, Goffin and colleagues estimated that $32 \%$ of 55-59 year-olds would be eligible for screening, decreasing to $30 \%$ for $60-64,23 \%$ for $65-69$ and $15 \%$ for $70-74{ }^{239}$

## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with screening for lung cancer in adults aged 55 to 74 years who have a 30 pack-year smoking history and currently smoke or have quit within the past 15 years, in a BC birth cohort of 40,000 .

In modelling CPB, we made the following assumptions:

- Based on BC life tables for 2010 to 2012, a total of 8,909 deaths would be expected between the ages of 55-79 in a BC birth cohort of 40,000 (see Table 2). Routine screening occurs to age 74 , but we have assumed the protective effect of routine screening continues to age 79 .
- Based on BC vital statistics data, there were 5,117 deaths between the ages of 45 and 64 in BC in 2012, with 544 ( $10.6 \%$ ) of these deaths due to lung cancer (ICD-10 codes C34). There were also 8,674 deaths between the ages of 65 and 79 that year, with $1,102(12.7 \%)$ of these deaths due to lung cancer. ${ }^{240}$ This suggests that 1,098 of the $8,909(12.3 \%)$ of the deaths in the BC birth cohort between the ages of 55 and 79 would be due to lung cancer (see Table 2).

[^58]| Table 2: Mortality Due to Lung Cancer Between the Ages of 55 and 79 <br> in a British Columbia Birth Cohort of 40,000 |  |  |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Age Group | Mean Survival Rate |  | Individuals in Birth Cohort |  |  | Life Years Lived | Deaths in Birth Cohort |  | Deaths due to Lung Cancer |  | Life Years Lost Per |  |
|  | Males | Females | Males | Females | Total |  | \% | \# | \% | \# | Death | Total |
| 50-54 | 0.950 | 0.969 | 19,003 | 19,375 | 38,378 | 191,890 |  |  |  |  |  |  |
| 55-59 | 0.931 | 0.956 | 18,619 | 19,118 | 37,737 | 188,686 | 1.7\% | 641 | 10.6\% | 68 | 27.7 | 1,882 |
| 60-64 | 0.902 | 0.936 | 18,041 | 18,726 | 36,767 | 183,834 | 2.6\% | 970 | 10.6\% | 103 | 23.4 | 2,407 |
| 65-69 | 0.858 | 0.906 | 17,164 | 18,113 | 35,277 | 176,387 | 4.2\% | 1,489 | 12.7\% | 189 | 19.2 | 3,632 |
| 70-74 | 0.792 | 0.857 | 15,837 | 17,144 | 32,981 | 164,903 | 7.0\% | 2,297 | 12.7\% | 292 | 15.3 | 4,463 |
| 75-79 | 0.693 | 0.780 | 13,861 | 15,608 | 29,469 | 147,346 | 11.9\% | 3,511 | 12.7\% | 446 | 11.8 | 5,262 |
|  |  |  |  |  |  |  |  | 8,909 | 12.3\% | 1,098 | 16.1 | 17,645 |

- In the National Lung Cancer Screening Trial (NLST), 53,454 persons at high risk of lung cancer were randomly assigned to undergo three annual screenings (see Table 4, row $j$ ) with low-dose computed tomography (LDCT group) or singleview posteroanterior chest radiography (X-ray group). Mortality from lung cancer was reduced by $19.6 \%$ ( RR of $0.804,95 \% \mathrm{CI}$ of 0.700 to 0.923 ) in the CT group (see Table 4, row $w$ ) compared to the X-ray group. Mortality from any cause was reduced by $6.1 \%$ (RR of $0.939,95 \% \mathrm{CI}$ of 0.884 to 0.998 ). Based on a nodule cut-off size of 4 mm (to be identified as a positive screen), $24.2 \%$ of all screens in the CT group were positive (see Table 4, row $m$ ). Of these positive screens, $96.4 \%$ were false positives (see Table 4, row $o$ ). ${ }^{241}$
- Three smaller, low quality RCTs have found no significant reduction in either lung cancer or all-cause mortality associated with screening with LDCT versus usual care (RR of $1.42,95 \% \mathrm{CI}$ of 0.91 to 2.22). ${ }^{242}$
- Compared with usual care, screening with LDCT detects lung cancers at an earlier stage. With LDCT, $66 \%$ of lung cancers at detected at Stage I or II, versus $40 \%$ with usual care (see Table 3). ${ }^{243,244}$

| Table 3: Stage of Lung Cancers: Screening with LDCT vs. Usual Care |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: |
| Usual Care Group |  |  | LDCT Group |  |
| Stage | \# | \% | \# | \% |
| I or II | 21 | 40.4\% | 83 | 65.9\% |
| III or IV | 31 | 59.6\% | 43 | 34.1\% |
| Total | 52 | 100.0\% | 126 | 100.0\% |
| Source: Canadian Cancer: Systema | k Forc eview | Preventive eta-analys | Care. | ening for |

[^59]- To date, the uptake of lung cancer screening has been less than optimal, with just $6.0 \%$ of the eligible US population being screened in 2015 (see Reference Document for more details). ${ }^{245}$ For modelling purposes we have assumed that screening rates of $60 \%$ (see Table 4 , row $k$ ) would eventually be achieved, with sensitivity analysis using a range from $50-70 \%$. The $60 \%$ is approximately halfway between current screening rates in BC for breast cancer ( $52 \%$ ) and cervical cancer (69\%) (see Reference Document).
- Screening with LDCT is also associated with a number of harms, including deaths following invasive follow-up testing, over diagnosis, major complications, false positive results and invasive procedures as a consequence of the false positive results. ${ }^{246}$
- Death from follow-up testing refers to "mortality that is the direct consequence of an invasive follow-up procedure (e.g., video-assisted thoracoscopic surgery, fine-needle aspiration biopsy or fine-needle aspiration cytology, thoracotomy, bronchoscopy, mediastinoscopy, surgical resection) initiated as a result of screening. ${ }^{, 247}$ Based upon a review of seven studies, the CTFPHC found that 20 of $1,502(1.33 \%)$ patients died as a result of follow-up testing after screening with LDCT (see Table 4, row $s$ ).
- "Overdiagnosis refers to the detection of a lung cancer that will not otherwise cause symptoms throughout the person's lifetime or result in death. ${ }^{248}$ Based upon a review of four studies, the CTFPHC found an overdiagnosis rate of between $11.0 \%$ and $25.8 \%$. The rate in the NLST was $11.0 \%$ ( $95 \%$ CI of $3.2 \%$ to $18.2 \%$ ).
- Major complications are defined as "requiring hospitalization or medical intervention (e.g., hemothorax and pneumothorax requiring tube placement, lung collapse, severe pain, cardiac arrhythmias and thromboembolic complications) that are the direct result of an invasive procedure (e.g., video-assisted thoracoscopic surgery, fine-needle aspiration biopsy or fine-needle aspiration cytology, thoracotomy, bronchoscopy, mediastinoscopy, surgical resection) initiated as a result of screening." ${ }^{249}$ Based upon a review of four studies, the CTFPHC found that 92 of 1,336 (1.33\%) patients had major complications as a result of follow-up testing after screening with LDCT.
- "A false positive refers to a screening test result that indicates the presence of lung cancer, when in fact no lung malignancy exists." ${ }^{250}$ Based upon a review of seven studies, the CTFPHC found that 8,290 of 42,774 (19.4\%) individuals who underwent screening with LDCT received at least one false positive result.
- Minor (e.g., fine-needle aspiration biopsy or fine-needle aspiration cytology, thoracic or lymph node biopsy, bronchoscopy) and major (e.g., video-assisted thoracoscopic surgery, thoracotomy, surgical resection) invasive procedures

[^60]initiated as a result of false positive screening tests. Based on a review of seven studies, the CTFPHC found that $0.72 \%$ ( $95 \% \mathrm{CI}$ of $0.33 \%$ to $1.11 \%$ ) of individuals with benign conditions underwent minor invasive procedures. Based on a further review of 17 studies, the CTFPHC found that $0.50 \%$ ( $95 \% \mathrm{CI}$ of $0.37 \%$ to $0.63 \%$ ) of individuals with benign conditions underwent major invasive procedures. ${ }^{251}$

- We have assumed a disutility of 0.05 associated with a false positive screen (see Table 4, row $q$ ). ${ }^{252,253}$
- Note that the NLTS (which the CTFPHC and our model follow) used a nodule cut-off size of 4 mm (to be identified as a positive screen). Significant analysis has since been completed to assess the pros and cons of moving to a larger nodule cut-off size as well as developing more advanced algorithms to fine-tune screening frequency.
- Gierada and colleagues re-examined the NLST results based on results associated with different size nodules. ${ }^{254}$ Moving the nodule cut-off size from 4 mm to 5 mm resulted in a $1.0 \%$ increase in missed or delayed lung cancer diagnosis but a $15.8 \%$ reduction in false positive results. With a cut-off of 8 mm , there would have been a $10.5 \%$ increase in missed or delayed lung cancer diagnosis but a $65.8 \%$ reduction in false positive results.
- Henschke et al. tested the effect of moving the nodule cut-off size to between 6 mm and 9 mm on false positive results and potential delays in detecting lung cancers. ${ }^{255}$ When alternative cut-offs of $6,7,8$ and 9 mm were used, the overall proportion of positive results declined to $10.2 \%, 7.1 \%, 5.1 \%$ and $4.8 \%$. The use of these alternative cut-offs would have reduced the work-up load by $36 \%, 56 \%$, $68 \%$ and $75 \%$ respectively. Concomitantly, a lung cancer diagnosis would have been delayed by at most 9 months in $0 \%, 5.0 \%, 5.9 \%$, and $6.7 \%$ of cases of cancer.
- The Pan-Canadian Early Detection of Lung Cancer Study (PAN-CAN) developed a more sophisticated approach to ascertaining the probability of lung cancer in pulmonary nodules detected on first screening CT, based on a combination of nodule size, age, sex, family history of lung cancer, emphysema location, type and count of the nodule and spiculation. ${ }^{256}$ Based on this approach, $80 \%$ of first screens placed patients in Category I ( $<1.5 \%$ lung cancer risk over the next 5.5 years), $12 \%$ in Category II ( $1.5 \%-<6 \%$ risk), $6 \%$ in Category 3 ( $6 \%$ $-<30 \%$ risk) and $2 \%$ in Category IV ( $\geq 30 \%$ risk). ${ }^{257}$

[^61]- The PAN-CAN lung cancer risk model has been validated in at least two studies. ${ }^{258,259}$ The results suggest that nodule size is still the most important predictor of lung cancer risk, with nodule spiculation, age and family history of lung cancer also being important predictive variables.
- The developers of the PAN-CAN lung cancer risk model suggest that patients in Category I require biennial screening, those in Category II require annual screening, those in Category III require rescreening in three months with annual screening thereafter if no growth in nodule size and those in Category IV should be referred for a definitive diagnosis. ${ }^{260}$
- A recent retrospective analysis of the NLST data suggests that annual screening might not be needed in individuals who have no abnormality identified on their initial screen and that a screening interval of at least two years could be considered on these individuals. ${ }^{261,262}$

Based on the above assumptions drawn from the NLST and the CTFPHC, the CPB is 1,745 quality-adjusted life years saved (see Table 4, row $z$ ). The CPB of 1,745 represents the gap between the existing coverage (no coverage) and $60 \%$.

$v=$ Estimates from the literature

[^62]We also modified a number of major assumptions and recalculated the CPB as follows:

- Assume the estimated effectiveness of lung cancer screening in reducing deaths due to lung cancers is reduced from $19.6 \%$ to $7.7 \%$ (Table 4, row $w$ ): $\mathrm{CPB}=485$.
- Assume the estimated effectiveness of lung cancer screening in reducing deaths due to lung cancers is increased from $19.6 \%$ to $30.0 \%$ (Table 4, row $w$ ): $\mathrm{CPB}=2,846$.
- Assume the adherence rate is reduced from $60 \%$ to $50 \%$ (Table 4 , row $k$ ): $\mathrm{CPB}=$ 1,454.
- Assume the adherence rate is increased from $60 \%$ to $70 \%$ (Table 4 , row $k$ ): $\mathrm{CPB}=$ 2,036.


## Modelling Cost-Effectiveness

In this section, we will calculate the CE associated with screening for lung cancer in adults aged 55 to 74 years who have a 30 pack-year smoking history and currently smoke or have quit within the past 15 years, in a BC birth cohort of 40,000 .

In modelling CE, we made the following assumptions:

- Assessment of patient risk - There are an expected 37,737 individuals in a BC birth cohort of 40,000 who are expected to survive to age 55 (see Table 2). Each of the 37,737 survivors would undergo a one-time screen by their primary care practitioner to determine if they were eligible for lung cancer screening. We assumed that $85 \%$ of individuals would agree to this screening and varied this in the sensitivity analysis from $75 \%$ to $95 \%$ (see Table 6, row $c$ ).
- Costs of screening - We assumed an annual LDCT screening exam would cost $\$ 198$ (2017 CAD) (see Table 6, row $i$ ). ${ }^{263}$
- Physician visits - LDCT screening results in an additional 14 physician visits per 100 persons screened (see Table 6, row $j$ ). ${ }^{264}$
- Positive findings on the screening CT result in the ensuing follow-up procedures (Table 5 rows $c$ to $k$ ): ${ }^{265}$
- Follow-up chest CT $-49.8 \%$
- Follow-up chest radiograph $-14.4 \%$
- Follow-up PET/CT scan - 8.3\%
- Percutaneous biopsy $-1.8 \%$
- Bronchoscopy without biopsy $-1.8 \%$
- Bronchoscopy with biopsy $-1.8 \%$
- Mediastinoscopy $-0.7 \%$
- Thoracoscopy - $1.3 \%$
- Thoracotomy - $2.9 \%$

By including all ensuing procedures following a positive screening CT result, we also include those procedures attributable to all identified harms, including deaths

[^63]following invasive follow-up testing, overdiagnosis, major complications, false positive results and invasive procedures as a consequence of the false positive results.

- The unit cost of the ensuing follow-up procedures is as follows (Table 5, rows $u$ to $a c):{ }^{266}$
- Follow-up chest radiograph - \$67
- Follow-up chest CT - \$164
- Follow-up PET/CT scan - \$1,399
- Percutaneous biopsy - CT-guided $=\$ 1,083$, US-guided $=\$ 682$
- Bronchoscopy without biopsy - $\$ 747$
- Bronchoscopy with biopsy - $\$ 804$
- Mediastinoscopy - $\$ 976$
- Thoracoscopy - $\$ 16,814$
- Thoracotomy - \$18,689
- Patient time and travel costs for follow-up procedures - We assumed 2 hours of patient time for a follow-up chest radiograph or chest CT, and 7.5 hours of patient time for a PET/CT scan, percutaneous biopsy or bronchoscopy. For a mediastinoscopy or a thoracoscopy we assumed a hospital stay of 3 days plus 4 weeks recovery (see Table 5, rows ae to $a m$ ).

[^64]| Row Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Number of positive screens | 1,297 | Table 4, row n |
| b | Number of false positive screens | 1,250 | Table 4, row p |
|  | Proportion of positive screens undergoing investigation |  |  |
| c | Follow-up chest radiograph | 14.4\% | $\checkmark$ |
| d | Follow-up chest CT | 49.8\% | $\checkmark$ |
| e | Follow-up PET/CT scan | 8.3\% | $\checkmark$ |
| f | Percutaneous biopsy | 1.8\% | $\checkmark$ |
| g | Bronchoscopy without biopsy | 1.8\% | $\checkmark$ |
| h | Bronchoscopy with biopsy | 1.8\% | $\checkmark$ |
| i | Mediastinoscopy | 0.7\% | $\checkmark$ |
| j | Thoracoscopy | 1.3\% | $\checkmark$ |
| k | Thoracotomy | 2.9\% | $\checkmark$ |
|  | Number of procedures following a positive screen |  |  |
| 1 | Follow-up chest CT | 187 | =a*c |
| m | Follow-up chest radiograph | 646 | =a*d |
| n | Follow-up PET/CT scan | 108 | =a*e |
| 0 | Percutaneous biopsy | 23 | = ${ }^{*} \mathrm{f}$ |
| p | Bronchoscopy without biopsy | 23 | = ${ }^{*} \mathrm{~g}$ |
| q | Bronchoscopy with biopsy | 23 | =a* h |
| r | Mediastinoscopy | 9 | =a*i |
| s | Thoracoscopy | 16 | = ${ }^{*} \mathrm{j}$ |
| t | Thoracotomy | 36 | = ${ }^{*}$ k |
|  | Unit cost of procedures following a positive screen |  |  |
| u | Follow-up chest radiograph | \$67 | $\checkmark$ |
| v | Follow-up chest CT | \$164 | $\checkmark$ |
| w | Follow-up PET/CT scan | \$1,399 | $\checkmark$ |
| x | Percutaneous biopsy | \$883 | $\checkmark$ |
| y | Bronchoscopy without biopsy | \$747 | $\checkmark$ |
| z | Bronchoscopy with biopsy | \$804 | $\checkmark$ |
| aa | Mediastinoscopy | \$976 | $\checkmark$ |
| ab | Thoracoscopy | \$16,814 | $\checkmark$ |
| ac | Thoracotomy | \$18,689 | $\checkmark$ |
| ad | Follow-up costs of positive screens | \$1,283,108 | $\begin{gathered} =\mathrm{I}^{*} \mathrm{u}+\mathrm{m}^{*} \mathrm{v}+\mathrm{n}^{*} \mathrm{w}+\mathrm{o}^{*} \mathrm{x} \\ +\mathrm{p}^{*} \mathrm{y}+\mathrm{q}^{*} \mathrm{z}+\mathrm{r}^{*} \mathrm{aa}+ \\ \mathrm{s}^{*} \mathrm{ab}+\mathrm{t}^{*} \mathrm{ac} \end{gathered}$ |
|  | Estimated patient time (in hours) per follow-up procedure |  |  |
| ae | Follow-up chest CT | 2.0 | Assumed |
| af | Follow-up chest radiograph | 2.0 | Assumed |
| ag | Follow-up PET/CT scan | 7.5 | Assumed |
| ah | Percutaneous biopsy | 7.5 | Assumed |
| ai | Bronchoscopy without biopsy | 7.5 | Assumed |
| aj | Bronchoscopy with biopsy | 7.5 | Assumed |
| ak | Mediastinoscopy | 7.5 | Assumed |
| al | Thoracoscopy | 172.5 | Assumed |
| am | Thoracotomy | 172.5 | Assumed |
| an | Hours of patient time associated with positive screens | 12,101 | $\begin{gathered} \hline=\text { l }^{*} a \mathrm{e}+\mathrm{m}^{*} \mathrm{af}+\mathrm{n}^{*} \mathrm{ag}+ \\ \text { o }^{*} \mathrm{ah}+\mathrm{p}^{*} \mathrm{ai}+\mathrm{q}^{*} \mathrm{aj}+ \\ \mathrm{r}^{*} \mathrm{ak}+\mathrm{s}^{*} \mathrm{al}+\mathrm{t}^{*} a \mathrm{~m} \end{gathered}$ |
| ao | Value of patient time per hour | \$29.69 | $\checkmark$ |
| ap | Total cost of patient time for follow-up procedures | \$359,290 | = ao * ap |
| aq | Cost of follow-up procedures | \$1,642,398 | = ad + ap |

- Costs avoided due to early detection of lung cancers - As noted in Table 3, screening with LDCT results in the earlier detection of lung cancers, thus potentially reducing the cost of treatment. Research by Cressman et al. suggests that the mean per person cost of treating stage I \& II lung cancer is \$34,267 (95\% CI of \$32,426$\$ 35,902) .{ }^{267}$ This increases to $\$ 49,115(95 \%$ CI of $\$ 44,451$ - $\$ 53,645)$ for stage III \& IV lung cancers. These costs include the diagnostic work-up, treatment and 2 years of follow-up. Based on the stage distribution noted in Table 3, the weighted cost would be $\$ 43,119$ for the usual care group and $\$ 37,288$ for the CT group, resulting in costs avoided of $\$ 5,831$ per lung cancer associated with LDCT screening (see Table 6 , row n).
- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the estimated cost per QALY would be $\$ 2,204$ (see Table 6, row $\mathrm{u})$.

[^65] Cancer Screening with Computed Tomography in Canada. Journal of Thoracic Oncology. 2014; 9(10): 1449-58.

Table 6. Summary of Cost Effectiveness (CE) Estimate for Lung Cancer Screening

| Row Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
|  | Assessment of patient risk |  |  |
| a | Proportion of cohort alive at age 55 | 94.3\% | $\checkmark$ |
| b | Total number of primary care provider screens (100\% adherence) | 37,737 | = a * 40,000 |
| c | Adherence with screening | 85\% | Assumed |
| d | Cost of 10-minute office visit | \$34.85 | Ref Doc |
| e | Value of patient time and travel for office visit | \$59.38 | Ref Doc |
| f | Portion of 10-minute office visit for screen | 50\% | Assumed |
| g | Cost of primary care provider screening | \$1,511,290 | $=\left(b^{*} \mathrm{c}\right) *((\mathrm{~d}+\mathrm{e}) * \mathrm{f})$ |
|  | Screening for Lung Cancer |  |  |
| h | Potential screens with 60\% adherence | 5,359 | =Table 4, row 1 |
| i | Cost per screen | \$198 | $\checkmark$ |
| j | Additional physician visits per screening exam | 0.14 | $\checkmark$ |
| k | Cost of screening | \$1,131,712 | $=(\mathrm{i} * \mathrm{~h})+((\mathrm{h} * \mathrm{j}) *(\mathrm{~d}+\mathrm{e}))$ |
| I | Costs Asspociated with Follow-up Procedures | \$1,642,398 | =Table 5, row aq |
| m | Total Costs of Screening and Follow-up | \$4,285,400 | $=\mathrm{g}+\mathrm{k}+\mathrm{l}$ |
|  | Costs Avoided |  |  |
| n | Treatment costs avoided with earlier detection, per cancer | -\$5,831 | $\checkmark$ |
| $\bigcirc$ | Number of incident lung cancers detected earlier | 112 | = Table 4, row y |
| p | Treatment costs avoided with earlier detection | -\$655,691 | = ${ }^{*}$ o |
| q | Net screening and patient costs (undiscounted) | \$3,629,710 | = $\mathrm{m}+\mathrm{p}$ |
| r | QALY s saved (undiscounted) | 1,745 | Table 4, row z |
| s | Net screening and patient costs (1.5\% discount) | \$3,140,279 | Calculated |
| t | QALYs saved (1.5\% discount) | 1,402 | Calculated |
| u | CE (\$/QALY saved) | \$2,240 | =s/t |

$\checkmark$ = Estimates from the literature

We also modified a number of major assumptions and recalculated the cost per QALY as follows:

- Assume the estimated effectiveness of lung cancer screening in reducing deaths due to lung cancers is reduced from $19.6 \%$ to $7.7 \%$ (Table 4, row $w$ ): $\mathrm{CE}=\$ 9,026$.
- Assume the estimated effectiveness of lung cancer screening in reducing deaths due to lung cancers is increased from $19.6 \%$ to $30.0 \%$ (Table 4, row $w$ ): $\mathrm{CE}=\$ 1,228$.
- Assume the adherence rate is reduced from $60 \%$ to $50 \%$ (Table 4 , row $k$ ): $\mathrm{CE}=$ \$2,425.
- Assume the adherence rate is increased from $60 \%$ to $70 \%$ (Table 4, row $k$ ): $\mathrm{CE}=$ \$2,107.
- Assume the adherence rate with the assessment of patient risk is reduced from $85 \%$ to $75 \%$ (Table 6, row $c$ ): $\mathrm{CE}=\$ 2,131$.
- Assume the adherence rate with the assessment of patient risk is increased from $85 \%$ to $95 \%$ (Table 6, row $c$ ): $\mathrm{CE}=\$ 2,349$.
- Assume that the portion of a 10-minute office visit for the assessment of patient risk is reduced from $50 \%$ to $33 \%$ (Table 6 , row $f$ ): $\mathrm{CE}=\$ 1,924$.
- Assume that the portion of a 10 -minute office visit for the assessment of patient risk is increased from $50 \%$ to $67 \%$ (Table 6, row $f$ ): $\mathrm{CE}=\$ 2,555$.


## Summary

| Cohort of 40,000 Between the Ages of 55 and 74 Summary |  |  |  |
| :---: | :---: | :---: | :---: |
|  | $\begin{aligned} & \text { Base } \\ & \text { Case } \end{aligned}$ | Range |  |
| CPB (Potential QALYs Gained) |  |  |  |
| Gap between B.C. Current (0\%) and 'Best in the World' (60\%) |  |  |  |
| 1.5\% Discount Rate | 1,402 | 390 | 2,287 |
| 3\% Discount Rate | 1,303 | 362 | 2,125 |
| 0\% Discount Rate | 1,745 | 485 | 2,846 |
|  |  |  |  |
| 1.5\% Discount Rate | \$2,240 | \$1,228 | \$9,206 |
| 3\% Discount Rate | \$2,296 | \$1,261 | \$9,239 |
| \%\% Discount Rate | \$2,080 | \$1,135 | \$8,419 |
| CE (\$/QALY) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | \$1,408 | \$718 | \$6,035 |
| 3\% Discount Rate | \$1,445 | \$739 | \$6,180 |
| 0\% Discount Rate | \$1,303 | \$658 | \$5,625 |

## Hypertension Screening and Treatment

## United States Preventive Services Task Force Recommendations (2015)

The USPSTF recommends screening for high blood pressure in adults age 18 years and older. (A recommendation).

The USPSTF recommends obtaining measurements outside of the clinical setting for diagnostic confirmation before starting treatment. ${ }^{268}$

## Canadian Task Force on Preventive Health Care Recommendations (2012)

We recommend blood pressure measurement at all appropriate primary care visits... (in) adults aged 18 years and older without previously diagnosed hypertension for the purpose of screening for hypertension. (Strong recommendation; moderate quality evidence)
We recommend that blood pressure be measured according to the current techniques described in the Canadian Hypertension Education Program (CHEP)

[^66]recommendations for office and out-of-office (ambulatory) blood pressure measurement). (Strong recommendation; moderate quality evidence)
For people who are found to have an elevated blood pressure during screening, the CHEP criteria for assessment and diagnosis of hypertension should be applied to determine whether the patient meets diagnostic criteria for hypertension. (Strong recommendation; moderate quality evidence) $)^{269}$

## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with screening for and treatment of hypertension in adults aged 18 and older in a BC birth cohort of 40,000 .

In modelling CPB, we made the following assumptions:

- Based on BC life tables for 2010 to 2012, there are a total of $2,436,832$ life years lived and 15,233 deaths between the ages of 18 and 84 in a BC birth cohort of 40,000 (see Table 1).
- Based on BC vital statistics data, 59 of 993 (5.9\%) deaths in 25-44 year olds in 2011 were due to cardiovascular disease (ICD-10 codes I00-I51). In 45-64 year olds, 601 of 5,076 (11.8\%) deaths were due to cardiovascular disease. In 65-84 year olds, 2,248 of $13,481(16.7 \%)$ deaths were due to cardiovascular disease. ${ }^{270}$
- Congestive heart failure deaths (ICD-10 codes I50) are a subset of cardiovascular disease (ICD-10 codes I50). In 2011, 23 of the 5,076 ( $0.45 \%$ ) deaths in 45-64 year olds was due to CHF. In 65-79 year olds, 88 of $8,600(1.02 \%)$ deaths were due to CHF. In the population ages 80 and older, 596 of $16,612(3.59 \%)$ deaths were due to CHF. ${ }^{271}$
- Based on BC vital statistics data, 31 of 993 (3.1\%) deaths in 25-44 year olds in 2011 were due to cerebrovascular disease (ICD-10 codes I60-I69). In 45-64 year olds, 191 of 5,076 (3.8\%) deaths were due to cerebrovascular disease. In 65-84 year olds, 905 of $13,481(6.7 \%)$ deaths were due to cerebrovascular disease. ${ }^{272}$
- This data was used to estimate that approximately 2,092 ( $13.7 \%$ ) of the 15,233 deaths in the birth cohort would be due to cardiovascular disease (excluding deaths due to CHF), 266 ( $1.74 \%$ ) due to CHF and 929 (6.1\%) due to cerebrovascular disease (see Table 1)

[^67]| Between the Ages of 18 and 84 <br> in a British Columbia Birth Cohort of 40,000 |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Age Group | Mean Survival Rate | Individuals in Birth Cohort | Life Years Lived | Death <br> Birth <br> \% | hs in Cohort \# | Cardiov Dise \% | ascular <br> \# | Conge <br> Heart F \% |  | ths due to <br> Cardio Disease \% | scular <br> xcl CHF) <br> \# | $\begin{gathered} \text { Cerebro } \\ \text { Dise } \\ \% \\ \hline \end{gathered}$ | ascular <br> \# | Life Expectancy | All Deaths | Life Yea <br> Cardio | rs Lost CHF | Cerebro |
| 18-19 | 0.994 | 39,744 | 79,488 | 0.1\% | 40 | 5.9\% | 2 | 0.0\% | 0 | 5.9\% | 2 | 3.1\% | 1 | 63.7 | 2,548 | 150 | 0 | 79 |
| 20-24 | 0.992 | 39,682 | 198,408 | 0.2\% | 62 | 5.9\% | 4 | 0.0\% | 0 | 5.9\% | 4 | 3.1\% | 2 | 60.8 | 3,794 | 224 | 0 | 118 |
| 25-29 | 0.989 | 39,570 | 197,850 | 0.3\% | 112 | 5.9\% | 7 | 0.0\% | 0 | 5.9\% | 7 | 3.1\% | 3 | 56.0 | 6,250 | 369 | 0 | 194 |
| 30-34 | 0.986 | 39,458 | 197,290 | 0.3\% | 112 | 5.9\% | 7 | 0.0\% | 0 | 5.9\% | 7 | 3.1\% | 3 | 51.1 | 5,723 | 338 | 0 | 177 |
| 35-39 | 0.983 | 39,310 | 196,550 | 0.4\% | 148 | 5.9\% | 9 | 0.0\% | 0 | 5.9\% | 9 | 3.1\% | 5 | 46.3 | 6,852 | 404 | 0 | 212 |
| 40-44 | 0.978 | 39,105 | 195,526 | 0.5\% | 205 | 5.9\% | 12 | 0.0\% | 0 | 5.9\% | 12 | 3.1\% | 6 | 41.5 | 8,499 | 501 | 0 | 263 |
| 45-49 | 0.970 | 38,814 | 194,070 | 0.8\% | 291 | 11.8\% | 34 | 0.45\% | 1 | 11.4\% | 33 | 3.8\% | 11 | 36.8 | 10,716 | 1,216 | 48 | 407 |
| 50-54 | 0.960 | 38,390 | 191,948 | 1.1\% | 424 | 11.8\% | 50 | 0.45\% | 2 | 11.4\% | 48 | 3.8\% | 16 | 32.2 | 13,666 | 1,551 | 61 | 519 |
| 55-59 | 0.944 | 37,757 | 188,786 | 1.7\% | 632 | 11.8\% | 75 | 0.45\% | 3 | 11.4\% | 72 | 3.8\% | 24 | 27.7 | 17,517 | 1,988 | 79 | 666 |
| 60-64 | 0.920 | 36,800 | 183,998 | 2.6\% | 958 | 11.8\% | 113 | 0.45\% | 4 | 11.4\% | 109 | 3.8\% | 36 | 23.4 | 22,408 | 2,543 | 101 | 851 |
| 65-69 | 0.883 | 35,332 | 176,658 | 4.2\% | 1,468 | 16.7\% | 245 | 1.02\% | 15 | 15.7\% | 230 | 6.7\% | 98 | 19.2 | 28,186 | 4,420 | 287 | 1,888 |
| 70-74 | 0.827 | 33,072 | 165,362 | 6.8\% | 2,259 | 16.7\% | 377 | 1.02\% | 23 | 15.7\% | 354 | 6.7\% | 151 | 15.3 | 34,566 | 5,420 | 353 | 2,316 |
| 75-79 | 0.741 | 29,628 | 148,142 | 11.6\% | 3,444 | 16.7\% | 575 | 1.02\% | 35 | 15.7\% | 540 | 6.7\% | 231 | 11.8 | 40,639 | 6,372 | 415 | 2,723 |
| 80-84 | 0.614 | 24,551 | 122,756 | 20.7\% | 5,077 | 16.7\% | 848 | 3.59\% | 182 | 13.1\% | 666 | 6.7\% | 340 | 8.7 | 44,172 | 5,791 | 1,586 | 2,959 |
| Total |  |  | 2,436,832 |  | 15,233 | 15.5\% | 2,358 | 1.74\% | 266 | 13.7\% | 2,092 | 6.1\% | 929 |  | 245,536 | 31,288 | 2,930 | 13,374 |

- An estimated $38.5 \%$ (Table 2, row $l$ ) of cerebrovascular deaths, $24.6 \%$ (Table 2, row $j$ ) of cardiovascular deaths and $33.0 \%$ (Table 2, row $k$ ) of CHF deaths are attributable to hypertension. ${ }^{273}$
- In a meta-analysis of 147 randomized trials, Law and colleagues found that lowering blood pressure by $10 / 5 \mathrm{~mm} \mathrm{Hg}$ (the equivalent of taking one drug at a standard dose) resulted in a $22 \%$ ( $95 \% \mathrm{CI}$ of $17 \%$ to $27 \%$ ) (Table 2 , rows $q \& r$ ) reduction in cardiovascular events and a $41 \%$ ( $95 \%$ CI of $33 \%$ to $48 \%$ ) (Table 2, row $s$ ) reduction in cerebrovascular events. ${ }^{274}$
- Other assumptions used in assessing the clinically preventable burden are detailed in the Reference Document.

Based on these assumptions, the CPB associated with screening for and treatment of hypertension in adults aged 18 and older is 11,587 QALYs saved (Table 2, row $a z$ ). The CPB of 11,587 QALYs saved represents the gap between no coverage and the 'best in the world' coverage estimated at $73 \%$.

We also modified several major assumptions and recalculated the CPB as follows:

- Assume that the proportion of the population with hypertension receiving drug treatment is decreased from $73 \%$ to $68 \%$ (Table 2, row $p$ ): $\mathrm{CPB}=10,523$.
- Assume that the proportion of the population with hypertension receiving drug treatment is increased from $73 \%$ to $78 \%$ (Table 2, row $p$ ): $\mathrm{CPB}=12,707$.
- Assume that the effectiveness of drug treatment in reducing cardiovascular disease events is decreased from $22 \%$ to $17 \%$ (Table 2 , rows $q \& r$ ) and the effectiveness of drug treatment in reducing cerebrovascular disease events is decreased from $41 \%$ to $33 \%$ (Table 2, row $s$ ): $\mathrm{CPB}=8,199$.

[^68]- Assume that the effectiveness of drug treatment in reducing cardiovascular disease events is increased from $22 \%$ to $29 \%$ (Table 2, rows $q \& r$ ) and the effectiveness of drug treatment in reducing cerebrovascular disease events is increased from $41 \%$ to $48 \%$ (Table 2, row $s$ ): $\mathrm{CPB}=15,792$.
- Assume that the disutility associated with living with a nonfatal cerebrovascular event is reduced from 0.264 to 0.177 (Table 2, row al): $\mathrm{CPB}=11,019$.
- Assume that the disutility associated with living with a nonfatal cerebrovascular event is increased from 0.264 to 0.350 (Table 2, row al): CPB 12,146.
- Assume that the disutility associated with taking pills for cardiovascular prevention is reduced from 0.0032 to 0.0 (Table 2, row $a x$ ): $\mathrm{CPB}=13,128$.
- Assume that the disutility associated with taking pills for cardiovascular prevention is increased from 0.0032 to 0.0044 (Table 2, row $a x$ ): $\mathrm{CPB}=11,009$.

| Row | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
|  | Estimated Current Status - Mortality |  |  |
| a | Total CHD (excluding CHF) mortality in the birth cohort | 2,092 | Table 1 |
| b | Total CHF mortality in the birth cohort | 266 | Table 1 |
| c | Total stroke mortality in the birth cohort | 929 | Table 1 |
| d | Life years lost per CHD death | 15.0 | Table 1 |
| e | Life years lost per CHF death | 11.0 | Table 1 |
| f | Life years lost per stroke death | 14.4 | Table 1 |
| g | Total life years lost due to CHD death | 31,288 | $=a^{*} \mathrm{~d}$ |
| h | Total life years lost due to CHF death | 2,930 | $={ }^{*}$ * |
| i | Total life years lost due to stroke death | 13,374 | = ${ }^{*} \mathrm{f}$ |
| j | \% CHD mortality attributable to hypertension | 24.6\% | $\checkmark$ |
| k | \% CHF mortality attributable to hypertension | 33.0\% | V |
| 1 | \% stroke mortality attributable to hypertension | 38.5\% | $\checkmark$ |
| m | Total CHD mortality in the birth cohort attributable to hypertension | 515 | $=a^{*} \mathrm{j}$ |
| n | Total CHF mortality in the birth cohort attributable to hypertension | 88 | $={ }^{*}$ * |
| 0 | Total stroke mortality in the birth cohort attributable to hypertension | 358 | = ${ }^{*}$ \\| |
| p | \% with hypertension receiving drug treatment | 73\% | Ref Doc |
| q | Effectiveness of drug treatment on CHD deaths | 22\% | $\checkmark$ |
| r | Effectiveness of drug treatment on CHF deaths | 22\% | V |
| s | Effectiveness of drug treatment on stroke deaths | 41\% | $\checkmark$ |
|  | Estimates in the Absence of Screening / Treatment |  |  |
|  | Mortality attributable to hypertension |  |  |
| t | Predicted hypertension-attributable CHD deaths in absence of screening | 613 | $=m /(1-(p * q))$ |
| $u$ | Predicted hypertension-attributable CHF deaths in absence of screening | 104 | $=\mathrm{n} /\left(1-\left(p^{*} r\right)\right)$ |
| v | Predicted hypertension-attributable stroke deaths in absence of screening | 511 | $=o /\left(1-\left(p{ }^{*} s\right)\right)$ |
| w | Predicted hypertension-attributable CHD life years lost in absence of screening | 9,169 | = ${ }^{*}$ d |
| x | Predicted hypertension-attributable CHF life years lost in absence of screening | 1,152 | $=u^{*} \mathrm{e}$ |
| y | Predicted hypertension-attributable stroke life years lost in absence of screening | 7,348 | = ${ }^{*} \mathrm{f}$ |
|  | Life years lost due to total deaths | 17,670 | $=w+x+y$ |
|  | Morbidity attributable to hypertension |  |  |
| z | Ratio of nonfatal cardiovascular events per fatal event | 5.09 | See Ref Doc |
| aa | \# of nonfatal cardiovascular events | 3,652 | $=(\mathrm{t}+\mathrm{u}) * \mathrm{z}$ |
| ab | Average age of individual with a cardiovascular event | 68.0 | See Ref Doc |
| ac | Life years lived with a nonfatal cardiovascular event | 12.1 | See Ref Doc |
| ad | Life years lost due to a nonfatal cardiovascular event | 6.3 | See Ref Doc |
| ae | QoL reduction living with a nonfatal cardiovascular event (for 1 month) | 0.125 | See Ref Doc |
| af | QALYs lost due to nonfatal cardiovascular events | 23,047 | $\begin{gathered} \hline=\mathrm{aa} *(\mathrm{ad}+(\mathrm{ae} / \\ 12)) \\ \hline \end{gathered}$ |
| ag | Ratio of nonfatal cerebrovascular events per fatal event | 4.58 | See Ref Doc |
| ah | \# of nonfatal cerebrovascular events | 2,339 | v*ag |
| ai | Average age of individual with a cerebrovascular event | 72.8 | See Ref Doc |
| aj | Life years lived with a nonfatal cerebrovascular event | 9.3 | See Ref Doc |
| ak | Life years lost due to a nonfatal cerebrovascular event | 5.5 | See Ref Doc |
| al | QoL reduction living with a nonfatal cerebrovascular event | 0.264 | See Ref Doc |
| am | QALYs lost due to nonfatal cerebrovascular events | 18,608 | $\begin{gathered} =\mathrm{ah} *(\mathrm{ak}+(\mathrm{aj} * \\ \mathrm{al})) \end{gathered}$ |
|  | Benefits if 73\% of individuals with hypertension are on drug treatment |  |  |
| ao | Number of CHD deaths prevented | 98 | $=t * p * q$ |
| ap | Number of CHF deaths prevented | 17 | $=u^{*} p^{*} r$ |
| aq | Number of stroke deaths prevented | 153 | =v*p*s |
| ar | Number of life years saved from CHD death prevented | 1,473 | $=w^{*} p^{*} q$ |
| as | Number of life years saved from CHF death prevented | 185 | $=x^{*} p^{*}{ }^{\text {r }}$ |
| at | Number of life years saved from stroke death prevented | 2,199 | =y*p*s |
| au | Total years of live saved from deaths prevented | 3,857 | = ar + as + at |
| av | QALY saved from prevented nonfatal cardiovascular disease events | 3,701 | $=\mathrm{af}$ * ${ }^{*}$ q |
| aw | QALY saved from prevented nonfatal cerebrovascular disease events | 5,569 | $=a m * p * s$ |
|  | Harms if 73\% of individuals with hypertension are on drug treatment |  |  |
| ax | Disutility per year associated with taking pills for cardiovascular prevention | -0.0032 | See Ref Doc |
| ay | Disutility associated with taking pills for cardiovascular prevention | -1,541 | $\begin{gathered} =\mathrm{p} \text { * Table 4, row } \\ \text { b*ax } \end{gathered}$ |
| az | Potential QALYs gained, screening and intervention from 0\% to 73\% | 11,587 | $=a u+a v+a w+a y$ |

[^69]
## Modelling Cost-Effectiveness

In this section, we will calculate the CE associated with screening for and treatment of hypertension in adults aged 18 and older in a BC birth cohort of 40,000 .

In modelling CE, we made the following assumptions:

- The proportion of the population with diagnosed hypertension is based on data provided by the BC Ministry of Health, Chronic Disease Management for fiscal 2002/03 (Table 3). ${ }^{275}$

| Table 3: Years Lived with Hypertension in a Birth Cohort of 40,000 |  |  |  |
| :---: | :---: | :---: | :---: |
| Age Group | $\begin{gathered} \% \\ \text { Hypertensive } \end{gathered}$ | Life Years | Life Years Lived with hypertension |
| 18-19 | 0.7\% | 79,488 | 544 |
| 20-24 | 1.5\% | 198,408 | 2,921 |
| 25-29 | 2.6\% | 197,850 | 5,16 |
| 30-34 | 4.0\% | 197,290 | 7,821 |
| 35-39 | 6.3\% | 196,550 | 12,359 |
| 40-44 | 10.7\% | 195,526 | 20,869 |
| 45-49 | 17.4\% | 194,070 | 33,803 |
| 50-54 | 26.3\% | 191,948 | 50,529 |
| 55-59 | 35.4\% | 188,786 | 66,816 |
| 60.64 | 43.9\% | 183,998 | 80,713 |
| 65-69 | 52.1\% | 176,658 | 92,077 |
| 70.74 | 59.6\% | 165,362 | 98,500 |
| 75-79 | 68.2\% | 148,142 | 101,101 |
| 80-84 | 75.3\% | 122,756 | 92,490 |
| Total |  | 2,436,832 | 665,769 |
| \% of years lived with hypertension 27.3\% |  |  |  |

- Costs of laboratory tests - The costs per diagnostic test (Table 4, rows $h$ to $o$ ) are based on information from the BC Medical Services Plan 2016/17 payment analysis. ${ }^{276}$
- Average annual cost of antihypertensive medication - Calculated based on an estimated average cost per day of treatment for antihypertensive medication in Canada of $\$ 0.53$ (Table 4, row p). ${ }^{277}$
- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the CE associated with universal screening for and treatment of hypertension in adults aged 18 and older is \$15,254 / QALY (Table 4, row $a v$ ).

[^70]Table 4: Summary of Cost-effectiveness Estimate for Hypertension in a Birth Cohort of 40,000

| (B.C.) |  |  |  |
| :---: | :---: | :---: | :---: |
| Row | Variable | Base Case | Data Source |
| a | Years of life in target population age range | 2,436,832 | Table 1 |
| b | Years of life lived with hypertension in target population age range | 665,769 | Table 3 |
| c | Portion of years eligible for screening | 1,771,063 | = $\mathrm{a}-\mathrm{b}$ |
|  | Costs of screening, lab monitoring and antihypertensive therapy |  |  |
| d | Cost of patient time and travel for office visit | \$59.38 | Ref Doc |
| e | Cost of office visit | \$34.85 | Ref Doc |
| f | Portion of 10 minute office visit used for screen | 50\% | Ref Doc |
| g | Portion of 10 minute office visit used for monitoring | 50\% | Ref Doc |
| h | 12-lead ECG | \$24.05 | $\checkmark$ |
| i | Urinalysis | \$7.42 | $\checkmark$ |
| j | Blood glucose | \$1.25 | $\checkmark$ |
| k | Hematocrit | \$3.22 | $\checkmark$ |
| I | Serum potassium | \$1.04 | $\checkmark$ |
| m | Creatinine | \$1.52 | V |
| n | Calcium | \$1.11 | $\checkmark$ |
| 0 | Lipid profile | \$6.87 | $\checkmark$ |
| p | Total costs for monitoring tests | \$46.48 | $=\mathrm{h}+\mathrm{i}+\mathrm{j}+\mathrm{k}+\mathrm{l}+\mathrm{m}+\mathrm{n}+\mathrm{o}$ |
| q | Average annual cost of antihypertensive, given current market share and adherence | \$193.45 | $\checkmark$ |
| $r$ | Average number of recommended hypertension screening tests per person year without diagnosis of hypertension | 0.5 | Ref Doc |
| t | Average number of recommended hypertension monitoring tests per person year of treatment | 2.0 | Assumed |
| u | Adherence with screening | 79\% | Ref Doc |
| v | Adherence with treatment | 73\% | Ref Doc |
| w | Lifetime screening costs | \$32,960,236 | $=\left(c^{*} u^{*}\right)^{*}((d+e) * f)$ |
| x | Lifetime non-screening monitoring costs | \$90,976,462 | $=\left(b *{ }^{*} \mathrm{t}\right) *(\mathrm{p}+((\mathrm{d}+\mathrm{e}) * \mathrm{~g}))$ |
| y | Lifetime anti-hypertensive therapy costs | \$94,018,893 | $=b^{*} q^{*} v$ |
| Estimated costs avoided due to intervention |  |  |  |
| z | Acute care costs avoided per avoided cardiovascular death | \$15,536 | Ref Doc |
| aa | Acute care costs avoided per avoided cerebrovascular death | \$9,583 | Ref Doc |
| ab | Costs avoided due to deaths avoided | \$1,725,327 | $\begin{gathered} \hline=\text { Table 2, row ao + (Table 2, row ap } \\ \quad{ }^{2} \text { z) + (Table 2, row aq * aa) } \\ \hline \end{gathered}$ |
| ac | First year costs avoided per nonfatal cardiovascular event avoided | \$33,934 | Ref Doc |
| ad | First year costs avoided per nonfatal cerebrovascular event avoided | \$21,139 | Ref Doc |
| ae | \# of cardiovascular events avoided | 587 | $\begin{gathered} \text { = Table } 2, \text { row aa } * \text { Table } 2 \text {, row } \mathrm{p}^{*} \\ \text { Table 2, row q } \end{gathered}$ |
| af | First-year acute care costs avoided / event | \$33,934 | Ref Doc |
| ag | Post-first-year annual costs avoided per nonfatal cardiovascular events avoided | \$2,278 | Ref Doc |
| ah | Number of years for which the costs are avoided | 12.1 | Ref Doc |
| ai | Total costs avoided for nonfatal cardiovascular events avoided | \$36,071,383 | = ae * (af + (ag * ah) ) |
| aj | \# of cerebrovascular events avoided | 700 | $\begin{gathered} =\text { Table } 2, \text { row ah * Table } 2 \text {, row } \mathrm{p}^{*} \\ \text { Table } 2 \text {, row s } \end{gathered}$ |
| ak | First-year acute care costs avoided / event | \$21,139 | Ref Doc |
| al | Post-first-year annual costs avoided per nonfatal cerebrovascular events avoided | \$6,246 | Ref Doc |
| am | Number of years for which the costs are avoided | 9.3 | Ref Doc |
| an | Post-first-year costs avoided for nonfatal cerebrovascular events avoided | \$55,453,391 | = aj * (ak + $\left.\mathrm{al}^{*} \mathrm{am}\right)$ ) |
| ao | Costs avoided due to intervention | \$93,250,100 | $=a b+a i+a n$ |
| Cost-effectiveness Calculation |  |  |  |
| ap | Cost of intervention over lifetime of birth cohort | \$217,955,592 | $=w+x+y$ |
| aq | Costs avoided due to intervention over lifetime of birth cohort | \$93,250,100 | ao |
| ar | QALYs saved | 11,587 | = Table 2, row az |
| as | Cost of intervention over lifetime of birth cohort (1.5\% discount) | \$140,544,975 | Calculated |
| at | Costs avoided due to intervention over lifetime of birth cohort (1.5\% discount) | \$48,541,462 | Calculated |
| au | QALYs saved (1.5\% discount) | 6,032 | Calculated |
| av | CE (\$/QALY saved) | \$15,254 | = (as - at) / au |

[^71]We also modified several major assumptions and recalculated the CE as follows:

- Assume that the proportion of the population with hypertension receiving drug treatment is decreased from $73 \%$ to $68 \%$ (Table 2, row $p$ ): $\mathrm{CE}=\$ 17,584$.
- Assume that the proportion of the population with hypertension receiving drug treatment is increased from $73 \%$ to $78 \%$ (Table 2, row $p$ ): $\mathrm{CE}=\$ 13,219$.
- Assume that the effectiveness of drug treatment in reducing cardiovascular disease events is decreased from $22 \%$ to $17 \%$ (Table 2 , rows $q \& r$ ) and the effectiveness of drug treatment in reducing cerebrovascular disease events is decreased from $41 \%$ to $33 \%$ (Table 2, row $s$ ): $\mathrm{CE}=\$ 24,485$.
- Assume that the effectiveness of drug treatment in reducing cardiovascular disease events is increased from $22 \%$ to $29 \%$ (Table 2, rows $q \& r$ ) and the effectiveness of drug treatment in reducing cerebrovascular disease events is increased from $41 \%$ to $48 \%$ (Table 2, row $s$ ): $\mathrm{CE}=\$ 9,314$.
- Assume that the disutility associated with living with a nonfatal cerebrovascular event is reduced from 0.264 to 0.177 (Table 2, row al): $\mathrm{CE}=\$ 16,036$.
- Assume that the disutility associated with living with a nonfatal cerebrovascular event is increased from 0.264 to 0.350 (Table 2, row al): $\mathrm{CE}=\$ 14,549$.
- Assume that the disutility associated with taking pills for cardiovascular prevention is reduced from 0.0032 to 0.0 (Table 2, row $a x$ ): $\mathrm{CE}=\$ 13,461$.
- Assume that the disutility associated with taking pills for cardiovascular prevention is increased from 0.0032 to 0.0044 (Table 2, row $a x$ ): $\mathrm{CE}=\$ 16,051$.
- Assume that the portion of a 10 -minute office visit for screening and/or monitoring is reduced from $50 \%$ to $33 \%$ (Table 4, rows $f$ \& $g$ ): $\mathrm{CE}=\$ 12,388$.
- Assume that the portion of a 10 -minute office visit for screening and/or monitoring is increased from $50 \%$ to $67 \%$ (Table 4, rows $f \& g$ ): $\mathrm{CE}=\$ 18,114$.


## Summary

| Summary |  |  |  |
| :---: | :---: | :---: | :---: |
|  | $\begin{aligned} & \text { Base } \\ & \text { Case } \end{aligned}$ | Range |  |
| CPB (Potential QALYs Gained) |  |  |  |
| Gap between no current service 'Best in the World' of 73\% |  |  |  |
| 1.5\% Discount Rate | 6,032 | 4,268 | 8,220 |
| 3\% Discount Rate | 3,088 | 2,185 | 4,208 |
| 0\% Discount Rate | 11,587 | 8,199 | 15,792 |
| CE ( $\$ /$ QALY) including patient time costs |  |  |  |
| 1.5\% Discount Rate | \$15,254 | \$9,314 | \$24,485 |
| 3\% Discount Rate | \$22,850 | \$14,890 | \$35,244 |
| 0\% Discount Rate | \$10,760 | \$6,019 | \$18,139 |
| CE (\$/QALY) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | \$9,945 | \$5,421 | \$16,987 |
| 3\% Discount Rate | \$15,814 | \$9,727 | \$25,281 |
| 0\% Discount Rate | \$6,476 | \$2,876 | \$12,086 |

## Screening for Cardiovascular Disease Risk and Treatment with Statins

## United States Preventive Services Task Force Recommendations (2016)

The USPSTF recommends initiating use of low- to moderate-dose statins in adults aged 40 to 75 years without a history of CVD who have 1 or more CVD risk factors (dyslipidemia, diabetes, hypertension, or smoking) and a calculated 10-year CVD event risk of $10 \%$ or greater. ( $B$ recommendation)

Identification of dyslipidemia and calculation of 10-year CVD event risk requires universal lipids screening in adults aged 40-74 years.

The USPSTF recommends using the ACC/AHA Pooled Cohort Equations to calculate the 10-year risk of CVD events. The calculator derived from these equations takes into account age, sex, race, cholesterol levels, blood pressure level, antihypertension treatment, presence of diabetes, and smoking status as risk factors. ${ }^{278}$

The CTFPHC has not completed a recent update due to the review completed by the Canadian Cardiovascular Society (CCS) in 2016. ${ }^{279}$ A number of the CCS recommendations, particularly those associated with screening and primary prevention, are highlighted below.

## Canadian Cardiovascular Society (2016)

## Screening

We recommend that a CV risk assessment be completed every 5 years for men and women aged 40 to 75 years using the modified FRS (Framingham Heart Study Risk Score) or CLEM (Cardiovascular Life Expectancy Model) to guide therapy to reduce major CV events. A risk assessment might also be completed whenever a patient's expected risk status changes. (Strong Recommendation; High Quality Evidence).

## Primary Prevention

We recommend management that does not include statin therapy for individuals at low risk (modified FRS $\leqslant 10 \%$ ) to decrease the risk of CVD events. (Strong Recommendation; High-Quality Evidence).

We recommend management that includes statin therapy for individuals at high risk (modified $F R S \geq 20 \%$ ) to decrease the risk of CVD events. (Strong Recommendation; High-Quality Evidence).

We recommend management that includes statin therapy for individuals at IR (intermediate risk: modified FRS $10 \%-19 \%$ ) with $L D L-C \geq 3.5 \mathrm{mmol} / \mathrm{L}$ to decrease the risk of CVD events. Statin therapy should also be considered for IR persons with LDL-C $<3.5 \mathrm{mmol} / \mathrm{L}$ but with apoB$\geq 1.2 \mathrm{~g} / \mathrm{L}$ or non-HDL-C $\geq 4.3 \mathrm{mmol} / \mathrm{L}$ or in men 50 years of age and older and women 60 years of age and older with $\geq 1$ CV risk factor. (Strong Recommendation; High-Quality Evidence). ${ }^{280}$

[^72]
## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB and CE associated with universal screening for and initiating use of low- to moderate-dose statins in adults aged 40 to 75 years without a history of CVD, who have 1 or more CVD risk factors, and a calculated 10-year CVD event risk of $10 \%$ or greater.

In estimating CPB, we made the following assumptions:

- Based on BC life tables for 2010 to 2012, there are a total of $1,296,348$ life years lived and 6,238 deaths between the ages of 40 and 74 in a BC birth cohort of 40,000 (see Table 1).

| Table 1: Deaths and Years of Life Lived and Lost <br> Between the Ages of 40 and 74 <br> in a British Columbia Birth Cohort of 40,000 |  |  |  |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Age Group | Mean Survival Rate | Individuals in Birth Cohort | Life Years <br> Lived |  | in Birth hort <br> \# | Cardiov <br> Dis \% | Death scular se \# | due to Cerebro Dis \% | scular <br> e <br> \# | Life Expectancy |  | Years Lost <br> Cardio | Cerebro |
| 35-39 | 0.983 | 39,310 |  |  |  |  |  |  |  |  |  |  |  |
| 40-44 | 0.978 | 39,105 | 195,526 | 0.5\% | 205 | 5.9\% | 12 | 3.1\% | 6 | 41.5 | 8,499 | 501 | 263 |
| 45-49 | 0.970 | 38,814 | 194,070 | 0.8\% | 291 | 11.8\% | 34 | 3.8\% | 11 | 36.8 | 10,716 | 1,265 | 407 |
| 50-54 | 0.960 | 38,390 | 191,948 | 1.1\% | 424 | 11.8\% | 50 | 3.8\% | 16 | 32.2 | 13,666 | 1,613 | 519 |
| 55-59 | 0.944 | 37,757 | 188,786 | 1.7\% | 632 | 11.8\% | 75 | 3.8\% | 24 | 27.7 | 17,517 | 2,067 | 666 |
| 60-64 | 0.920 | 36,800 | 183,998 | 2.6\% | 958 | 11.8\% | 113 | 3.8\% | 36 | 23.4 | 22,408 | 2,644 | 851 |
| 65-69 | 0.883 | 35,332 | 176,658 | 4.2\% | 1,468 | 16.7\% | 245 | 6.7\% | 98 | 19.2 | 28,186 | 4,707 | 1,888 |
| 70-74 | 0.827 | 33,072 | 165,362 | 6.8\% | 2,259 | 16.7\% | 377 | 6.7\% | 151 | 15.3 | 34,566 | 5,772 | 2,316 |
| Total |  |  | 1,296,348 |  | 6,238 | 14.5\% | 907 | 5.5\% | 344 |  | 135,558 | 18,569 | 6,911 |

- Based on BC vital statistics data, 59 of 993 (5.9\%) deaths in 25-44 year olds in 2011 were due to cardiovascular disease (ICD-10 codes I00-I51) and 31 of 993 ( $3.1 \%$ ) deaths were due to cerebrovascular disease (ICD-10 codes I60-I69). In 45-64 year olds, 601 of $5,076(11.8 \%)$ deaths were due to cardiovascular disease, and 191 of $5,076(3.8 \%)$ deaths were due to cerebrovascular disease. In 65-84 year olds, 2,248 of $13,481(16.7 \%)$ deaths were due to cardiovascular disease while 905 of 13,481 $(6.7 \%)$ deaths were due to cerebrovascular disease. ${ }^{281}$ This data was used to estimate that approximately 907 ( $14.5 \%$ ) of the 6,238 deaths in the birth cohort would be due to cardiovascular disease and 344 ( $5.5 \%$ ) due to cerebrovascular disease (see Table 1 and Table 3, rows $f, g, h \& i$ ).
- We are not aware of any information which indicates the proportion of adults aged 40 to 74 years in BC who have had a cardiovascular risk assessment within the past five years. Nor are we aware of BC-specific data on the proportion of adults at intermediate or higher risk of CVD who are taking statins over the longer term for primary prevention purposes. Research suggests that $54.8 \%$ of Canadians between the ages of 40 and 79 are at low risk (defined as a mean 10-year risk of a CVD event of less than $10 \%$ ), $14.4 \%$ are at intermediate risk (mean 10 -year risk of a CVD event of $10 \%-19 \%$ ) and $30.9 \%$ are at high risk (mean 10-year risk of a CVD event of $\geq 20 \%)^{282}$ (see Table 2 below and Table 3, row $b$ ).

[^73]| By CVD Risk Status, 2007 to 2011 |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Age |  | Estimat | by CVD | Status | Estima Ris | ted \% b <br> sk Statu | oy CVD |
| Group | Population | Low | Int. | High | Low | Int. | High |
| 20-39 | 8,983,467 | 8,893,999 | 4,335 | 85,133 | 99.0\% | 0.05\% | 0.95\% |
| 40-59 | 9,863,690 | 7,231,730 | 1,014,437 | 1,617,523 | 73.3\% | 10.3\% | 16.4\% |
| 60-79 | 5,186,843 | 1,011,071 | 1,148,828 | 3,026,944 | 19.5\% | 22.1\% | 58.4\% |
| Total | 24,034,000 | 17,136,800 | 2,167,600 | 4,729,600 | 71.3\% | 9.0\% | 19.7\% |
| 40-79 | 15,050,533 | 8,242,801 | 2,163,265 | 4,644,467 | 54.8\% | 14.4\% | 30.9\% |

- In a systematic review for the USPSTF, Chou et al. included 19 randomized control trials (RCTs) with 71,344 participants with a mean age between 51 and 66 years and an average of 4.1 years of follow-up. They conclude that statin therapy is associated with a decreased risk of the following: ${ }^{283}$
- All-cause mortality (RR, 0.86 [ $95 \% \mathrm{CI}, 0.80$ to 0.93 ]) (Table 3, row $y$ )
- Cardiovascular mortality (RR, 0.69 [ $95 \% \mathrm{CI}, 0.54$ to 0.88$]$ )
- Myocardial infarction (RR, 0.64 [ $95 \%$ CI, 0.57 to 0.71 ]) (Table 3, row $a b$ )
- Stroke (RR, 0.71 [ $95 \%$ CI, 0.62 to 0.82 ]) (Table 3, row ae)
- Based on the review for the USPSTF, statin therapy (when compared with a placebo) is not associated with an increased risk of withdrawal due to adverse events, serious adverse events, any cancer, fatal cancer, myalgias or elevated aminotransferase levels, rhabdomyolysis or myopathy, renal dysfunction, cognitive harms or new-onset diabetes following initiation of statin therapy. ${ }^{284}$
- The review for the USPSTF by Chou et al. has been criticized on several fronts. Redberg and Katz note that the review did not exclude studies that included patients taking statins for secondary prevention. ${ }^{285}$ A 2010 review by Ray and colleagues, which included only studies of patients receiving statins for primary prevention, did not find a benefit of statin use and all-cause mortality (RR, $0.91 ; 95 \% \mathrm{CI}$ of 0.83 to 1.01). ${ }^{286}$ In addition, Redberg and Katz note that the most commonly reported side effect of muscle weakness and pain is not included in the review by Chou et al. Clinical trials suggest that statin myopathy occurs in 1-5\% of patients while it may range as high as $20-30 \%$ based on observations in clinical practice. ${ }^{287,288}$
- In a 2016 review of the available evidence on the safety of statin therapy, Collins and colleagues note that " $(\mathrm{t})$ he only serious adverse events that have been shown to be caused by long-term statin therapy - i.e., adverse effects of the statin, are myopathy

[^74](defined as muscle pain or weakness combined with large increases in blood concentrations of creatine kinase), new-onset diabetes mellitus, and, probably, haemorrhagic stroke. Typically, treatment of 10000 patients for 5 years with an effective regimen (e.g., atorvastatin 40 mg daily) would cause about 5 cases of myopathy (one of which might progress, if the statin therapy is not stopped, to the more severe condition of rhabdomyolysis), 50-100 new cases of diabetes, and 5-10 haemorrhagic strokes. However, any adverse impact of these side-effects on major vascular events has already been taken into account in the estimates of the absolute benefits. Statin therapy may cause symptomatic adverse events (e.g., muscle pain or weakness) in up to about $50-100$ patients (i.e., $0.5-1.0 \%$ absolute harm) per 10000 treated for 5 years. However, placebo-controlled randomised trials have shown definitively that almost all of the symptomatic adverse events that are attributed to statin therapy in routine practice are not actually caused by it (i.e., they represent misattribution)....It is, therefore, of concern that exaggerated claims about side-effect rates with statin therapy may be responsible for its under-use among individuals at increased risk of cardiovascular events. For, whereas the rare cases of myopathy and any muscle-related symptoms that are attributed to statin therapy generally resolve rapidly when treatment is stopped, the heart attacks or strokes that may occur if statin therapy is stopped unnecessarily can be devastating."289

- The controversy over side-effects continues, especially regarding muscle problems, as evidenced by the series of letters in the March 18, 2017 issue of The Lancet responding to the Collins et al. review. In our sensitivity analysis, we have included an assumption that $5 \%^{290,291}$ of patients taking statins would develop muscle problems and that their QoL would be reduced by $53 \%{ }^{292}$ during the estimated 3 months it would take for the statin withdrawal and rechallenge process ${ }^{293,294}$ to determine that the muscle problem is associated with the use of statins.
- Other assumptions used in assessing the clinically preventable burden are detailed in the Reference Document.

Based on these assumptions, the CPB associated with universal CVD risk-factor screening and initiating use of low- to moderate-dose statins in adults aged 40 to 74 years without a history of CVD who have 1 or more CVD risk factors and a calculated 10-year CVD event risk of $10 \%$ or greater is 9,370 QALYs (see Table 3, row $a p$ ). This is based on the assumption of moving from no statin use in this intermediate or high risk cohort, to $30 \%$ of this cohort initiating and sustaining statin use.

[^75]| Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
|  | Estimated current status |  |  |
| a | \# of life years lived between the ages of 40-74 in birth cohort | 1,296,348 | Table 1 |
| b | \% of life years at intermediate or high risk | 45.2\% | Table 2 |
| c | \# of life years at intermediate or high risk | 586,371 | = (a*b) |
| d | \% of life years at intermediate or high risk on statins | 30.0\% | See Ref Doc |
| e | \# of life years at intermediate or high risk on statins | 175,911 | = (c*d) |
| f | Total deaths in birth cohort between the ages of 40-74 | 6,238 | Table 1 |
| g | Cardiovascular deaths in birth cohort between the ages of 40-74 | 907 | Table 1 |
| h | Cerebrovascular deaths in birth cohort between the ages of 40-74 | 344 | Table 1 |
| i | Life years lost due to total deaths | 135,558 | Table 1 |
| j | Life years lost per death | 21.7 | = (i/f) |
| k | \# of nonfatal cardiovascular events per fatal event | 5.09 | See Ref Doc |
| I | \# of nonfatal cardiovascular events | 4,615 | = ( $\mathrm{g}^{*}$ k) |
| m | Average age of individual with a cardiovascular event | 68.0 | See Ref Doc |
| n | Life years lived with a nonfatal cardiovascular event | 12.1 | See Ref Doc |
| 0 | Life years lost due to a nonfatal cardiovascular event | 6.3 | See Ref Doc |
| p | QoL reduction living with a nonfatal cardiovascular event (for 1 month) | 0.125 | See Ref Doc |
| q | QALYs lost due to nonfatal cardiovascular events | 29,120 | $=(1 * o)+(1 * p / 12)$ |
| r | Ratio of nonfatal cerebrovascular events per fatal event | 4.58 | See Ref Doc |
| s | \# of nonfatal cerebrovascular events | 1,574 | = ( $\mathrm{r}^{*} \mathrm{~h}$ ) |
| t | Average age of individual with a cerebrovascular event | 72.8 | See Ref Doc |
| u | Life years lived with a nonfatal cerebrovascular event | 9.3 | See Ref Doc |
| v | Life years lost due to a nonfatal cerebrovascular event | 5.5 | See Ref Doc |
| w | QoL reduction living with a nonfatal cerebrovascular event | 0.264 | See Ref Doc |
| x | QALYs lost due to nonfatal cerebrovascular events | 12,525 | $=(s * v)+\left(s^{*} u^{*} w\right)$ |
|  | Benefits if 30\% of intermediate or high risk individuals were on statins |  |  |
| y | \% reduction in all cause mortality associated with statin use | 14\% | $\checkmark$ |
| z | Deaths avoided with statin usage | 262 | $=\left(f{ }^{*}{ }^{*} \mathrm{y}\right)$ |
| aa | QALYs gained due to a reduction in all cause mortality | 5,693 | $=\left(z{ }^{*}\right)^{\text {) }}$ |
| ab | \% reduction in cardiovascular events associated with statin use | 36\% | $\checkmark$ |
| ac | Cardiovascular events avoided with $30 \%$ statin usage | 498 | $=(1 * d * a b)$ |
| ad | QALYs gained due to a reduction in nonfatal cardiovascular events associated with statin use | 3,145 | $=\left(q^{*} d^{*} a b\right)$ |
| ae | \% reduction in cerebrovascular events associated with statin use | 29\% | $\checkmark$ |
| af | Cerebrovascular events avoided with $30 \%$ statin usage | 137 | $=\left(s^{*} d^{*} \mathrm{ae}\right)$ |
| ag | QALYs gained due to a reduction in nonfatal cerebrovascular events associated with statin use | 1,090 | $=(\mathrm{af} * \mathrm{t} * \mathrm{u})$ |
| ah | Total QALYs gained if $\mathbf{3 0 \%}$ of intermediate or high risk individuals were on statins | 9,928 | $=(a a+a d+a g)$ |
|  | Harms if 30\% of intermediate or high risk individuals were on statins |  |  |
| ai | Disutility per year associated with taking pills for cardiovascular prevention | -0.0032 | See Ref Doc |
| aj | Disutility associated with taking pills for cardiovascular prevention | -558 | $=(\mathrm{e}$ * ai ) |
| ak | Proportion of individuals taking statins who experience muscle problems | 0.0\% | $\checkmark$ |
| al | Length of time for muscle problems to be indentified and resolved (in years) | 0.25 | $\checkmark$ |
| am | Disutilty per year associated with muscle problems | -0.53 | $\checkmark$ |
| an | Disutility associated with muscle problems | 0 | $\begin{gathered} \hline \text { Table } 1 *{ }^{*} * \mathrm{ak}^{*} \mathrm{al} \\ * \mathrm{am} \\ \hline \end{gathered}$ |
| ao | QALYs lost if 30\% of intermediate or high risk individuals were on statins | -558 | = (aj + an) |
| ap | Potential QALYs gained, Screening \& Intervention from 0\% to 30\% | 9,370 | $=(\mathrm{ah}+\mathrm{ao}$ ) |

$V=$ Estimates from the literature
For our sensitivity analysis, we modified a number of major assumptions and recalculated the CPB as follows:

- Assume that the QoL reduction associated with a stroke is reduced from 0.264 to 0.177 (Table 3, row w): $\mathrm{CPB}=9,259$.
- Assume that the QoL reduction associated with a stroke is increased from 0.264 to 0.350 (Table 3, row $w$ ): $\mathrm{CPB}=9,480$.
- Assume that decreased risk of all-cause mortality associated with statin therapy is reduced from $14 \%$ to $7 \%$ (Table 3, row $y$ ), the decreased risk of a myocardial infarction is reduced from $36 \%$ to $29 \%$ (Table 3, row $a b$ ) and the decreased risk of stroke is reduced from $29 \%$ to $18 \%$ (Table 3, row $a e$ ): $\mathrm{CPB}=5,499$.
- Assume that decreased risk of all-cause mortality associated with statin therapy is increased from $14 \%$ to $20 \%$ (Table 3, row $y$ ), the decreased risk of a myocardial infarction is increased from $36 \%$ to $43 \%$ (Table 3, row $a b$ ) and the decreased risk of stroke is increased from $29 \%$ to $38 \%$ (Table 3, row $a e$ ): $\mathrm{CPB}=12,760$.
- Assume that the disutility per year associated with taking pills for cardiovascular prevention is reduced from -0.0032 to 0.0 (Table 3, row ai): $\mathrm{CPB}=9,928$.
- Assume that the disutility per year associated with taking pills for cardiovascular prevention is increased from -0.0032 to -0.0044 (Table 3, row ai): $\mathrm{CPB}=9,161$.
- Assume that the percent of life years at intermediate risk on statins is reduced from $30 \%$ to $25 \%$ (Table 3, row $d$ ): CPB $=7,809$.
- Assume that the percent of life years at intermediate risk on statins is increased from $30 \%$ to $40 \%$ (Table 3, row $d$ ): CPB $=12,494$.
- Assume that statin use is associated with muscle problems in 5\% of users (Table 3, row $a k$ ): CPB $=9,259$.


## Modelling Cost-Effectiveness

In this section, we will calculate the CE associated with universal screening for and initiating use of low- to moderate-dose statins in adults aged 40 to 74 years without a history of CVD, who have 1 or more CVD risk factors, and a calculated 10 -year CVD event risk of $10 \%$ or greater.

In estimating CE, we made the following assumptions:

## Cost of Screening for CVD Risk

- The USPSTF recommends using the ACC/AHA Pooled Cohort Equations to calculate the 10 -year risk of CVD events. ${ }^{295}$
- The 2013 ACC/AHA Guideline on the Assessment of Cardiovascular Risk indicate that "it is reasonable to ...estimate 10-year ASCVD risk every 4-6 years in adults 4079 years of age who are free from ASCVD."296
- The ACC-AHA-ASCVD score, however, overestimates the 10-year ASCVD risk. The USPSTF recognizes this. "The reasons for this possible overestimation are still unclear. The Pooled Cohort Equations were derived from prospective cohorts of volunteers from studies conducted in the 1990s and may not be generalizable to a

[^76]more contemporary and diverse patient population seen in current clinical practice. ${ }^{י 297}$

- Cook and Ridker, using the Women's Health Study, found that the ACC-AHAASCVD score overestimated the actual 10-year ASCVD risk in women by $43 \%$ to $90 \%$ in women, depending on their baseline risk. ${ }^{298}$ DeFilippis and colleagues compared the performance of five risk assessment tools in a community-based, sexbalanced, multiethnic cohort. The ACC-AHA-ASCVD score overestimated the $10-$ year ASCVD risk by $78 \%$. They found that the best risk assessment tool was the Reynolds Risk Score. ${ }^{299}$ Rana and co-authors used a large contemporary, multi-ethnic population to assess the ACC-AHA-ASCVD score. They found that the ACC-AHAASCVD score substantially overestimated the actual 5-year ASCVD risk and that this overestimation was similar in both males and females and in four major ethnic groups (black, Asian/Pacific Islander, Hispanic and white). ${ }^{300}$ In a commentary, Nissen notes that "the extent of miscalibration is substantial.... This is not a trivial problem.... Overestimation by the guideline risk equations would likely add millions of Americans to the roles of patients for whom statins are recommended., ${ }^{301}$
- The USPSTF notes that "because the Pooled Cohort Equations lack precision, the risk estimation tool should be used as a starting point to discuss with patients their desire for lifelong statin therapy., ${ }^{302}$
- For screening purposes, we have assumed that $54.8 \%$ of the BC population ages 4075 is at a low risk for CVD (Table 4, row $b$ ), $14.4 \%$ is at an intermediate risk (Table 4 , row $d$ ) and $30.9 \%$ is at a high risk (Table 4, row $f$ ) (see also Table 2).
- We have assumed that the CVD screening would take place once every five years and modified this to once every two years in the sensitivity analysis (Table 4, row $h$ ).
- Completion of a risk assessment includes a clinician visit and a full lipid profile (total cholesterol [TC]; high density lipoprotein cholesterol [HDL-C]; low-density lipoprotein cholesterol [LDL-C], non-HDL-C; and triglycerides [TG]). The full lipid profile costs $\$ 21.31$ (Table 4, row $p$ ). ${ }^{303}$
- We assumed that a 10 -minute office visit would be required for the initial screening. If the results indicate a low risk of CVD, then the follow-up would consist of a phone call to the patient. If the results indicate an intermediate or high risk of CVD, then a

[^77]follow-up visit would be required to discuss the results and the possibility of taking statins (Table 4, row $l$ ).

## Costs of the Intervention

- Adherence with statin therapy in the real world is relatively poor. Benner and colleagues found that early and frequent follow-up by physicians (including cholesterol retesting) improves long-term adherence by approximately 45\% (OR $1.45 ; 95 \% \mathrm{CI}$ of $1.34-1.55) .{ }^{304}$
- Brookhart et al., in a study based on BC data, found that a return to adherence after a period of nonadherence was associated with a return visit to the physician who initially prescribed the statin and a retest of cholesterol. "Our results suggest that continuity of care combined with increased follow-up and cholesterol testing could promote long-term adherence. ${ }^{י 305}$
- Pandya and colleagues estimated one additional physician visit per year for individuals in a disease-free state taking statins (i.e., for primary prevention). ${ }^{306}$
- The BC Guidelines for the primary prevention of cardiovascular disease suggest a follow-up physician visit 4-6 months after the initiation of statin which includes the measuring of lipid levels with a non-HDL-C or an apolipoprotein B (apoB) test, to assess patient adherence to statin therapy and any response to statin therapy, with further follow-ups as clinically indicated. The cost of a non-HDL-C test is $\$ 12.20$ while that of an apoB test is $\$ 16.60 .{ }^{307}$ For modelling purposes, we used the midpoint cost of these two tests (Table 4, row $a b$ ).
- For modelling purposes, we have assumed that $30 \%$ of intermediate and high risk patients would adhere to long-term statin therapy and modified this from $25 \%$ to $40 \%$ in the sensitivity analysis (Table 3, row $d$ ). We further assumed, based on expert input, that one annual follow-up office visit per year (Table 4, row $y$ ) is required for patients on statin therapy, that $100 \%$ of this office visit (Table 4, row $z$ ) is allocated to discussing the statin therapy and that a follow-up lipid test (non-HDL-C or apoB) would be required once every five years (Table 4, row $a a$ ).
- The BC Reference Drug Pricing program fully covers the costs of two statins, atorvastatin and rosuvastatin. ${ }^{308}$ The cost of 10 mg rosuvastatin, taken by the majority of patients, is $\$ 95$ plus four dispensing fees of $\$ 10$ each, for an annual cost of $\$ 135$ (Table 4, row $w$ ). The cost of 80 mg atorvastatin is $\$ 206$ plus four dispensing fees of $\$ 10$ each, for an annual cost of $\$ 246$. We have used this higher cost in the sensitivity analysis.


## Costs Avoided due to the Intervention

- For modelling purposes, we assumed that the acute care costs avoided per death avoided would be $\$ 13,929$ (Table 4, row $a h$ ). This is based on the mix of

[^78]cardiovascular and cerebrovascular deaths in the cohort ( $73 \%$ and $27 \%$, respectively) (see Table 1) and the estimated cost of the acute care phase associated with a fatal myocardial infarction $(\$ 15,536)$ and a fatal stroke $(\$ 9,583)$.

- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the CE associated with universal screening for and initiating use of low- to moderate-dose statins in adults aged 40 to 74 years without a history of CVD, who have 1 or more CVD risk factors, and a calculated 10-year CVD event risk of $10 \%$ or greater is $\$ 3,223 /$ QALY (Table 4, row ay).

Table 4: CE of Universal Screening for and Initiating Use of Statins in Adults Aged 40 to 74 Years with an Intermediate or High Risk of CVD in a Birth Cohort of 40,000

| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | \# of life years lived between the ages of 40-74 in birth cohort | 1,296,348 | Table 1 |
| b | \% of life years at low risk | 54.8\% | Table 2 |
| c | \# of life years at low risk | 709,977 | = (a*b) |
| d | \% of life years at intermediate risk | 14.4\% | Table 2 |
| e | \# of life years at intermediate risk | 186,329 | = (a*d) |
| f | \% of life years at high risk | 30.9\% | Table 2 |
| g | \# of life years at high risk | 400,042 | = (a*f) |
| h | Annual frequency of screening | 0.20 | $\checkmark$ |
| i | Adherence with offers to receive screening | 48\% | See Ref Doc |
| j | Total \# of screens in birth cohort | 124,449 | = (a*h*i) |
|  | Estimated cost of screening |  |  |
| k | Number of office visits associated with screening - low risk | 1.0 | Expert Opinion |
| I | Number of office visits associated with screening - medium or high risk | 2.0 | Expert Opinion |
| m | Cost of 10-minute office visit | \$34.85 | See Ref Doc |
| n | Cost of a follow-up phone call | \$15.00 | See Ref Doc |
| o | Cost to measure cholesterol | \$21.31 | $\checkmark$ |
| p | Health care costs of screening - low risk | \$4,850,111 | $=(\mathrm{j} * \mathrm{~b}) * \mathrm{k} *(\mathrm{~m}+\mathrm{n}+\mathrm{o})$ |
| q | Health care costs of screening - intermediate and high risk | \$5,123,096 | $\begin{gathered} =((\mathrm{d}+\mathrm{f}) * \mathrm{j} * \mathrm{I}) *(\mathrm{~m}+ \\ (\mathrm{o} / 2)) \end{gathered}$ |
| r | Patient time required / office visit (hours) | 2.0 | $\checkmark$ |
| s | Value of patient time (per hour) | \$29.69 | $\checkmark$ |
| t | Value of patient time and travel for screening | \$7,389,806 | $=(\mathrm{j} * \mathrm{r} * \mathrm{~s})$ |
|  | Estimated cost of intervention |  |  |
| u | Adherence with long-term statin therapy in intermediate and high risk cohort | 30\% | Table 3, row d |
| v | Years on statin therapy | 175,911 | $=(\mathrm{e}+\mathrm{g}) * \mathrm{u}$ |
| w | Cost of statin therapy / year | \$135 | $\checkmark$ |
| X | Cost of statin therapy | \$23,748,009 | $=\left(v^{*} \mathrm{w}\right)$ |
| y | \# of follow-up office visits per year re: statin therapy | 1.0 | Expert Opinion |
| z | Portion of 10-minute office visit for follow-up re: statin therapy | 100\% | Expert Opinion |
| aa | \# of lab tests (non-HDL-C or apoB) per year re: statin therapy | 0.2 | Expert Opinion |
| ab | Cost per lab test | \$14.40 | $v$ |
| ac | Follow-up costs | \$6,637,129 | $\begin{gathered} =\left(v^{*} y^{*} z * m\right)+(v * \\ \text { aa *ab) } \end{gathered}$ |
| ad | Value of patient time and travel for intervention | \$10,445,606 | $=(v * y * s * r)$ |
|  | Estimated costs avoided due to intervention |  |  |
| ae | \# of deaths avoided | 262.0 | Table 3, row z |
| af | \# of nonfatal cardiovascular events avoided | 498.4 | Table 3, row ac |
| ag | \# of nonfatal cerebrovascular events avoided | 136.9 | Table 3, row af |
| ah | Acute care costs avoided per avoided death | -\$13,929 | See Ref Doc |
| ai | First year costs avoided per nonfatal cardiovascular event avoided | -\$33,934 | See Ref Doc |
| aj | First year costs avoided per nonfatal cerebrovascular event avoided | -\$21,139 | See Ref Doc |
| ak | First-year acute care costs avoided | -\$23,455,536 | $\begin{gathered} =(\mathrm{ae} * a h)+(a f * a i)+ \\ (a g * a j) \end{gathered}$ |
| al | Post-first-year annual costs avoided for nonfatal cardiovascular events avoided | -\$2,278 | See Ref Doc |
| am | Number of years for which the costs are avoided | 12.1 | See Ref Doc |
| an | Post-first-year costs avoided for nonfatal cardiovascular events avoided | -\$13,736,935 | = af * $\mathrm{am}^{*} \mathrm{al}$ ) |
| ao | Post-first-year annual costs avoided for nonfatal cerebrovascular events avoided | -\$6,246 | See Ref Doc |
| ap | Number of years for which the costs are avoided | 9.3 | See Ref Doc |
| aq | Post-first-year costs avoided for nonfatal cerebrovascular events avoided | -\$7,954,795 | = (ag * ap * ao) |
| ar | Costs avoided due to intervention | -\$45,147,265 | = $\mathrm{ak}+\mathrm{an}+\mathrm{aq}$ |
|  | CE Calculation |  |  |
| as | Cost of intervention over lifetime of birth cohort | \$58,193,757 | $=p+q+t+x+a c+a d$ |
| at | Costs avoided due to intervention over lifetime of birth cohort | -\$45,147,265 | = ar |
| au | QALYs saved | 9,370 | Table 3, row ap |
| av | Cost of intervention over lifetime of birth cohort (1.5\% discount) | \$45,893,093 | Calculated |
| aw | Costs avoided due to intervention over lifetime of birth cohort (1.5\% discount) | -\$28,135,568 | Calculated |
| ax | QALYs saved (1.5\% discount) | 5,510 | Calculated |
| ay | CE (\$/QALY saved) | \$3,223 | = (av + aw) / ax |

For our sensitivity analysis, we modified a number of major assumptions and recalculated the CE as follows:

- Assume that the QoL reduction associated with a stroke is reduced from 0.264 to 0.177 (Table 3, row w): $\mathrm{CE}=\$ 3,261$.
- Assume that the QoL reduction associated with a stroke is increased from 0.264 to 0.350 (Table 3, row $w$ ): $\mathrm{CE}=\$ 3,186$.
- Assume that decreased risk of all-cause mortality associated with statin therapy is reduced from $14 \%$ to $7 \%$ (Table 3, row $y$ ), the decreased risk of a myocardial infarction is reduced from $36 \%$ to $29 \%$ (Table 3, row $a b$ ) and the decreased risk of stroke is reduced from $29 \%$ to $18 \%$ (Table 3, row $a e$ ): $\mathrm{CE}=\$ 7,849$.
- Assume that decreased risk of all-cause mortality associated with statin therapy is increased from $14 \%$ to $20 \%$ (Table 3, row $y$ ), the decreased risk of a myocardial infarction is increased from $36 \%$ to $43 \%$ (Table 3, row $a b$ ) and the decreased risk of stroke is increased from $29 \%$ to $38 \%$ (Table 3, row ae): $\mathrm{CE}=\$ 1,458$.
- Assume that the disutility per year associated with taking pills for cardiovascular prevention is reduced from -0.0032 to 0.0 (Table 3, row ai): $\mathrm{CE}=\$ 2,996$.
- Assume that the disutility per year associated with taking pills for cardiovascular prevention is increased from -0.0032 to -0.0044 (Table 3, row ai): $\mathrm{CE}=\$ 3,317$.
- Assume that the percent of life years at intermediate risk on statins is reduced from $30 \%$ to $25 \%$ (Table 3, row $d$ ): $\mathrm{CE}=\$ 3,720$.
- Assume that the percent of life years at intermediate risk on statins is increased from $30 \%$ to $40 \%$ (Table 3, row $d$ ): $\mathrm{CE}=\$ 2,601$.
- Assume that statin use is associated with muscle problems in 5\% of users (Table 3, row $a k$ ): $\mathrm{CE}=\$ 3,272$.
- Assume that the annual frequency of screening is increased from once every five years to once every two years (Table 4 , row $i$ ): $\mathrm{CE}=\$ 6,950$.
- Assume that the cost of statin therapy in increased from $\$ 135$ per year to $\$ 246$ per year (Table 4, row $w$ ): $\mathrm{CE}=\$ 6,017$.
- Assume that the first-year costs avoided following a nonfatal cerebrovascular are decreased from $\$ 21,139$ to $\$ 16,642$ (Table 4 , row $a j$ ) and the post-first-year annual costs avoided decreased from $\$ 6,246$ to $\$ 4,930$ (Table 4, row $a o$ ): $\mathrm{CE}=\$ 3,471$.
- Assume that the first-year costs avoided following a nonfatal cerebrovascular are increased from $\$ 21,139$ to $\$ 25,635$ (Table 4 , row $a j$ ) and the post-first-year annual costs avoided increased from \$6,246 to \$7,562 (Table 4, row ao): CE = \$2,974.


## Summary

| Statins in Adults aged 40 to 74 years with an Intermediate or High Risk of CVD in a Birth Cohort of 40,000 Summary |  |  |  |
| :---: | :---: | :---: | :---: |
|  | Base <br> Case | Range |  |
| CPB (Potential QALYs Gained) |  |  |  |
| Gap between No Service and 'Best in the World' (30\%) |  |  |  |
| 1.5\% Discount Rate | 5,510 | 3,204 | 7,531 |
| 3\% Discount Rate | 3,144 | 1,800 | 4,322 |
| 0\% Discount Rate | 9,370 | 5,499 | 12,760 |
| CE (\$/QALY) including patient time costs |  |  |  |
| 1.5\% Discount Rate | \$3,223 | \$1,458 | \$7,849 |
| 3\% Discount Rate | \$6,222 | \$3,567 | \$13,376 |
| 0\% Discount Rate | \$1,392 | \$169 | \$4,537 |
| CE (\$/QALY) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | \$1,174 | -\$409 | \$3,459 |
| 3\% Discount Rate | \$2,634 | \$958 | \$7,109 |
| 0\% Discount Rate | -\$511 | -\$1,229 | \$1,293 |

## Screening for Type 2 Diabetes Mellitus

## Canadian Task Force on Preventive Health Care (2012)

The CTFPHC suggests a two-phase approach to screening. ${ }^{309}$ First, it recommends screening all adults ages 18 and older using a validated risk calculator such as FINDRISC (Finnish Diabetes Risk Score) or CANRISK (Canadian Diabetes Risk Assessment Questionnaire). This first level of screening should be completed once every 3-5 years. Those with a FINDRISC score of 15 to 20 are considered to be at high risk of diabetes (an individual's risk of developing type 2 diabetes within 10 years is between $33 \%$ and $49 \%$ ) and those with a score greater than 21 are at very high risk (an individual's risk of developing diabetes within 10 years is $50 \%$ or higher). The second phase of screening involves either an A1C, fasting glucose or oral glucose tolerance test. The CTFPHC recommends the use of the A1C test given its "convenience for patients." Individuals at high risk are to be screened every 3-5 years while individuals at very high risk are to be screened every year. The CTFPHC considers these recommendations to be "weak" based on "low-quality evidence". ${ }^{310}$

## United States Preventive Services Task Force Recommendations (2015)

The USPSTF recommends screening for abnormal blood glucose in all adults ages 40 to 70 who are overweight or obese as part of a cardiovascular risk assessment. This recommendation receives a " $B$ " grade from the USPSTF. ${ }^{311}$

## Modelling the Clinically Preventable Burden

In this section, we model the CPB associated with the two-phase approach to screening for type 2 diabetes, recommended by the CTFPHC, in a British Columbia birth cohort of 40,000.

In modelling CPB, we made the following assumptions:

- $35 \%$ of the population aged 40 or older would have a FINDRISC score of 15-19 (high risk) and $10 \%$ would have a score of $20+$ (very high risk) (see Table 1 and 2 below). ${ }^{312}$
- Detailed information on the prevalence of diagnosed diabetes in Canada in 2008/09 by age group and sex is provided by the CTFPHC. Overall, rates for Canadian females and males were $6.4 \%$ and $7.2 \%$, respectively. ${ }^{313}$ Rates of diagnosed diabetes in British Columbia in 2007/08 were $6.0 \%$ for females and $6.9 \%$ for males. ${ }^{314}$ This data was not stratified by age. In estimating the age and sex specific prevalence rates for diagnosed diabetes in BC , we adjusted the Canadian age and sex specific rates downwards by the difference between the Canadian and British Columbian rates (see Figure 1).

[^79]

- Estimates of the proportion of diabetes cases that are undiagnosed by age group and sex are as follows: ${ }^{315}$

| Age Group | Males | Females |
| :--- | :---: | :---: |
| $40-49$ | $44 \%$ | $24 \%$ |
| $\mathbf{5 0 - 5 9}$ | $21 \%$ | $15 \%$ |
| $\mathbf{6 0 - 6 9}$ | $17 \%$ | $16 \%$ |
| $\mathbf{7 0 - 7 9}$ | $19 \%$ | $14 \%$ |
| $\mathbf{8 0 +}$ | $16 \%$ | $14 \%$ |

- A total of 798,605 years would be lived by males from age $40-89$ in a BC birth cohort of 40,000 (see Table 1). The equivalent number for females would be 857,481 (see Table 2). Among males, 279,512 of these years would be spent at high risk for type 2 diabetes, and 79,861 would be spent at very high risk. Among females, 300,118 would be spent at high risk and 85,748 at very high risk.

[^80]Table 1: Prevalence and Increased Risk for Type 2 Diabetes in a Male Birth Cohort of 20,000

|  | Mean Survival | Individuals in Birth | Years of Life in Birth |  | mated SC Status Very High | Prev <br> Diagn <br> \% | lence osed | of Diab Undia \% | nosed | Years | ife with tes |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Group | Rate |  |  | High |  | \% | \# | \% | \# | Diagnos | diagnosed |
| 40-44 | 0.972 | 19,442 | 97,211 | 34,024 | 9,721 | 3.9\% | 764 | 1.7\% | 336 | 3,820 | 1,681 |
| 45-49 | 0.963 | 19,263 | 96,314 | 33,710 | 9,631 | 5.9\% | 1,145 | 2.6\% | 504 | 5,723 | 2,518 |
| 50-54 | 0.950 | 19,003 | 95,017 | 33,256 | 9,502 | 9.1\% | 1,730 | 1.9\% | 363 | 8,651 | 1,817 |
| 55-59 | 0.931 | 18,619 | 93,095 | 32,583 | 9,310 | 13.4\% | 2,498 | 2.8\% | 525 | 12,490 | 2,623 |
| 60-64 | 0.902 | 18,041 | 90,204 | 31,571 | 9,020 | 18.3\% | 3,302 | 3.1\% | 561 | 16,511 | 2,807 |
| 65-69 | 0.858 | 17,164 | 85,820 | 30,037 | 8,582 | 22.7\% | 3,898 | 3.9\% | 663 | 19,492 | 3,314 |
| 70-74 | 0.792 | 15,837 | 79,183 | 27,714 | 7,918 | 26.0\% | 4,113 | 4.9\% | 781 | 20,564 | 3,907 |
| 75-79 | 0.693 | 13,861 | 69,305 | 24,257 | 6,931 | 27.3\% | 3,786 | 5.2\% | 719 | 18,929 | 3,596 |
| 80-84 | 0.553 | 11,053 | 55,266 | 19,343 | 5,527 | 24.4\% | 2,697 | 3.9\% | 432 | 13,485 | 2,158 |
| 85-89 | 0.372 | 7,438 | 37,190 | 13,017 | 3,719 | 24.4\% | 1,815 | 3.9\% | 290 | 9,074 | 1,452 |
| Total Ages 40-89 |  | 798,605 |  | 279,512 79,861 |  |  |  |  |  | 128,739 | 25,872 |


|  | Mean | Individuals | Years of | Estimated FINDRISC Status |  | Prevalence of Diabetes <br> Diagnosed Undiagnosed |  |  |  | Years of Life with Diabetes Diagnosed Undiagnosed |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Age | Survival | in Birth | Life in Birth |  |  |  |  |  |  |  |  |
| Group | Rate | Cohort | Cohort | High | Very High | \% | \# | \% | \# |  |  |
| 40-44 | 0.984 | 19,672 | 98,358 | 34,425 | 9,836 | 3.5\% | 682 | 0.8\% | 164 | 3,412 | 819 |
| 45-49 | 0.978 | 19,560 | 97,800 | 34,230 | 9,780 | 4.8\% | 935 | 1.1\% | 224 | 4,676 | 1,122 |
| 50-54 | 0.970 | 19,395 | 96,977 | 33,942 | 9,698 | 6.9\% | 1,346 | 1.0\% | 202 | 6,728 | 1,009 |
| 55-59 | 0.957 | 19,150 | 95,748 | 33,512 | 9,575 | 10.0\% | 1,921 | 1.5\% | 288 | 9,605 | 1,441 |
| 60-64 | 0.939 | 18,774 | 93,872 | 32,855 | 9,387 | 13.3\% | 2,499 | 2.1\% | 400 | 12,497 | 1,999 |
| 65-69 | 0.909 | 18,190 | 90,948 | 31,832 | 9,095 | 16.7\% | 3,035 | 2.7\% | 486 | 15,177 | 2,428 |
| 70-74 | 0.863 | 17,265 | 86,325 | 30,214 | 8,633 | 20.0\% | 3,448 | 2.8\% | 483 | 17,238 | 2,413 |
| 75-79 | 0.790 | 15,799 | 78,995 | 27,648 | 7,900 | 21.7\% | 3,421 | 3.0\% | 479 | 17,107 | 2,395 |
| 80-84 | 0.676 | 13,517 | 67,587 | 23,655 | 6,759 | 20.3\% | 2,744 | 2.8\% | 384 | 13,720 | 1,921 |
| 85-89 | 0.509 | 10,174 | 50,871 | 17,805 | 5,087 | 20.3\% | 2,065 | 2.8\% | 289 | 10,327 | 1,446 |
| Total Ag | es 40-89 |  | 857,481 | 300,118 | 85,748 |  |  |  |  | 110,486 | 16,994 |

- Screening of the entire target population every 3-5 years starting at age 40 is associated with the following benefits over a 50 year period: ${ }^{316}$
$\checkmark 5.2$ (range of $2.7-7.5$ ) myocardial infarction events prevented per 1,000 people screened (Table 3, row $d$ ).
$\checkmark 8.0$ (range of $6.2-9.5$ ) microvascular events (foot amputations/ulcers, end-stage renal disease or blindness) prevented per 1,000 people screened (Table 3, row $h$ ).
$\checkmark 3.2$ (range of $1.0-5.8$ ) premature deaths prevented per 1,000 people screened (Table 3, row $l$ ).
- We have assumed that each event would be prevented, on average, half way through the 50 year follow-up period.
- A myocardial infarction reduces a person's quality of life by $12.6 \%$ for a period of one month or a 0.0105 reduction in QoL (Table 3, row $f$ ).

[^81]- End-stage renal disease (ESRD) reduces a person's quality of life by $20 \%$, foot amputation by $10.5 \%$ and blindness by $16 \% .{ }^{317}$ For microvascular events prevented, we assumed an overall quality of life reduction of $15.8 \%$ based on a 40:33:27 distribution for incidence of ESRD, foot amputation or blindness (Table 3, row $j$ ). ${ }^{318}$
- Other assumptions used in assessing the clinically preventable burden are detailed in the Reference Document.

Based on these assumptions, the CPB associated with screening for type 2 diabetes is 3,494 QALYs (Table 3, row $p$ ).

| Table 3: CPB of Screening for Type 2 Diabetes in a Birth Cohort of 40,000 |  |  |  |
| :---: | :--- | :---: | :---: |
| Row <br> Label | Variable | Base Case | Data Source |
| a | Individuals in birth cohort at age 40 | 39,114 | Tables 1 and 2 |
| b | Adherence with screening | $80 \%$ | Ref Doc |
| c | Individuals screened | 31,291 | $=\mathrm{a} * \mathrm{~b}$ |
|  | Benefits Associated with Screening |  |  |
| d | Myocardial infarction events prevented / 1,000 people screened | 5.2 | V |
| e | Myocardial infarction events prevented | 163 | $=(\mathrm{c} / 1,000)^{*} \mathrm{~d}$ |
| f | Quality of life adjustment per myocardial event | 0.0105 | Ref Doc |
| g | QALYs gained | 1.7 | $=e^{*} \mathrm{f}$ |
| h | Microvascular events prevented / 1,000 people screened | 8.0 | V |
| i | Microvascular events prevented | 250 | $=(\mathrm{c} / 1,000)^{*} \mathrm{~h}$ |
| j | Quality of life adjustment | $15.8 \%$ | V |
| k | QALYs gained | 989 | $=\mathrm{i} * 25 * \mathrm{j}$ |
| l | Premature deaths averted / 1,000 people screened | 3.2 | V |
| m | Premature deaths averted | 100 | $=(\mathrm{c} / 1,000)^{*} \mathrm{~m}$ |
| n | Life-years gained / death averted | 25 | V |
| o | Life-years gained | 2,503 | $=\mathrm{m} * \mathrm{n}$ |
| p | Potential QALYs gained, Screening increasing from 0\% to 80\% | 3,494 | $=\mathrm{g}+\mathrm{k}+\mathrm{o}$ |

V = Estimates from the literature
We also modified a major assumption and recalculated the CPB as follows:

- Assume the number of myocardial infarction events prevented per 1,000 people screened is reduced from 5.2 to 2.7 (Table 3, row $d$ ), the number of microvascular events prevented per 1,000 people screened is reduced from 8.0 to 6.2 (Table 3, row $h$ ) and the number of premature deaths prevented per 1,000 people screened is reduced from 3.2 to 1.0 (Table 3, row $l$ ): $\mathrm{CPB}=1,549$ QALYs.
- Assume the number of myocardial infarction events prevented per 1,000 people screened is increased from 5.2 to 7.5 (Table 3, row $d$ ), the number of microvascular events prevented per 1,000 people screened is increased from 8.0 to 9.5 (Table 3, row $h$ ) and the number of premature deaths prevented per 1,000 people screened is increased from 3.2 to 5.8 (Table 3, row $l$ ): CPB = 5,714 QALYs.

[^82]
## Modelling Cost-Effectiveness

In this section, we model the CE associated with the two-phase approach to screening for type 2 diabetes, recommended by the CTFPHC, in a British Columbia birth cohort of 40,000.

In modelling CE, we made the following assumptions:

- Laboratory screening tests - The cost of an A1C test (MSP fee item 91745) in BC is $\$ 6.09$ (Table 4, row $l$ ). ${ }^{319}$
- The typical event (i.e., first year) cost for an acute myocardial infarction is $\$ 33,934$, with annual costs thereafter of $\$ 1,193$ (see Reference Document).
- The annual costs for blindness are $\$ 2,330$ (see Reference Document).
- The annual costs for end-stage renal disease are $\$ 86,278$ (see Reference Document).
- The typical event cost for a lower extremity amputation is $\$ 33,642$ with annual costs thereafter of \$1,396 (see Reference Document).
- We have assumed that each event and the resulting costs would be prevented, on average, half way through the 50 year follow-up period.
- Screening detects diabetes, on average, 5.3 years earlier than no screening. ${ }^{320}$
- Average costs avoided per acute myocardial infarction event would therefore be $\$ 6,323(\$ 1,193 * 5.3)$ (Table 4, row $t$ ).
- For microvascular events prevented, we assumed a 40:33:27 distribution for ESRD, foot amputation or blindness. ${ }^{321}$ Average costs avoided per microvascular event would therefore be $\$ 188,685$ (Table 4, row $w$ ).
- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the CE associated with screening for type 2 diabetes is $-\$ 3,121$ per QALY (Table 4, row ee).

[^83]| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Individuals in birth cohort at age 40 | 39,114 | Table 3, row a |
| b | Life years at increased risk for diabetes | 1,656,086 | Tables 1 and 2 |
| c | Life years at high risk for diabetes | 579,630 | Tables 1 and 2 |
| d | Life years at very high risk for diabetes | 165,609 | Tables 1 and 2 |
|  | Costs of intervention |  |  |
| e | Frequency of screening with FINDRISC/CANRISK (every x years) | 4 | $\checkmark$ |
| f | Total number of screens with FINDRISC/CANRISK (100\% adherence) | 414,022 | $=\mathrm{b} / \mathrm{e}$ |
| g | Adherence with screening | 80\% | Ref Doc |
| h | Cost of 10-minute office visit | \$34.85 | Ref Doc |
| i | Value of patient time and travel for office visit | \$59.38 | Ref Doc |
| j | Portion of 10-minute office visit for screen | 50\% | Ref Doc |
| k | Cost of screening with FINDRISC/CANRISK | \$15,605,298 | $=(\mathrm{f} * \mathrm{~g}) *(\mathrm{~h}+\mathrm{i}) * \mathrm{j}$ |
| I | Lab cost of A1C test | \$6.09 | $\checkmark$ |
| m | Value of patient time and travel for lab test | \$29.69 | Ref Doc |
| n | Frequency of lab testing for high risk patients (every x years) | 4 | $\checkmark$ |
| 0 | \# of lab tests high risk patients | 115,926 | $=(\mathrm{c} / \mathrm{n}) * \mathrm{~g}$ |
| p | Frequency of lab testing for very high risk patients (every x years) | 1 | $v$ |
| q | \# of lab tests for very high risk patients | 132,487 | $=d^{*} p^{*} g$ |
| $r$ | Cost of lab testing | \$20,592,187 | $\begin{gathered} =((\mathrm{o}+\mathrm{q}) *(\mathrm{l}+\mathrm{m}))+((\mathrm{o}+ \\ \mathrm{q}) *(\mathrm{~h}+\mathrm{i}) * \mathrm{j}) \end{gathered}$ |
|  | Cost avoided |  |  |
| 5 | Myocardial infarction events prevented | 163 | Table 3, row e |
| t | Cost avoided per event avoided | \$6,323 | $\checkmark$ |
| u | Total costs avoided | \$1,028,837 | = ${ }^{*}$ t |
| v | Microvascular events prevented | 250 | Table 3, row i |
| w | Cost avoided per event avoided | \$188,685 | $\checkmark$ |
| X | Total costs avoided | \$47,233,248 | $=v^{*} \mathrm{w}$ |
|  | CE calculation |  |  |
| y | Cost of intervention over lifetime of birth cohort | \$36,197,486 | $=\mathrm{k}+\mathrm{r}$ |
| z | Costs avoided | \$48,262,085 | = $\mathrm{u}+\mathrm{x}$ |
| aa | QALYs saved | 3,494 | Table 3, row p |
| bb | Cost of intervention over lifetime of birth cohort (1.5\% discount) | \$25,566,103 | Calculated |
| cc | Costs avoided (1.5\% discount) | \$31,908,799 | Calculated |
| dd | QALYs saved (1.5\% discount) | 2,032 | Calculated |
| ee | CE (\$/QALY saved) | -\$3,121 | $=(\mathrm{bb}-\mathrm{cc}) / \mathrm{dd}$ |

V = Estimates from the literature
We also modified several major assumptions and recalculated the cost per QALY as follows:

- Assume the number of myocardial infarction events prevented per 1,000 people screened is reduced from 5.2 to 2.7 (Table 3, row $d$ ), the number of microvascular events prevented per 1,000 people screened is reduced from 8.0 to 6.2 (Table 3 , row $h$ ) and the number of premature deaths prevented per 1,000 people screened is reduced from 3.2 to $1.0($ Table 3 , row $l)$ : $\mathrm{CE}=\$ 1,121$
- Assume the number of myocardial infarction events prevented per 1,000 people screened is increased from 5.2 to 7.5 (Table 3, row $d$ ), the number of microvascular events prevented per 1,000 people screened is increased from 8.0 to 9.5 (Table 3, row $h$ ) and the number of premature deaths prevented per 1,000 people screened is increased from 3.2 to 5.8 (Table 3, row $l$ ): $\mathrm{CE}=-\$ 3,761$
- Assume the frequency of screening with FINDRISC is increased from every 4 years to every 3 years (Table 4, row $e$ ): $\mathrm{CE}=-\$ 1,313$
- Assume the frequency of screening with FINDRISC is decreased from every 4 years to every 5 years (Table 4, row $e$ ): $\mathrm{CE}=-\$ 4,206$
- Assume that the portion of a 10 -minute office visit for the assessment of patient risk is reduced from $50 \%$ to $33 \%$ (Table 4 , row $j$ ): $\mathrm{CE}=-\$ 6,348$
- Assume that the portion of a 10 -minute office visit for the assessment of patient risk is increased from $50 \%$ to $67 \%$ (Table 4, row $j$ ): $\mathrm{CE}=\$ 106$


## Summary

| Table 5: Screening for Type 2 Diabetes in a Birth Cohort <br> of 40,000 <br> Summary |  |  |  |
| :---: | :---: | :---: | :---: |
|  | $\begin{aligned} & \text { Base } \\ & \text { Case } \end{aligned}$ | Range |  |
| CPB (Potential QALYs Gained) |  |  |  |
| Gap between No Service and 'Best in the World' (80\%) |  |  |  |
| 1.5\% Discount Rate | 2,032 | 901 | 3,324 |
| 3\% Discount Rate | 1,162 | 515 | 1,901 |
| 0\% Discount Rate | 3,494 | 1,459 | 5,714 |
| CE (\$/QALY) including patient time costs |  |  |  |
| 1.5\% Discount Rate | -\$3,121 | -\$6,348 | \$1,121 |
| 3\% Discount Rate | -\$1,879 | -\$5,990 | \$5,067 |
| 0\% Discount Rate | -\$3,453 | -\$6,111 | -\$608 |
| CE ( $\$ / \mathrm{QALY}$ ) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | -\$11,666 | -\$12,859 | -\$18,145 |
| 3\% Discount Rate | -\$12,764 | -\$14,285 | -\$19,477 |
| 0\% Discount Rate | -\$10,490 | -\$11,473 | -\$16,475 |

## Screening for Depression in the General Adult Population

## Canadian Task Force on Preventive Health Care (2013) ${ }^{322}$

Recommendations on screening for depression in primary care settings are provided for people 18 years of age or older who present at a primary care setting with no apparent symptoms of depression. These recommendations do not apply to people with known depression, with a history of depression or who are receiving treatment for depression.

For adults at average risk of depression, ${ }^{323}$ we recommend not routinely screening for depression. (Weak recommendation; very-low-quality evidence)

For adults in subgroups of the population who may be at increased risk of depression, ${ }^{324}$ we recommend not routinely screening for depression. ${ }^{325}$ (Weak recommendation; very-low-quality evidence)

Note that the 2013 recommendations from the CTFPHC are different than their 2005 recommendations. In 2005, the CTFPHC recommended the following:

There is fair evidence to recommend screening adults in the general population for depression in primary care settings that have integrated programs for feedback to patients and access to case management or mental health care (grade B recommendation).

This is insufficient evidence to recommend for or against screening adults in the general; population for depression in primary care settings where effective follow-up and treatment are not available (grade I recommendation). ${ }^{326}$

United States Preventive Services Task Force Recommendations (2016)
The USPSTF recommends screening for depression in the general adult population, including pregnant and postpartum women. Screening should be implemented with adequate systems in place to ensure accurate diagnosis, effective treatment, and appropriate follow-up. (B recommendation) ${ }^{327}$

[^84]
## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with screening non-pregnant adults ages 18 and older for depression when staff-assisted depression care supports are in place to assure accurate diagnosis, effective treatment, and follow-up in a BC birth cohort of 40,000 .
In modelling CPB, we made the following assumptions:

- In BC in 2012, $4.6 \%$ of the population aged $\geq 15$ had a major depressive episode (MDE) within the previous 12 months ( $4.0 \%$ for males and $5.2 \%$ for females). The lifetime risk for an MDE is $11.6 \%$ ( $9.3 \%$ for males and $13.9 \%$ for females). ${ }^{328}$
- The average duration of a first episode of a MDE is 71.0 weeks ( 1.37 years) for males and 75.9 weeks ( 1.46 years) for females (see Table 1 ). ${ }^{329}$

| Table 1: Length of First Major Depression Episode British Columbia in 2012 by Sex |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | Males |  |  |  | Femal |  |
| Episode duration (as reported) | Episode duration (in weeks) | Number | Percent | Cumulative percent | Episode duration (in weeks) | Number | Percent | Cumulative percent |
| 2 weeks | 2.0 | 8 | 6.1\% | 6.1\% | 2.0 | 10 | 4.0\% | 4.0\% |
| 3 weeks | 3.0 | 5 | 3.8\% | 9.9\% | 3.0 | 4 | 1.6\% | 5.6\% |
| 1 month | 4.3 | 11 | 8.4\% | 18.3\% | 4.3 | 33 | 13.1\% | 18.7\% |
| 2 months | 8.7 | 9 | 6.9\% | 25.2\% | 8.7 | 19 | 7.6\% | 26.3\% |
| 3 months | 13.0 | 16 | 12.2\% | 37.4\% | 13.0 | 17 | 6.8\% | 33.1\% |
| 4 months | 17.3 | 5 | 3.8\% | 41.2\% | 17.3 | 7 | 2.8\% | 35.9\% |
| 5 months | 21.7 | 1 | 0.8\% | 42.0\% | 21.7 | 9 | 3.6\% | 39.4\% |
| 6 months | 26.0 | 15 | 11.5\% | 53.4\% | 26.0 | 31 | 12.4\% | 51.8\% |
| 7 months | 30.3 | 1 | 0.8\% | 54.2\% | 30.3 | 0 | 0.0\% | 51.8\% |
| 8 months | 34.7 | 4 | 3.1\% | 57.3\% | 34.7 | 5 | 2.0\% | 53.8\% |
| 9 months | 39.0 | 2 | 1.5\% | 58.8\% | 39.0 | 4 | 1.6\% | 55.4\% |
| 10 months | 43.3 | 3 | 2.3\% | 61.1\% | 43.3 | 2 | 0.8\% | 56.2\% |
| 11 months | 47.7 | 0 | 0.0\% | 61.1\% | 47.7 | 2 | 0.8\% | 57.0\% |
| 1 year | 52.0 | 17 | 13.0\% | 74.0\% | 52.0 | 40 | 15.9\% | 72.9\% |
| 2 years* | 156.0 | 25 | 19.1\% | 93.1\% | 156.0 | 48 | 19.1\% | 92.0\% |
| 5 years* | 364.0 | 9 | 6.9\% | 100.0\% | 364.0 | 20 | 8.0\% | 100.0\% |
| Total | 71.0 | 131 |  |  | 75.9 | 251 |  |  |
| * Reponses were categorized as ranges: 2-4 years and 5 or more years. Assume a duration of 3 years for the first category and 7 years for the second. |  |  |  |  |  |  |  |  |

- Depression is a highly recurrent disorder. ${ }^{330}$ On average, half of individuals experiencing at least one MDE during their lifetime will experience between 5-9 recurrent episodes during their lifetime.${ }^{331,332,333}$ For modelling purposes, we assumed

[^85]that $50 \%$ of individuals experiencing an initial MDE would experience 7 recurrent episodes during their lifetime.

- The above information was used to generate the expected number of life years lived with depression by males and females in a BC birth cohort of 40,000 . For males, an estimated $0.95 \%$ of life years lived between the age of 18 and death would be with diagnosed depression (see Tables 2). For females, an estimated $1.33 \%$ of life years lived between the age of 18 and death would be with diagnosed depression (see Tables 3).

| Table 2: Years of Life Lived with Depression in a British Columbia Male Birth Cohort of 20,000 |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Age Group | Mean Survival Rate | Individuals in Birth Cohort | Estimated First MDE | Males <br> Estimated Subsequent MDE | Years of Life with Depression in Birth Cohort | Years of Life in Birth Cohort | \% of Life <br> Years with <br> Depression |
| 18-19 | 0.993 | 19,862 | 58.6 | 205.2 | 376.8 | 39,724 | 0.95\% |
| 20-24 | 0.991 | 19,821 | 146.3 | 512.0 | 940.0 | 99,106 | 0.95\% |
| 25-29 | 0.987 | 19,742 | 145.7 | 510.0 | 936.2 | 98,709 | 0.95\% |
| 30-34 | 0.983 | 19,666 | 145.2 | 508.0 | 932.6 | 98,332 | 0.95\% |
| 35-39 | 0.979 | 19,571 | 144.5 | 505.6 | 928.1 | 97,854 | 0.95\% |
| 40-44 | 0.972 | 19,442 | 143.5 | 502.3 | 922.0 | 97,211 | 0.95\% |
| 45-49 | 0.963 | 19,263 | 142.2 | 497.6 | 913.5 | 96,314 | 0.95\% |
| 50-54 | 0.950 | 19,003 | 140.3 | 490.9 | 901.2 | 95,017 | 0.95\% |
| 55-59 | 0.931 | 18,619 | 137.4 | 481.0 | 883.0 | 93,095 | 0.95\% |
| 60-64 | 0.902 | 18,041 | 133.2 | 466.1 | 855.5 | 90,204 | 0.95\% |
| 65-69 | 0.858 | 17,164 | 126.7 | 443.4 | 814.0 | 85,820 | 0.95\% |
| 70-74 | 0.792 | 15,837 | 116.9 | 409.1 | 751.0 | 79,183 | 0.95\% |
| 75-79 | 0.693 | 13,861 | 102.3 | 358.1 | 657.3 | 69,305 | 0.95\% |
| 80+ | 0.296 | 5,918 | 17.5 | 61.2 | 112.3 | 11,836 | 0.95\% |
| Total Ag | es 18+ |  | 1,700 | 5,950 | 10,923 | 1,151,710 | 0.95\% |


| Table 3: Years of Life Lived with Depression in a British Columbia Female Birth Cohort of 20,000 |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Age Group | Mean Survival Rate | Individuals <br> in Birth <br> Cohort | Estimated <br> First MDE | Females <br> Estimated <br> Subsequent <br> MDE | Years of Life with Depresion in Birth Cohort | Years of Life in Birth Cohort | \% of Life <br> Years with <br> Depression |
| 18-19 | 0.994 | 19,887 | 82.5 | 288.8 | 530.2 | 39,775 | 1.33\% |
| 20-24 | 0.993 | 19,868 | 206.1 | 721.3 | 1,324.1 | 99,339 | 1.33\% |
| 25-29 | 0.992 | 19,836 | 205.8 | 720.2 | 1,322.0 | 99,179 | 1.33\% |
| 30-34 | 0.990 | 19,799 | 205.4 | 718.8 | 1,319.6 | 98,997 | 1.33\% |
| 35-39 | 0.987 | 19,748 | 204.8 | 717.0 | 1,316.1 | 98,738 | 1.33\% |
| 40-44 | 0.984 | 19,672 | 204.1 | 714.2 | 1,311.1 | 98,358 | 1.33\% |
| 45-49 | 0.978 | 19,560 | 202.9 | 710.1 | 1,303.6 | 97,800 | 1.33\% |
| 50-54 | 0.970 | 19,395 | 201.2 | 704.2 | 1,292.7 | 96,977 | 1.33\% |
| 55-59 | 0.957 | 19,150 | 198.6 | 695.2 | 1,276.3 | 95,748 | 1.33\% |
| 60-64 | 0.939 | 18,774 | 194.7 | 681.6 | 1,251.3 | 93,872 | 1.33\% |
| 65-69 | 0.909 | 18,190 | 188.7 | 660.4 | 1,212.3 | 90,948 | 1.33\% |
| 70-74 | 0.863 | 17,265 | 179.1 | 626.8 | 1,150.7 | 86,325 | 1.33\% |
| 75-79 | 0.790 | 15,799 | 163.9 | 573.6 | 1,053.0 | 78,995 | 1.33\% |
| 80+ | 0.384 | 7,677 | 95.6 | 334.5 | 614.0 | 46,063 | 1.33\% |
| Total Ag | es 18+ |  | 2,533 | 8,867 | 16,277 | 1,221,114 | 1.33\% |

- Depression increases an individual's mortality risk. Males living with depression are 21 times as likely to commit suicide as males without depression. For females, this ratio increases to 27 times. ${ }^{334}$ Individuals living with depression also have higher rates of overall excess mortality with an early meta-analysis suggesting a RR of 1.81 ( $95 \%$ CI of 1.58 to 2.07 ). ${ }^{335}$ This review, however, did not adjust for confounding variables such as chronic illness and lifestyle. After adjusting for tobacco smoking and heavy alcohol use, Murphy et al. found a non-significant increase in mortality associated with depression in men (RR 1.6, 95\% CI of 0.8 to 3.1). ${ }^{336}$ Other research has found that the effect of depression on mortality is independent of chronic illnesses such as diabetes ${ }^{337}$ and congestive heart failure. ${ }^{338}$ After adjusting for a number of potentially confounding covariates, including the presence of chronic disease, Schoevers, et al. found a $41 \%$ higher mortality rate associated with chronic depression. ${ }^{339}$ A more recent meta-analysis of excess mortality associated with depression found a RR of 1.52 ( $95 \%$ CI of 1.45 to 1.59 ). ${ }^{340}$ For modelling purposes we calculated the number of deaths occurring for males and females between the ages of 20 and 74 in our birth cohort and then estimated how many of these deaths would be in individuals living with depression. We assumed that depression would increase the premature mortality rate by $52 \%$ and varied this in the sensitivity analysis from $45 \%$ to $59 \%$. In males, 20 deaths and 477 life years lost in the cohort are attributable to depression (see Table 4). In females, 18 deaths and 444 life years lost are attributable to depression (see Table 5).

[^86]Table 4: Deaths and Life Years Lost Attributable to Depression in a British Columbia Male Birth Cohort of 20,000

| Age Group | Individuals <br> in Birth <br> Cohort | Deaths | Proportion with <br> Depression | Unadjusted Deaths in Pop. With Depression | Adjusted <br> Deaths in <br> Pop. With <br> Depression | Deaths Attributable to Depression | Average Life Years Lived | Life Years <br> Lost to <br> Depression |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 18-19 | 19,862 |  |  |  |  |  |  |  |
| 20-24 | 19,821 | 41 | 0.95\% | 0.4 | 0.6 | 0.2 | 58.9 | 12 |
| 25-29 | 19,742 | 79 | 0.95\% | 0.8 | 1.1 | 0.4 | 56.0 | 22 |
| 30-34 | 19,666 | 75 | 0.95\% | 0.7 | 1.1 | 0.4 | 51.1 | 19 |
| 35-39 | 19,571 | 96 | 0.95\% | 0.9 | 1.4 | 0.5 | 46.3 | 22 |
| 40-44 | 19,442 | 129 | 0.95\% | 1.2 | 1.9 | 0.6 | 41.5 | 26 |
| 45-49 | 19,263 | 179 | 0.95\% | 1.7 | 2.6 | 0.9 | 36.8 | 33 |
| 50-54 | 19,003 | 259 | 0.95\% | 2.5 | 3.7 | 1.3 | 32.2 | 41 |
| 55-59 | 18,619 | 384 | 0.95\% | 3.6 | 5.5 | 1.9 | 27.7 | 53 |
| 60-64 | 18,041 | 578 | 0.95\% | 5.5 | 8.3 | 2.9 | 23.4 | 67 |
| 65-69 | 17,164 | 877 | 0.95\% | 8.3 | 12.6 | 4.3 | 19.2 | 83 |
| 70-74 | 15,837 | 1,327 | 0.95\% | 12.6 | 19.1 | 6.5 | 15.3 | 100 |
| Total |  | 4,025 |  | 38 | 58 | 20 |  | 477 |


| Age Group | Individuals <br> in Birth Cohort | Female Deaths | Proportion with <br> Depression | Unadjusted <br> Deaths in <br> Pop. With <br> Depression | Adjusted <br> Deaths in <br> Pop. With <br> Depression | Deaths Attributable to Depression | Average Life Years Lived | Life Years <br> Lost to <br> Depression |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 18-19 | 19,887 |  |  |  |  |  |  |  |
| 20-24 | 19,868 | 20 | 1.33\% | 0.3 | 0.4 | 0.1 | 62.7 | 9 |
| 25-29 | 19,836 | 32 | 1.33\% | 0.4 | 0.6 | 0.2 | 57.8 | 13 |
| 30-34 | 19,799 | 36 | 1.33\% | 0.5 | 0.7 | 0.3 | 52.9 | 13 |
| 35-39 | 19,748 | 52 | 1.33\% | 0.7 | 1.0 | 0.4 | 48.1 | 17 |
| 40-44 | 19,672 | 76 | 1.33\% | 1.0 | 1.5 | 0.5 | 43.2 | 23 |
| 45-49 | 19,560 | 112 | 1.33\% | 1.5 | 2.3 | 0.8 | 38.5 | 30 |
| 50-54 | 19,395 | 165 | 1.33\% | 2.2 | 3.3 | 1.1 | 33.8 | 39 |
| 55-59 | 19,150 | 246 | 1.33\% | 3.3 | 5.0 | 1.7 | 29.2 | 50 |
| 60-64 | 18,774 | 375 | 1.33\% | 5.0 | 7.6 | 2.6 | 24.7 | 64 |
| 65-69 | 18,190 | 585 | 1.33\% | 7.8 | 11.8 | 4.1 | 20.4 | 83 |
| 70-74 | 17,265 | 925 | 1.33\% | 12.3 | 18.7 | 6.4 | 16.3 | 104 |
| Total |  | 2,622 |  | 35 | 53 | 18 |  | 444 |

- Diagnosing depression is challenging. "The diagnosis of a mental health disorder is a process that often takes time and develops in a context of trust. Both patient and doctor may need to be sure that the somatic symptoms of depression are exactly that, and not the symptoms of an underlying physical illness." ${ }^{341}$
- Based on a meta-analysis of 41 studies including 50,371 patients, for every 100 patients, GPs identify 10 true positive cases of depression, diagnose 15 patients with depression who do not have depression (false positives) and miss 10 cases of depression (false negatives). Accuracy is improved with prospective examination

[^87]over an extended period of time (3-12 months) rather than relying on a one-time assessment or case-note records. ${ }^{342}$

- Those who meet screening criteria and were previously undiagnosed by their primary care physician tend to be less severely ill than those who were previously diagnosed. ${ }^{343,344}$ Approximately half ( $52 \%$ ) of primary care patients identified by screening have transient symptoms (possibly related to life events) lasting less than two weeks and do not require treatment. ${ }^{345}$
- Zimmerman et al. found that $71 \%$ of patients diagnosed with major depressive disorder in their outpatient practice had a Hamilton Depression Rating Scale (HDRS) score of less than $22 .{ }^{346}$ Scores on the HDRS can be interpreted as follows: no depression (0-7), mild depression (8-16), moderate depression (17-23) and severe depression $(\geq 24) .{ }^{347}$
- When a longitudinal perspective is taken, $30 \%$ of patients with depression remain undetected at 1 year and only $14 \%$ at the end of 3 years, or approximately one out of seven patients with treatable depression. ${ }^{348,349,350}$ For modelling purposes, we assumed that $14 \%$ of depression is undiagnosed treatable depression (see Table 6 , row $i$ ) and increased this to $30 \%$ in the sensitivity analysis.
- $85 \%$ of patients diagnosed with depression were prescribed anti-depressant medication (ADM) in 2011/12 in Canada. ${ }^{351}$
- Approximately $60 \%$ of patients stay on ADM for at least 3 months and $45 \%$ for at least 6 months. ${ }^{352,353}$

[^88]- The use of ADM for major depression is associated with a $64 \% ~(\mathrm{OR}=0.36,95 \% \mathrm{CI}$ of 0.15 to 0.88 ) reduced risk of recurrent depression eight years later ${ }^{354}$ and a $70 \%$ ( $\mathrm{OR}=0.30,95 \% \mathrm{CI}$ of 0.1 to 1.0) reduced risk after 10 years. ${ }^{355}$
- The theoretical cumulative effectiveness of achieving remission through four levels of treatment (primarily medication switching or augmentation) based on the Sequenced Treatment Alternatives to relieve Depression (STAR*D) trial is $36.8 \%$ at Level 1, 56.1\% at Level 2, $62.1 \%$ at Level 3 and $67.1 \%$ at Level 4. ${ }^{356,357}$ For modelling purposes we used Level $2(56.1 \%$ ) results as the base with sensitivity analysis using Level 1 and Level 4 results (see Table 6, row $n$ ).
- Depression has an important influence on a person's QoL. Studies have also shown that individuals with current or treated depression report lower preference scores for depression health states that the general population. ${ }^{358,359}$ Pyne and colleagues suggest that "public stigma may result in the general population being less sympathetic to the suffering of individuals with depression and less willing to validate the impact of depression symptoms.. ${ }^{360}$ Revicki and Wood, based on input from patients with depression who had completed at least eight weeks of ADM, identified the following health state utilities: severe depression $=0.30$, moderate depression $=0.55$ to 0.63 , mild depression $=0.64$ to 0.73 and antidepressant maintenance therapy $=0.72$ to $0.83{ }^{361}$ Whiteford and colleagues ${ }^{362}$ suggest the following health utilities:
- Severe depression $=0.35$ ( $95 \%$ CI of $0.18-0.53$ )
- Moderate depression $=0.59(95 \% \mathrm{CI}$ of $0.45-0.72)$
- Mild depression $=0.84$ ( $95 \% \mathrm{CI}$ of $0.78-0.89$ )

For modelling purposes we assumed an equal proportion of individuals with mild, moderate and severe depression and used the average health utilities provided by Whiteford and colleagues ( $0.59,95 \% \mathrm{CI}$ of $0.47-0.72$ ) adjusted for a general population QoL of 0.848 (see Reference Document) resulting in a QoL reduction of 0.30 (see Table 6 , row $p$ ), ranging from 0.16 to 0.45 .

- Other assumptions used in assessing the clinically preventable burden are detailed in the Reference Document.

Based on these assumptions, screening for depression results in a CPB of 92 quality-adjusted life years saved (see Table 6, row s). The CPB of 92 represents the gap between existing coverage (no coverage) and the 'best in the world' coverage estimated at $12 \%$.

[^89]Table 6: CPB of Screening for Depression in a Birth Cohort of 40,000

| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Life years lived from age 18 to death in a birth cohort of 20,000 males | 1,151,710 | Table 2 |
| b | Life years lived from age 18 to death in a birth cohort of 20,000 females | 1,221,114 | Table 3 |
| c | Life years lived with depression in a birth cohort of 20,000 males | 10,923 | Table 2 |
| d | Life years lived with depression in a birth cohort of 20,000 females | 16,277 | Table 3 |
| e | Proportion of life years lived with depression in a birth cohort of 20,000 males | 0.95\% | = $/$ / a |
| f | Proportion of life years lived with depression in a birth cohort of 20,000 females | 1.33\% | = d/b |
| g | Life years lost attributable to depression in a birth cohort of 20,000 males | 477 | Table 4 |
| h | Life years lost attributable to depression in a birth cohort of 20,000 females | 444 | Table 5 |
| i | Proportion of treatable depression undiagnosed | 14\% | $\checkmark$ |
| j | Life years lived with undiagnosed treatable depression in a birth cohort of 20,000 males | 1,529 | = ${ }^{*}$ i |
| k | Life years lived with undiagnosed treatable depression in a birth cohort of 20,000 females | 2,279 | = ${ }^{*}$ i |
| 1 | Adherence with screening | 12\% | $\checkmark$ |
| m | Life years lived with undiagnosed treatable depression identified by screening | 457 | $=(\mathrm{j}+\mathrm{k}) *$ I |
| n | Effectiveness of ADM in achieving remission | 56\% | $\checkmark$ |
| - | Life years lived in remission with treated depression identified by screening | 256 | $=\mathrm{m}$ * |
| p | Quality of life reduction | 30\% | $\checkmark$ |
| q | QALYs gained | 77 | = * ${ }^{\text {p }}$ |
| r | Life-years gained / death averted | 15 | $=(\mathrm{g}+\mathrm{h}){ }^{\text {i }}$ * ${ }^{\text {l }}$ |
| s | Potential QALYs gained, Screening increasing from 0\% to 12\% | 92 | $=q+r$ |

V = Estimates from the literature
We also modified a number of major assumptions and recalculated the CPB as follows:

- Assume that the RR of excess mortality associated with depression is reduced from 1.52 to 1.45 (Table 4 and 5): CPB $=90$.
- Assume that the RR of excess mortality associated with depression is increased from 1.52 to 1.59 (Table 4 and 5): CPB $=94$.
- Assume the proportion of treatable depression that is undiagnosed is increased from $14 \%$ to $30 \%$ (Table 6, row $i$ ): $\mathrm{CPB}=198$.
- Assume the effectiveness of ADM in achieving remission is reduced from $56 \%$ to $37 \%$ (Table 6, row $l$ ): $\mathrm{CPB}=66$.
- Assume the effectiveness of ADM in achieving remission is increased from $56 \%$ to $67 \%$ (Table 6, row $n$ ): $\mathrm{CPB}=107$.
- Assume the QoL adjustment is reduced from $30 \%$ to $16 \%$ (Table 6, row $p$ ): $\mathrm{CPB}=$ 55.
- Assume the QoL adjustment is increased from $30 \%$ to $45 \%$ (Table 6, row $p$ ): $\mathrm{CPB}=$ 130.

To this point we have not considered some of the potential harms associated with screening for depression, including the negative side-effects of ADM or the possibility that individuals may be diagnosed with depression who do not have depression (false positives).

- There is a side effect burden associated with taking ADM: $48.7 \%$ of individuals taking ADM experienced side effects at least $50 \%$ of the time, with the maximum side effect burden being at least moderate $34.2 \%$ of the time. ${ }^{333}$ Based on input from patients with depression who had completed at least eight weeks of ADM, Revicki and Wood identified a health state utility of between 0.72 and 0.83 associated with antidepressant maintenance therapy. ${ }^{364}$ With an average population health state utility of 0.848 (see Reference Document), this represents a disutility of between 0.02 (or $2.4 \%$ ) and 0.13 ( $15.3 \%$ ). For modelling purposes we assumed a disutility of $8.8 \%$ (the midpoint) and varied this assumption from $2.4 \%$ and $15.3 \%$ in the sensitivity analysis (Table 7, row $t$ ).
- Screening for depression may result in 15 patients being diagnosed with depression who do not have depression (false positives) for every 10 patients who are true positive cases of depression. ${ }^{365}$ For modelling purposes, we have assumed a ratio of 1.5 to 1 false positives to true positives (Table 7, row $n$ ) and that false positive patients will be prescribed ADM the same as true positive patients.
- One of the harms associated with a diagnosis of depression is being rated (i.e. charged a higher life insurance premium) or being refused insurance coverage when the diagnosis of depression is included in the patient's medical chart. Bell suggests that this is one reason why underdiagnoses may be by design rather than accident. ${ }^{366}$ We have not included this potential harm in the modelling.

Based on these additional assumptions, the calculation of CPB is reduced from 92 to -8 quality-adjusted life years saved (see Table 7, row $v$ ). That is, when these harms are taken into account, screening for depression does more harm than good.

[^90]Table 7: CPB of Screening for Depression in a Birth Cohort of 40,000

| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Life years lived from age 18 to death in a birth cohort of 20,000 males | 1,151,710 | Table 2 |
| b | Life years lived from age 18 to death in a birth cohort of 20,000 females | 1,221,114 | Table 3 |
| c | Life years lived with depression in a birth cohort of 20,000 males | 10,923 | Table 2 |
| d | Life years lived with depression in a birth cohort of 20,000 females | 16,277 | Table 3 |
| e | Proportion of life years lived with depression in a birth cohort of 20,000 males | 0.95\% | = $/$ / ${ }^{\text {a }}$ |
| f | Proportion of life years lived with depression in a birth cohort of 20,000 females | 1.33\% | $=d / b$ |
| g | Life years lost attributable to depression in a birth cohort of 20,000 males | 477 | Table 4 |
| h | Life years lost attributable to depression in a birth cohort of 20,000 females | 444 | Table 5 |
| i | Proportion of treatable depression undiagnosed | 14\% | $\checkmark$ |
| j | Life years lived with undiagnosed treatable depression in a birth cohort of 20,000 males | 1,529 | = ${ }^{*} \mathrm{i}$ |
| k | Life years lived with undiagnosed treatable depression in a birth cohort of 20,000 females | 2,279 | $=\mathrm{d}^{*} \mathrm{i}$ |
| 1 | Adherence with screening | 12\% | $\checkmark$ |
| m | Life years lived with undiagnosed treatable depression identified by screening | 457 | $=(j+k) * 1$ |
| n | Life years treated for depression - false positives | 685 | = m 1.5 |
| 0 | Effectiveness of ADM in achieving remission | 56\% | $\checkmark$ |
| p | Life years lived in remission with treated depression identified by screening | 256 | = $\mathrm{m}^{*}$ o |
| q | Quality of life adjustment | 30\% | $\checkmark$ |
| r | QALYs gained | 77 | $=\mathrm{p}^{*} \mathrm{q}$ |
| s | Life-years gained / death averted | 15 | $=(\mathrm{g}+\mathrm{h}) *{ }_{\mathrm{i}}{ }^{\prime} \mathrm{l}$ |
| t | Disutility associated with ADM | -8.8\% |  |
| $u$ | QALYs lost associated with ADM | -101 | $=(\mathrm{m}+\mathrm{n}) * \mathrm{t}$ |
| v | Potential QALYs gained, Screening increasing from 0\% to 12\% | -8 | $=r+s+u$ |

$\checkmark$ = Estimates from the literature

## Modelling Cost-Effectiveness

In this section, we will calculate the CE associated with screening non-pregnant adults ages 18 and older for depression when staff-assisted depression care supports are in place to assure accurate diagnosis, effective treatment, and follow-up in a BC birth cohort of 40,000.

In modelling CE, we made the following assumptions:

- We did not include false positives or the potential disutility associated with taking ADM, as identified in Table 7.
- We assumed that screening would occur annually (Table 8, row $c$ ).
- For patient time and travel costs, we estimated two hours of patient time required per screening visit (Table 8, row $g$ ).
- We assumed that diagnosed depression results in an additional 6 physician visits per year and modified this assumption from 4 to 8 in the sensitivity analysis (see Table 8, row $m$ ).
- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the estimated cost per QALY would be $\$ 148,602$ (see Table 8, row $s$ ).

| Table 8: CE of Screening for Depression in a Birth Cohort of 40,000 |  |  |  |
| :---: | :---: | :---: | :---: |
| Row <br> Label | Variable | Base Case | Data Source |
| a | Life years lived from age 18 to death without diagnosed depression in a birth cohort of 20,000 males | 1,140,786 | Table 6, row a - row C |
| b | Life years lived from age 18 to death without diagnosed depression in a birth cohort of 20,000 females | 1,204,837 | Table 6, row b-row d |
|  | Costs of intervention |  |  |
| c | Frequency of screening (every x years) | 1 | Assumed |
| d | Total number of screens (100\% adherence) | 2,345,623 | $=(a+b) / c$ |
| e | Adherence with screening | 12\% | Table 6, row 1 |
| f | Cost of 10-minute office visit | \$34.85 | Ref Doc |
| g | Value of patient time and travel for office visit | \$59.38 | Ref Doc |
| h | Portion of 10-minute office visit for screen | 50\% | Assumed |
| i | Cost of screening | \$13,261,683 | $=\left(d^{*} e\right) *(f+g) * h$ |
| j | Life years treated for depression | 457 | Table 6, row m |
| k | Annual cost of ADM | \$438 | Ref Doc |
| 1 | Cost of ADM | \$200,150 | $=j^{*} \mathrm{k}$ |
| m | Annual \# of additional visits to a clinician associated with treatment for depression | 6 | Assumed |
| n | Cost of additional follow-up office visits to a clinician | \$258,358 | $=\left(\mathrm{m}^{*} \mathrm{j}\right) *(\mathrm{f}+\mathrm{g})$ |
|  | CE calculation |  |  |
| 0 | Cost of intervention over lifetime of birth cohort | \$13,720,192 | $=(i+1+n)$ |
| p | QALYs saved | 92 | Table 6, row s |
| q | Cost of intervention over lifetime of birth cohort (1.5\% discount) | \$8,692,068 | Calculated |
| $r$ | QALYs saved (1.5\% discount) | 58 | Calculated |
| s | CE (\$/QALY saved) | \$148,602 | $=q / r$ |

$V=$ Estimates from the literature
We also modified a number of major assumptions and recalculated the CE as follows:

- Assume the proportion of treatable depression that is undiagnosed is increased from $14 \%$ to $30 \%$ (Table 6, row $i$ ): $\mathrm{CE}=\$ 71,996$.
- Assume the effectiveness of ADM in achieving remission is reduced from $56 \%$ to $37 \%$ (Table 6, row $n$ ): $\mathrm{CE}=\$ 207,084$.
- Assume the effectiveness of ADM in achieving remission is increased from $56 \%$ to $67 \%$ (Table 6, row $n$ ): $\mathrm{CPB}=\mathrm{CE}=\$ 127,720$.
- Assume the QoL adjustment is reduced from $30 \%$ to $16 \%$ (Table 6 , row $p$ ): $\mathrm{CE}=$ \$248, 053 .
- Assume the QoL adjustment is increased from $30 \%$ to $45 \%$ (Table 6, row $p$ ): $\mathrm{CE}=$ \$105,909.
- Assume that the proportion of an office visit required for screening is reduced from $50 \%$ to $33 \%$ (Table 8 , row $h$ ): $\mathrm{CE}=\$ 99,776$.
- Assume that the proportion of an office visit required for screening is increased from $50 \%$ to $67 \%$ (Table 8 , row $h$ ): $\mathrm{CE}=\$ 197,438$.
- Assume that diagnosed depression results in an additional 4 physician visits per year rather than 6 (see Table 8 , row $m$ ): $\mathrm{CE}=\$ 147,669$.
- Assume that diagnosed depression results in an additional 8 physician visits per year rather than 6 (see Table 8, row $m$ ): $\mathrm{CE}=\$ 149,535$.


## Summary - Excluding Harms

| $40,000$ <br> Summary Excluding Harms |  |  |  |
| :---: | :---: | :---: | :---: |
|  |  |  |  |
| $\begin{aligned} & \text { Base } \\ & \text { Case } \end{aligned}$ |  | Range |  |
| CPB (Potential QALYs Gained) |  |  |  |
| Gap between B.C. Current (0\%) and 'Best in the World' (12\%) |  |  |  |
| 1.5\% Discount Rate | 58 | 35 | 125 |
| 3\% Discount Rate | 39 | 23 | 84 |
| 0\% Discount Rate | 92 | 55 | 198 |
| CE (\$/QALY) including patient time costs |  |  |  |
| 1.5\% Discount Rate | \$148,602 | \$71,996 | \$207,084 |
| 3\% Discount Rate | \$148,602 | \$11,996 | \$207,084 |
| 0\% Discount Rate | \$148,602 | \$11,996 | \$207,084 |
| CE (\$/QALY) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | \$56,325 | \$27,993 | \$78,492 |
| 3\% Discount Rate | \$56,325 | \$27,993 | \$78,492 |
| 0\% Discount Rate | \$56,325 | \$27,993 | \$78,492 |

## Summary - Including Harms

| $40,000$ |  |  |  |
| :---: | :---: | :---: | :---: |
| Summary Including Harms |  |  |  |
|  | $\begin{aligned} & \text { Base } \\ & \text { Case } \\ & \hline \end{aligned}$ | Range |  |
| CPB (Potential QALYs Gained) |  |  |  |
| Gap between B.C. Current (0\%) and 'Best in the World' (12\%) |  |  |  |
| 1.5\% Discount Rate | -5 | -29 | 18 |
| 3\% Discount Rate | -3 | -19 | 12 |
| 0\% Discount Rate | -8 | -45 | 29 |
| CE (\$/QALY) including patient time costs |  |  |  |
| 1.5\% Discount Rate |  |  | \$472,872 |
| 3\% Discount Rate | Dominated | Dominated | \$472,872 |
| 0\% Discount Rate |  |  | \$472,872 |
| CE (\$/QALY) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate |  |  | \$179,234 |
| 3\% Discount Rate | Dominated | Dominated | \$179,234 |
| 0\% Discount Rate |  |  | \$179,234 |

## Screening for Depression in Pregnant and Postpartum Women

## Canadian Task Force on Preventive Health Care (2013)

For adults in subgroups of the population who may be at increased risk of depression, including pregnant and postpartum women, ${ }^{367}$ we recommend not routinely screening for depression. ${ }^{368}$ (Weak recommendation; very-low-quality evidence) ${ }^{369}$

## United States Preventive Services Task Force Recommendations (2016)

The USPSTF recommends screening for depression in the general adult population, including pregnant and postpartum women. Screening should be implemented with adequate systems in place to ensure accurate diagnosis, effective treatment, and appropriate follow-up. (B recommendation) ${ }^{370}$

The Lifetime Prevention Schedule Expert Oversight Committee acknowledges the conflict between the two recommendations. Upon further examination, the USPSTF review included literature investigating screening and treatment of depression in perinatal and postpartum women. The CTFPHC included literature examining screening only, which was sparse; literature examining screening and treatment was excluded. In BC, the current standard for delivery of public health services is offering the Edinburgh Postnatal Depression Scale (EPDS) by eight weeks postpartum, with education/intervention/referral for treatment as needed. The USPSTF review includes a number of validation studies on perinatal and postpartum depression screening tools (including the Edinburgh Postnatal Depression Scale) in a variety of settings. These do not appear in the CTFPHC review. Finally, there are several studies on perinatal and postpartum depression screening and treatment that were published after the CTFPHC review in 2013, but were included in the more recent USPSTF review. Therefore, the LPS will use the USPSTF recommendation as the most current evidence of clinical effectiveness and proceed with the modeling of population health impact and cost effectiveness of screening and treatment for depression in perinatal and postpartum women.

## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with screening pregnant and postpartum women for depression in a BC birth cohort of 40,000 .

In modelling CPB, we made the following assumptions:

- On average, each female in a BC birth cohort would be expected to birth 1.42 children over their lifetime (Table 1, row $a$ ). ${ }^{371}$

[^91]- In 2003/04, $11.9 \%$ of pregnant women in BC visited a physician at least once for depression services during the 27 month time period surrounding their child's birth ( 9 months before conception to 9 months after giving birth). ${ }^{372}$
- A 2004 systematic review found prevalence rates of depression of $7.4 \%, 12.8 \%$ and $12.0 \%$ during the first, second and third trimesters. ${ }^{373}$
- A 2005 systematic review found that the point prevalence of minor and major depressions ranged from approximately $8-11 \%$ during pregnancy, peaked at approximately $13 \%$ three months after giving birth and then fell to about $6 \%$ eight months after giving birth. Less than half of the depressive episodes are MDE. ${ }^{374}$ MDE is a distinct clinical syndrome for which treatment is clearly indicated. ${ }^{375}$
- The majority of depressive episodes resolve within three to six months postpartum. A subset of new mothers (approximately $30 \%$ ), however, remain chronically depressed after this time period. ${ }^{376}$
- For modelling purposes we assumed that screening would occur at 7 weeks post birth (Table 1, row $d$ ) and modified this to screen at 30 weeks pregnancy in the sensitivity analysis (Table 1, row e).
- For modelling purposes we assumed a prevalence of depression of $7.4 \%$ during the first trimester, $12.8 \%$ during the second trimester, $12.0 \%$ during the third trimester and $13 \%$ during the eight months after giving birth. We also assumed an equal distribution between mild, moderate and severe depression, yielding a weighted average prevalence of $7.9 \%$ for moderate to severe depression (Table 1, row $v$ ). If we screen at 7 weeks post birth, a potential total of 1,274 years lived with moderate to severe depression between 7 weeks and eight months post birth would be identified in the cohort (Table 1, row $d$ ). If we screen at 30 weeks pregnant, a potential total of 1,996 years lived with moderate to severe depression between 30 weeks pregnant and eight months post birth would be identified in the cohort (Table 1, row $e$ ).
- Depression is associated with the following disutility: ${ }^{377}$
- Severe depression $=0.65(95 \% \mathrm{CI}$ of $0.47-0.82)$
- Moderate depression $=0.41$ ( $95 \%$ CI of 0.28-0.55)
- Mild depression $=0.16$ ( $95 \% \mathrm{CI}$ of 0.11-0.22)

We assumed an equal distribution between mild, moderate and severe depression, yielding an average disutility of $0.53(95 \% \mathrm{CI}$ of $0.38-0.69)$ for moderate to severe

[^92]depression. The average QoL for a 18-39 year old is 0.90 (see Reference Document), resulting in a \% reduction in QoL of $59 \%$ ( 0.53 / 0.90) (Table 1, row $f$ ).

- Suicide during the perinatal period is rare, with estimates between one and five per 100,000 live births in high income settings. For modelling purposes we have used a rate of $3 / 100,000$ as the base case and modified this from 1 to $5 / 100,000$ in the sensitivity analysis (Table 1, row $h$ ). When suicides do occur during this period, the mean age of the mother is 30.5 years, resulting in a loss of 55 QALYs per suicide (Table 1, row $j$ ). ${ }^{378}$ Women who commit suicide during the perinatal period are twice as likely ( RR of $2.19,95 \% \mathrm{CI}$ of 1.43 to 3.34 ) to have a diagnosis of depression as women who commit suicide outside of the perinatal period (Table 1, row $k$ ). ${ }^{379}$
- Mothers with a high level of depressive symptoms report significantly poorer adherence with childhood safety prevention practices such as the consistent use of car seats, covering electrical plugs, and having syrup of ipecac in the home. ${ }^{380}$
- Postpartum depression does not appear to influence the number of well-baby visits or the likelihood of immunization but it may increase the likelihood of infant hospitalization and sick/emergency visits during the first year of life. ${ }^{381,382}$
- Postpartum depression is associated with a $59 \%$ (OR of $1.59,95 \%$ CI of 1.24 to 2.04) increase in unintentional injury (Table 1, row $o$ ) and a $41 \%$ (OR of $1.41,95 \%$ CI of 1.02 to 1.95 ) increase in falls in infants. ${ }^{383}$
- In BC, the rate of hospital separations due to unintentional injuries in children less than 5 years of age is 671 per 100,000 (Table 1, row $m$ ). The rate of deaths due to unintentional injuries is 10.7 per 100,000 (Table 1 , row $n$ ). ${ }^{384}$ If we assume that the average death occurs at age 2, then each death results in 80 years of life lost (Table 1, row $r$ ). ${ }^{385}$
- Pregnancy and postpartum depression are associated with a shorter duration of breastfeeding. ${ }^{386}$ An Australian study found the median duration of breastfeeding to be 26-28 weeks in women with depression and 39 weeks in women without depression. ${ }^{387}$ Maternal depressive symptoms at 2 to 4 months postpartum are associated with a $27 \%$ ( $95 \%$ CI of $12 \%$ to $39 \%$ ) reduced odds of continuing

[^93]breastfeeding. ${ }^{388}$ For modelling purposes, we assumed a $27 \%$ reduction of exclusive breastfeeding to six months associated with maternal depression (Table 1, row $u$ ) and varied this from $12 \%$ to $39 \%$ in the sensitivity analysis.

- Breastfeeding is associated with a reduced risk of excess weight, otitis media, atopic dermatitis, gastrointestinal infection, lower respiratory tract infection, asthma, type 1 diabetes, childhood leukemia and sudden infant death syndrome in infants and breast and ovarian cancers in the mother. ${ }^{389,390}$ In a previous analysis of the promotion of breastfeeding, we calculated that exclusive breastfeeding to six months is associated with an increase of 0.40 QALYs per infant/mother pair (Table 1, row $t$ ). ${ }^{391}$
- Depression in the year before birth is independently associated with an increase in the risk of Sudden Infant Death Syndrome (SIDS) (OR of 4.9, 95\% CI of 1.1 to 22.1). Depression during pregnancy or after birth is not significantly associated with SIDS. ${ }^{392}$ Since the proposed screening for depression would take place during pregnancy or shortly after birth, we have not included SIDS in this analysis.
- An increased risk of preterm birth is associated with antenatal depression and has been estimated at $37 \%$ (OR of $1.37,95 \%$ CI of 1.04 to 1.81 ) and $39 \%$ (OR of 1.39 , $95 \% \mathrm{CI}$ of 1.19 to 1.61) in two meta-analyses. ${ }^{393,394}$
- Preterm births, including late preterm births, are associated with a greater risk of developmental delay, mental retardation, cerebral palsy, and poor health related outcomes (and utilization) during their first year. ${ }^{395,396,397}$
- Children born preterm tend to have a lower overall QoL than their full term counterparts. The difference in QoL decreases with age (a disutility of 0.13 from birth to age 12 and a disutility of 0.06 from age 13 to 19) and tends to disappear when they become adults. ${ }^{398}$

[^94]- Screening and treatment for depression starting late in pregnancy or shortly after birth, however, is unlikely to have an impact on pre-term birth rates and has not been included in this analysis.
- Maternal depressive symptoms at 2 to 4 months postpartum are associated with a $19 \%$ reduced odds of showing books, $30 \%$ reduced odds of playing with the infant, $26 \%$ reduced odds of talking to the infant and $39 \%$ reduced odds of following routines, compared to mothers without depressive symptoms. ${ }^{399}$
- Few studies have assessed the benefits of treating depression during the perinatal period and the subsequent well-being of the child. The limited research available "has yielded a mixed pattern of results suggesting additional investigations are needed." ${ }^{400}$
- A commonly used depression screening instrument in postpartum and pregnant women is the Edinburgh Postnatal Depression Scale (EPDS). The sensitivity of the EPDS is 0.79 ( $95 \%$ CI of 0.72 to 0.85 ) and the specificity is always higher than $0.87 .{ }^{401}$ This means that the test would identify $79 \%$ of true positive cases (women with perinatal depression) and would falsely identify $13 \%$ of cases as positive (the false positive rate) (Table 1, row $y$ ).
- Involvement in screening programs, with or without additional treatment components, is associated with an $18 \%$ to $59 \%$ (weighted mean of $32 \%$ ) reduced risk of depression (Table 1, row $a b$ ). ${ }^{402}$
- The use of second generation antidepressants during pregnancy may be associated with increased risk of some serious side-effects, ${ }^{403}$ although the research remains unclear. ${ }^{404,405}$
- Cognitive behavioural therapy (CBT) is associated with a $34 \%$ (RR of $1.34,95 \%$ CI of 1.19 to 1.50 ) increase in the likelihood of remission. ${ }^{406}$
- Other assumptions used in assessing the clinically preventable burden are detailed in the Reference Document.

Based on these assumptions, the CPB is 109 quality-adjusted life years saved (see Table 1, row $a e$ ). The CPB of 109 represents the gap between no coverage and the 'best in the world' coverage estimated at $40 \%$.

[^95]Table 1: Calculation of Clinically Preventable Burden (CPB) Estimate for Screening Pregnant and Postpartum Women for Depression in a Birth Cohort of 40,000 (B.C.)

| Row Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Lifetime live births per female | 1.42 | $\checkmark$ |
| b | Proportion of females surviving to age 20 in the cohort | 99.39\% | $\checkmark$ |
| C | Number of pregnancies in the birth cohort | 28,226 | $=(b * 20,000) * a$ |
| d | Estimated years lived with moderate to severe perinatal depression -7 weeks post birth to 34 weeks post birth | 1,274 | $\checkmark$ |
| e | Estimated years lived with moderate to severe perinatal depression - 30 weeks pregnant to 34 weeks post birth | 1,996 | $\checkmark$ |
| f | Disutility associated with moderate to severe depression | 0.59 | $\checkmark$ |
| g | QALYs lost due to moderate to severe perinatal depression | 750 | $=\mathrm{d} * \mathrm{f}$ |
| h | Rate of suicide in perinatal women without depression | 0.00003 | $\checkmark$ |
| I | Suicides in perinatal women without depression | 0.85 | = ${ }^{*} \mathrm{~h}$ |
| j | Years of life lost due to suicide | 55 | V |
| k | Increase in risk of suicide in perinatal women with depression | 119\% | $\checkmark$ |
| I | QALYs lost due to suicide attributable to perinatal depression | 55.4 | $=\left(i^{*}\right)^{*}{ }^{\text {j }}$ |
| m | Rate of hospitalizations due to unintentional injuries in children age 0-4; mothers without depression | 0.0067 | $\checkmark$ |
| n | Mortality rate due to unintentional injuries in children age 0-4; mothers without depression | 0.00011 | $\checkmark$ |
| 0 | Increased risk of unintentional injuries; mothers with depression | 59\% | $\checkmark$ |
| p | Hospitalizations due unintentional injuries in children age 0-4 attributable to mothers with depression | 112 | $=\left(r^{*}\right)^{*} \mathrm{t}$ |
| q | Deaths due to unintentional injuries in children age 0-4 attributable to mothers with depression | 1.8 | $=(\mathrm{s} * \mathrm{c}) * \mathrm{t}$ |
| $r$ | Years of life lost due to death of child from unintentional injury | 80 | $V$ |
| S | QALYs lost due to unintentional injury attributable to perinatal depression | 143 | $=q^{*} r$ |
| t | QALYs lost per mother/infant pair due to not exclusively breastfeeding to six months | 0.40 | $\checkmark$ |
| u | Reduced risk of exclusive breastfeeding to six months associated with maternal depression | 27\% | $\checkmark$ |
| $v$ | Estimated prevalence of moderate to severe perinatal depression | 7.9\% | $\checkmark$ |
| w | QALYs lost due to shorter duration of breastfeeding | 241 | $=v^{*} c^{*} t^{*} u$ |
| x | Total QALYs lost due to moderate to severe perinatal depression | 1,189 | $=\mathrm{g}+\mathrm{j}+\mathrm{s}+\mathrm{w}$ |
| y | Proportion of true positive cases identified by using the EPDS | 79\% | $\checkmark$ |
| z | Adherence with screening | 39\% | Ref Doc |
| aa | Years lived with moderate to severe perinatal depression identified | 366 | $=(w * z) * y$ |
| ab | Effectiveness of screening in reducing the risk of moderate to severe depression | 32\% | $\checkmark$ |
| ac | Years lived with moderate to severe perinatal depression reduced by | 117 | = aa * ab |
| ad | \% of years lived with moderate to severe perinatal depression reduced by screening | 9.2\% | = ac/d |
| ae | Potential QALYs saved (CPB) - Screening increasing from 0\% to 40\% | 109 | = x * ad |

$v=$ Estimates from the literature
We also modified a number of major assumptions and recalculated the CPB as follows:

- Assume that screening would occur at 30 weeks pregnant and again at 7 weeks post birth instead of just at 7 weeks post birth (Table 1, row $e$ ): CPB $=202$.
- Assume that the disutility associated with moderate to severe depression is reduced from 0.59 to 0.42 (Table 1, row $f$ ): $\mathrm{CPB}=73$.
- Assume that the disutility associated with moderate to severe depression is increased from 0.59 to 0.76 (Table 1, row $f$ ): $\mathrm{CPB}=153$.
- Assume that the increased risk of unintentional injuries in children (mothers with depression) is reduced from $59 \%$ to $24 \%$ (Table 1, row $o$ ): $\mathrm{CPB}=94$.
- Assume that the increased risk of unintentional injuries in children (mothers with depression) is increased from $59 \%$ to $104 \%$ (Table 1, row o): CPB $=130$.
- Assume that the effectiveness of screening in reducing the risk of moderate to severe depression is reduced from $32 \%$ to $18 \%$ (Table 1, row $a b$ ): CPB $=62$.
- Assume that the effectiveness of screening in reducing the risk of moderate to severe depression is increased from $32 \%$ to $59 \%$ (Table 1, row $a b$ ): $\mathrm{CPB}=202$.
- Assume that the reduced risk of exclusive breastfeeding to six months associated with maternal depression is reduced from $27 \%$ to $12 \%$ (Table 1, row $u$ ): CPB $=86$.
- Assume that the reduced risk of exclusive breastfeeding to six months associated with maternal depression is increased from $27 \%$ to $39 \%$ (Table 1, row $u$ ): CPB = 130.


## Modelling Cost-Effectiveness

In this section, we will calculate the CE associated with screening pregnant and postpartum women for depression in a BC birth cohort of 40,000 .

In modelling CE, we made the following assumptions:

- Expected screens - We assumed that screening would occur once per pregnancy (Table 2, row $a$ ) and modified this to twice in the sensitivity analysis. ${ }^{407,408}$
- Cost of office visit - Screening with the EPDS takes approximately 5 minutes. ${ }^{409}$ We therefore assumed that $50 \%$ of a 10 -minute office visit would be required for the screening and varied this from $33 \%$ to $67 \%$ in the sensitivity analysis (Table 2, row h).
- Evaluation of women with positive screens - Women who test positive for depression on the EPDS should be offered a psychiatric diagnostic assessment. ${ }^{410} \mathrm{We}$ assumed a cost of $\$ 237.95$ for this assessment, based on fee code 00610 - full diagnostic interview by a psychiatrist in the BC MSC Payment Schedule (Table 2, row $o$ ). ${ }^{411}$ The assessment and fee applies to all true and false positive cases.
- Treatment for depression - For the base model, we assumed that women with severe depression would be treated with CBT rather than antidepressant medication, due to potential safety concerns. CBT can be provided in a group or to an individual. Individual therapy consists of 12 - 90 minute sessions with 1-2 follow-up sessions

[^96]lasting from 10-30 minutes for a total therapy time of approximately 19 hours. ${ }^{412}$ The cost of psychiatric treatment in BC is $\$ 169.75$ per hour ${ }^{413}$ for a total cost of $\$ 3,225$ per individual. Group therapy general consists of 1 initial individual session lasting 90 minutes, eight individuals receiving 12 - 120 minute sessions with 1-2 follow-up sessions lasting from $10-30$ minutes. ${ }^{414}$ The cost of group therapy in BC with eight clients is $\$ 269$ per hour. ${ }^{415}$ The cost of group therapy would therefore be $\$ 1,231$ per person (Table 2, row $q$ ). For modelling purposes, we assumed in the base model that CBT would be provided as group therapy and then included the costs for individual therapy in the sensitivity analysis. For patient time and travel costs associated with CBT we assumed 26.5 hours in therapy plus 1 hour travel for each session for a total of 41 hours. If antidepressant medication is used, the cost/day for antidepressant prescriptions in BC ranges from $\$ 1.00$ for prescriptions paid by the provincial government to $\$ 1.19$ for prescription paid for by uninsured patients and $\$ 1.27$ paid for by private insurers. ${ }^{466}$ The weighted average is $\$ 1.15 /$ day or $\$ 420 /$ year.

- Hospitalizations avoided due to unintentional injury - We assumed that the hospital costs per unintentional injury would be $\$ 20,524$ (Table 2, row $u$ ). ${ }^{417}$
- Costs avoided due to increased duration of breastfeeding - In a previous analysis of the promotion of breastfeeding, we calculated that exclusive breastfeeding to six months is associated with costs avoided of $\$ 2,067$ per infant/mother pair (Table 2, row $w$ ). ${ }^{418}$
- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the estimated cost per QALY would be $\$ 23,042$ (Table 2, row $a d$ ).

[^97]Table 2. Calculation of Cost-effectiveness (CE) for Screening Pregnant and Postpartum Women for Depression in a Birth Cohort of 40,000

| Row Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Number of screens per pregnancy | 1 | $\checkmark$ |
| b | Number of pregnancies in the birth cohort | 28,226 | = Table 1, row c |
| c | Total \# of screens in birth cohort - 100\% adherence | 28,226 | = a * b |
| d | Adherence with screening | 39\% | = Table 1, row z |
| e | Total \# of screens in birth cohort - 40\% adherence | 11,008 | $=c^{*} \mathrm{~d}$ |
| $f$ | Cost of 10-minute office visit | \$34.85 | Ref Doc |
| g | Value of patient time and travel for office visit | \$59.38 | Ref Doc |
| h | Portion of 10-minute office visit for screen | 50\% | $\checkmark$ |
| I | Cost of screening | \$518,652 | $=e^{*}(\mathrm{f}+\mathrm{g})^{*} \mathrm{~h}$ |
| j | Estimated prevalence of perinatal depression | 7.9\% | = Table 1, row v |
| k | EPDS true positive \% | 79\% | = Table 1, row y |
| 1 | EPDS false positive \% | 13\% | $\checkmark$ |
| m | \# of true positive screens | 688 | $=b^{*} d^{*}{ }^{*} k$ |
| n | \# of false positive screens | 113 | $=b^{*} \mathrm{~d}^{*} \mathrm{j}^{*} \mathrm{l}$ |
| 0 | Cost per psychiatric assessment | \$237.95 | $\checkmark$ |
| p | Cost of psychiatric assessment | \$238,068 | $=(\mathrm{m}+\mathrm{n}) * \mathrm{o}+(\mathrm{m}+\mathrm{n}) * \mathrm{~g}$ |
| q | Cost of CBT / ADM per individual | \$1,231 | $\checkmark$ |
| $r$ | Costs of patient time for CBT per individual | \$1,217 | $=41 *(\mathrm{~g} / 2)$ |
| S | Cost of CBT | \$1,683,308 | $=(q+r) * m$ |
| t | Hospitalizations due to unintentional injuries avoided with screening | 10.3 | = Table 1, row p * Table 1, row ad |
| u | Cost of hospital treatment | \$20,524 | V |
| v | Costs avoided due to unintentional injury hospitalizations avoided | -\$211,015 | $=t^{*} u$ |
| w | Costs avoided due to exclusive breastfeeding to six months per mother / infant pair | -\$2,067 | $\checkmark$ |
| X | Reduced risk of exclusive breastfeeding associated with maternal depression | 27\% | = Table 1, row u |
| Y | Costs avoided due to longer duration of breastfeeding | -\$114,588 | = Table 1, row v * Table 1, row c * Table 1, row ad * w ${ }^{*} x$ |
| z | Net screening and patient costs (undiscounted) | \$2,114,425 | $=i+p+s+v+y$ |
| aa | QALYs saved (undiscounted) | 109 | = Table 1, row ae |
| ab | Net screening and patient costs (1.5\% discount) | \$2,131,450 | Calculated |
| ac | QALYs saved (1.5\% discount) | 93 | Calculated |
| ad | CE (\$/QALY saved) | \$23,042 | = ab / ac |

$\checkmark=$ Estimates from the literature
We also modified a number of major assumptions and recalculated the CE as follows:

- Assume that screening would occur at 30 weeks pregnant and again at 7 weeks post birth instead of just at 7 weeks post birth (Table 1, row $e$ ): $\mathrm{CE}=\$ 28,566$.
- Assume that the disutility associated with moderate to severe depression is reduced from 0.59 to 0.42 (Table 1, row $f$ ): $\mathrm{CE}=\$ 36,843$.
- Assume that the disutility associated with moderate to severe depression is increased from 0.59 to 0.76 (Table 1, row $f$ ): $\mathrm{CE}=\$ 15,632$.
- Assume that the increased risk of unintentional injuries in children (mothers with depression) is reduced from $59 \%$ to $24 \%$ (Table 1, row $o$ ): $\mathrm{CE}=\$ 27,714$.
- Assume that the increased risk of unintentional injuries in children (mothers with depression) is increased from $59 \%$ to $104 \%$ (Table 1, row $o$ ): $\mathrm{CE}=\$ 18,030$.
- Assume that the effectiveness of screening in reducing the risk of depression is reduced from $32 \%$ to $18 \%$ (Table 1, row $a b$ ): $\mathrm{CE}=\$ 43,255$.
- Assume that the effectiveness of screening in reducing the risk of depression is increased from $32 \%$ to $59 \%$ (Table 1, row $a b$ ): $\mathrm{CE}=\$ 11,149$.
- Assume that the portion of a 10 -minute office visit required for screening is reduced from $50 \%$ to $33 \%$ (Table 2, row $h$ ): $\mathrm{CE}=\$ 21,163$.
- Assume that the portion of a 10 -minute office visit required for screening is increased from $50 \%$ to $67 \%$ (Table 2, row $h$ ): CE $=\$ 24,920$.
- Assume that the cost of CBT per individual is increased from $\$ 1,231$ to $\$ 3,225$ (Table 2, row $q$ ): $\mathrm{CE}=\$ 37,644$.
- Assume that $50 \%$ of individuals use group CBT and $50 \% \mathrm{ADM}$ (Table 2, row $q$ ): CE $=\$ 20,072$.
- Assume that the reduced risk of exclusive breastfeeding to six months associated with maternal depression is reduced from $27 \%$ to $12 \%$ (Table 1 , row $u$ ): $\mathrm{CE}=$ \$29,016.
- Assume that the reduced risk of exclusive breastfeeding to six months associated with maternal depression is increased from $27 \%$ to $39 \%$ (Table 1, row $u$ ): $\mathrm{CE}=$ \$19,357.


## Summary

| Table 3: Offer of Screening Pregnant and Postpartum Women for Depression in a Birth Cohort of 40,000 Summary |  |  |  |
| :---: | :---: | :---: | :---: |
|  | Base <br> Case | Range |  |
| CPB (Potential QALYs Gained) |  |  |  |
| Gap between 0\% and 'Best in the World' (39\%) |  |  |  |
| 1.5\% Discount Rate | 93 | 52 | 171 |
| 3\% Discount Rate | 79 | 45 | 146 |
| 0\% Discount Rate | 109 | 62 | 202 |
| CE (\$/QALY) including patient time costs |  |  |  |
| 1.5\% Discount Rate | \$23,042 | \$11,149 | \$43,255 |
| 3\% Discount Rate | \$26,846 | \$13,163 | \$50,109 |
| 0\% Discount Rate | \$19,334 | \$9,124 | \$36,688 |
| CE (\$/QALY) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | \$10,140 | \$4,151 | \$20,319 |
| 3\% Discount Rate | \$12,002 | \$5,110 | \$23,715 |
| 0\% Discount Rate | \$8,258 | \$3,116 | \$16,997 |

## Screening for Osteoporosis to Prevent Fractures

## United States Preventive Services Task Force Recommendations ${ }^{419}$

The USPSTF recommends screening for osteoporosis with bone measurement testing to prevent osteoporotic fractures in women 65 years and older. ( $B$ recommendation)

The USPSTF recommends screening for osteoporosis with bone measurement testing to prevent osteoporotic fractures in postmenopausal women younger than 65 years at increased risk of osteoporosis, as determined by a formal clinical risk assessment tool. (B recommendation)

The USPSTF concludes that the current evidence is insufficient to assess the balance of benefits and harms of screening for osteoporosis to prevent osteoporotic fractures in men. (I statement)

In discussing the limitations of their recommendation, the USPSTF states that "...evidence is limited on the direct question of the benefits and harms of screening for elevated osteoporotic fracture risk. The indirect evidence pathway rests on studies evaluating (1) the accuracy of screening approaches in identifying osteoporosis and predicting fractures and (2) the benefits of treatment among those with osteoporosis or at high risk for fractures. Other limitations of the evidence base relate to underlying heterogeneity in baseline risk, prior fractures, prior treatment, and duration of follow-up. ${ }^{420}$

## Canadian Task Force on Preventive Health Care Recommendations

The CTFPHC does not have a current published recommendation on screening for osteoporosis. ${ }^{421}$

We will follow the approach of the USPSTF and model the path of indirect evidence.

## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with screening for osteoporosis in females ages 65 and older.

In modelling CPB, we made the following assumptions:

- Using longitudinal peak bone mineral data from the Canadian multicentre osteoporosis study (CaMos), Berger et al. estimate the prevalence of osteoporosis in Canadian women over 65 years old to be $37.1 \% ~(95 \%$ CI $33.6 \%-42.7 \%) .{ }^{422}$

[^98]- Cheng et al. evaluated Medicare claims in the US and estimated the following prevalence of osteoporosis in women by age: ${ }^{423} 65-69$ (29.8\%), $70-74$ (33.7\%), and $75-79$ ( $41.8 \%$ ), $80+(48.3 \%)$.
- The prevalence of osteoporosis in BC women by age, based on data from BBC's Chronic Disease Registry between 2001 and 2017, is as follows: $65-69$ (19.2\%), 70 $-74(25.3 \%)$, and $75-79(30.7 \%), 80+(37.1 \%) .{ }^{424}$

| Table 1: Screening for Osteoporosis in Women Ages 65 and Older |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Prevalence of Osteoporosis |
| In a BC Birth Cohort of 40,000 |

- We used the age-specific estimates of prevalence from the BC Chronic Disease Registry applied to our BC cohort of women starting at age 65 and continuing to age 86 (based on the average life expectancy of 22 years for a 65 year old women) and estimated a prevalence in BC of $28.3 \%$ (see Table 1), lower than the $37.1 \%$ identified by Berger et al. ${ }^{425}$

[^99]- A study by Hopkins and colleagues calculated the total number of patients with fractures in Canada between April 1, 2010 and March 31, 2011, by sex, age and type of fracture using data from the Canadian Institute for Health Information (CIHI). ${ }^{426}$ The various types of fractures were identified based on International Statistical Classification of Diseases and Related Health Problems, Tenth Revision, Canada (ICD-10-CA) codes. We compiled the relevant data for women ages 60-89 and calculated the incidence rate per 100,000 by age group ( $60-69,70-79$ and $80-89$ ) by fracture type (see Table 2).

|  | Age Group |  |  |  |
| :---: | :---: | :---: | :---: | :---: |
|  | 60-69 | 70-79 | 80-89 | Total |
| Female Population in 2011 | 1,760,036 | 1,085,293 | 681,159 | 3,526,488 |
| Number of Fractures in Canada in 2011 |  |  |  |  |
| Hip | 1,826 | 4,238 | 9,612 | 15,676 |
| Vertebral | 904 | 1,673 | 2,540 | 5,117 |
| All Other |  |  |  |  |
| Wrist | 7,584 | 5,131 | 4,486 | 17,201 |
| Humerus | 1,844 | 2,015 | 2,423 | 6,282 |
| Other | 8,867 | 8,055 | 11,779 | 28,701 |
| Multiple | 1,271 | 1,835 | 2,769 | 5,875 |
| Subtotal All Other | 19,566 | 17,036 | 21,457 | 58,059 |
| Total | 22,296 | 22,947 | 33,609 | 78,852 |
| Fracture Rate per 100,000 person years |  |  |  |  |
| Hip | 104 | 390 | 1,411 | 445 |
| Vertebral | 51 | 154 | 373 | 145 |
| All Other |  |  |  |  |
| Wrist | 431 | 473 | 659 | 488 |
| Humerus | 105 | 186 | 356 | 178 |
| Other | 504 | 742 | 1,729 | 814 |
| Multiple | 72 | 169 | 407 | 167 |
| Subtotal All Other | 1,112 | 1,570 | 3,150 | 1,646 |
| Total | 1,267 | 2,114 | 4,934 | 2,236 |

- For modelling purposes, we assumed a hip fracture rate of 104 / 100,000 person years in women ages 65-69, $390 / 100,000$ person years in women ages 70-79 and 1,411 / 100,000 person years in women ages $80-86$. Furthermore, we assumed a vertebral fracture rate of 51 / 100,000 person years in women ages $65-69,154$ / 100,000 person years in women ages 70-79 and 373 / 100,000 person years in women ages 80-86. Finally, we assumed a non-hip, non-vertebral fracture rate of 1,112 / 100,000 person years in women ages $65-69,1,570 / 100,000$ person years in women ages 70-79 and $3,150 / 100,000$ person years in women ages 80-86.

[^100]- Lippuner and colleagues estimated that 71\% of hip fractures in 65-74 year olds are attributable to osteoporosis. ${ }^{427}$ This increases to $91 \%$ in 74-84 year olds. Similarly, approximately $81 \%$ of vertebral fractures in 65-84 year olds are attributable to osteoporosis. Finally, non-hip, non-vertebral, non-stress fractures attributable to osteoporosis ranged from 50-78\% for ages 65-74 and between $60-91 \%$ for ages $75+$.
- In their economic modelling, Hopkins et al. assumed that $100 \%$ of hip and vertebral fractures are attributable to osteoporosis while $81.5 \%$ of all other fractures in women are attributable to osteoporosis. ${ }^{428}$
- For modelling purposes, we assumed that $71 \%$ of hip fractures in 65-74 year olds are attributable to osteoporosis, increasing to $91 \%$ at age 75 , that $81 \%$ of vertebral fractures are attributable to osteoporosis and $81.5 \%$ of all other fractures are attributable to osteoporosis (see Table 3).
- In Table 3, we show that for the 22 years modelled for the cohort beginning at age 65 , the total number of osteoporosis-attributable fractures is 7,379 . Of these, 1,708 are hip fractures, 507 are vertebral fractures and 5,164 are other fractures.

- In their meta-analysis on morbidity associated with hip fractures, Haentjen and colleagues calculated a hazard ratio of 2.87 ( $95 \%$ CI $2.52-3.27$ ) of death in the first year for women 50 and older with a hip fracture compared to those without. ${ }^{429} \mathrm{~A}$

[^101]hazard ratio of 1.00 suggests that the death rate in the group of interest is the same as that in the general population.

- Tran and colleagues report that for women over 50 the hazard ratio (of excess mortality) of any fragility fracture is 1.51 ( $95 \%$ CI $1.31-1.75$ ), 2.13 (1.58-2.87) for hip fractures, 1.82 (1.28-2.57) for vertebral fractures and 1.38 (1.18-1.62) for nonhip, non-vertebral fractures. ${ }^{430}$
- In his commentary on mortality after osteoporotic fractures, Schousboe discusses some of the links between fracture and mortality. He notes that "...after adjustment for comorbidity, and/or functional status, some studies report longer-term excess mortality after hip fracture and others do not. ${ }^{3431}$
- We will model the risk of excess mortality for women with a hip fracture using a hazard ratio of 2.87 in the first year after hip fracture (and vary this from 2.52 to 3.27 in our sensitivity analysis). We will model the risk of excess mortality for women with vertebral fractures at 1.82 (varied between 1.28 and 2.57) and for all other fractures (i.e. non-hip, non-vertebral) we use a hazard ratio of 1.38 (varied between 1.18 and 1.62 ). We conservatively apply the excess mortality only in the year of the incident fracture.
- Based on the number of osteoporotic fractures calculated in Table 3, we calculate the number of deaths and life years lost attributable to osteoporotic fractures (see Table 4).
- In Table 4, we show that 181 excess deaths are attributable to osteoporosis-related fractures, 113 of which are due to hip fractures, 13 to vertebral fractures and 55 to other fractures. Combining the year when the deaths occur with life expectancy at the time of death, we further show that a total of 1,000 life years are lost due to osteoporosis-related fractures. Of these, 571 life years lost are due to hip fractures, 75 are due to vertebral fractures and 354 are due to other fractures.

[^102]| Table 4: Screening for Osteoporosis in Women Ages 65 and Older Number of Deaths Attributable to Osteoporotic Fracture <br> In a BC Birth Cohort of 40,000 |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Age | \# in Cohort | Deaths <br> in Cohort | Years <br> Lived | Death <br> Rate / 100,000 | Fractur <br> Hip <br> Fracture | ures Attribut Osteoporosi <br> Vertebral Fracture | table to is <br> All Other Fractures | Hazard R Due to Hip Fracture | Ratio of Exc Incident F <br> Vertebral Fracture | cess Death Fracture <br> All Other Fractures | Excess De <br> Hip <br> Fracture | Deaths Due to Fracture <br> Vertebral Fracture | Incident <br> All Other Fractures | Life Expectancy | Life Y Osteo Hip Fracture | Years Lost Du porotic Fra <br> Vertebral Fracture | bue to ctures <br> All Other Fractures |
| 64 | 18,572 |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| 65 | 18,456 | 116 | 18,392 | 629 | 14 | 8 | 167 | 2.87 | 1.82 | 1.38 | 0.16 | 0.04 | 0.40 | 22 | 4 | 1 | 9 |
| 66 | 18,329 | 127 | 18,259 | 692 | 13 | 8 | 165 | 2.87 | 1.82 | 1.38 | 0.17 | 0.04 | 0.43 | 21 | 4 | 1 | 9 |
| 67 | 18,190 | 139 | 18,113 | 765 | 13 | 8 | 164 | 2.87 | 1.82 | 1.38 | 0.19 | 0.05 | 0.48 | 20 | 4 | 1 | 10 |
| 68 | 18,037 | 152 | 17,954 | 845 | 13 | 7 | 163 | 2.87 | 1.82 | 1.38 | 0.21 | 0.05 | 0.52 | 19 | 4 | 1 | 10 |
| 69 | 17,870 | 167 | 17,778 | 936 | 13 | 7 | 161 | 2.87 | 1.82 | 1.38 | 0.23 | 0.06 | 0.57 | 18 | 4 | 1 | 10 |
| 70 | 17,687 | 183 | 17,586 | 1,036 | 49 | 22 | 225 | 2.87 | 1.82 | 1.38 | 0.94 | 0.19 | 0.89 | 17 | 16 | 3 | 15 |
| 71 | 17,486 | 201 | 17,375 | 1,151 | 48 | 22 | 222 | 2.87 | 1.82 | 1.38 | 1.04 | 0.20 | 0.97 | 16 | 17 | 3 | 16 |
| 72 | 17,265 | 221 | 17,144 | 1,278 | 48 | 21 | 219 | 2.87 | 1.82 | 1.38 | 1.14 | 0.22 | 1.06 | 15 | 17 | 3 | 16 |
| 73 | 17,023 | 242 | 16,890 | 1,422 | 47 | 21 | 216 | 2.87 | 1.82 | 1.38 | 1.24 | 0.25 | 1.17 | 14 | 17 | 3 | 16 |
| 74 | 16,758 | 265 | 16,612 | 1,584 | 46 | 21 | 213 | 2.87 | 1.82 | 1.38 | 1.36 | 0.27 | 1.28 | 13 | 18 | 4 | 17 |
| 75 | 16,467 | 291 | 16,307 | 1,766 | 58 | 20 | 209 | 2.87 | 1.82 | 1.38 | 1.91 | 0.29 | 1.40 | 12 | 23 | 4 | 17 |
| 76 | 16,148 | 319 | 15,973 | 1,974 | 57 | 20 | 204 | 2.87 | 1.82 | 1.38 | 2.10 | 0.32 | 1.53 | 11 | 23 | 4 | 17 |
| 77 | 15,799 | 349 | 15,608 | 2,209 | 55 | 19 | 200 | 2.87 | 1.82 | 1.38 | 2.29 | 0.35 | 1.68 | 10 | 23 | 4 | 17 |
| 78 | 15,418 | 381 | 15,209 | 2,474 | 54 | 19 | 195 | 2.87 | 1.82 | 1.38 | 2.50 | 0.39 | 1.83 | 9 | 23 | 3 | 16 |
| 79 | 15,001 | 417 | 14,774 | 2,777 | 52 | 18 | 189 | 2.87 | 1.82 | 1.38 | 2.73 | 0.42 | 1.99 | 8 | 22 | 3 | 16 |
| 80 | 14,547 | 454 | 14,300 | 3,121 | 184 | 43 | 367 | 2.87 | 1.82 | 1.38 | 10.72 | 1.11 | 4.35 | 7 | 75 | 8 | 30 |
| 81 | 14,053 | 494 | 13,785 | 3,514 | 177 | 42 | 354 | 2.87 | 1.82 | 1.38 | 11.63 | 1.20 | 4.73 | 6 | 70 | 7 | 28 |
| 82 | 13,517 | 536 | 13,228 | 3,964 | 170 | 40 | 340 | 2.87 | 1.82 | 1.38 | 12.59 | 1.30 | 5.12 | 5 | 63 | 6 | 26 |
| 83 | 12,938 | 579 | 12,626 | 4,477 | 162 | 38 | 324 | 2.87 | 1.82 | 1.38 | 13.57 | 1.40 | 5.51 | 4 | 54 | 6 | 22 |
| 84 | 12,314 | 624 | 11,980 | 5,066 | 154 | 36 | 308 | 2.87 | 1.82 | 1.38 | 14.57 | 1.50 | 5.92 | 3 | 44 | 5 | 18 |
| 85 | 11,645 | 669 | 11,288 | 5,747 | 145 | 34 | 290 | 2.87 | 1.82 | 1.38 | 15.58 | 1.61 | 6.33 | 2 | 31 | 3 | 13 |
| 86 | 10,931 | 714 | 10,553 | 6,532 | 136 | 32 | 271 | 2.87 | 1.82 | 1.38 | 16.55 | 1.71 | 6.72 | 1 | 17 | 2 | 7 |
| Total |  | 7,640 | 341,738 |  | 1,708 | 507 | 5,164 |  |  |  | 113 | 13 | 55 |  | 571 | 75 | 354 |

- In Table 5, we subtract the number of deaths from the number of osteoporosis fracture events to determine the number of people still living after osteoporosisrelated fractures. This comes to 7,198 people in total, 1,594 of whom have had hip fractures, 494 of whom have had vertebral fractures and 5,110 of whom have had other fractures.

|  | Table 5: Screening for Osteoporosis in Women Ages 65 and Older <br> Number Living with Fracture <br> In a BC Birth Cohort of 40,000 |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Fractures Attributable to Osteoporosis |  |  | Excess Deaths Due to Incident Fracture |  |  | Number Living with Fractures |  |  |
| Age | Hip <br> Fracture | Vertebral Fracture | All Other Fractures | Hip <br> Fracture | Vertebral Fracture | All Other Fractures | Hip <br> Fracture | Vertebral Fracture | All Other Fractures |
| 64 |  |  |  |  |  |  |  |  |  |
| 65 | 14 | 8 | 167 | 0.2 | 0.0 | 0.4 | 13 | 8 | 166 |
| 66 | 13 | 8 | 165 | 0.2 | 0.0 | 0.4 | 13 | 8 | 165 |
| 67 | 13 | 8 | 164 | 0.2 | 0.0 | 0.5 | 13 | 7 | 164 |
| 68 | 13 | 7 | 163 | 0.2 | 0.1 | 0.5 | 13 | 7 | 162 |
| 69 | 13 | 7 | 161 | 0.2 | 0.1 | 0.6 | 13 | 7 | 161 |
| 70 | 49 | 22 | 225 | 0.9 | 0.2 | 0.9 | 48 | 22 | 224 |
| 71 | 48 | 22 | 222 | 1.0 | 0.2 | 1.0 | 47 | 21 | 221 |
| 72 | 48 | 21 | 219 | 1.1 | 0.2 | 1.1 | 46 | 21 | 218 |
| 73 | 47 | 21 | 216 | 1.2 | 0.2 | 1.2 | 46 | 21 | 215 |
| 74 | 46 | 21 | 213 | 1.4 | 0.3 | 1.3 | 45 | 20 | 211 |
| 75 | 58 | 20 | 209 | 1.9 | 0.3 | 1.4 | 56 | 20 | 207 |
| 76 | 57 | 20 | 204 | 2.1 | 0.3 | 1.5 | 55 | 20 | 203 |
| 77 | 55 | 19 | 200 | 2.3 | 0.4 | 1.7 | 53 | 19 | 198 |
| 78 | 54 | 19 | 195 | 2.5 | 0.4 | 1.8 | 52 | 19 | 193 |
| 79 | 52 | 18 | 189 | 2.7 | 0.4 | 2.0 | 50 | 18 | 187 |
| 80 | 184 | 43 | 367 | 10.7 | 1.1 | 4.4 | 173 | 42 | 363 |
| 81 | 177 | 42 | 354 | 11.6 | 1.2 | 4.7 | 165 | 40 | 349 |
| 82 | 170 | 40 | 340 | 12.6 | 1.3 | 5.1 | 157 | 39 | 334 |
| 83 | 162 | 38 | 324 | 13.6 | 1.4 | 5.5 | 149 | 37 | 319 |
| 84 | 154 | 36 | 308 | 14.6 | 1.5 | 5.9 | 139 | 35 | 302 |
| 85 | 145 | 34 | 290 | 15.6 | 1.6 | 6.3 | 129 | 32 | 283 |
| 86 | 136 | 32 | 271 | 16.6 | 1.7 | 6.7 | 119 | 30 | 264 |
| Total | 1,708 | 507 | 5,164 | 113 | 13 | 55 | 1,594 | 494 | 5,110 |

- Betram et al. use a hip-fracture disability weight of 0.272 based on Global Burden of Disease data to model hip-fracture health burden. The authors state that " $29 \%$ of hip fracture cases in the elderly do not reach their pre-fracture levels 1 year post-fracture. Those who do recover tend to reach their pre-fracture levels of functioning at around 6 months." ${ }^{432}$
- Vertebral fracture patients are often advised that it will be a full year before they reach their pre-fracture levels of functioning. ${ }^{433}$
- Kanis and colleagues ${ }^{434}$ assign different disability weights based on expert opinion derived from a 1998 National Osteoporosis Foundation paper. ${ }^{435}$ They suggest a firstyear utility loss with vertebral, rib and pelvis fractures of 0.0502 , with humerus, clavicle, scapula, sternum and distal forearm fractures of 0.0464 and hip, other femoral fractures and tibia and fibula fractures of 0.4681.436

[^103]- The USPSTF found no harms of screening in terms of anxiety or quality of life. ${ }^{437}$
- We model that $29 \%$ of hip fracture patients do not recover their pre-fracture functioning, and have a reduced quality of life for their remaining years of life. We model that the remaining hip fracture patients recover within an average of 6 months. We model vertebral fracture patients recover to pre-fracture levels of functioning in one year and assume that all other fracture types recover in an average of 6 months. We model a 0.27 reduction in QoL following a hip fracture, a 0.050 reduction in QoL for vertebral fractures and a 0.046 QoL reduction for other fractures. Compared to an average quality of life of 0.76 of a 70-79 year old (see Reference Document), this results in a $35.5 \%$ ( $0.27 / 0.76$ ) reduction in QoL due to hip fracture, a $6.6 \%$ reduction due to vertebral fracture and a $6.0 \%$ reduction due to other fractures.
- We apply our assumptions to the individuals living with fractures and calculate QALYs lost attributable to osteoporotic fractures in Table 6. For example, at age 65, $29 \%$ of the 13.4 hip fractures (3.9) will have a lifelong quality decrement. The QALYs lost in this group is the number multiplied by the decrement multiplied by the number of life years remaining, and comes to $30.5(=3.9 * 0.355 * 22)$. The remaining 9.5 hip fractures have the decrement applied for half a year, resulting in 1.7 QALY lost ( $9.5 * 0.355 * 0.5$ ). The total QALYs lost due to hip fracture is the sum of these two, or 32 QALYs.
- Table 6 shows that the total QALYs lost due to osteoporosis-related fractures is 1,606 . Hip fractures account for 1,420 of the QALYs lost, with vertebral fractures and other fractures accounting for 33 and 153 QALYs lost respectively.

[^104]| Table 6: Screening for Osteoporosis in Women Ages 65 and Older Quality Adjusted Life Years for those Living with Fracture In a BC Birth Cohort of 40,000 |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Age | Number <br> Hip <br> Fracture | Living with <br> Vertebral Fracture | Fractures <br> All Other <br> Fractures | Lifetime Dis Hip Fractu <br> Percentage | ability in e Cases <br> Number | Hip <br> Fracture | L Decremen <br> Vertebral <br> Fracture | nt <br> All Other Fractures | Length of <br> Lifetime Hip Cases | Tme for QoL <br> Vertebral <br> Fracture <br> Cases | Decrement <br> All other cases | Quality Adj <br> Due to <br> Hip <br> Fracture | djusted Life steoporotic <br> Vertebral Fracture | Years Lost Fractures <br> All Other <br> Fractures |
| 64 |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| 65 | 13 | 8 | 166 | 29\% | 3.9 | 0.36 | 0.07 | 0.06 | 22 | 1.0 | 0.5 | 32 | 0.5 | 5.0 |
| 66 | 13 | 8 | 165 | 29\% | 3.9 | 0.36 | 0.07 | 0.06 | 21 | 1.0 | 0.5 | 30 | 0.5 | 4.9 |
| 67 | 13 | 7 | 164 | 29\% | 3.8 | 0.36 | 0.07 | 0.06 | 20 | 1.0 | 0.5 | 29 | 0.5 | 4.9 |
| 68 | 13 | 7 | 162 | 29\% | 3.8 | 0.36 | 0.07 | 0.06 | 19 | 1.0 | 0.5 | 27 | 0.5 | 4.9 |
| 69 | 13 | 7 | 161 | 29\% | 3.7 | 0.36 | 0.07 | 0.06 | 18 | 1.0 | 0.5 | 25 | 0.5 | 4.8 |
| 70 | 48 | 22 | 224 | 29\% | 13.9 | 0.36 | 0.07 | 0.06 | 17 | 1.0 | 0.5 | 90 | 1.4 | 6.7 |
| 71 | 47 | 21 | 221 | 29\% | 13.7 | 0.36 | 0.07 | 0.06 | 16 | 1.0 | 0.5 | 84 | 1.4 | 6.6 |
| 72 | 46 | 21 | 218 | 29\% | 13.5 | 0.36 | 0.07 | 0.06 | 15 | 1.0 | 0.5 | 77 | 1.4 | 6.5 |
| 73 | 46 | 21 | 215 | 29\% | 13.2 | 0.36 | 0.07 | 0.06 | 14 | 1.0 | 0.5 | 71 | 1.4 | 6.4 |
| 74 | 45 | 20 | 211 | 29\% | 13.0 | 0.36 | 0.07 | 0.06 | 13 | 1.0 | 0.5 | 65 | 1.4 | 6.3 |
| 75 | 56 | 20 | 207 | 29\% | 16.3 | 0.36 | 0.07 | 0.06 | 12 | 1.0 | 0.5 | 76 | 1.3 | 6.2 |
| 76 | 55 | 20 | 203 | 29\% | 15.9 | 0.36 | 0.07 | 0.06 | 11 | 1.0 | 0.5 | 69 | 1.3 | 6.1 |
| 77 | 53 | 19 | 198 | 29\% | 15.4 | 0.36 | 0.07 | 0.06 | 10 | 1.0 | 0.5 | 61 | 1.3 | 5.9 |
| 78 | 52 | 19 | 193 | 29\% | 14.9 | 0.36 | 0.07 | 0.06 | 9 | 1.0 | 0.5 | 54 | 1.2 | 5.8 |
| 79 | 50 | 18 | 187 | 29\% | 14.4 | 0.36 | 0.07 | 0.06 | 8 | 1.0 | 0.5 | 47 | 1.2 | 5.6 |
| 80 | 173 | 42 | 363 | 29\% | 50.1 | 0.36 | 0.07 | 0.06 | 7 | 1.0 | 0.5 | 146 | 2.8 | 10.9 |
| 81 | 165 | 40 | 349 | 29\% | 48.0 | 0.36 | 0.07 | 0.06 | 6 | 1.0 | 0.5 | 123 | 2.7 | 10.5 |
| 82 | 157 | 39 | 334 | 29\% | 45.6 | 0.36 | 0.07 | 0.06 | 5 | 1.0 | 0.5 | 101 | 2.6 | 10.0 |
| 83 | 149 | 37 | 319 | 29\% | 43.1 | 0.36 | 0.07 | 0.06 | 4 | 1.0 | 0.5 | 80 | 2.4 | 9.6 |
| 84 | 139 | 35 | 302 | 29\% | 40.4 | 0.36 | 0.07 | 0.06 | 3 | 1.0 | 0.5 | 61 | 2.3 | 9.0 |
| 85 | 129 | 32 | 283 | 29\% | 37.5 | 0.36 | 0.07 | 0.06 | 2 | 1.0 | 0.5 | 43 | 2.1 | 8.5 |
| 86 | 119 | 30 | 264 | 29\% | 34.5 | 0.36 | 0.07 | 0.06 | 1 | 1.0 | 0.5 | 27 | 2.0 | 7.9 |
| Total | 1,594 | 494 | 5,110 |  | 462 |  |  |  |  |  |  | 1,420 | 33 | 153 |

- The USPSTF found convincing evidence that "...screening can detect osteoporosis and that treatment of women with osteoporosis can provide at least a moderate benefit in preventing fractures. ${ }^{43}{ }^{43}$
- We have assumed a potential screening rate of $57.8 \%$ (Table 7, row $p$ ). ${ }^{439}$ We assume that all persons with a positive screen for osteoporosis are prescribed medication.
- Fraser and colleagues report on the accuracy of a Canadian modification of the FRAX ${ }^{\circledR}$ fracture prediction screening tool. Combining FRAX ${ }^{\circledR}$ with BMD (bone mineral density) testing resulted in an area under the receiver operator curve of 0.69 (a poor to fair score) for predicting major osteoporotic fractures and 0.80 (a good score) for predicting hip fractures. When just the BMD testing results are used, the equivalent results are 0.66 for major osteoporotic fractures and 0.76 for hip fractures. ${ }^{440}$

[^105]- For women over 65 , the USPSTF ${ }^{441}$ does not explicitly recommend a risk assessment, only DXA screening. ${ }^{42}$ We model accordingly.
- We model a single screening at age 65 to detect osteoporosis and assume that $76 \%$ of hip fractures and $66 \%$ of all other fractures could be predicted by screening with DXA alone (Table 7, rows $q \& r$ ).
- Bisphosphonates have been shown effective in building back bone mineral density and were the most frequently studied medication referenced by the USPSTF. ${ }^{443}$ We therefore model treatment as being carried out with bisphosphonates.
- The review by the USPSTF found that bisphosphonates were found to significantly reduce vertebral fractures ( RR of $0.57,95 \% \mathrm{CI}, 0.41-0.78$ ) and nonvertebral fractures (RR of $0.84,95 \% \mathrm{CI}, 0.76-0.92$ ) but not hip fractures (RR of $0.70,95 \% \mathrm{CI}, 0.44-$ 1.11). ${ }^{444}$
- Long-term treatment compliance is critical in achieving a reduced risk of fracture. In a study of 19,987 (mostly [ $97 \%$ ]) females ages 65 and older, Patrick et al. calculated that $36.5 \%$ of the study cohort took their medication between $80 \%$ and $100 \%$ of the time during the 300 -day medication study compliance period. ${ }^{445}$ A further $31.8 \%$ of the cohort were in the $0-19 \%$ compliance group, $11.3 \%$ were in the $20-39 \%$ compliance group, $8.8 \%$ were in the $40-59 \%$ compliance group and $11.5 \%$ in the $60-$ $79 \%$ compliance group.
- It was in the high compliance group $(80-100 \%)$ that Patrick et al. found a statistically significant 5 -year reduction of $23 \%$ ( $95 \%$ CI of $8 \%$ to $36 \%$ ) in hip fractures, $26 \%$ ( $95 \%$ CI of $12 \%$ to $38 \%$ ) reduction in vertebral fractures and a $20 \%$ ( $95 \%$ CI of $9 \%$ to $29 \%$ ) reduction in other non-hip fractures when compared to the group with poor or no compliance. ${ }^{446}$ The only other compliance group that saw a significant reduction in hip fractures was the $60-79 \%$ group ( $24 \%, 95 \%$ CI of $1 \%$ to $42 \%$ ).
- For the $36.5 \%$ of patients in the high compliance group (the $80-100 \%$ group) (Table 7 , row $s$ ), we model a $23 \%$ reduction in hip fractures, a $26 \%$ reduction in vertebral fractures and a $20 \%$ reduction in all other fractures (Table 7, rows $t$ to $v$ ).
- Shepstone and colleagues ${ }^{477}$ recently published an RCT investigating the potential benefits of a fracture-risk based, community screening program in older women (ages 70-85) in the UK. BMD measurement was only applied to a selected subgroup of these women based on their risk assessment using the Fracture Risk Assessment Tool (FRAX). They found that this screening approach, followed by appropriate osteoporosis medication, did not reduce the overall incidence of osteoporosis-related

[^106]fractures (hazard ratio [HR] 0.94, 95\% CI $0.85-1.03$ ), nor the overall incidence of all clinical fractures ( $0.94,0.86-1.03$ ). It did, however, reduce the incidence of hip fractures ( $0.72,0.59-0.89$ ). As noted previously, we do not assume any risk stratification in our modelling.

Based on these assumptions, the CPB associated with screening for osteoporosis in females ages 65 and older is 91 QALYs (see Table 7, row $a f$ ).

| Table 7: CPB of Screening for Osteoporosis in Women 65+ In a BC Birth Cohort of 40,000 |  |  |  |
| :---: | :---: | :---: | :---: |
| Row Label | Variable | Base case | Data Source |
| a | Expected life-years between age 65 and 86 | 341,738 | Table 1 |
| b | Prevalence of osteoporosis | 38.5\% | Table 1 |
| c | Years lived with osteoporosis | 131,418 | = a*b |
| d | Expected number of hip fractures | 1,971 | Table 3 |
| e | Expected number of vertebral fractures | 626 | Table 3 |
| f | Expected number of all other fractures | 6,337 | Table 3 |
| g | Expected number of hip fractures attributable to osteoporosis | 1,708 | Table 3 |
| h | Expected number of vertebral fractures attributable to osteoporosis | 507 | Table 3 |
| i | Expected number of all other fractures attributable to osteoporosis | 5,164 | Table 3 |
| j | Life years lost due death from to osteoporotic hip fractures | 571 | Table 4 |
| k | Life years lost due to death from osteoporotic vertebral fractures | 75 | Table 4 |
| 1 | Life years lost due to death from all other osteoporotic fractures | 354 | Table 4 |
| m | QALYs lost due to living with osteoporotic hip fractures | 1,420 | Table 6 |
| n | QALYs lost due to living with osteoporotic vertebral fractures | 33 | Table 6 |
| 0 | QALYs lost due to living with osteoporotic other fractures | 153 | Table 6 |
| p | Screening Rate | 57.8\% | $\checkmark$ |
| q | Accuracy of bone density screening to predict hip fractures | 76\% | $\checkmark$ |
| r | Accuracy of bone density screening to predict non-hip fractures | 66\% | $\checkmark$ |
| s | Long term compliance rate with medical treatment | 36.5\% | $\checkmark$ |
| t | Hip fracture reduction rate due to treatment | 23.0\% | $\checkmark$ |
| u | Vertebral fracture reduction rate due to treatment | 26.0\% | $\checkmark$ |
| v | Other fracture reduction rate due to treatment | 20.0\% | $\checkmark$ |
| w | Hip fractures avoided due to treatment | 63 | $=\mathrm{g} * \mathrm{p}^{*} \mathrm{q}^{*} \mathrm{~s}^{*} \mathrm{t}$ |
| x | Vertebral fractures avoided due to treatment | 18 | $=h^{*} \mathrm{p}^{*} \mathrm{r}^{*} \mathrm{~s}^{*} \mathrm{u}$ |
| y | Other fractures avoided due to treatment | 144 | =i*p*r*s*v |
| z | Life years gained (deaths avoided) due to screening, osteoporotic hip fractures | 21 | $=j^{*} p^{*} q^{*} s^{*} t$ |
| aa | Life years gained (deaths avoided) due to screening, osteoporotic vertebral fractures | 2.7 | =k*p*r*s*u |
| ab | Life years gained (deaths avoided) due to screening, osteoporotic other fractures | 10 | $=l^{*} p^{*} r^{*}{ }^{*}{ }^{*}$ |
| ac | QALYs gained due to screening in those living with osteoporotic hip fractures | 52.4 | $=m * p * q * s * t$ |
| ad | QALYs gained due to screening in those living with osteoporotic vertebral fractures | 1.2 | $=n * p * r * s * u$ |
| ae | QALYs gained due to screening in those living with osteoporotic other fractures | 4.3 | =o*p*r*s*v |
| af | Total QALYs gained due to screening (going from 0\% to 57.8\%) | 91 | $=z+a a+a b+a c+a d+a e$ |

$V=$ Estimates from the literature
For the sensitivity analysis, we modified a number of major assumptions and recalculated the CPB as follows:

- Assume that the hazard ratio (HR) for death after hip fracture is reduced from 2.87 to 2.52, the HR for death after vertebral fractures is reduced from 1.82 to 1.28 and the HR for death after other fractures is reduced from 1.38 to 1.18 (Table 4): $\mathrm{CPB}=88$
- Assume that the hazard ratio (HR) for death after hip fracture is increased from 2.87 to 3.27 , the HR for death after vertebral fractures is increased from 1.82 to 2.57 and the HR for death after other fractures is increased from 1.38 to 1.62 (Table 4): $\mathrm{CPB}=$ 96
- Assume that the hip fracture reduction rate is reduced from $23 \%$ to $8 \%$ (Table 7 , row $t$ ), the vertebral fracture reduction rate is reduced from $26 \%$ to $12 \%$ (Table 7 , row $u$ ) and the other fracture reduction rate is reduced from $20 \%$ to $9 \%$ (Table 7 , row $v$ ): $\mathrm{CPB}=34$
- Assume that the hip fracture reduction rate is increased from $23 \%$ to $36 \%$ (Table 7, row $t$ ), the vertebral fracture reduction rate is increased from $26 \%$ to $38 \%$ (Table 7, row $u$ ) and the other fracture reduction rate is increased from $20 \%$ to $29 \%$ (Table 7, row $v$ ): $\mathrm{CPB}=141$


## Modelling Cost-Effectiveness

In this section, we will calculate the CE associated with screening for osteoporosis in women ages 65 and older.

In modelling CE, we made the following assumptions:

- We model that $57.8 \%{ }^{448}$ of 65 year old women are referred to and receive a bone density (DXA) scan (Table 8 , row $b$ ). This rate takes into account both physician adherence (willingness to make the referral) and patient adherence (willingness to get the scan done).
- The cost of each 10 minute primary care provider office visit is $\$ 34.85$ (Reference Document) (Table 8, row $d$ )
- The value of patient time for each visit to a primary care office and for bone density scanning is $\$ 59.38$ (Reference Document) (Table 8, row $e$ ).
- The proportion of each office visit attributable to screening is $50 \%$ (Reference Document) (Table 8, row $f$ ).
- We model that all those who receive a DXA scan have also visited their primary care provider to receive the referral for the scan. During this appointment, a risk assessment (e.g. FRAX ${ }^{\circledR}$ ) could be conducted within the portion of the office visit attributable to screening. The FRAX ${ }^{\circledR}$ tool adapted for the Canadian population can be found online at no cost. ${ }^{449}$
- According to the BC Medical Services Plan Fee-For-Service Payment Analysis for 2012/13 - 2016/17, a single area bone density scan (fee item 8688) averages $\$ 66.94$ per scan. Adding a second area (fee item 8689 ) costs an additional $\$ 45.88$ per scan. A second area scan occurred at a rate of approximately $95.2 \%$ of single area scans. ${ }^{450}$
- We assume that bone scans to determine bone mineral density are conducted by means of DXA and model the cost of the average bone scan as $\$ 66.94+(0.952 *$ $45.88)=\$ 110.62($ Table 8 , row $h)$.
- In the study by Patrick et al. ${ }^{451}$ of 19,987 individuals initiating treatment with bisphosphonates, they found that $31.8 \%$ had a cumulative proportion of days covered

[^107](i.e. the proportion of days taking medication) between $0-19 \%, 11.3 \%$ had a proportion of days covered (PDC) between $20-39 \%, 8.8 \%$ had a PDC between $40-$ $59 \%, 11.5 \%$ had a PDC between $60-79 \%$ and $36.5 \%$ had a PDC between $80-100 \%$. (Table 8, rows $l$ to $p$ ). Their study assessed medication compliance rates over a 300day period.

- For modelling purposes, we assume that each PDC group has a compliance rate at the midpoint of their range. Groups with a PDC of between $0-79 \%$ stop taking medication after 300 days. For the high compliance group, we assume that the medication is taken for 5 years in the base model (Table 8 , row $y$ ). In the sensitivity analysis, we model 5 years of taking medication, followed by a 5-year medication 'holiday' followed by a further 5 years of taking medication.
- Alendronate is the most commonly prescribed bisphosphonate in BC and is typically prescribed to be taken orally once per week at a dose of 70 mg . ${ }^{452}$
- We model weekly treatment with 70mg alendronate. The cost per 70mg pill ranges from $\$ 2.17$ - $\$ 13.88$ in BC. ${ }^{453}$ Only two records for BC, however, showed a price above $\$ 3.21$. We assume pricing above $\$ 3.21$ per 70 mg are outliers and model using the mid-point of the $\$ 2.17$ - $\$ 3.21$ range for the pills, or $\$ 2.69$. The dispensing fee ranges from $\$ 4.49-\$ 13.99$, with only a single dispensing fee below $\$ 9.95$. We assume a dispensing fee at the midpoint of \$9.95-\$13.99 (or \$11.97) and assume a 3-month dose is dispensed each time.
- We model the annual cost of treatment as $\$ 187.76((\$ 2.69 * 52)+(4 * \$ 11.97))$. Translating this into a daily cost results in $\$ 0.51$ / day ( $\$ 187.76 / 365$ ). Using the low and high numbers of the ranges above (excluding outliers), we use a range of between $\$ 0.42$ and $\$ 0.62$ / day in the sensitivity analysis (Table 8 , row $v$ ).
- A December 20, 2018 publication by Reid and colleagues assessed the efficacy of 4 infusions of 5 mg zoledronate (or zoledronic acid) at 18-month intervals vs. placebo in older women (mean age of 71) with osteopenia. ${ }^{454}$ They noted a $37 \%$ (HR of 0.63 , $94 \%$ CI $0.50-0.79$ ) reduction in fragility fractures in women receiving zoledronate. The efficacy of such a reduction in medication dose and frequency is encouraging for the potential compliance with and cost of treatment.
- In comparing less-frequent zoledronic acid infusions with more frequent bisphosphonate treatment regimes, Lozano and Sanchez-Fidalgo report that "patients appear to have (a) preference for less frequent dosing. Switching from oral to intravenous therapy...may allow obtaining better outcomes in adherence to osteoporosis treatment. ${ }^{" 455}$
- Potential changes in adherence and the costs associated with zoledronic acid infusions are two important variables that should be considered in future updates of

[^108]this model, should the results observed by Reid and colleagues ${ }^{456}$ be confirmed for patients with osteoporosis.

- We model one additional visit to a primary care provider for monitoring medication for those with low compliance (PDC of $0-79 \%$ ) (Table 8, row ab) and one annual visit to a primary care provider for monitoring medication for those with high compliance (PDC of $80-100 \%$ ) (Table 8, row $i$ ).
- A recent Canadian study by Hopkins et al. estimated the annual direct medical costs of a hip fracture to be $\$ 61,540$, the cost of a vertebral fracture to be $\$ 25,965$ and the cost of "other" fractures to be $\$ 13,579$ (all in 2014 CAD). ${ }^{457}$ Costs included acute care, rehabilitation care, long term care, home care, outpatient physician services and mobility devices.
- We adjusted the costs calculated by Hopkins et al. to 2017 CAD and use $\$ 62,152$ for the total cost per hip fracture (Table 8, row ai), \$26,223 (Table 8, row $a j$ ) per vertebral fracture and $\$ 13,714$ for all other fractures (Table 8, row $a k$ ).

Based on these assumptions, the CE associated with screening for osteoporosis in women ages 65 and older is cost saving ( $-\$ 29,412 /$ QALY) (see Table 8, row $a r$ ).

[^109]Table 8: Cost Effectiveness of Osteoporosis Screening in Women 65+

| Row Label | Variable | Base case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Population in cohort, age 65 | 18,456 | BC Life Table |
| b | Proportion screened for osteoporosis | 0.578 | Table 7, row p |
| c | Number in cohort receiving bone density screen (DXA) | 10,667 | $=\mathrm{a}$ * b |
| d | Cost of 10 minute office visit | \$34.85 | Ref Doc |
| e | Value of patient time and travel for office visit | \$59.38 | Ref Doc |
| f | Portion of 10-minute visit for screening | 50\% | Ref Doc |
| g | Cost of initial screening visit | \$502,592 | = ${ }^{*} \mathrm{f}^{*}(\mathrm{~d}+\mathrm{e})$ |
| h | Bone density screening cost, per screen | \$110.62 | $\checkmark$ |
| i | Cost of bone density screening | \$1,813,447 | $c^{*}(\mathrm{e}+\mathrm{h})$ |
| j | Number of osteoporotic patients at age 65 | 3,543 | Table 1 |
| k | Number of osteoporotic patients identified via screening | 2,048 | = ${ }^{*}$ b |
| I | Percent of patients with proportion of days covered (PDC) 0-19\% | 31.8\% | V |
| m | Percent of patients with PDC of 20-39\% | 11.3\% | $\checkmark$ |
| n | Percent of patients with PDC of 40-59\% | 8.8\% | $\checkmark$ |
| 0 | Percent of patients with PDC of 60-79\% | 11.5\% | $\checkmark$ |
| p | Percent of patients with PDC of $80-100 \%$ | 36.5\% | Table 7, row s |
| q | Average days taking medication - PDC 0-19\% group | 30 | $=300 * 0.10$ |
| r | Average days taking medication - PDC 20-39\% group | 90 | $=300 * 0.30$ |
| s | Average days taking medication - PDC 40-59\% group | 150 | $=300 * 0.50$ |
| t | Average days taking medication - PDC 60-79\% group | 210 | $=300 * 0.70$ |
| u | Total days taking medication-PDC 0-79\% group | 116,866 | $\begin{gathered} =(\mathrm{k} * \mathrm{I} * \mathrm{q})+(\mathrm{k} * \mathrm{~m} * \mathrm{r})+(\mathrm{k} * \mathrm{n} \\ * \mathrm{~s})+\left(\mathrm{k} * \mathrm{o}^{*} \mathrm{t}\right) \end{gathered}$ |
| v | Daily cost of medication | \$0.51 | $\checkmark$ |
| w | Total cost of medication - PDC 0-79\% | \$60,117 | = ${ }^{*} \mathrm{v}$ |
| x | Average days taking medication - PDC 80-100\% group | 329 | $=365$ * 0.90 |
| y | Years of treatment - PDC 80-100\% group | 5 | $\checkmark$ |
| z | Total days taking medication - PDC 80-100\% group | 1,227,879 | =k*p*x*y |
| aa | Total cost of medication - PDC 80-100\% group | \$631,634 | $=z^{*} \mathrm{v}$ |
| ab | Annual office visits required to monitor medication | 1 | Assumption |
| ac | Cost of annual visits to monitor medication - PDC 0-79\% group | \$61,276 | $=(1-p) * k * a b *(d+e) * f$ |
| ad | Cost of annual visits to monitor medication - PDC 80-100\% group | \$176,108 | = ${ }^{*} \mathrm{k}^{*} \mathrm{y}^{*} \mathrm{ab}^{*}(\mathrm{~d}+\mathrm{e}){ }^{*} \mathrm{f}$ |
| ae | Total cost of screening and treatment | \$3,245,174 | = $\mathrm{g}+\mathrm{i}+\mathrm{w}+\mathrm{aa}+\mathrm{ac}+\mathrm{ad}$ |
|  | Potential Costs Avoided |  |  |
| af | Total hip fractures avoided | 63 | Table 7, row w |
| ag | Total vertebral fractures avoided | 18 | Table 7, row x |
| ah | Other fractures avoided | 144 | Table 7, row y |
| ai | Average cost per hip fracture in the year following the fracture | \$62,152 | V |
| aj | Average cost per vertebral fracture in the year following the fracture | \$26,223 | $\checkmark$ |
| ak | Average cost per other fracture in the year following the fracture | \$13,714 | $\checkmark$ |
| al | Total costs avoided | \$6,367,537 | $=\left(a f\right.$ * ai) $+\left(a g^{*}\right.$ aj) + (ah * ak) |
| am | Net cost of intervention | -\$3,122,363 | =ae - al |
| an | QALYs gained | 91 | Table 7, row af |
| ao | Cost effectiveness (CE) of intervention, \$/QALY | -\$34,145 | $=\mathrm{am} / \mathrm{an}$ |
| ap | Net Cost of Intervention (1.5\% Discount) | -\$2,248,682 | Calculated |
| aq | Net QALYs Gained (1.5\% Discount) | 76 | Calculated |
| ar | Cost Effectiveness (CE) of Intervention, \$/QALY (1.5\% Discount) | -\$29,412 | = ap / aq |

V $=$ Estimates from the literature
For the sensitivity analysis, we modified a number of major assumptions and recalculated the CE as follows:

- Assume that the hazard ratio (HR) for death after hip fracture is reduced from 2.87 to 2.52, the HR for death after vertebral fractures is reduced from 1.82 to 1.28 and the HR for death after other fractures is reduced from 1.38 to 1.18 (Table 4): CE $=-\$ 30,527$
- Assume that the hazard ratio (HR) for death after hip fracture is increased from 2.87 to 3.27 , the HR for death after vertebral fractures is increased from 1.82 to 2.57 and the HR for death after other fractures is increased from 1.38 to 1.62 (Table 4):
CE $=-\$ 28,234$
- Assume that the hip fracture reduction rate is reduced from $23 \%$ to $8 \%$ (Table 7, row $t$ ), the vertebral fracture reduction rate is reduced from $26 \%$ to $12 \%$ (Table 7 , row $u$ )
and the other fracture reduction rate is reduced from $20 \%$ to $9 \%$ (Table 7, row $v$ ): CE $=\$ 38,997$
- Assume that the hip fracture reduction rate is increased from $23 \%$ to $36 \%$ (Table 7, row $t$ ), the vertebral fracture reduction rate is increased from $26 \%$ to $38 \%$ (Table 7, row $u$ ) and the other fracture reduction rate is increased from $20 \%$ to $29 \%$ (Table 7, row $v$ ): $\mathrm{CE}=-\$ 43,257$
- Assume that the cost of treatment is increased from $\$ 0.51$ / day to $\$ 0.61$ / day (Table 8, row $v$ ): $\mathrm{CE}=-\$ 27,765$
- Assume that the cost of treatment is reduced from $\$ 0.51$ / day to $\$ 0.42$ / day (Table 8 , row $v$ ): $\mathrm{CE}=-\$ 31,060$
- Assume that treatment pattern for the PDC $80-100 \%$ group changes from five years of treatment to five years of treatment followed by five years untreated followed by another five years of treatment, for a total treatment time of 10 years (Table 8, row $y$ ): CE $=-\$ 20,574$
A number of others have calculated the cost-effectiveness of screening and treatment options for osteoporosis in women ages 65 and older. ${ }^{458,459,460,461}$ In a Canadian cost-effectiveness analysis published in 2006, Goeree and colleagues estimated a CE of $\$ 32,571$ / QALY for etidronate when compared with no intervention. ${ }^{462}$ The CE / QALY was $\$ 38,623$ for alendronate and $\$ 114,070$ for raloxifene. Their study made a number of different key assumptions than we have. First, they assumed that $100 \%$ of patients with osteoporosis would adhere to medication regimens for a five-year period. Based on a large real-world adherence study published in $2010,{ }^{463}$ we assume that just $36.5 \%$ of patients with osteoporosis would adhere to medication regimens for a five year period. In addition, their estimated annual cost of drugs was between $\$ 546$ and $\$ 969$ compared to our base case scenario of $\$ 188$. Applying an annual drug cost of $\$ 546$ to our model results in a cost / QALY of $-\$ 12,608$. An annual drug cost of $\$ 969$ would increase the cost / QALY to $\$ 7,234$.


## Summary

The clinically preventable burden ( CPB ) associated with screening for, and treatment of, osteoporosis in females ages 65 and older in order to prevent fractures is 91 quality-adjusted life years (QALYs) while the cost-effectiveness (CE) is estimated at a saving of \$29,412 per QALY (see Table 9).

[^110]| Birth Cohort of 40,000 Summary |  |  |  |
| :---: | :---: | :---: | :---: |
|  | Base <br> Case | Ran |  |
| CPB (Potential QALYs Gained) |  |  |  |
| Assume No Current Service |  |  |  |
| 1.5\% Discount Rate | 76 | 28 | 118 |
| 3\% Discount Rate | 65 | 24 | 100 |
| 0\% Discount Rate | 91 | 34 | 141 |
| CE (\$/QALY) including patient time costs |  |  |  |
| 1.5\% Discount Rate | -\$29,412 | -\$43,257 | \$38,997 |
| 3\% Discount Rate | -\$24,048 | -\$40,489 | \$57,000 |
| 0\% Discount Rate | -\$34,145 | -\$45,672 | \$22,976 |
| CE (\$/QALY) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | -\$43,755 | -\$52,552 | \$81 |
| 3\% Discount Rate | -\$40,996 | -\$51,474 | \$11,028 |
| 0\% Discount Rate | -\$46,171 | -\$53,466 | -\$9,663 |

## Screening for Abdominal Aortic Aneurysms

## United States Preventive Services Task Force Recommendations ${ }^{464}$

The USPSTF recommends 1-time screening for AAA with ultrasonography in men aged 65 to 75 years who have ever smoked. (B recommendation).

## Canadian Task Force on Preventive Health Care Recommendations ${ }^{465}$

We recommend one-time screening with ultrasonography for AAA of men aged 65 to 80 years (weak recommendation; moderate quality of evidence).

We recommend not screening men older than 80 years of age for AAA (weak recommendation; low quality of evidence).

The Canadian Task force acknowledged "evidence showing increased risk of AAA among smokers" but did not make a separate recommendation on screening this population "because there is no evidence on outcomes of screening smokers for AAA." ${ }^{466}$

## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with screening for abdominal aortic aneurysms in males ages 65 to 75 who have ever smoked.

An abdominal aortic aneurysm is conventionally diagnosed when the diameter of the aorta below the kidneys is $30 \mathrm{~mm}(3.0 \mathrm{~cm})$ or greater. ${ }^{467}$

The USPSTF considers an "ever-smoker" someone who has smoked at least 100 cigarettes in their lifetime. ${ }^{468}$

Unless otherwise noted, we apply these conventions and definitions in our modeling.
In modelling CPB, we made the following assumptions:

- The single screen recommended by the USPSTF is conducted at age 65.
- Jacomelli and colleagues report that the National Health Service in England's AAA screening programme had mean uptake across the country of $78.1 \%$, but varied regionally between $61.7-85.8 \%{ }^{469} \mathrm{We}$ use $85.8 \%$ as the best in the world screening rate for AAA.

[^111]- The large, population-based randomized controlled trials (RCTs) used by the USPSTF in making their recommendation found an abdominal aortic aneurysm (AAA) in $4.0-7.7 \%$ of male screening participants. ${ }^{470}$
- Citing more recent epidemiologic evidence from Europe and New Zealand, the USPSTF acknowledged a "substantial decrease in AAA prevalence in men aged 65 years or older in the past 2 decades" ${ }^{" 471}$ and referenced a study by Svensjö et al. citing an AAA prevalence rate of $1.7 \%$ in Sweden. ${ }^{472}$
- In the UK, the AAA prevalence rate in 65 -year old men has decreased from $5.0 \%$ in 1991 to $1.3 \%$ in 2015. ${ }^{473}$ In Denmark, the prevalence rate in 65 -year old men was $2.6 \%$ during 2008-2011. ${ }^{474}$
- For modelling purposes, we use an AAA prevalence rate in 65-year old men of $2.35 \%$ (Table 5, row $e$ ). Using $2.35 \%$ prevalence in our model brings the model results with screening reasonably close to actual BC results. The $2.35 \%$ prevalence rate used is between the values reported for the UK and Denmark.
- The USPSTF rated the quality of the population-based randomized controlled trials (RCTs) used by the USPSTF in making their recommendation. The USPSTF considered the Multicentre Aneurysm Screening Study (MASS) and the Viborg AAA studies as "good-quality", and the Chichester and Western Australia AAA studies as "fair-quality". ${ }^{475}$ Neither good-quality study included men over the age of 74 . On the other hand, both fair-quality studies included older men up to ages 80 (Chichester) and 83 (Western Australia).
- The prevalence of AAA increases with increasing age. ${ }^{476}$
- In the MASS study, $4.9 \%$ of screened men were diagnosed with AAA and the total AAA-related death rate was 109 per 100,000 person years in the control group. ${ }^{477}$ In the Viborg study, $4.0 \%$ of screened men were diagnosed with AAA and the total AAA-related death rate was 87 per 100,000 person years in the control group. ${ }^{478}$

[^112]- Based on 25 years of experience with an ultrasound screening program for AAA in the UK, Oliver-Williams and colleagues report that while the "prevalence of screendetected small and medium AAAs has decreased over the past 25 years, ...growth rates have remained similar. Men with a subaneurysmal aorta at age 65 years have a substantial risk of developing a large AAA by the age of 80 years. ${ }^{\prime 479}$
- For modelling purposes, we assume that the death rate / 100,000 person years of 98.0 observed in the control groups of the MASS and Viborg studies would be reduced linearly to 51.7 / 100,000 person years due to the lower estimated prevalence of AAA ( $2.35 \%$ ) used in our model (see Table 1).

Table 1: Screening for Abdominal Aortic Aneurysm Men Ages 65+ Adjusted Study Results Based on Lower AAA Prevalence

| Study | USPSTF Study Rating | Study Prevalence of AAA | Study Death Rate in Control Group per 100,000 person years | Model <br> Prevalence <br> of AAA | Adjusted <br> Death Rate per 100,000 person years |
| :---: | :---: | :---: | :---: | :---: | :---: |
| MASS (Thompson et al., 2012) | Good | 4.9\% | 109 | 2.35\% | 52.3 |
| Viborg (Lindholt et al.) | Good | 4.0\% | 87 | 2.35\% | 51.1 |
| Average of Good Quality Studies |  |  | 98.0 |  | 51.7 |

- As early as 1998 , Semmens et al. reported a decline in AAA-related emergency and elective procedures in Western Australia, ahead of similar results being reported in Europe and theorized that this may be due to "significant changes in the health of the Australian community" including "the success of the anti-smoking movement". ${ }^{480}$
- In Sweden, Johansson and colleagues observed that AAA mortality declined from 36 to 10 deaths per 100,000 for men aged 65-74 between the early 2000s and 2015. ${ }^{481}$ They note, however, that only an estimated $30 \%$ of this reduction was associated with the introduction of screening for AAA and that $70 \%$ is due to other factors, most notably a reduction in smoking. Between 1970 and 2010, the prevalence of smoking in Sweden decreased from $44 \%$ to $15 \%$. ${ }^{482}$
- In a 2018 systematic review and meta-analysis of tobacco smoking and AAA, Aune and colleagues report that the relative risk of AAA in current smokers is 4.87 ( $95 \%$ CI $3.93-6.02$ ) and in former smokers is $2.10(95 \%$ CI $1.76-2.50)$ compared to never smokers. ${ }^{483}$

[^113]- The Canadian Tobacco, Alcohol and Drugs Survey, 2017 indicated that $16.8 \%$ (95\% CI 11.6 - 22.0\%) of men 45+ in BC are current smokers, $36.3 \%$ ( $95 \%$ CI 29.6 $43.0 \%$ ) are former smokers and $47 \%(95 \%$ CI $39.6-54.3)$ have never smoked. ${ }^{484}$
- Based on Canadian Community Health Survey data from 2014, $12.9 \%$ of BC men ages 65-69 are daily or occasional smokers. ${ }^{485}$
- For modelling purposes, we assume that $12.9 \%$ of men 65 years of age are current smokers (Table 5, row $d$ ), $47 \%$ are never smokers (Table 5, row $b$ ) and the balance $(40.1 \%)$ are former smokers (Table 5, row $c$ ).
- In Table 2 we combine the estimated AAA-related death rate for the population as a whole (51.7 / 100,000 person years, see Table 1), the proportion of 65 year old BC men by smoking category and the relative risk of AAA for current-smokers, formersmokers and never-smokers. At the same time, we calculated the prevalence of AAA in each group, using our model prevalence of $2.35 \%$ for the whole population (Table 5 , row $e$ ).
- The results suggest a prevalence of $1.21 \%$ (Table 5 , row $f$ ) and an AAA-related death rate of 26.6 / 100,000 in never-smokers, a prevalence of $2.54 \%$ (Table 5, row $g$ ) and an AAA-related death rate of 55.9 / 100,000 in former-smokers and a prevalence of $5.90 \%$ (Table 5, row $h$ ) and an AAA-related death rate of 129.7 / 100,000 in currentsmokers.

Table 2: Screening for Abdominal Aortic Aneurysm Men 65+
AAA Prevalence and Death Rates by Smoking Category

|  | Total | Never-Smoker | Former-Smoker | Current-Smoker |
| :---: | :---: | :---: | :---: | :---: |
| Proportion of Population | 1.00 | 0.470 | 0.401 | 0.129 |
| Relative Risk of AAA |  | 1.00 | 2.10 | 4.87 |
| Prevalence of AAA | 2.35\% | 1.21\% | 2.54\% | 5.90\% |
| Death Rate per 100,000 | 51.7 | 26.6 | 55.9 | 129.7 |

- Howard et al. report the incidence of acute AAA events to be 55 / 100,000 per year in 65-74 year olds and 112 / 100,000 per year in 75-84 year olds. Of these acute AAA events, $59.2 \%$ were fatal within 30 days. ${ }^{486}$ This works out to AAA-related death rates of $32.6(55 * 0.592)$ and $66.3(112 * 0.592) / 100,000$ for 65-74 and 75-84 year olds respectively.
- Howard and colleagues also report that $22.3 \%$ of incident AAA-events took place in $65-74$ year olds, with only $13.1 \%$ of AAA-related deaths occurring in this age group. ${ }^{487}$
- We adjust the rates for age groups from $65-74$ and $75-84$ to reflect that $86.9 \%$ of AAA-related deaths are in the $75+$ age group, while ensuring the total population rates still reflect what was calculated in Table 2. The deaths and life-years lost in a

[^114]cohort of BC men 65+ due to AAA is shown in Table 3. We model from AAA screening at age 65 through to age 84 , in keeping with the average life expectancy of 19.5 years for a 65 year old male from the BC Life Table.

- AAA is usually asymptomatic prior to rupture, ${ }^{488}$ therefore reduced quality of life in those living with AAA is not presented in Table 3 or considered in our model.
- Table 3 indicates that, in our birth cohort, we would expect 36 AAA-related deaths in male never-smokers (Table 5, row $p$ ), 65 AAA-related deaths in former-smokers (Table 5, row $q$ ) and 48 AAA-related deaths in current-smokers (Table 5, row $r$ ). These 149 AAA-related deaths represent $1.90 \%$ of the total 7,872 deaths in the cohort between the ages of 65 and 84 . Research from other jurisdictions suggests an AAArelated death rate of between $1-2 \%$ of total deaths. ${ }^{499,490}$ These 149 deaths would result in the loss of $1,068(259+464+346)$ QALYs in our cohort.
- BC Vital Statistics annual reports provide a detailed listing (by ICD-10 code) of annual deaths by age and sex. ICD-10 code I71 is for deaths due to "aortic aneurysm \& dissection." If we combine deaths due to ICD-10 code I71 from the $2013^{491}$, $2014^{492}$ and $2015^{493}$ BC Vital Statistics annual reports, $0.78 \%$ of deaths in males 65 79 and $0.72 \%$ of deaths in males 80 and over were attributed to ICD-10 code I71. In males over $65,0.74 \%$ of deaths were attributed to ICD-10 code I71. This proportion of deaths attributable to ICD-10 code I71 is considerably lower than our modelled estimate of $1.90 \%$. Using cause of death data from vital statistics can be somewhat challenging as research has indicted that at least $15 \%$ of all deaths are miscoded in vital statistics data in the US and Canada. ${ }^{494}$ It is possible, therefore, that the $0.74 \%$ is an underrepresentation of the actual proportion of deaths due to AAA in BC males 65 years of age and older due to AAA. We include the $0.74 \%$ in our sensitivity analysis.

[^115]\begin{tabular}{|c|c|c|c|c|c|c|c|c|c|c|c|c|c|c|c|}
\hline \multicolumn{16}{|c|}{\begin{tabular}{l}
Table 3: Screening for Abdominal Aortic Aneurysm in Men 65+ \\
Deaths and Life Years Lost Due to Abdominal Aortic Aneurysm In a BC Birth Cohort of 40,000
\end{tabular}} \\
\hline Age \& \# in Cohort \& Proportion of Population \& \begin{tabular}{l}
ever Smokers \\
AAA-Related \\
Deaths per 100,000 person years
\end{tabular} \& \begin{tabular}{l}
AAA- \\
Related Deaths
\end{tabular} \& Proportion of Population \& \begin{tabular}{l}
rmer Smokers \\
AAA-Related \\
Deaths per 100,000 person years
\end{tabular} \& \begin{tabular}{l}
AAA- \\
Related \\
Deaths
\end{tabular} \& Proportion of Population \& \begin{tabular}{l}
rrent Smokers \\
AAA-Related \\
Deaths per 100,000 person years
\end{tabular} \& \begin{tabular}{l}
AAA- \\
Related \\
Deaths
\end{tabular} \& AAADeaths in Ever Smokers \& Life Expectancy \& Life Years

Never
Smokers \& Lost Due to

Former

Smokers \& | o Death |
| :--- |
| Current |
| Smokers | <br>

\hline 65 \& 17,559 \& 47.0\% \& 6.1 \& 0.5 \& 40.1\% \& 12.9 \& 0.9 \& 12.9\% \& 29.8 \& 0.7 \& 1.6 \& 20 \& 10.1 \& 18.1 \& 13.5 <br>
\hline 66 \& 17,370 \& 47.0\% \& 6.1 \& 0.5 \& 40.1\% \& 12.9 \& 0.9 \& 12.9\% \& 29.8 \& 0.7 \& 1.6 \& 19 \& 9.5 \& 17.0 \& 12.7 <br>
\hline 67 \& 17,164 \& 47.0\% \& 6.1 \& 0.5 \& 40.1\% \& 12.9 \& 0.9 \& 12.9\% \& 29.8 \& 0.7 \& 1.5 \& 18 \& 8.9 \& 15.9 \& 11.9 <br>
\hline 68 \& 16,940 \& 47.0\% \& 6.1 \& 0.5 \& 40.1\% \& 12.9 \& 0.9 \& 12.9\% \& 29.8 \& 0.7 \& 1.5 \& 17 \& 8.3 \& 14.8 \& 11.1 <br>
\hline 69 \& 16,697 \& 47.0\% \& 6.1 \& 0.5 \& 40.1\% \& 12.9 \& 0.9 \& 12.9\% \& 29.8 \& 0.6 \& 1.5 \& 16 \& 7.7 \& 13.8 \& 10.3 <br>
\hline 70 \& 16,434 \& 47.0\% \& 6.1 \& 0.5 \& 40.1\% \& 12.9 \& 0.8 \& 12.9\% \& 29.8 \& 0.6 \& 1.5 \& 15 \& 7.1 \& 12.7 \& 9.5 <br>
\hline 71 \& 16,147 \& 47.0\% \& 6.1 \& 0.5 \& 40.1\% \& 12.9 \& 0.8 \& 12.9\% \& 29.8 \& 0.6 \& 1.5 \& 14 \& 6.5 \& 11.7 \& 8.7 <br>
\hline 72 \& 15,837 \& 47.0\% \& 6.1 \& 0.5 \& 40.1\% \& 12.9 \& 0.8 \& 12.9\% \& 29.8 \& 0.6 \& 1.4 \& 13 \& 5.9 \& 10.6 \& 7.9 <br>
\hline 73 \& 15,500 \& 47.0\% \& 6.1 \& 0.4 \& 40.1\% \& 12.9 \& 0.8 \& 12.9\% \& 29.8 \& 0.6 \& 1.4 \& 12 \& 5.4 \& 9.6 \& 7.2 <br>
\hline 74 \& 15,136 \& 47.0\% \& 6.1 \& 0.4 \& 40.1\% \& 12.9 \& 0.8 \& 12.9\% \& 29.8 \& 0.6 \& 1.4 \& 11 \& 4.8 \& 8.6 \& 6.4 <br>
\hline 75 \& 14,743 \& 47.0\% \& 53.9 \& 3.7 \& 40.1\% \& 113.1 \& 6.7 \& 12.9\% \& 262.3 \& 5.0 \& 11.7 \& 10 \& 37.3 \& 66.9 \& 49.9 <br>
\hline 76 \& 14,318 \& 47.0\% \& 53.9 \& 3.6 \& 40.1\% \& 113.1 \& 6.5 \& 12.9\% \& 262.3 \& 4.8 \& 11.3 \& 9 \& 32.6 \& 58.4 \& 43.6 <br>
\hline 77 \& 13,861 \& 47.0\% \& 53.9 \& 3.5 \& 40.1\% \& 113.1 \& 6.3 \& 12.9\% \& 262.3 \& 4.7 \& 11.0 \& 8 \& 28.1 \& 50.3 \& 37.5 <br>
\hline 78 \& 13,370 \& 47.0\% \& 53.9 \& 3.4 \& 40.1\% \& 113.1 \& 6.1 \& 12.9\% \& 262.3 \& 4.5 \& 10.6 \& 7 \& 23.7 \& 42.4 \& 31.7 <br>
\hline 79 \& 12,844 \& 47.0\% \& 53.9 \& 3.3 \& 40.1\% \& 113.1 \& 5.8 \& 12.9\% \& 262.3 \& 4.3 \& 10.2 \& 6 \& 19.5 \& 35.0 \& 26.1 <br>
\hline 80 \& 12,283 \& 47.0\% \& 53.9 \& 3.1 \& 40.1\% \& 113.1 \& 5.6 \& 12.9\% \& 262.3 \& 4.2 \& 9.7 \& 5 \& 15.5 \& 27.9 \& 20.8 <br>
\hline 81 \& 11,686 \& 47.0\% \& 53.9 \& 3.0 \& 40.1\% \& 113.1 \& 5.3 \& 12.9\% \& 262.3 \& 4.0 \& 9.3 \& 4 \& 11.8 \& 21.2 \& 15.8 <br>
\hline 82 \& 11,053 \& 47.0\% \& 53.9 \& 2.8 \& 40.1\% \& 113.1 \& 5.0 \& 12.9\% \& 262.3 \& 3.7 \& 8.8 \& 3 \& 8.4 \& 15.0 \& 11.2 <br>
\hline 83 \& 10,386 \& 47.0\% \& 53.9 \& 2.6 \& 40.1\% \& 113.1 \& 4.7 \& 12.9\% \& 262.3 \& 3.5 \& 8.2 \& 2 \& 5.3 \& 9.4 \& 7.0 <br>
\hline 84 \& 9,688 \& 47.0\% \& 53.9 \& 2.5 \& 40.1\% \& 113.1 \& 4.4 \& 12.9\% \& 262.3 \& 3.3 \& 7.7 \& 1 \& 2.5 \& 4.4 \& 3.3 <br>
\hline Total \& \& \& 26.6 \& 36 \& \& 55.9 \& 65 \& \& 129.7 \& 48 \& 113 \& \& 259 \& 464 \& 346 <br>
\hline
\end{tabular}

- There are three primary AAA-related modes of death considered by the randomized controlled trials: death as a result of AAA rupture before receiving emergency surgery at a hospital, death as a result of AAA rupture after receiving emergency surgery, and death due to complications following elective surgery.
- Only one good quality USPSTF referenced study reported on rates of elective and emergency surgery in the control and screening intervention groups; the Viborg study reported by Lindholt and colleagues. ${ }^{495}$ They report an elective surgery rate of 70 / 100,000 and an emergency surgery rate of $70 / 100,000$ in the control population at a reported AAA prevalence of $4.0 \%$.
- We model that these rates would be reduced linearly to 41 / 100,000 person years (Table 5, row $v$ ) and $41 / 100,000$ person years (Table 5, row $a c$ ) for elective and emergency procedures respectively due to the lower estimated prevalence of AAA (2.35\%) used in our model (see Table 4).

| Adjusted Surgery Rates Based on Lower AAA Prevalence ${ }^{1}$ |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: |
| Variable | Study Prevalence of AAA | Incidence per 100,000 person years | Model Prevalence of AAA | Adjusted Incidence per 100,000 person years |
| Elective Operations, Control | 4.0\% | 70 | 2.35\% | 41 |
| Acute Operation, with Rupture, Control | 4.0\% | 57 | 2.35\% | 33 |
| Acute Operation, without rupture, Control | 4.0\% | 13 | 2.35\% | 8 |
| Total for Acute Operations, Control | 4.0\% | 70 | 2.35\% | 41 |

${ }^{1}$ Source: Lindholt et al. (2010)

[^116]- Guirguis-Blake and colleagues conducted a pooled analysis of RCTs reporting 13-15 year follow up results and calculated the following relative risks in the screening group: ${ }^{496}$
- RR of elective operations for AAA: 2.15 ( $95 \%$ CI, $1.89-2.44$ )
- RR of emergency operations for AAA: 0.52 ( $95 \%$ CI, $0.40-0.66$ )
- RR of AAA-related mortality: 0.58 ( $95 \% \mathrm{CI}, 0.39-0.88$ )
- We model the RR after the pooled analysis by Guirguis-Blake et al. with a relative risk of elective operations of 2.15 (Table 5, row $a l$ ), a relative risk of emergency operations of 0.52 (Table 5 , row $a u$ ), and an overall relative risk of AAA-related death of 0.58 in the screening group (Table 5, row $a z$ ).
- There are a number of cases of asymptomatic AAA that could be found without screening. This number ranges from $7-25 \%$ in economic analyses and studies reporting this variable. ${ }^{497,498,499,500,501}$
- For modelling purposes, we use the mid-point between $7 \%$ and $25 \%$ ( $13 \%$ ) and vary this from $7-25 \%$ in our sensitivity analysis (Table 5, row $a k$ ).
- Reporting on the years 2003 - 2004 for Canada, Forbes et al. reported that $8.9 \%$ of elective AAA-repair was carried out by endovascular surgery, with the balance being open surgery. ${ }^{502}$
- Jetty and Husereau reported on Canadian trends from 2004-2009 and reported that endovascular aneurysm repair (EVAR) rates rose from $11.5 \%$ to $35.5 \%$ in Canada during that time. They also report substantial regional differences in elective endovascular repair rates, from a low of $15.8 \%$ in Manitoba to a high of $45.0 \%$ in BC in 2009. BC's rate increased each year from $7.5 \%$ in 2005 to $45.0 \%$ in 2009. ${ }^{503}$
- Of the 1,958 surgeries for AAA in BC between 2013/14 and 2017/18, 1,142 were EVAR (58\%) and 816 were open ( $42 \%$ ). ${ }^{504}$

[^117]- Recent evidence from the UK and Sweden also indicate a rate for elective EVAR of $59 \%$. ${ }^{505,506}$
- We model an EVAR rate of $58 \%$ in BC (Table 5, rows $x \& a p$ ).
- The USPSTF referenced two key studies comparing early open surgery with surveillance in their analysis of the harms of screening. ${ }^{507}$ One study was conducted in the UK (UKSAT) ${ }^{508}$ and the other in the US (ADAM). ${ }^{509}$
- Greenhalgh and colleagues reported a 30-day mortality rate of $5.8 \%$ in patients receiving open surgery in the UK Small Aneurysm Trial (UKSAT). The authors acknowledge that this rate was "about half the national in-hospital mortality rate for elective repair" of AAA. ${ }^{510}$ This study was conducted at a time when endovascular surgery was "still under development".
- Lederle and colleagues reported a 30-day mortality rate of $2.0 \%$ in patients receiving open surgery in the Aneurysm Detection and Management (ADAM) study. ${ }^{511}$
- Thompson and colleagues reported a 30 -day mortality of $1.8 \%$ and $4.6 \%$ for elective endovascular and elective open AAA surgeries respectively (MASS study in UK). ${ }^{512}$
- Several studies published since the USPSTF recommendation in 2014 have reported on elective surgery mortalities. A study of Medicare beneficiaries in the US reported a perioperative (within 30-days of surgery) mortality rate of $1.6 \%$ for endovascular repair of AAA and $5.2 \%$ for open repair. The mean age was 75.6 for those receiving surgery and the data used was from $2001-2008 .{ }^{513}$
- More recent European studies report ranges of $0.3 \%-0.7 \%$ and $0.9 \%-1.3 \%$ for $30-$ day mortality following endovascular repair and open surgery respectively. ${ }^{514,515}$ Neither study explicitly states the mean age of patients receiving surgery, but

[^118]Jacomelli et al. ${ }^{516}$ report on screening of 65 year-old men and Wanhainen et al. ${ }^{517}$ on $65-74$ year old men, so it can be inferred that their results are taken from a younger cohort than is reported by Schermerhorn and colleagues. ${ }^{518}$

- In a report using Ontario data, de Mestral and colleagues report a 90 -day mortality rate following endovascular repair of $1.6 \%$. ${ }^{519}$
- Reporting on outcomes of open repair of AAA in Ontario, Dubois and colleagues report a 30 -day mortality for open repair of $3 \% .{ }^{520}$
- We model a 30 -day mortality of $1.0 \%$ and $3.0 \%$ for elective endovascular and open surgery respectively (Table 5, rows $z \& a a$ and $a r \& a s$ ).
- In their evidence synthesis for the USPSTF, Guirguis-Blake and colleagues report an estimate of $41 \%$ mortality (either in hospital or 30-day) associated with emergency surgery for AAA. ${ }^{521}$
- We model an emergency surgery 30-day mortality of $41 \%$ (Table 5, row ae \& ax).

Based on these assumptions, the CPB associated with screening for abdominal aortic aneurysms in males aged 65 who have ever smoked is 340 QALYs (see Table 5, row $b k$ ).

## Comparison to Actual BC Data

Analysis from the discharge abstract database in BC from 2013/14-2017/18 indicates that 77.8 / 100,000 men over 65 years old had elective AAA surgery and 24.8 / 100,000 men over 65 years old had emergency and / or ruptured AAA surgery, a ratio of 3.14. ${ }^{522}$ Our model calculates these rates at $88.4 / 100,000$ and 21.4 / 100,000 respectively, a difference of approximately $14 \%$ from the actuals in both cases. With no screening (i.e. in the control group), the Viborg study reported the same rates of elective and emergency surgery (see Table 4). If there was no screening in BC, we might expect a similar ratio as the unscreened population in the Viborg study. The fact that there are more than three times as many elective as emergency surgeries in $B C$ suggests that $B C$ physicians are already opportunistically screening their patients in the province. In the fully screened population analysed by the USPSTF, ${ }^{523}$ the ratio of elective to emergency surgeries was 4.13 , indicating that while

[^119]opportunistic screening is occurring in BC , it has not yet reached a level in which the majority of eligible males (we model a 'best in the world' rate of $85.8 \% \%^{524}$ ) are screened.

Table 5: CPB of Abdominal Aortic Aneurysm Screening in Ever-Smoking Men 65+ In a BC Birth Cohort of 40,000

| Row Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
|  | Deaths and Life-Years Lost due to AAA in an Unscreened Cohort |  |  |
| a | Number of 65-year old men in cohort | 17,559 | BC Life Table |
| b | Proportion of population, never-smokers | 47.0\% | $\checkmark$ |
| C | Proportion of population, former smokers | 40.1\% | $\checkmark$ |
| d | Proportion of population, current smokers | 12.9\% | $\checkmark$ |
| e | Prevalence of AAA in population | 2.35\% | $\checkmark$ |
| $f$ | Prevalence of AAA in never-smokers | 1.21\% | Table 2 |
| g | Prevalence of AAA in former smokers | 2.54\% | Table 2 |
| h | Prevalence of AAA in current smokers | 5.90\% | Table 2 |
| i | Life years for cohort from 65-84 | 289,017 | Table 3 |
| j | Life years, ever-smokers for cohort from 65-84 | 153,179 | $=i^{*}(c+d)$ |
| k | Number with AAA in cohort at age 65, never-smokers | 100 | $=a * b * f$ |
| 1 | Number with AAA in cohort at age 65, former smokers | 179 | $=a * c * g$ |
| m | Number with AAA in cohort at age 65, current smokers | 134 | $={ }^{*} d^{*}$ h |
| n | Number of AAA-related deaths over cohort lifetime | 149 | Table 3 |
| 0 | Fraction of those with AAA dying over cohort lifetime, total population | 36.2\% | = $\mathrm{n} /(\mathrm{k}+\mathrm{l}+\mathrm{m})$ |
| p | Number of deaths over cohort lifetime, never-smokers | 36 | = ${ }^{*}$ o |
| q | Number of deaths over cohort lifetime, former smokers | 65 | $={ }^{*}$ o |
| $r$ | Number of deaths over cohort lifetime, current smokers | 48 | $=\mathrm{m}^{*} \mathrm{o}$ |
| s | Life years lost over cohort lifetime, never-smokers | 259 | Table 3 |
| t | Life years lost over cohort lifetime, former smokers | 464 | Table 3 |
| $u$ | Life years lost over cohort lifetime, current smokers | 346 | Table 3 |
|  | AAA-related deaths in an Unscreened Cohort of Ever-Smokers |  |  |
| v | Rate of elective surgery per 100,000, unscreened population | 41 | Table 4 |
| w | Number of elective surgeries in cohort | 63 | $=(\mathrm{v} / 100,000) * \mathrm{j}$ |
| x | Proportion of elective surgeries that are endovascular | 58\% | $\checkmark$ |
| y | Proportion of elective surgeries that are open | 42\% | $=(1-\mathrm{ag})$ |
| z | 30-day mortality for elective endovascular AAA surgery | 1.0\% | $\checkmark$ |
| aa | 30-day mortality for elective open AAA surgery | 3.0\% | $\checkmark$ |
| ab | Number of deaths associated with elective surgeries | 1.2 | $=w^{*}\left(\left(x^{*} z\right)+\left(y^{*} \mathrm{aa}\right)\right)$ |
| ac | Rate of emergency surgery per 100,000, unscreened population | 41 | Table 4 |
| ad | Number of emergency surgeries in cohort | 63 | $=(\mathrm{ac} / 100,000){ }^{*} \mathrm{j}$ |
| ae | Death rate, emergency surgery | 41\% | $\checkmark$ |
| af | Number of deaths associated with emergency surgeries | 25.8 | = ad * ae |
| ag | Number of deaths prior to arriving at hospital for surgery | 86.2 | $=(q+r)-a b-a f$ |

${ }^{524}$ Jacomelli J, Summers L, Stevenson A et al. Impact of the first 5 years of a national abdominal aortic aneurysm screening programme. British Journal of Surgery. 2016; 103(9): 1125-31.

Table 5: CPB of Abdominal Aortic Aneurysm Screening in Ever-Smoking Men 65+

| Row Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
|  | AAA-related deaths in a Screened Cohort of Ever-Smokers |  |  |
| ah | Number targeted for screening, base case: ever-smokers (current + former) | 9,306 | $=a^{*}(c+d)$ |
| ai | Screening Rate | 85.8\% | $\checkmark$ |
| aj | Total Number screened | 7,985 | $=v^{*} \mathrm{w}$ |
| ak | Proportion of AAA opportunistically detected without screening | 13\% | $\checkmark$ |
| al | Relative risk of elective surgery, screened vs. unscreened population | 2.15 | $\checkmark$ |
| am | Rate of elective surgery per 100,000, screened population | 88.4 | $=\mathrm{al}^{*} \mathrm{v}$ |
| an | Number of elective surgeries in cohort | 135 | $=((\mathrm{am} / 100,000) * \mathrm{j})$ |
| ao | Number of elective surgeries in cohort, due to screening alone | 63 | = an * (1-ak) |
| ap | Proportion of elective surgeries that are endovascular | 58\% | = x |
| aq | Proportion of elective surgeries that are open | 42\% | = y |
| ar | 30-day mortality for elective endovascular AAA surgery | 1.0\% | = |
| as | 30-day mortality for elective open AAA surgery | 3.0\% | = aa |
| at | Number of deaths associated with elective surgeries | 2.5 | = an * ( $\left.a p^{*} \mathrm{ar}\right)+(\mathrm{aq}$ * as$)$ ) |
| au | Relative risk of emergency surgery, screened vs. unscreened population | 0.52 | $\checkmark$ |
| av | Rate of emergency surgery per 100,000, unscreened population | 21.4 | = au *ac |
| aw | Number of emergency surgeries in cohort | 33 | $=(a u / 100,000) *$ j |
| ax | Death rate, emergency surgery | 41\% | $\checkmark$ |
| ay | Number of deaths associated with emergency surgeries | 13.4 | = aw * ax |
| az | Relative risk of AAA-related death, overall, screened vs. unscreened population | 0.58 | $\checkmark$ |
| ba | AAA-related deaths in screened cohort | 66 | $=(q+r) * a z$ |
| bb | Number of deaths prior to arriving at hospital for surgery | 49.7 | = ba - ay - at |
|  | Difference in AAA-related deaths in a Screened vs. Unscreened Cohort of EverSmokers |  |  |
| bc | Deaths due to elective surgeries, screened vs. unscreened | 1.3 | = at - ab |
| bd | Deaths due to emergency surgeries, screened vs. unscreened | -12.4 | = ay - af |
| bf | Deaths prior to hospital arrival, screened vs. unscreened | -36.5 | $=\mathrm{bb}-\mathrm{ag}$ |
| bg | Difference in total AAA-related deaths, screened vs. unscreened | -47.6 | $=b c+b d+b f$ |
| bh | Total AAA-related deaths in unscreened cohort | 113 | $=q+r$ |
| bi | Fraction of deaths avoided as a result of screening | 42\% | $=(-b g) / \mathrm{bh}$ |
|  | Difference in Life Years, Screened vs. Unscreened Cohort of Ever-Smokers |  |  |
| bj | Life years lost due to death from AAA in unscreened ever-smoking group | 810 | Table 3 |
| bk | QALYs saved by screening | 340 | $=\mathrm{bi}$ * bj |

V = Estimates from the literature
For the sensitivity analysis, we modified the relative risk assumptions and recalculated the CPB as follows:

- Assume that the relative risk of overall death is increased from 0.58 to 0.88 (Table 5, row $a z$ ), the relative risk of elective surgery in screened individuals is decreased from 2.15 to 1.89 (Table 5 , row $a l$ ) and the relative risk of emergency surgery is increased from 0.52 to 0.66 (Table 5 , row $a u$ ): $\mathrm{CPB}=97$
- Assume that the relative risk of overall death is decreased from 0.58 to 0.39 (Table 5, row az), the relative risk of elective surgery in screened individuals is increased from 2.15 to 2.44 (Table 5 , row al) and the relative risk of emergency surgery is decreased from 0.52 to 0.40 (Table 5, row au): $\mathrm{CPB}=494$
- Offer screening to all 65 year old males, rather than to just 65 year old male eversmokers (Table 5, rows $b, c$ and $d$ ): $\mathrm{CPB}=449$
- Assume vital statistics death rate of $0.74 \%$ in population 65 and older due to abdominal aortic aneurysm, rather than the $1.90 \%$ calculated in the model: $\mathrm{CPB}=$ 133


## Modelling Cost-Effectiveness

In this section, we will calculate the CE associated with screening for abdominal aortic aneurysms in males ages 65 to 75 who have ever smoked.

In modelling CE , we made the following assumptions:

- The single screen recommended by the USPSTF is conducted at age 65 .
- The screen targets only the population of ever-smokers (i.e. current and former smokers). We assess the benefits of screening the whole population in our sensitivity analysis.
- For modelling purposes, we assume that $12.9 \%$ of men 65 years of age are current smokers (Table 6, row $d$ ) and $40.1 \%$ are former smokers (Table 6, row $c$ ).
- We assume that all 65 year old males will have at least one visit to their GP each year.
- We model a best in the world screening acceptance rate of $85.8 \%$ (Table 6 , row $e$ ). ${ }^{525}$
- The cost of each 10 minute primary care provider office visit is $\$ 34.85$ (Reference Document) (Table 6, row $g$ )
- The value of patient time (based on 2 hours, including travel time) for each visit to a primary care office and for abdominal ultrasound screening is $\$ 59.38$ (Reference Document) (Table 6, row $h$ ).
- The proportion of each office visit attributable to recommending screening is $50 \%$ (Reference Document) (Table 8, row $i$ ).
- The average service fee cost of an abdominal B-scan (ultrasound - fee item 8648) in BC between 2012 and 2016 was $\$ 106.81$ (Table 6, row $k$ ). ${ }^{526}$
- Visser reported elective endovascular surgery costs at $€ 20,767$ (2003) or $\$ 38,084$ ( 2017 CAD), with those costs rising to $€ 23,588$ (2003) or $\$ 43,257$ (2017 CAD) if one-year follow-up costs were included. ${ }^{527}$
- Matsumura and colleagues reported elective endovascular surgery costs between $\$ 34,800-38,900$ USD (2008) or $\$ 33,750-37,726$ (2017CAD), depending on which device was used in the surgery. ${ }^{528}$
- Similarly, in their cost-effectiveness analysis, Svensjo and colleagues use an elective endovascular surgery cost of $€ 24,493$ (2012), with that cost rising to $€ 29,758$ if post-

[^120]operative costs were included as well. ${ }^{529}$ Converted to 2017 CAD, the amounts are $\$ 40,778$ and $\$ 49,544$ respectively.

- For elective endovascular surgery, Burgers and colleagues reported surgery costs of $€ 14,690$ (2013) or $\$ 22,534$ (2017 CAD). ${ }^{530}$
- Elective endovascular surgery costs, adjusted to 2017 CAD, range between $\$ 22,534$ (Burgers et al.) and \$49,544 (Svensjö et al.). We model elective endovascular AAArepair surgery costs at $\$ 36,039$ (the mid-point of this) and vary this to $\$ 22,534$ and $\$ 49,544$ in our sensitivity analysis (Table 6 , row $s$ ).
- We noted previously that we assume a 30 -day mortality of $1.0 \%$ and $3.0 \%$ for elective endovascular and open surgery respectively. This early mortality advantage associated with EVAR erodes over time, with no survival advantage after 4 to 5 years of follow-up. ${ }^{531,532,533}$
- Based on 15 years of follow-up results from the UK EVAR trial, graft-related reinterventions remained higher in patients with endovascular repair compared with open repair. Overall, any graft-related re-intervention occurred in $26 \%$ of EVAR vs. $12 \%$ of open patients. Serious graft-related re-interventions occurred in $22 \%$ of EVAR vs. $9 \%$ of open patients while life-threatening re-interventions occurred in $14 \%$ of EVAR vs. $7 \%$ of open patients. The authors note that "there is no time to assume that it is safe to discontinue surveillance in patients who have had EVAR". ${ }^{534}$
- Studies assessing the long-term cost-effectiveness of EVAR vs. open surgery that take into account the changing survival profile following EVAR and open surgery, as well as differential graft-related intervention rates, have found no differences in costeffectiveness. Epstein and colleagues "did not find that EVAR is cost-effective compared with open repair in the long term in trials conducted in European centres. ${ }^{,{ }^{535}}$ Lederle and co-authors conclude that, based on follow-up of 9 years, "survival, quality of life, costs and cost-effectiveness did not differ between elective open and endovascular repair of AAA. ${ }^{" 536}$ Cost-effectiveness studies with a followup period of less than 4 years, on the other hand, find EVAR to be cost-effective

[^121]compared with open surgery, largely due to the early survival advantages associated with EVAR. ${ }^{537}$

- Because of this long term convergence in the benefits and costs between EVAR and open surgery, we have not taken into account the longer-term benefits or costs of EVAR or open surgery in our modelling.
- Visser reported elective open surgery costs at $€ 35,470$ (2003) or $\$ 65,047$ (2017 CAD), with those costs rising to $€ 36,448$ (2003) or $\$ 66,840$ (2017 CAD) if one-year follow-up costs were included. ${ }^{538}$
- Matsumura and colleagues reported elective open surgery costs between $\$ 38,900$ 45,100 USD (2008) or $\$ 37,726-43,739$ (2017 CAD), depending on which device was used in the surgery. ${ }^{539}$
- Similarly, in their cost-effectiveness analysis, Svensjo and colleagues use an elective open surgery cost of $€ 30,099$ (2012), with that cost rising to $€ 35,615$ if post-operative costs were included as well. ${ }^{540}$ Converted to 2017 CAD, the amounts are $\$ 50,112$ and $\$ 59,295$ respectively.
- For elective open surgery, Burgers and colleagues reported surgery costs of $€ 16,399$ (2013) or $\$ 25,156$ (2017 CAD). ${ }^{541}$
- In papers not reporting on the specific type of elective surgery, the elective surgery costs ranged from $\$ 14,075$ - $\$ 44,388$ (2017 CAD). ${ }^{542,543,544,545,546,547,548,549}$

[^122]- Elective open surgery costs, adjusted to 2017 CAD, range between $\$ 25,156$ (Burgers et al.) and $\$ 66,840$ (Visser et al.). We model elective open AAA-repair surgery costs at $\$ 45,998$ (open surgery mid-point) and vary this to $\$ 25,156$ and $\$ 66,840$ in our sensitivity analysis (Table 6 , row $t$ ).
- Chew and colleagues reported that emergency AAA-repair surgery costs in Nova Scotia were $\$ 18,899$ ( 1998 CAD), including overhead. This is equivalent to $\$ 27,500$ ( 2017 CAD). ${ }^{550}$
- In a Swedish cost analysis, Wanhainen and colleagues used $€ 32,183$ (2003) for emergency AAA-repair with rupture or $\$ 50,301$ (2017 CAD). ${ }^{551}$
- In a model of US costs, Silverstein and colleagues used \$60,000 (2003) USD to account for emergency surgery and emergency care costs. Adjusted to 2017 CAD, this comes to $\$ 66,582 .{ }^{552}$
- Montreuil and colleagues conducted a Monte Carlo analysis of screening Canadian men for AAA and used $\$ 35,982$ (2005 CAD) for emergency AAA-repair surgery costs, equivalent to $\$ 43,494$ (2017 CAD). ${ }^{553}$
- Lindholt and colleagues reported an emergency AAA-repair surgery cost of $€ 35,928$ (2007) in Denmark or $\$ 63,497$ (2017 CAD). ${ }^{554}$
- Reporting on the cost-effectiveness of screening using the MASS results, Thompson and colleagues used an emergency AAA-repair cost of $£ 14,825$ (2008) or $\$ 29,935$ ( 2017 CAD). ${ }^{555}$
- Giardina and colleagues report an emergency AAA-repair cost of $€ 15,602$ (2009) in Italy, or $\$ 27,123$ (2017 CAD). ${ }^{556}$
- Emergency AAA-repair surgery costs, adjusted to 2017 CAD, range between $\$ 27,123$ (Giardina et al.) and $\$ 66,582$ (Silverstein et al.). We model the cost of emergency surgery as $\$ 46,853$ (mid-point of emergency surgery range) and vary this from $\$ 27,123$ to $\$ 66,582$ in our sensitivity analysis (Table 6, row ao).
- Chew et al. reported a mean length of stay in Nova Scotia of 19.57 days in hospital for emergency surgery survivors and 9.22 days in hospital for emergency surgery patients who died. ${ }^{557}$ We model accordingly (Table 6, rows aq \& ar)

[^123]- The Canadian Society for Vascular Surgery (CSVS) and HealthLinkBC agree that hospital stays for elective endovascular AAA-repair surgery will range between $1-3$ days. ${ }^{58,559}$
- The Canadian Society for Vascular Surgery suggests that elective open AAA-repair surgery will require 5-7 days in hospital. ${ }^{560}$
- Analysis from the discharge abstract database in BC from 2013/14-2017/18 indicates the average length of stay for elective endovascular AAA repair in BC is no less than 4 days, while the average length of stay for elective open AAA repair is 10 days. ${ }^{561}$
- HealthLinkBC states that patients will typically fully recover 4 weeks after endovascular AAA-repair surgery and suggests planning to take $1-2$ weeks off work. ${ }^{562}$ The CSVS reports a full recovery time between $2-4$ weeks. ${ }^{563}$
- HealthLinkBC states that patients will typically resume "usual activities" $4-6$ weeks after open AAA-repair surgery and that full recovery will take $2-3$ months. ${ }^{564}$ The CSVS reports a full recovery time between $1-3$ months. ${ }^{565}$
- For the purposes of calculating patient time costs, we model 4 days and 10 days in hospital for elective endovascular and open AAA-repair surgeries respectively (Table 6 , rows $v \& w$ ). We model time off work at 10 days (midpoint of $1-2$ weeks) and 35 days (midpoint of $4-6$ weeks) for endovascular and open AAA-repair surgeries respectively (Table 6 , rows $x \& y$ ). In our sensitivity analysis we range the days off work between $7-14$ for endovascular and $28-42$ for open surgery.
- Emergency ground transport in BC costs $\$ 530$ for non-MSP beneficiaries. ${ }^{566}$ This can be considered the unsubsidized cost of emergency ground transportation.
- We model that the difference in the sum of emergency surgeries and deaths prior to hospitalization for AAA between the unscreened and screened cohort is equivalent to the number of avoided emergency transports (Table 6, row ay). These emergency transports each cost $\$ 530$ (Table 6, row $a z$ ).

Based on these assumptions, the CE associated with screening for abdominal aortic aneurysms in males ages 65 to 75 who have ever smoked is $\$ 11,995$ / QALY (see Table 6, row $b g$ ).

[^124]Table 6: Cost Effectiveness of Abdominal Aortic Aneurysm Screening in Ever-Smoking Men 65+ In a BC Birth Cohort of 40,000

| Row Label | Variable | Base case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Number of 65-year old men in cohort | 17,559 | BC Life Table |
| b | Proportion who are former smokers | 40.1\% | $\checkmark$ |
| c | Proportion who are current smokers | 12.9\% | $\checkmark$ |
| d | Number targeted for screening | 9,306 | $=a^{*}(\mathrm{~d}+e)$ |
| e | Screening Rate | 85.8\% | $\checkmark$ |
| f | Total Number screened | 7,985 | = f * g |
| g | Cost of 10 minute office visit | \$34.85 | Ref Doc |
| h | Value of patient time and travel for office visit | \$59.38 | Ref Doc |
| i | Portion of 10-minute office visit for screening | 50\% | Ref Doc |
| j | Cost of initial primary care visit for cohort | \$376,207 | $=\mathrm{f}^{*}(\mathrm{~g}+\mathrm{h}) * \mathrm{i}$ |
| k | Cost of ultrasonic screening session | \$107 | $\checkmark$ |
| I | Cost of ultrasonic screening for cohort | \$1,327,006 | = $\mathrm{f}^{*}(\mathrm{~h}+\mathrm{k})$ |
| m | Number of elective surgeries in ever-smokers, unscreened | 63 | Table 5, row w |
| n | Number of elective surgeries in ever-smokers, screened | 135 | Table 5, row an |
| 0 | Rate of opportunistically detected AAA | 13\% | Table 5, row ak |
| p | Number of additional elective surgeries attributable to screening alone | 63 | $=((\mathrm{n}-\mathrm{m}) *(1-\mathrm{o})$ ) |
| q | Proportion of surgeries that are endoscopic surgeries | 58\% | Table 5, row ap |
| r | Proportion of surgeries that are open surgeries | 42\% | $=1-\mathrm{q}$ |
| s | Cost per elective surgery, endoscopic AAA repair | \$36,039 | $\checkmark$ |
| t | Cost per elective surgery, open AAA repair | \$45,998 | $\checkmark$ |
| $u$ | Cost of additional elective surgery due to screening | \$2,533,146 | $=p^{*}\left(\left(q^{*} s\right)+\left(r^{*} t\right)\right)$ |
| v | Time in hospital, days, endovascular AAA repair | 4 | V |
| w | Time in hospital, days, open AAA repair | 10 | $\checkmark$ |
| x | Recovery time, days, endovascular AAA repair | 10 | $\checkmark$ |
| y | Recovery time, days, open AAA repair | 35 | $\checkmark$ |
| z | Cost per day of patient time in hospital | \$223 | Ref Doc |
| aa | Patient time cost for additional elective AAA surgeries | \$377,903.66 | $=p^{*}\left(\left(q^{*}(v+x)\right)+\left(r^{*}(w+y)\right)^{*} z\right.$ |
| ab | Number of elective surgeries, endoscopic | 37 | = ${ }^{*}$ q |
| ac | Cost of CT Scan | \$223.50 | $\checkmark$ |
| ad | Cost of office visit, 100\% for AAA follow-up | \$94 | $=\mathrm{g}+\mathrm{h}$ |
| ae | Average life expectancy of 65-year old man | 20 | BC Life Table |
| af | Estimated compliance with annual follow-up protocol | 70\% | V |
| ag | Cost of CT Scans | \$114,973 | = ab *ac*ae * f |
| ah | Cost of follow-up office visits | \$48,474 | $=a b * a d$ *e * $\mathrm{ff}^{\text {f }}$ |
| ai | Lifetime failure rates of EVAR | 10\% | $\checkmark$ |
| aj | Cost to correct EVAR failure with open surgery | \$169,017 | $=\mathrm{ab} * \mathrm{ai}{ }^{*} \mathrm{t}$ |
| ak | Total cost due to additional elective AAA surgery in cohort | \$3,243,513 | $=u+a a+a g+a h+a j$ |
| al | Number of emergency surgeries in ever-smokers, unscreened | 63.0 | Table 5, row ad |
| am | Number of emergency surgeries in ever-smokers, screened | 32.8 | Table 5, row aw |
| an | Reduction in emergency surgeries in screened population | 30.2 | = al - am |
| ao | Cost of emergency surgery, AAA rupture repair | \$46,853 | $\checkmark$ |
| ap | Cost reduction due to avoided surgery | \$1,416,717 | = an * $\mathrm{a}^{\text {o }}$ |
| aq | Time in hospital, emergency AAA repair, survivors | 19.57 | $\checkmark$ |
| ar | Time in hospital, emergency AAA repair, patients who die | 9.22 | V |
| as | Death rate, emergency surgery | 41\% | $\checkmark$ |
| at | Average time in hospital, emergency AAA repair | 15.3 | $=((a q *(1-\mathrm{as}))+(\mathrm{ar} * \mathrm{as}))$ |
| au | Patient time cost avoided due to avoided emergency surgery | \$103,195 | an *at * |
| av | Total cost reduction due to avoided surgeries | \$1,519,913 | $=a p+a v$ |
| aw | Number of emergency surgeries and pre-hospital deaths, unscreened cohort | 149 | Table 5, row ad + Table 5, row ag |
| ax | Number of emergency surgeries and pre-hospital deaths, screened cohort | 83 | Table 5, row aw + Table 5, row bb |
| ay | Number of avoided emergency transports due to screening | 67 | =aw -ax |
| az | Average cost of emergency transport | \$530 | $\checkmark$ |
| ba | Avoided emergency transportation cost | \$35,361 | = ay *az |
| bb | Net cost of intervention | \$3,391,452 | $=j+l+a k-a v-b a$ |
| bc | QALYs saved | 340 | Table 5, row bk |
| bd | Cost effectiveness (CE) of intervention, \$/QALY | \$9,973 | $=\mathrm{bb} / \mathrm{bc}$ |
| be | Net Cost of Intervention (1.5\% Discount) | \$3,512,843 | Calculated |
| bf | Net QALYs Gained (1.5\% Discount) | 293 | Calculated |
| bg | Cost Effectiveness (CE) of Intervention, \$/QALY (1.5\% Discount) | \$11,995 | = be / bf |

For the sensitivity analysis, we modified a number of major assumptions and recalculated the CE as follows:

- Assume that the relative risk of overall death moves from 0.58 to 0.88 (Table 5, row $a z$ ), the relative risk of elective surgery in screened individuals is decreased from 2.15 to 1.89 (Table 5, row al) and the relative risk of emergency surgery moves from 0.52 to 0.66 (Table 5, row $a u$ ): $\mathrm{CE}=\$ 38,251$
- Assume that the relative risk of overall death moves from 0.58 to 0.39 (Table 5, row $a z$ ), the relative risk of elective surgery in screened individuals is increased from 2.15 to 2.44 (Table 5, row al) and the relative risk of emergency surgery moves from 0.52 to 0.40 (Table 5, row au): $\mathrm{CE}=\$ 9,328$
- Assume the rate of opportunistically detected AAA in the population increases from $13 \%$ to $25 \%$ (Table 5, row $a k$ ): $\mathrm{CE}=\$ 10,512$
- Assume the rate of opportunistically detected AAA in the population decreases from $13 \%$ to $7 \%$ (Table 5, row $a k$ ): $\mathrm{CE}=\$ 12,736$
- Assume the cost of elective endovascular surgery increases from $\$ 36,039$ to $\$ 49,544$ (Table 6 , row $s$ ), the cost of elective open endovascular surgery increases from $\$ 45,998$ to $\$ 66,840$ (Table 6, row $t$ ), and the cost of emergency AAA-repair surgery increases from $\$ 46,853$ to $\$ 66,582$ (Table 6 , row $a f$ ): $\mathrm{CE}=\$ 13,955$
- Assume the cost of elective endovascular surgery decreases from $\$ 36,039$ to $\$ 22,534$ (Table 6, row $s$ ), the cost of elective open endovascular surgery decreases from $\$ 45,998$ to $\$ 25,156$ (Table 6 , row $t$ ), and the cost of emergency AAA-repair surgery decreases from $\$ 46,853$ to $\$ 27,123$ (Table 6 , row $a f$ ): $\mathrm{CE}=\$ 10,034$
- Assume that the time off work for elective endovascular surgery increases from 10 to 14 days (Table 6 , row $x$ ) and the time off work for elective open surgery increases from 35 to 42 days (Table 6, row $y$ ): $\mathrm{CE}=\$ 12,239$
- Assume that the time off work for elective endovascular surgery decreases from 10 to 7 days (Table 6 , row $x$ ) and the time off work for elective open surgery increases from 35 to 28 days (Table 6, row $y$ ): $\mathrm{CE}=\$ 11,778$
- Assume vital statistics death rate of $0.74 \%$ in population 65 and older due to abdominal aortic aneurysm, rather than the $1.90 \%$ calculated in the model: $\mathrm{CE}=$ \$21,015
- Offer screening to all 65 year old males, rather than to just 65 year old male eversmokers (Table 5, rows $b, c$ and $d$ ): $\mathrm{CE}=\$ 17,293$


## Summary

## Ever-Smoking Males Ages 65 and Older

The clinically preventable burden (CPB) associated with screening for, and treatment of, abdominal aortic aneurysm in ever-smoking males ages 65 and older is 293 quality-adjusted life years (QALYs) while the cost-effectiveness (CE) is estimated at $\$ 11,995$ per QALY (see Table 7).

Table 7: Abdominal Aortic Aneurysm Screening in EverSmoking Men 65+ in a BC Birth Cohort of 40,000 Summary

| Base <br> Case$\quad$ Range |
| :--- |

CPB (Potential QALYs Gained)
Assume No Current Service

| 1.5\% Discount Rate | 293 | 84 | 425 |
| :--- | :--- | :--- | :--- |
| 3\% Discount Rate | 254 | 73 | 369 |
| 0\% Discount Rate | 340 | 97 | 494 |

CE (\$/QALY) including patient time costs

| $\mathbf{1 . 5 \%}$ Discount Rate | $\$ 11,995$ | $\$ 9,328$ | $\$ 38,251$ |
| :--- | :--- | :--- | :--- |
| $3 \%$ Discount Rate | $\$ 14,175$ | $\$ 11,053$ | $\$ 44,859$ |
| $0 \%$ Discount Rate | $\$ 9,973$ | $\$ 7,725$ | $\$ 32,136$ |

CE (\$/QALY) excluding patient time costs

| $1.5 \%$ Discount Rate | $\$ 8,516$ | $\$ 6,750$ | $\$ 26,836$ |
| :--- | :---: | :---: | :---: |
| $3 \%$ Discount Rate | $\$ 10,162$ | $\$ 8,079$ | $\$ 31,705$ |
| $0 \%$ Discount Rate | $\$ 6,984$ | $\$ 5,511$ | $\$ 22,315$ |

## All Males Ages 65 and Older

The clinically preventable burden (CPB) associated with screening for, and treatment of, abdominal aortic aneurysm in all males ages 65 and older is 386 quality-adjusted life years (QALYs) while the cost-effectiveness (CE) is estimated at $\$ 17,293$ per QALY (see Table 8).


## Screening for Sexually Transmitted Infections and Blood Borne Pathogens

Human Immunodeficiency Virus

## United States Preventive Services Task Force Recommendations (2013)

An estimated 1.2 million persons in the United States are currently living with HIV infection, and the annual incidence of the disease is approximately 50000 cases. Since the first cases of AIDS were reported in 1981, more than 1.1 million persons have been diagnosed and nearly 595000 have died from the condition.
Approximately 20\% to $25 \%$ of individuals living with HIV infection are unaware of their positive status.

The USPSTF recommends that clinicians screen adolescents and adults aged 15 to 65 years for HIV infection. Younger adolescents and older adults who are at increased risk should also be screened. (A recommendation)

The USPSTF recommends that clinicians screen all pregnant women for HIV, including those who present in labor who are untested and whose HIV status is unknown. (A recommendation) $)^{567}$

## Canadian Task Force on Preventive Health Care Recommendations (2016)

The CTFPHC has reviewed the USPSTF guideline on screening for HIV infection and conclude that it "is a high-quality guideline, but the CTFPHC does not recommend its use in Canada. In the opinion of the CTFPHC, available evidence does not justify routinely screening all adult Canadians for HIV." Instead, the focus should be on screening high-risk groups and pregnant women. ${ }^{568}$

## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with screening adolescents and adults aged 15 to 65 years for HIV infection in a BC birth cohort of 40,000.

In modelling CPB, we made the following assumptions:

- The total number of individuals living with HIV infections in BC is estimated to be 12,100 (with a range from 9,700 to 14,500 ) (see Table 1). ${ }^{569}$

[^125]

- $20 \%$ of HIV-infected men who have sex with men (MSM), $24 \%$ of HIV-infected injection drug users (IDU) and $34 \%$ of HIV-infected heterosexuals (HET) are unaware of their HIV status (Table 2, rows $c, f \& i)$. ${ }^{570}$
- Adherence with universal screening was assumed to be $83 \%$ for MSM, $45 \%$ for HET and $60 \%$ for IDU (Table 2, rows $u, v \& w$ ) (see Reference Document).
- $4.56 \%$ of HIV infected individuals die prematurely without early initiation of antiretroviral therapy (ART) (deferring initiation of ART to CD4 levels of 200 cells $/ \mu \mathrm{L}$ ). This can be reduced to $1.11 \%$ with early initiation of ART (Table 2 , rows $y$ $\& z) .{ }^{511}$
- The average age at which undiagnosed HIV is detected is 40 (Table 2, row $b b$ ). ${ }^{572}$
- The gain in quality of life associated with early detection and treatment of an HIV infection is 0.11 (Table 2, row ee). ${ }^{573}$
- Antiretroviral therapy is a potent intervention for prevention of HIV in discordant couples. The RCT by Cohen, et al. found that just 1 of 28 transmissions occurred in a serodiscordant couple in which the infected partner received early initiation of antiretroviral therapy (a hazard ratio of 0.04; 95\% CI from 0.01 to 0.27). ${ }^{574}$ The 2013 Cochrane review by Anglemyer and colleagues noted the RCT study by Cohen, et al. as well as nine observational studies. Results from the observational studies suggested that treating the HIV-infected partner in a serodiscordant couple reduces

[^126]the risk of transmission by $64 \%$ (a relative risk of $0.36 ; 95 \% \mathrm{CI}$ from 0.17 to $0.75) .{ }^{575,576}$ In BC, the expanded utilization of highly active antiretroviral therapy (HAART) between 1996 and 2012 is associated with a $66 \%$ decrease in new diagnoses of HIV. ${ }^{577}$ To incorporate this information into our model, we first calculated the rate per person year of HIV transmission in HIV-discordant couples if the HIV-positive partner is not treated with ART. This is based on the results from the control arms of the 1 RCT and 9 observational studies included in the Cochrane review by Anglemyer et al. (1,094 transmissions during 42,917 person-years, a transmission rate of 0.0255 per person-year, Table 2 , row $g g$ ). We then assumed a $64 \%$ reduction in the transmission rate per person-year if the HIV-positive partner is treated with ART. This results in an annual transmission rate of 0.0092 per personyear (Table 2 , row $h h$ ). In the sensitivity analysis we used results from the Cohen et al. study ( $96 \%$ reduction) as the upper bounds and the $95 \%$ CI from the 9 observational studies reviewed by Anglemyer et al. (RR of 0.75 or a $25 \%$ reduction) as the lower bounds.

- We assumed that the 16.58 infections avoided associated with screening and the early treatment with ART (Table 2, row $k k$ ) would lead to an additional 11.91 infections avoided (Table 2, row $n n$ ), due to second order transmission benefits.
- The difference in quality of life between avoided infection and symptomatic HIV treated with ART is 0.17 (Table 2, row oo). ${ }^{578}$
- Other assumptions used in assessing the clinically preventable burden are detailed in the Reference Document.
Based on these assumptions, the calculation of CPB (Table 2, row $q q$ ) is 360 QALYs. This represents the potential CPB of moving from no screening to $45 \%$ in the heterosexual population, $60 \%$ in people who inject drugs and $83 \%$ in men who have sex with men.

[^127]Table 2: CPB of Screening to Detect and Treat HIV in a BC Birth Cohort of
40,000

| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Prevalence of HIV Infections in B.C. | 12,100 | Table 1 |
| b | Prevalence of HIV Infections in MSM | 5,500 | $\checkmark$ |
| c | \% Undiagnosed in MSM | 20\% | $\checkmark$ |
| d | Undiagnosed HIV in MSM | 1,100 | $=b^{*} \mathrm{c}$ |
| e | Prevalence of HIV Infections in PWID | 3,785 | V |
| f | \% Undiagnosed in PWID | 24\% | $\checkmark$ |
| g | Undiagnosed HIV in PWID | 908 | $=e^{*} \mathrm{f}$ |
| h | Prevalence of HIV Infections in HET | 2,690 | $\checkmark$ |
| i | \% Undiagnosed in HET | 34\% | $\checkmark$ |
| j | Undiagnosed HIV in HET | 915 | $=h^{*}$ |
| k | Undiagnosed HIV in BC | 2,923 | $=d+g+j$ |
| I | Diagnosed HIV in BC | 9,177 | = a-k |
| m | BC Population Ages 15-65 | 3,239,000 | $\checkmark$ |
| n | Prevalence / 100,000 Diagnosed HIV | 283 | =1/(m/100,000) |
| 0 | Prevalence / 100,000 Undiagnosed HIV | 90 | $=\mathrm{k} /(\mathrm{m} / 100,000)$ |
| p | Est. diagnosed HIV in BC birth cohort of 40,000 | 113 | = ${ }^{*} 0.4$ |
| q | Est. undiagnosed HIV in BC birth cohort of 40,000 | 36 | = o*0.4 |
| r | Est. undiagnosed HIV in BC birth cohort of 40,000-MSM | 14 | $=(\mathrm{d} / \mathrm{k})^{*} \mathrm{q}$ |
| s | Est. undiagnosed HIV in BC birth cohort of 40,000- PWID | 11 | $=(\mathrm{g} / \mathrm{k}) * \mathrm{q}$ |
| t | Est. undiagnosed HIV in BC birth cohort of 40,000-HET | 11 | $=(\mathrm{j} / \mathrm{k})^{*} \mathrm{q}$ |
| u | Adherence with screening - MSM | 83.0\% | Ref Doc |
| v | Adherence with screening - PWID | 60.0\% | $\checkmark$ |
| w | Adherence with screening - HET | 45.0\% | Ref Doc |
| x | Previously undiagnosed HIV infections detected by universal screening | 23.09 | $=r * u+s^{*} v+t^{*}$ w |
| y | \% early death without early initiation of antiretroviral therapy (ART) | 4.56\% | $\checkmark$ |
| z | \% early death with early initiation of ART | 1.11\% | $\checkmark$ |
| aa | Early deaths avoided with early initiation of ART | 0.80 | $=\left(x^{*} y\right)-\left(x^{*} z\right)$ |
| bb | Average age at which undiagnosed HIV infection detected | 40 | $\checkmark$ |
| cc | Life expectancy of a 40 year-old | 44 | $\checkmark$ |
| dd | QALYs gained - premature death avoided | 35.0 | =aa*cc |
| ee | Gain in QoL associated with early detection and treatment of HIV | 0.11 | $\checkmark$ |
| ff | QALYs gained - early detection and treatment | 112 | =x*cc*ee |
| gg | HIV transmission in HIV-discordant couples, HIV positive partner untreated with ART - rate/person year | 0.0255 | $\checkmark$ |
| hh | HIV transmission in HIV-discordant couples, HIV positive partner treated with ART - rate/person year | 0.0092 | $\checkmark$ |
| ii | Potential HIV transmissions, HIV positive partner untreated with ART | 25.91 | $=x^{*} \mathrm{cc}{ }^{*} \mathrm{gg}$ |
| jj | Potential HIV transmissions, HIV positive partner treated with ART | 9.33 | =x*cc*hh |
| kk | Infections avoided per early detection associated with ARTfirst order | 16.58 | =ii-jj |
| II | Potential HIV transmissions, HIV positive partner untreated with ART | 18.60 | =kk*gg*cc |
| mm | Potential HIV transmissions, HIV positive partner treated with ART | 6.70 | =kk*hh*cc |
| $n \mathrm{n}$ | Infections avoided per early detection associated with ARTsecond order | 11.91 | =II-mm |
| OO | Difference in QoL associated with no infection vs. symptomatic infection treated with ART | 0.17 | $\checkmark$ |
| pp | QALYs gained - infections avoided due to ART | 213 | $=(\mathrm{kk}+\mathrm{nn})^{*} \mathrm{cc} * \mathrm{oo}$ |
| qq | Total QALYs gained, Utilization increasing from 0\% to 45\% for HET, 60\% for PWID and 83\% for MSM | 360 | $=d d+f f+p p$ |

$v=$ Estimates from the literature

We also modified several major assumptions and recalculated the CPB as follows:

- Assume the prevalence of individuals living with HIV infections in BC is decreased from 12,100 to 9,700 (Table 2, row $a$ ): CPB $=288$.
- Assume the prevalence of individuals living with HIV infections in BC is increased from 12,100 to 14,500 (Table 2, row $a$ ): $\mathrm{CPB}=431$.
- Assume that the early initiation of antiretroviral therapy is associated with a $96 \%$ reduction (from 64\%) in the transmission rate per person-year (Table 2, row $h h$ ): $\mathrm{CPB}=533$.
- Assume that the early initiation of antiretroviral therapy is associated with a $25 \%$ reduction (from 64\%) in the transmission rate per person-year (Table 2, row $h h$ ): $\mathrm{CPB}=209$.


## Modelling Cost-Effectiveness

In this section, we will calculate the CE associated with screening adolescents and adults aged 15 to 65 years for HIV infection in a BC birth cohort of 40,000 .

In modelling CE, we made the following assumptions:

- Number of screens - We have assumed screening between the ages of 15-65 would occur every year in high risk populations and once every 5 years in low-risk populations. ${ }^{579}$ Long and colleagues estimated the high-risk population to be $2.85 \%$ of the total population ages $15-65$ in the $\mathrm{US}^{580}$ and $1.62 \%$ in the UK. ${ }^{581}$ We assumed $2.85 \%$ for BC (Table 3, row $a$ ). In the sensitivity analysis, we adjusted screening once every five years in the low-risk population to once every 10 years and once per lifetime.
- True / false positive screens - The ratio of true to false positive test results is 1:1 (Table 3, row $i$ ). ${ }^{582}$
- Laboratory cost per screen - The estimated cost per screen is $\$ 7$ (with a range from $\$ 5$ to $\$ 9$ ). The estimated cost of confirming true / false positive results is $\$ 400$ (with a range from $\$ 300$ to $\$ 500$ ) (Table 3, rows $m \& n$ ). ${ }^{583}$
- Cost of a counselling session - We estimated the average cost of a counselling session associated with a true / false positive result to be $\$ 84.45$, based on MSP fee item 13015 (HIV/AIDS Primary Care Management - in or out of office - per half hour or major portion thereof) (Table 3, row o). ${ }^{584}$

[^128]- Average annual cost of antiretrovirals for HIV - Calculated based on an estimated average cost per day of treatment in Canada of $\$ 26.00^{585}$ (Table 3, row $s$ ). Costs in BC may be as high as $\$ 47.00$ per day. ${ }^{586} \mathrm{We}$ have used this higher estimate in our sensitivity analysis.
- Direct medical costs avoided - The annual direct medical costs (excluding medications) associated with HIV/AIDS in Canada have been estimated by stage of infection at \$1,684 for asymptomatic HIV, \$2,534 for symptomatic HIV and \$9,715 for AIDS (in 2009 CAD ). ${ }^{587}$ We modelled avoided cost using the annual direct medical costs associated with symptomatic HIV, updated to 2017 CAD of $\$ 2,843$ (Table 3, row $w$ ).
- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the estimated cost per QALY would be $\$ 16,434$ (see Table 3, row $g g$ ).

[^129]| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Proportion of population high risk | 2.85\% | $\checkmark$ |
| b | Proportion of population low risk | 97.15\% | =1-a |
| c | Screening rate in high risk populations | Annual | $\checkmark$ |
| d | Screening rate in low risk populations | Every 5 years | $\checkmark$ |
| e | Lifetime screens in high risk populations | 45,583 | Calculated |
| f | Lifetime screens in low risk populations | 170,778 | Calculated |
| g | Total screens | 216,361 | =e+f |
| h | \# of true positive screens | 23.09 | Table 2, row x |
| i | Estimated \# of false positive screens | 23.09 | =h |
|  | Costs of screening and counseling |  |  |
| j | Cost of 10-minute office visit | \$34.85 | Ref Doc |
| k | Value of patient time and travel for office visit | \$59.38 | Ref Doc |
| I | Proportion of office visit required | 0.50 | Assumed |
| m | Cost per screen | \$7 | $\checkmark$ |
| n | Cost per true/false positive screen | \$400 | $\checkmark$ |
| o | Cost per counselling session | \$84.45 | $\checkmark$ |
| p | Cost of screening | \$5,303,081 | $=\left(\mathrm{g}^{*} \mathrm{j}^{*} \mathrm{l}\right)+\left(\mathrm{g}^{*} \mathrm{~m}\right)+(\mathrm{h}+\mathrm{i})^{*} \mathrm{n}$ |
| q | Cost of counselling | \$3,900 | $=(\mathrm{h}+\mathrm{i}) * \mathrm{O}$ |
| r | Patient time costs | \$6,423,750 | = g*k* |
|  | Costs of antiretrovirals |  |  |
| S | Cost per day of treatment | \$26 | $\checkmark$ |
| t | Cost of antiretrovirals | \$9,640,931 | $\begin{gathered} \text { =Table } 2, \text { row x * Table 2, } \\ \text { row cc *365 *s } \end{gathered}$ |
|  | Costs avoided |  |  |
| u | HIV infections avoided - treatment with ART | 28.49 | Table 2, row kk + Table 2, row nn |
| v | Cost of antiretrovirals avoided | -\$11,894,198 | $\begin{gathered} =-\mathrm{u} * \text { Table 2, row } \\ \text { cc*365*s } \end{gathered}$ |
| w | Annual direct medical costs (excluding medications) associated with symptomatic HIV | \$2,843 | $\checkmark$ |
| X | Direct medical costs avoided | -\$3,563,246 | = -u* Table 2, row cc*w |
|  | CE calculation |  |  |
| y | Cost of screening and counseling (undiscounted) | \$11,730,731 | = $p+q+r$ |
| z | Cost of antiretrovirals (undiscounted) | \$9,640,931 | = t |
| aa | Costs avoided (undiscounted) | -\$15,457,444 | = $\mathrm{v}+\mathrm{x}$ |
| bb | QALYs saved (undiscounted) | 360 | Table 2, row qq |
| cc | Cost of screening and counseling (1.5\% discount rate) | \$8,603,838 | Calculated |
| dd | Cost of antiretrovirals (1.5\% discount rate) | \$7,071,086 | Calculated |
| ee | Costs avoided (1.5\% discount rate) | -\$11,337,175 | Calculated |
| ff | QALYs saved (1.5\% discount rate) | 264 | Calculated |
| gg | CE (\$/QALY saved) | \$16,434 | =(cc+dd+ee)/ff |

$V=$ Estimates from the literature
We also modified several major assumptions and recalculated the cost per QALY as follows:

- Assume the prevalence of individuals living with HIV infections in BC is decreased from 12,100 to 9,700 (Table 2, row $a$ ): $\mathrm{CE}=\$ 24,483$.
- Assume the prevalence of individuals living with HIV infections in BC is increased from 12,100 to 14,500 (Table 2, row a): CE = \$11,049.
- Assume that the early initiation of antiretroviral therapy is associated with a $96 \%$ reduction (from 64\%) in the transmission rate per person-year (Table 2, row hh): CE $=-\$ 12,463$.
- Assume that the early initiation of antiretroviral therapy is associated with a $25 \%$ reduction (from 64\%) in the transmission rate per person-year (Table 2, row $h h$ ): CE $=\$ 80,739$.
- Assume screening once every 10 years rather than once every 5 years in the low-risk population (Table 3, row $d$ ): $\mathrm{CE}=\$ 3,521$.
- Assume screening once per lifetime rather than once every 5 years in the low-risk population (Table 3, row $d$ ): $\mathrm{CE}=-\$ 6,669$.
- Assume the cost of screening is reduced from $\$ 7$ and $\$ 400$ to $\$ 5$ and $\$ 300$ (Table 3, rows $m \& n$ ): $\mathrm{CE}=\$ 15,218$.
- Assume the cost of screening is increased from $\$ 7$ and $\$ 400$ to $\$ 9$ and $\$ 500$ (Table 3, rows $m \& n$ ): $\mathrm{CE}=\$ 17,649$.
- Assume the proportion of an office visit required is reduced from 0.50 to 0.33 (Table 3 , row $l$ ): $\mathrm{CE}=\$ 6,803$.
- Assume the proportion of an office visit required is increased from 0.50 to 0.67 (Table 3, row $l$ ): $\mathrm{CE}=\$ 26,084$.
- Assume the average annual cost of antiretrovirals for HIV is increased from $\$ 26$ to $\$ 47$ per day (Table 3, row $s$ ): $\mathrm{CE}=\$ 11,377$.


## Summary



## Chlamydia / Gonorrhea

There is a strong overlap in the at-risk populations for chlamydia and gonorrhea with both STIs often seen in the same individual. Indeed, the USPSTF recommends "chlamydia and gonorrhea screening for all sexually active women younger than 25 years (including adolescents), even if they are not engaging in high-risk sexual behaviours." ${ }^{1888}$ They further note that younger women tend to be at higher risk as they tend to have more new sex partners, their immune system tends to be relatively immature and the presence of "columnar epithelium on the adolescent exocervix." ${ }^{589}$

Following are the specific recommendations from the USPSTF and the CTFPHC with respect to screening for chlamydia and gonorrhea.

## USPSTF Recommendations (2014)

The USPSTF recommends screening for chlamydia in sexually active females aged 24 years or younger and in older women who are at increased risk for infection. (B recommendation)

The USPSTF recommends screening for gonorrhea in sexually active females aged 24 years or younger and in older women who are at increased risk for infection. (B recommendation $)^{590}$

## CTFPHC Recommendations (1994)

The CTFPHC recommendations have not been updated since 1994.
Although there is sufficient evidence linking chlamydial infections to many complications, there is currently insufficient evidence in males and non-pregnant females to show that screening is effective in preventing these complications. Thus routine screening is not recommended in the general population ( $D$ Recommendation). ${ }^{591}$

The low prevalence rate of infection with N. gonorrheae would make mass screening of the general population an inefficient intervention (D Recommendation). However, screening should be performed in certain populations: 1) individuals under 30 years, particularly adolescents, with at least 2 sexual partners in the previous year; 2) prostitutes; 3) sexual contacts of individuals known to have a sexually transmitted disease; and 4) age $\leq 16$ years at first intercourse (A Recommendation). ${ }^{592}$

## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with screening females less than 30 years of age at increased risk for infection with chlamydia and gonorrhea.

The USPSTF recommends that screening be performed in all sexually active females younger than 25. The CTFPHC also recommends screening in individuals under 30 years with at least

[^130]2 sexual partners in the previous year. This means that approximately 189,099 females would be eligible for screening in BC in 2017 (see Table 1).

| Table 1: Relevant Female Population for |
| :--- | :--- | :--- | :--- |
| Chlamydia/Gonorrhea Screening in B.C. |

In estimating CPB, we used the results based on a state transition simulation model developed by Hu and colleagues. ${ }^{593}$ They found the most cost-effective approach to screening included annual screening in at-risk women ages 15 to 29 years of age followed by semi-annual screening for those with a history of infection. Our analysis is based on the assumption that this screening approach would be followed. Unless otherwise noted, the following assumptions are based on their analysis.

- In the absence of screening, the lifetime risk of chronic pelvic pain, infertility and ectopic pregnancy is $3.44 \%, 3.88 \%$ and $1.74 \%$, respectively (Table 2 , rows $d, e \& f$ ).
- With the screening protocol noted above, the lifetime risk of chronic pelvic pain, infertility and ectopic pregnancy is reduced by $41 \%$ (Table 2 , row $g$ ).
- The quality of life impact estimates for chronic pelvic pain, infertility and ectopic pregnancy can have a significant impact on model results. ${ }^{594}$
- Hu and colleagues suggest that chronic pelvic pain is associated with a 0.40 reduction in quality of life for a period of 5 years. ${ }^{595}$ The GBD study, however, found that

[^131]moderate pelvic pain is associated a disability weight of 0.114 ( $95 \% \mathrm{CI}$ of 0.078 to 0.159 ). ${ }^{596}$ Given the average QoL of women ages less than 30 of 0.914 (see Reference Document), the 0.114 disability weight results in a reduced QoL of $12.5 \%$ ( $95 \%$ CI of $8.5 \%$ to $17.4 \%$ ) (Table 2, row $n$ ).

- Hu and colleagues suggest that infertility is associated with a 0.18 reduction in quality of life up until age $50 .{ }^{597}$ The GBD study, however, found that primary infertility ("wants to have a child and has a fertile partner but the couple cannot conceive") is associated with a disability weight of just 0.008 ( $95 \% \mathrm{CI}$ of 0.003 to $0.015) .{ }^{598}$ Given the average QoL of women ages less than 50 of approximately 0.886 (see Reference Document), the 0.008 disability weight results in a reduced QoL of $0.9 \%$ ( $95 \% \mathrm{CI}$ of $0.3 \%$ to $1.7 \%$ ). We assumed the average infection would occur at age $21^{599}$ with 29 potential years of infertility (Table 2 , rows $o$ ).
- Hu and colleagues suggest that ectopic pregnancy is associated with a 0.42 reduction in quality of life for a period of 4 weeks. ${ }^{600}$ The GBD study, however, found that an ectopic pregnancy is associated a disability weight of 0.114 ( $95 \% \mathrm{CI}$ of 0.078 to $0.159) .{ }^{601}$ Given the average QoL of women ages less than 30 of 0.914 (see Reference Document), the 0.114 disability weight results in a reduced QoL of $12.5 \%$ ( $95 \%$ CI of $8.5 \%$ to $17.4 \%$ ) (Table 2, rows $p$ ).
- Other assumptions used in assessing the clinically preventable burden are detailed in the Reference Document.
Based on these assumptions, the calculation of CPB (Table 2, row $t$ ) is 143 QALYs. This represents the potential CPB moving from no screening to approximately $55 \%$ screening uptake.

[^132]Table 2: CPB of Screening to Detect and Treat Chlamydia/Gonorrhea in a Birth Cohort of 40,000 (B.C.)

| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | At-risk population in B.C. birth cohort of 40,000 | 20,000 | $\checkmark$ |
| b | Potential adherence with screening | 55\% | Ref Doc |
| c | At-risk population screened | 11,000 | = a*b |
| d | Lifetime risk of chronic pelvic pain (CPP) without screening | 3.44\% | $\checkmark$ |
| e | Lifetime risk of infertility without screening | 3.88\% | $\checkmark$ |
| f | Lifetime risk of ectopic pregnancy (EP) without screening | 1.74\% | $\checkmark$ |
| g | Effectiveness of screening in reducing CPP, infertility and EP | 41\% | $\checkmark$ |
| h | Lifetime risk of chronic pelvic pain with screening | 2.03\% | $=(1-\mathrm{g})^{*} \mathrm{~d}$ |
| i | Lifetime risk of infertility with screening | 2.29\% | $=(1-\mathrm{g})^{*} \mathrm{e}$ |
| j | Lifetime risk of ectopic pregnancy with screening | 1.03\% | $=(1-\mathrm{g})^{*} \mathrm{f}$ |
| k | Cases of chronic pelvic pain avoided with screening | 155 | =(c*d)-(c*h) |
| 1 | Cases of infertility avoided with screening | 175 | $=\left(c^{*} e\right)-\left(c^{*} \mathrm{i}\right)$ |
| m | Cases of ectopic pregnancy avoided with screening | 79 | $=\left(c^{*} \mathrm{f}\right)-\left(\mathrm{c}^{*} \mathrm{j}\right)$ |
| n | QALYs parameters - chronic pelvic pain (5 years) | 0.125 | $\checkmark$ |
| 0 | QALYs parameters - infertility (to age 50) | 0.009 | $\checkmark$ |
| p | QALYs parameters - ectopic pregnancy (4 weeks) | 0.125 | $\checkmark$ |
| q | QALYs gained with screening - chronic pelvic pain | 97 | =k*n*5 |
| $r$ | QALYs gained with screening - infertility | 46 | =1*o*29 |
| S | QALYs gained with screening - ectopic pregnancy | 0.8 | =m*p*0.077 |
| t | Total QALYs gained, $55 \%$ adherence with screening | 143 | =q+r+s |

$v=$ Estimates from the literature
As noted by Hu and colleagues, the effectiveness and cost-effectiveness associated with their modelling is highly sensitive to a number of key assumptions. ${ }^{602}$ Furthermore, there is significant debate about these key assumptions. For example, Hu and colleagues assumed that $30 \%$ of infections with chlamydia would lead to acute pelvic inflammatory disease (PID), with a range from $10-40 \%$. Subsequent research suggests that the rate might be much lower, resulting in a change in the lower end of the range from $10 \%$ to just $0.43 \%{ }^{603,604}$ Others indicate that we simply do not know very much about the natural progression from infection with either chlamydia or gonorrhea to PID. ${ }^{605}$

There is also significant debate about whether screening is associated with any significant reduction in PID and its sequelae. In a seminal article published in the New England Journal of Medicine in 1996, Scholes et al. present the results of a randomized controlled clinical trial in which they observed a significant reduction in PID in women screened for chlamydia (relative risk of $0.44 ; 95 \% \mathrm{CI}$ of 0.20 to 0.90 ). ${ }^{606}$ Subsequent research, however, has not been able to replicate these results. The Prevention of Pelvic Infection (POPI) trial in the UK, also

[^133]a randomized controlled trail, found a non-significant reduction in PID associated with screening (relative risk of $0.65 ; 95 \% \mathrm{CI}$ of 0.34 to 1.22 ). ${ }^{607}$

Assumptions about the proportion of women with an infection that progresses to PID and the effectiveness of screening (and early treatment) in reducing the proportion of women with an infection who progress to PID are critical to any analysis about the effectiveness and costeffectiveness of screening. In fact, Low notes that "under realistic assumptions, introducing a chlamydia screening programme is likely to be an expensive intervention". ${ }^{608}$ She further notes that many chlamydia screening programs have been uncritically accepted as being effective.

With these caveats in mind, we modified the following major assumptions and recalculated the CPB as follows:

- Assume the potential adherence rate with screening is reduced from $55 \%$ to $45 \%$ (Table 2, row $b$ ): $\mathrm{CPB}=117$.
- Assume the potential adherence rate with screening is increased from $55 \%$ to $65 \%$ (Table 2, row $b$ ): $\mathrm{CPB}=169$.
- Assume the effectiveness of screening in reducing chronic pelvic pain, infertility and ectopic pregnancies is reduced from $41 \%$ to $10 \%$ (Table 2, rows $g$ ): $\mathrm{CPB}=35$.
- Assume that the QoL reduction associated with chronic pelvic pain is reduced from $12.5 \%$ to $8.5 \%$ (Table 2 - row $n$ ), the QoL reduction associated with infertility is reduced from $0.9 \%$ to $0.3 \%$ (Table 2 - row $o$ ) and the QoL reduction associated with ectopic pregnancy is reduced from $12.5 \%$ to $8.5 \%$ (Table 2 - row $p$ ): CPB $=84$.
- Assume that the QoL reduction associated with chronic pelvic pain is increased from $12.5 \%$ to $17.4 \%$ (Table 2 - row $n$ ), the QoL reduction associated with infertility is increased from $0.9 \%$ to $1.7 \%$ (Table 2 - row $o$ ) and the QoL reduction associated with ectopic pregnancy is increased from $12.5 \%$ to $17.4 \%$ (Table 2 - row $p$ ): CPB $=$ 222.


## Modelling Cost-Effectiveness

In this section, we will calculate the CE associated with screening females less than 30 years of age at increased risk for infection with chlamydia and gonorrhea.

In modelling CE, we made the following assumptions:

- Proportion of at-risk population with infection - We assumed that $5.68 \%$ of the atrisk population would test positive for either chlamydia or gonorrhea (Table 3, row f). ${ }^{609}$ This assumption was varied between $2 \%$ and $33 \%$ in the sensitivity analysis. ${ }^{610}$

[^134]- Screening protocol - We assumed that screening included annual screening in atrisk women ages 15 to 29 years of age followed by semi-annual screening for those with a history of infection (Table 3, rows $g, h$ and $i$ ). ${ }^{611}$
- Costs of screening tests -Hu et al. estimated the cost of a urine nucleic acid amplification test to be $\$ 13$ (2000 USD) ${ }^{612}$ or $\$ 15.28$ in 2017 CAD. Robinson et al. estimated the costs to be $£ 7.35$ (in 2005) ${ }^{613}$ or $\$ 16.17$ in 2017 CAD. We used an estimate of $\$ 15.73$ (the midpoint between the two estimates) per screening test in the model (Table 3, row $m$ ).
- Average cost of antibiotic treatment - The recommended drug regimen for chlamydia is doxycycline 100 mg PO bid for 7 days (estimated cost of $\$ 22.18$ including dispensing fee ${ }^{614}$ ) or azithromycin 1 g PO in a single dose (estimated cost of $\$ 18.10$ including dispensing fee ${ }^{615}$ ) while the recommended drug regimen for gonorrhea is cefixime 800 mg PO in a single dose (estimated cost of $\$ 19.04$ including dispensing fee ${ }^{616}$ ) or ceftriaxone 250 mg in a single dose plus azithromycin 1 g PO in a single dose. ${ }^{617}$ We used an average cost of $\$ 19.77$ (Table 3, row $p$ ) with a range from \$18.10 to \$22.18.
- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the estimated cost per QALY would be $\$ 57,174$ (see Table 3, row $v$ ).

[^135]Table 3: CE of Screening to Detect and Treat Chlamydia/Gonorrhea in a Birth Cohort of 40,000 (B.C.)

| Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | At-risk population screened | 11,000 | Table 2, row c |
| b | \# of annual screens between age 15 and 24 | 10 | $\checkmark$ |
| C | Total \# of screens, 15-24 | 110,000 | =a*b |
| d | \% Population at-risk between 25-29 | 6\% | $\checkmark$ |
| e | Total \# of screens, 25-29 | 3,300 | =d*a*5 |
| f | \% with chlamydia/gonorrhea infection | 5.68\% | $\checkmark$ |
| g | Total screens - positive | 6,435 | $=(\mathrm{c}+\mathrm{e}) * \mathrm{~d}$ |
| h | Total screens - negative | 106,865 | $=\mathrm{c}+\mathrm{e}-\mathrm{g}$ |
| i | Additional follow-up screens in positive women | 6,435 | = g |
|  | Costs of screening |  |  |
| j | Cost of 10-minute office visit | \$34.85 | Ref Doc |
| k | Cost of patient time and travel for office visit | \$59.38 | Ref Doc |
| 1 | Portion of office visit needed | 50\% | Ref Doc |
| m | Cost per screening test | \$15.73 | $\checkmark$ |
| n | Costs of screening | \$7,524,774 | $=(\mathrm{g}+\mathrm{h}+\mathrm{i}) *(((\mathrm{j}+\mathrm{k}) * \mathrm{l}) * \mathrm{~m})$ |
| 0 | Costs of antibiotics |  |  |
| p | Cost per treatment | \$19.77 | $\checkmark$ |
| q | Cost of antibiotics | \$127,218 | $=g^{*} p$ |
|  | CE calculation |  |  |
| $r$ | Costs (undiscounted) | \$7,651,992 | $=\mathrm{n}+\mathrm{q}$ |
| S | QALYs saved (undiscounted) | 143 | Table 2, row t |
| t | Costs (1.5\% discount rate) | \$6,813,920 | Calculated |
| u | QALYs saved (1.5\% discount rate) | 119 | Calculated |
| V | CE (\$/QALY saved) | \$57,174 | = t/u |

$V=$ Estimates from the literature
We also modified several major assumptions and recalculated the cost per QALY as follows:

- Assume the effectiveness of screening in reducing chronic pelvic pain, infertility and ectopic pregnancies is reduced from $41 \%$ to $10 \%$ (Table 2, row $b$ ): $\mathrm{CE}=\$ 234,414$.
- Assume that the QoL reduction associated with chronic pelvic pain is reduced from $12.5 \%$ to $8.5 \%$ (Table 2 - row $n$ ), the QoL reduction associated with infertility is reduced from $0.9 \%$ to $0.3 \%$ (Table 2 - row $o$ ) and the QoL reduction associated with ectopic pregnancy is reduced from $12.5 \%$ to $8.5 \%$ (Table 2 , row $p$ ): $\mathrm{CE}=\$ 96,519$.
- Assume that the QoL reduction associated with chronic pelvic pain is increased from $12.5 \%$ to $17.4 \%$ (Table 2 - row $n$ ), the QoL reduction associated with infertility is increased from $0.9 \%$ to $1.7 \%$ (Table 2 - row $o$ ) and the QoL reduction associated with ectopic pregnancy is increased from $12.5 \%$ to $17.4 \%$ (Table 2 , row $p$ ): $\mathrm{CE}=$ \$37,189.
- Assume that the proportion of the at-risk population who would test positive for either chlamydia or gonorrhea is reduced from $5.68 \%$ to $2.0 \%$ (Table 3, row $f$ ): $\mathrm{CE}=$ \$54,601.
- Assume that the proportion of the at-risk population who would test positive for either chlamydia or gonorrhea is increased from $5.68 \%$ to $33.0 \%$ (Table 3, row $f$ ): CE $=\$ 76,281$.
- Assume the portion of an office visit required is decreased from 50 to $33 \%$ (Table 3, row $l$ ): $\mathrm{CE}=\$ 42,843$.
- Assume the portion of an office visit required is increased from $50 \%$ to $67 \%$ (Table 3, row $l$ ): $\mathrm{CE}=\$ 71,506$.
- Assume the cost for antibiotic treatment is decreased from $\$ 19.77$ to $\$ 18.10$ (Table 3 , row $p$ ): $\mathrm{CE}=\$ 57,094$.
- Assume the cost for antibiotic treatment is increased from $\$ 19.77$ to $\$ 22.18$ (Table 3, row $p$ ): $\mathrm{CE}=\$ 57,290$.


## Summary

| Table 4: Screening to Diagnose and Treat Chlamydia/Gonorrhea Infections in a Birth Cohort of 40,000 Summary |  |  |  |
| :---: | :---: | :---: | :---: |
|  | Base Case | Range |  |
| CPB (Potential QALYs Gained) |  |  |  |
| 1.5\% Discount Rate | 119 | 29 | 183 |
| 3\% Discount Rate | 100 | 24 | 153 |
| 0\% Discount Rate | 143 | 35 | 222 |
| CE (\$/QALY) including patient time costs |  |  |  |
| 1.5\% Discount Rate | \$57,174 | \$37,189 | \$234,414 |
| 3\% Discount Rate | \$60,733 | \$39,750 | \$249,007 |
| 0\% Discount Rate | \$53,410 | \$34,494 | \$218,983 |
| CE (\$/QALY) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | \$30,612 | \$19,912 | \$125,511 |
| 3\% Discount Rate | \$32,518 | \$21,283 | \$133,324 |
| 0\% Discount Rate | \$28,597 | \$18,469 | \$117,248 |

## Hepatitis C Virus

## United States Preventive Services Task Force Recommendations (2013)

Hepatitis C virus is the most common chronic bloodborne pathogen in the United States and a leading cause of complications from chronic liver disease. The prevalence of the anti-HCV antibody in the United States is approximately $1.6 \%$ in noninstitutionalized persons. According to data from 1999 to 2008, about three fourths of patients in the United States living with HCV infection were born between 1945 and 1965, with a peak prevalence of $4.3 \%$ in persons aged 40 to 49 years from 1999 to 2002. The most important risk factor for HCV infection is past or current injection drug use, with most studies reporting a prevalence of $50 \%$ or more. The incidence of HCV infection was more than 200000 cases per year in the 1980s but decreased to 25000 cases per year by 2001. According to the Centers for Disease Control and Prevention (CDC), there were an estimated 16000 new cases of HCV infection in 2009 and an estimated 15000 deaths in 2007. Hepatitis C-related endstage liver disease is the most common indication for liver transplants among U.S. adults, accounting for more than 30\% of cases. Studies suggest that about one half of the recently observed 3-fold increase in incidence of hepatocellular carcinoma is related to acquisition of HCV infection 2 to 4 decades earlier.

The USPSTF recommends screening for HCV infection in persons at high risk for infection. The USPSTF also recommends offering 1-time screening for HCV infection to adults born between 1945 and 1965. (B recommendation) ${ }^{618}$

## Canadian Task Force on Preventive Health Care Recommendations (2017)

The task force recommends against screening for HCV in asymptomatic Canadian adults (including baby boomers) who are not at elevated risk of HCV infection. Strong recommendation based on very low-quality evidence.

A strong recommendation against screening is warranted given its uncertain benefits but the certainty that it would lead to high levels of resource consumption. Referring individuals with screen-detected HCV for assessment would reduce access to assessment and treatment for people with clinically evident HCV. ${ }^{619}$

In 2016, BC had an HCV infection rate of 48.6 per 100,000 population, ranging from 18.7 in the Richmond HSDA to 74.2 in the Fraser East HSDA. The rate in BC is significantly higher than the Canadian average of 30.4 / 100,000 (in 2015). ${ }^{220}$ As a result, the Lifetime Prevention Schedule Expert Committee has recommended that the analysis of CPB and CE be completed following the USPSTF recommendation to offer one-time screening for HCV infection to adults born between 1945 and 1965.

## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with one-time screening for HCV infection in BC adults born between 1945 and 1965.

[^136]In modelling CPB, we made the following assumptions:

- There are an estimated 1,301,000 individuals in BC born between 1945 and 1965 (ages 52 to 72 in 2017) or $27.0 \%$ of BC's population of 4.82 million. This translates into an at-risk population of 11,604 in a birth cohort of 40,000 (27.0\%) (Table 1, row $a)$.
- The estimated prevalence of HCV infection in this at-risk population is $3.60 \%{ }^{621}$ (Table 1, row $e$ ).
- The probability of cirrhosis in individuals with HCV infection is $15 \%{ }^{622}$ (Table 1 , row $h$ ).
- The annual probability of transitioning from cirrhosis to decompensated cirrhosis is $3.90 \%$. The annual probability of transitioning from cirrhosis to liver cancer is $2.50 \%{ }^{623}$ (Table 1, rows $j \& k$ ).
- The annual probability of a liver transplant following decompensated cirrhosis or liver cancer is $3.10 \%{ }^{624}$ (Table 1, row $l$ ).
- The annual probability of death due to decompensated cirrhosis is $13.5 \%$. The annual probability of death due to liver cancer is $40.9 \%^{625}$ (Table 1 , rows $n \& o$ ).
- Quality of life losses associated with cirrhosis, decompensated cirrhosis and liver cancer are $0.19,0.30$ and 0.33 , respectively ${ }^{626}$ (Table 1 , rows $p, q \& r$ ).
- The average age at which an individual is identified with HCV infection and subsequent cirrhosis is 62 , the mid-point between 52 and 72 (Table 1 , row $s$ ).
- The effectiveness of antiviral therapy in producing a sustained viral response (i.e. a cure) is $95 \%{ }^{627,628,629,630}$ (Table 1, row $x$ ).
- Other assumptions used in assessing the CPB are detailed in the Reference Document.

Based on these assumptions, the calculation of CPB is 3,920 QALYs (Table 1, row $y$ ). This represents the potential CPB of moving from no screening to screening uptake of $48 \%$.

[^137]Table 1: CPB of Screening to Detect and Treat Hepatitis C Infection in a Birth Cohort of 40,000 (B.C.)

| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | At-risk population in B.C. | 1,301,000 | $\checkmark$ |
| b | B.C. population | 4,817,160 | $\checkmark$ |
| C | \% of B.C. population at risk | 27.0\% | = a/b |
| d | At-risk population in B.C. birth cohort of 40,000 | 10,803 | = c *40,000 |
| e | Estimated prevalence of HCV in at-risk population | 3.60\% | $\checkmark$ |
| f | Adherence with screening | 48\% | Ref Doc |
| g | Cases of HCV infection detected through screening | 187 | $=d^{*} e^{*} \mathrm{f}$ |
| h | Probability of cirrhosis in HCV positive individuals | 15.0\% | $\checkmark$ |
| i | Cases of cirrhosis detected through screening | 28 | $=h^{*}{ }^{\text {i }}$ |
| j | Annual probability of decompensated cirrhosis with cirrhosis | 3.9\% | $\checkmark$ |
| k | Annual probability of liver cancer with cirrhosis | 2.5\% | $\checkmark$ |
| I | Annual probability of liver transplantation with decompensated cirrhosis or liver cancer | 3.1\% | $\checkmark$ |
| m | \# of liver transplants | 0.72 | Calculated |
| n | Annual probability of death - decompensated cirrhosis | 13.5\% | $\checkmark$ |
| 0 | Annual probability of death - liver cancer | 40.9\% | $\checkmark$ |
| p | Reduction in QoL associated with cirrhosis | 0.19 | $\checkmark$ |
| q | Reduction in QoL associated with decompensated cirrhosis | 0.30 | $\checkmark$ |
| r | Reduction in QoL associated with liver cancer | 0.33 | $\checkmark$ |
| S | Average age | 62 | $\checkmark$ |
| t | QALYs Lost - Cirrhosis | 703 | Calculated |
| u | QALYs Lost - Decompensated cirrhosis | 1,956 | Calculated |
| V | QALYs Lost - Liver cancer | 1,595 | Calculated |
| w | \% Eligible for and accepting treatment | 97\% | $\checkmark$ |
| X | Effectiveness of antiviral therapy in producing a sustained viral response (i.e. a cure) | 95\% | V |
| y | Total QALYs gained, Utilization increasing from 0\% to 48\% | 3,920 | $=(t+u+v)^{*} w^{*} x$ |

$V=$ Estimates from the literature

We also modified several major assumptions and recalculated the CPB as follows:

- Assume the prevalence of HCV infection in the at-risk population is reduced from $3.60 \%$ to $1.60 \%$ (Table 1, row $e$ ): $\mathrm{CPB}=1,742$.
- Assume the prevalence of HCV infection in the at-risk population is increased from $3.60 \%$ to $5.60 \%$ (Table 1, row $e$ ): $\mathrm{CPB}=6,097$.
- Assume the probability of cirrhosis in HCV positive individuals is decreased from $15 \%$ to $10 \%$ (Table 1, row $h$ ): $\mathrm{CPB}=2,613$.
- Assume the probability of cirrhosis in HCV positive individuals is increased from $15 \%$ to $20 \%$ (Table 1, row $h$ ): $\mathrm{CPB}=5,226$.


## Modelling Cost-Effectiveness

In this section, we will calculate the CE associated with screening for HCV infection in BC adults born between 1945 and 1965 .

In modelling CE, we made the following assumptions:

- Costs of screening tests - we estimated the cost of a hepatitis C antibody EIA test to be $\$ 24.28$ (Table 2, row $g$ ). ${ }^{631}$ A positive screening test would be followed by a hepatitis C RNA amp probe and a hepatitis C RNA quant test to confirm RNA detection and quantify RNA for a total cost per positive screening test of $\$ 234.62$ (Table 2, row $h$ ). ${ }^{632}$
- Cost of treatment - the price for HCV direct-acting antivirals is estimated at approximately $\$ 55,000$ per treatment (Table 2, row $l$ ). ${ }^{633,634}$ In the sensitivity analysis, this cost was increased/decreased by $25 \%$.
- Follow-up - Patients on antiviral treatment would require an average of 9 follow-up visits to their physician, at weeks $2,4,8,12,16,24,32,40$ and $48 .{ }^{635}$ Each visit would include three lab tests (CBC, Renal panel and TSH). The costs of the lab tests are estimated at $\$ 10.94, \$ 12.22$ and $\$ 23.64$, respectively (Table 2, row $o$ ). ${ }^{636}$
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.
- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.

Based on these assumptions, the estimated cost per QALY would be $\$ 3,427$ (Table 2, row $u$ ).

[^138]Table 2: CE of Screening to Detect and Treat Hepatitis C Infection in a Birth Cohort of 40,000 (B.C.)

| Row Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | At-risk Population in a BC birth cohort of 40,000 | 10,803 | Table 1, row d |
| b | Estimated prevalence of HCV in at-risk population | 3.60\% | Table 1, row e |
| C | Cases of HCV infection detected through screening | 187 | Table 1, row g |
| d | \% Eligible for and accepting treatment | 97\% | Table 1, row w |
|  | Costs of screening |  |  |
| e | Cost of 10-minute office visit | \$34.85 | Ref Doc |
| f | Portion of office visit needed | 50\% | Ref Doc |
| g | Cost per negative screening test | \$24.28 | $\checkmark$ |
| h | Cost per positive screening tests | \$234.62 | $\checkmark$ |
| i | Costs of screening | \$497,592 | $\begin{gathered} =\left(a^{*} e^{* f}\right)+\left(a^{*} \mathrm{~g}\right)+\left(c^{*} \mathrm{~h}\right)+ \\ \left(\mathrm{c}^{*} \mathrm{e}^{*} \mathrm{f}\right) \end{gathered}$ |
| j | Cost of patient time and travel for office visit | \$59.38 | Ref Doc |
| k | Patient time costs - screening | \$326,285 | $=\left(j^{*} \mathrm{f}\right) *(\mathrm{a}+\mathrm{c})$ |
|  | Cost of treatment |  |  |
| 1 | Drug costs per treatment - antiviral therapy | \$55,000 | $\checkmark$ |
| m | Costs of antiviral therapy | \$9,959,198 | $=\left(c^{*} d\right) * \mid$ |
| n | Follow-up visits during treatment | 9 | $\checkmark$ |
| 0 | Cost of lab tests/follow-up | \$46.80 | $\checkmark$ |
| p | Follow-up costs | \$229,835 | $=\left(c^{*} d\right) *(e+j+o) * n$ |
|  | CE calculation |  |  |
| q | Costs (undiscounted) | \$11,012,910 | = i $+\mathrm{k}+\mathrm{m}+\mathrm{p}$ |
| $r$ | QALYs saved (undiscounted) | 3,920 | Table 1, row y |
| S | Costs (1.5\% discount rate) | \$11,012,910 | Calculated |
| t | QALYs saved (1.5\% discount rate) | 3,213 | Calculated |
| u | CE (\$/QALY saved) | \$3,427 | = s/t |

$V=$ Estimates from the literature

We also modified several major assumptions and recalculated the cost per QALY as follows:

- Assume the prevalence of HCV infection in the at-risk population is reduced from $3.60 \%$ to $1.60 \%$ (Table 1, row $e$ ): $\mathrm{CE}=\$ 3,727$.
- Assume the prevalence of HCV infection in the at-risk population is increased from $3.60 \%$ to $5.60 \%$ (Table 1, row $e$ ): $\mathrm{CE}=\$ 3,342$.
- Assume the probability of cirrhosis in HCV positive individuals is decreased from $15 \%$ to $10 \%$ (Table 1, row $h$ ): $\mathrm{CE}=\$ 5,141$.
- Assume the probability of cirrhosis in HCV positive individuals is increased from $15 \%$ to $20 \%$ (Table 1, row $h$ ): $\mathrm{CE}=\$ 2,570$.
- Assume the portion of an office visit needed is decreased from $50 \%$ to $33 \%$ (Table 2, row $f$ ): $\mathrm{CE}=\$ 3,482$.
- Assume the portion of an office visit needed is increased from $50 \%$ to $67 \%$ (Table 2, row $f$ ): $\mathrm{CE}=\$ 3,373$.
- Assume the cost of antiviral treatment in increased from $\$ 55,000$ to $\$ 68,750$ (Table 2 , row $l$ : $\mathrm{CE}=\$ 4,202$.
- Assume the cost of antiviral treatment in decreased from \$55,000 to \$41,250 (Table 2 , row $l$ ): $\mathrm{CE}=\$ 2,652$.


## Summary

| in a Birth Cohort of 40,000 (B.C.) |  |  |  |
| :---: | :---: | :---: | :---: |
| Summary |  |  |  |
| Base |  |  |  |
| CPB (Potential QALYs Gained) |  |  |  |
| Gap between 0\% and 'Best in the World' (48\%) |  |  |  |
| 1.5\% Discount Rate | 3,213 | 1,428 | 4,998 |
| 3\% Discount Rate | 2,661 | 1,183 | 4,139 |
| 0\% Discount Rate | 3,920 | 1,742 | 6,097 |
| Gap between B.C. Current (33\%) and 'Best in the World' (48\%) |  |  |  |
| 1.5\% Discount Rate | 1,004 | 446 | 1,562 |
| 3\% Discount Rate | 832 | 370 | 1,293 |
| 0\% Discount Rate | 1,225 | 544 | 1,905 |
| CE (\$/QALY) including patient time costs |  |  |  |
| 1.5\% Discount Rate | \$3,427 | \$2,570 | \$5,141 |
| 3\% Discount Rate | \$4,139 | \$3,104 | \$6,209 |
| 0\% Discount Rate | \$2,810 | \$2,107 | \$4,214 |
| CE ( $\$ / \mathrm{QALY})$ excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | \$3,296 | \$2,472 | \$4,944 |
| 3\% Discount Rate | \$3,980 | \$2,985 | \$5,970 |
| 0\% Discount Rate | \$2,702 | \$2,026 | \$4,052 |

## Behavioural Counselling Interventions

## Definition

In 2002, the USPSTF published an article outlining its vision for a broader appreciation of the importance of behavioural counselling interventions in clinical care. ${ }^{637}$ The paper includes important definitional and context information for this area and we have thus quoted liberally from the paper below.

Behavioral counseling interventions address complex behaviors that are integral to daily living; they vary in intensity and scope from patient to patient; they require repeated action by both patient and clinician, modified over time, to achieve health improvement; and they are strongly influenced by multiple contexts (family, peers, worksite, school, and community). Further, "counseling" is a broadly used but imprecise term that covers a wide array of preventive and therapeutic activities, from mental health or marital therapy to the provision of health education and behavior change support. Thus, we have chosen to use the term "behavioral counseling interventions" to describe the range of personal counseling and related behaviorchange interventions that are effectively employed in primary care to help patients change health-related behaviors. (p.270)

Behavioral counseling interventions in clinical care are those activities delivered by primary care clinicians and related healthcare staff to assist patients in adopting, changing, or maintaining behaviors proven to affect health outcomes and health status. Common health promoting behaviors include smoking cessation, healthy diet, regular physical activity, appropriate alcohol use, and responsible use of contraceptives. (p. 269-70)

The strongest evidence for the efficacy of primary care behavior-change interventions comes from tobacco-cessation research and, to a lesser extent, problem drinking. Accumulating evidence also shows the effectiveness of similar interventions for other behaviors. These interventions often provide more than brief clinician advice. Effective interventions typically involve behavioral counseling techniques and use of other resources to assist patients in undertaking advised behavior changes. For example, intervention adjuncts to brief clinician advice may involve a broader set of healthcare team members (e.g., nurses, other office staff, health educators, and pharmacists), a number of complementary communication channels (e.g., telephone counseling, video or computer assisted interventions, self-help guides, and tailored mailings), and multiple contacts with the patient. (p. 268)

In 2014, the USPSTF published an article discussing challenges it encounters in aggregating the behavioural counselling intervention literature, including clear descriptions of the study population, intervention protocols, assessment of outcomes, and linking behaviour changes to health outcomes. ${ }^{638}$ Researchers are encouraged to pay closer attention to these issues in designing and writing up their behavioural intervention research.

[^139]
## Prevention of Sexually Transmitted Diseases

## Canadian Task Force on Preventive Health Care (2001)

A 2001 report from the CTFPHC titled "Counseling for Risky Health Habits: A Conceptual Framework for Primary Care Practitioners" noted that,

Risky lifestyle choices contribute to many contemporary health conditions. Primary care practitioners have frequent opportunities to help patients clarify issues and alter adverse behaviour patterns....The six risky behaviours addressed in this paper are appropriate targets for counseling. Some situations respond to brief on-the-spot advice, others require a few repeated counseling sessions utilizing concepts from behavioural theory, and certain ones need referral to a structured counseling program that employs a longer time-frame and allows for the opportunity to use a range of methods. ${ }^{639}$

The "six risky behaviours" include dietary patterns, unintentional injury, problem drinking, physical inactivity patterns, risky sexual patterns and cigarette smoking.

## United States Preventive Services Task Force Recommendations (2014)

The USPSTF recommends intensive behavioral counseling for all sexually active adolescents and for adults who are at increased risk for STIs. (B recommendation)

All sexually active adolescents are at increased risk for STIs. Other risk groups include adults with current STIs or other infections within the past year, adults who have multiple sex partners, and adults who do not consistently use condoms.

Clinicians should be aware of populations with a particularly high prevalence of STIs. African Americans have the highest STI prevalence of any racial/ethnic group, and prevalence is higher in American Indians, Alaska Natives, and Latinos than in white persons. Increased STI prevalence rates are also found in men who have sex with men (MSM), persons with low incomes living in urban settings, current or former inmates, military recruits, persons who exchange sex for money or drugs, persons with mental illness or a disability, current or former intravenous drug users, persons with a history of sexual abuse, and patients at public STI clinics.

Behavioral counseling interventions can reduce a person's likelihood of acquiring an STI. Interventions ranging in intensity from 30 min to $\geq 2 \mathrm{~h}$ of contact time are beneficial; evidence of benefit increases with intervention intensity. Interventions can be delivered by primary care clinicians or through referral to trained behavioral counselors. Most successful approaches provide basic information about STIs and STI transmission; assess risk for transmission; and provide training in pertinent skills, such as condom use, communication about safe sex, problem solving, and goal setting. ${ }^{640}$

[^140]
## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with behavioural counselling interventions for the prevention of sexually transmitted diseases in a British Columbia birth cohort of 40,000 .

In estimating CPB, we made the following assumptions:

- The age and sex specific incidence rates per 100,000 for acute hepatitis B are taken from the BCCDC Annual Summary of Reportable Diseases 2016. ${ }^{641}$ The age and sex specific incidence rates per 100,000 for human immunodeficiency virus (HIV) are taken from the BCCDC HIV Annual Annual Report 2015. ${ }^{642}$ The age and sex specific incidence rates per 100,000 for chlamydia, gonorrhea and syphilis infections are taken from the BCCDC Annual Report 2015. ${ }^{643}$ The incidence of human papillomavirus (HPV) infection in females is taken from an Ontario study. ${ }^{644}$ We have assumed that the age specific incidence rate for males is the same as for females. ${ }^{645}$ We calculated the incidence of herpes simplex virus type 2 (HSV-2) infection based on the number of patients within each age group who had their first herpes-related physician billings in 2006, as reported by the BC Centre for Disease Control. ${ }^{646}$ We reduced the rates of first herpes-related visits proportional to the percentage of age-specific laboratory-diagnosed HSV infections in BC that were from genital specimens and were confirmed HSV-2. In 2005, approximately $31 \%$ of HSV-2 cases were identified in males and 69\% percent in females; therefore, new cases were distributed between sexes according to these proportions (see Table 1).

| Table 1: Sexually Transmitted Infections in British Columbia Rate per 100,000 by Sex and Age Group |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | HIV |  | Chlamydia |  | Gonorrhea |  | Hepatitis B - Acute |  | Syphilis |  | HPV |  | HSV-2 |  |
|  | Female | Male | Female | Male | Female | Male | Female | Male | Female | Male | Female | Male | Female | Male |
| 10-14 | - | - | 40 | 2 | 4 | - | - | - | - | - | NA | NA | 2.8 | 1.3 |
| 15-19 | 2 | 1 | 1,433 | 322 | 121 | 64 | - | - | 1 | 6 | 25,000 | 25,000 | 140.1 | 63.3 |
| 20-24 | 1 | 11 | 1,993 | 961 | 195 | 219 | - | - | 5 | 35 | 8,800 | 8,800 | 209.6 | 94.7 |
| 25-29 | 1 | 23 | 1,111 | 895 | 162 | 281 | - | - | 3 | 64 | 8,300 | 8,300 | 222.9 | 100.7 |
| 30-39 | 4 | 14 | 427 | 395 | 76 | 202 | - | 0.3 | 2 | 61 | 13,000 | 13,000 | 248.0 | 112.2 |
| 40-59 | 2 | 13 | 86 | 103 | 17 | 69 | 0.2 | 0.3 | 1 | 49 | 7,600 | 7,600 | 164.9 | 74.5 |
| 60+ | 1 | 3 | 6 | 17 | 2 | 15 | - | 0.2 | 0 | 10 | NA | NA | 113.0 | 51.6 |
| NA = not available |  |  |  |  |  |  |  |  |  |  |  |  |  |  |

[^141]- The age- and sex- specific incidence rates were combined with years of life in a given age group by sex in the BC birth cohort to calculate the expected number of STIs by age and sex (see Tables 2 and 3).


| Table 3: Estimated Number of Sexually Transmitted Infections in a Female Birth Cohort of 20,000 |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Age Group | Mean <br> Survival <br> Rate | Individuals in Birth Cohort | Years of Life in Birth Cohort | Chlamydia | HIV | Gonorrhea | Hepatitis <br> B - Acute | Syphilis | HPV | HSV-2 |
| 15-19 | 0.995 | 19,897 | 99,484 | 1,425 | 2 | 120 | 0 | 1 | 24,871 | 139 |
| 20-24 | 0.993 | 19,865 | 99,323 | 1,979 | 1 | 193 | 0 | 4 | 8,740 | 208 |
| 25-29 | 0.992 | 19,833 | 99,163 | 1,102 | 1 | 161 | 0 | 3 | 8,231 | 221 |
| 30-34 | 0.990 | 19,795 | 98,975 | 423 | 4 | 76 | 0 | 2 | 12,867 | 245 |
| 35-39 | 0.987 | 19,741 | 98,706 | 422 | 4 | 75 | 0 | 2 | 12,832 | 245 |
| 40-44 | 0.983 | 19,662 | 98,311 | 85 | 2 | 17 | 0 | 1 | 7,472 | 162 |
| 45-49 | 0.977 | 19,546 | 97,730 | 84 | 2 | 16 | 0 | 1 | 7,427 | 161 |
| 50-54 | 0.969 | 19,375 | 96,873 | 83 | 2 | 16 | 0 | 1 | 7,362 | 160 |
| 55-59 | 0.956 | 19,118 | 95,591 | 82 | 2 | 16 | 0 | 1 | 7,265 | 158 |
| Total Ag | s 15-59 |  | 884,156 | 5,685 | 21 | 691 | 1 | 17 | 97,067 | 1,699 |

- The data in Tables 2 and 3 was used to populate rows $a-n$ in Table 4 .
- High intensity (> 2 hours) behavioural counselling interventions are associated with a $62 \%(\mathrm{OR}=0.38,95 \% \mathrm{CI}$ of $0.24-0.60)$ reduction in STI incidence in adolescents and a $30 \%(\mathrm{OR}=0.70,95 \% \mathrm{CI}$ of $0.56-0.87$ ) reduction in STI incidence in adults (Table 4 , rows $o \& p) .{ }^{647}$
- Reductions in quality of life attributable to an infection with chlamydia, gonorrhea, HPV and HSV-2 are based on data provided in the relevant appendixes of the document Vaccines for the $21^{\text {st }}$ Century: A Tool for Decision Making (Table 4, rows $y, a a, d d \& e e) .{ }^{648}$ These appendixes include an estimated rate for all sequelae following the infection, together with the time in a given state and the relevant change in quality of life over that time period.

[^142]- Vaccines for the $21^{s t}$ Century: A Tool for Decision Making suggest that chronic pelvic pain is associated with a 0.40 reduction in quality of life for a period of 22.73 years. The GBD study, however, found that moderate pelvic pain is associated a disability weight of 0.114 ( $95 \% \mathrm{CI}$ of 0.078 to 0.159 ). ${ }^{.49}$ Given the average QoL of women ages less than 30 of 0.914 (see Reference Document), the 0.114 disability weight results in a reduced QoL of $12.5 \%$ ( $95 \% \mathrm{CI}$ of $8.5 \%$ to $17.4 \%$ ). We therefore modified the assumption in Vaccines for the $21^{\text {st }}$ Century: A Tool for Decision Making from 0.40 reduction in quality of life associated with chronic pelvic pain to 0.125 .
- Vaccines for the $21^{s t}$ Century: A Tool for Decision Making suggest that infertility is associated with a 0.18 reduction in quality of life for 22.73 years. The GBD study, however, found that primary infertility ("wants to have a child and has a fertile partner but the couple cannot conceive") is associated with a disability weight of just $0.008(95 \% \mathrm{CI}$ of 0.003 to 0.015$) .{ }^{650}$ Given the average QoL of women ages less than 50 of approximately 0.886 (see Reference Document), the 0.008 disability weight results in a reduced QoL of $0.9 \%$ ( $95 \% \mathrm{CI}$ of $0.3 \%$ to $1.7 \%$ ). We therefore modified the assumption in Vaccines for the $21^{\text {st }}$ Century: A Tool for Decision Making from 0.18 reduction in quality of life associated with infertility to 0.009 .
- We assumed that the average HIV infection would occur at age $40^{651}$ with 44 years of life remaining at a $17 \%$ reduced quality of life (Table 4 , row $z$ ). ${ }^{652}$ We assumed a reduction of 0.05 QALYs per infection with syphilis (Table 4, row $c c$ ), roughly equivalent to the calculated reductions for chlamydia ( 0.049 , Table 4 , row $y$ ) and gonorrhea ( 0.055 , Table 4 , row $a a$ ). We assumed an $18.5 \%$ reduction in quality of life attributable to a hepatitis $\mathrm{B}-$ acute infection (Table 4, row $b b$ ). ${ }^{653}$
- Other assumptions used in assessing the clinically preventable burden are detailed in the Reference Document.

Based on these assumptions, the CPB associated with behavioural counselling interventions for the prevention of sexually transmitted diseases is 3,285 QALYs (Table 4, row ff).

[^143]Table 4 CPB of Behavioural Counselling Interventions for the Prevention of Sexually Transmitted Infections in a Birth Cohort of 40,000

| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Estimated number of STIs in birth cohort as adolescents - Chlamydia | 1,745 | Tables 2 and 3 |
| b | Estimated number of STIs in birth cohort as adults - Chlamydia | 7,263 | Tables 2 and 3 |
| c | Estimated number of STIs in birth cohort as adolescents - HIV | 4 | Tables 2 and 3 |
| d | Estimated number of STIs in birth cohort as adults - HIV | 128 | Tables 2 and 3 |
| e | Estimated number of STIs in birth cohort as adolescents - Gonorrhea | 183 | Tables 2 and 3 |
| f | Estimated number of STIs in birth cohort as adults - Gonorrhea | 1,722 | Tables 2 and 3 |
| g | Estimated number of STIs in birth cohort as adolescents - Hep B-Acute | 0 | Tables 2 and 3 |
| h | Estimated number of STIs in birth cohort as adults - Hep B-Acute | 2 | Tables 2 and 3 |
| i | Estimated number of STIs in birth cohort as adolescents - Syphilis | 7 | Tables 2 and 3 |
| j | Estimated number of STIs in birth cohort as adults - Syphilis | 418 | Tables 2 and 3 |
| k | Estimated number of STIs in birth cohort as adolescents - HPV | 49,715 | Tables 2 and 3 |
| I | Estimated number of STIs in birth cohort as adults - HPV | 143,554 | Tables 2 and 3 |
| m | Estimated number of STIs in birth cohort as adolescents - HSV-2 | 202 | Tables 2 and 3 |
| n | Estimated number of STIs in birth cohort as adults - HSV-2 | 2,257 | Tables 2 and 3 |
|  | Benefits Associated with Behavioural Counselling |  |  |
| 0 | Effectiveness of high intensity behavioural counselling in reducing STI incidence in adolescents | 62\% | $\checkmark$ |
| p | Effectiveness of high intensity behavioural counselling in reducing STI incidence in adults | 30\% | $\checkmark$ |
| q | Adherence with behavioural counselling | 29\% | Ref Doc |
| $r$ | Estimated \# of chlamydia infections avoided | 946 | $=\left(\left(a^{*} o\right)+\left(b^{*} \mathrm{p}\right)\right)^{*} \mathrm{q}$ |
| s | Estimated \# of HIV infections avoided | 12 | $=\left(\left(c^{*} o\right)+\left(d^{*} p\right)\right)^{*} q$ |
| t | Estimated \# of gonorrhea infections avoided | 183 | $=((\mathrm{e} * \mathrm{o})+(\mathrm{f} * \mathrm{p}))^{*} \mathrm{q}$ |
| $u$ | Estimated \# of Hep B-Acute infections avoided | 0.2 | $=\left(\left(g^{*} \mathrm{o}\right)+(\mathrm{h} * \mathrm{p})\right)^{*} \mathrm{q}$ |
| v | Estimated \# of syphilis infections avoided | 38 | $=\left(\left(i^{*} o\right)+\left(j^{*} \mathrm{p}\right)\right)^{*} \mathrm{q}$ |
| w | Estimated \# of HPV infections avoided | 21,428 |  |
| x | Estimated \# of HSV-2 infections avoided | 233 | $=\left((m * o)+\left(n^{*} \mathrm{p}\right)\right)^{*} \mathrm{q}$ |
| y | Reduction in QALYs per infection - Chlamydia | 0.049 | $\checkmark$ |
| z | Reduction in QALYs per infection - HIV | 7.48 | $\checkmark$ |
| aa | Reduction in QALYs per infection - Gonorrhea | 0.055 | $\checkmark$ |
| bb | Reduction in QALYs per infection - Hep B - Acute | 0.185 |  |
| cc | Reduction in QALYs per infection - Syphilis | 0.050 | Assumed |
| dd | Reduction in QALYs per infection - HPV | 0.146 | $\checkmark$ |
| ee | Reduction in QALYs per infection - HSV-2 | 0.0028 | $\checkmark$ |
| ff | Potential QALYs gained, Behavioural Counseling increasing from 0\% to 29\% | 3,285 | $\begin{aligned} = & r^{*} y+s^{*} z+t^{*} a a+u * b b \\ & +v^{*} c c+w^{*} d d^{*} x * e e \end{aligned}$ |

$V=$ Estimates from the literature
We also modified a number of major assumptions and recalculated the CPB as follows:

- Assume the effectiveness of high intensity behavioural counselling interventions in reducing the incidence of STIs is reduced from $62 \%$ to $40 \%$ in adolescents and from $30 \%$ to $13 \%$ in adults (Table 4, rows $o \& p$ ): CPB $=1,706$ QALYs.
- Assume the effectiveness of high intensity behavioural counselling interventions in reducing the incidence of STIs is increased from $62 \%$ to $74 \%$ in adolescents and from $30 \%$ to $44 \%$ in adults (Table 4, rows $o \& p$ ): CPB $=4,498$ QALYs.


## Modelling Cost-Effectiveness

In this section, we will calculate the CE associated with behavioural counselling interventions for the prevention of sexually transmitted diseases in a British Columbia birth cohort of 40,000.

In estimating CE, we made the following assumptions:

- We have assumed that all individuals between the ages of 15 and 59 who had sexual intercourse within the past 12 months would be eligible for this intervention. Rates of sexually transmitted diseases are relatively rare before age 15 and after age 60 (see Table 1 above). The rates by sex and age group for those who have 'ever had sexual intercourse' and 'had sexual intercourse in the past 12 months' are taken from the 2010 Canadian Community Health Survey Public Use Microdata File. ${ }^{654}$ Based on this data, approximately $81 \%$ of individuals between the ages of 15 and 59 have been sexually active within the past 12 months (see Table 5).

| By Age and Sex, 2010 |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Age | Ever had interco | sexual ourse |  | sual <br> se in past onths | BC Popul 201 | ation in 10 | BC Popul Ris | lation at sk |
| Group | Males | Females | Males | Females | Males | Females | Males | Females |
| 15-17 | 31.9\% | 19.3\% | 28.4\% | 17.7\% | 87,147 | 78,702 | 24,774 | 13,932 |
| 18-19 | 70.0\% | 63.3\% | 61.8\% | 59.9\% | 59,622 | 54,725 | 36,876 | 32,794 |
| 20-24 | 84.4\% | 87.5\% | 74.6\% | 77.7\% | 154,199 | 150,826 | 114,961 | 117,200 |
| 25-29 | 91.9\% | 91.2\% | 87.0\% | 84.1\% | 158,599 | 158,757 | 138,019 | 133,532 |
| 30-34 | 99.3\% | 96.6\% | 93.6\% | 93.2\% | 146,617 | 146,738 | 137,211 | 136,730 |
| 35-39 | 95.7\% | 96.7\% | 89.1\% | 91.1\% | 148,222 | 151,380 | 132,139 | 137,833 |
| 40-44 | 99.5\% | 97.9\% | 91.4\% | 85.6\% | 158,902 | 162,455 | 145,166 | 139,097 |
| 45-49 | 99.5\% | 95.9\% | 86.1\% | 82.7\% | 178,859 | 182,002 | 154,079 | 150,497 |
| 50-59 | 99.5\% | 95.9\% | 86.1\% | 82.7\% | 328,360 | 331,907 | 282,868 | 274,454 |
| Total |  |  | 82.1\% | 80.1\% | 1,420,527 | 1,417,492 | 1,166,093 | 1,136,069 |

- Frequency of screening - We assumed that a general practitioner would enquire about a patient's sexual behaviours once every four years (Table 7, row $c$ ).
- Patient time costs for behavioural counselling intervention - We assumed three hours of patient time would be required (including travel to and from the session) (Table 7, row o).
- Costs of a behavioural counselling intervention - We assumed that a clinical nurse specialist with a wage rate of $\$ 53.42$ per hour ( $\$ 100,000$ per year) would lead the session. ${ }^{655}$ Their direct time involvement would be 3.5 hours ( 2.5 for the session and 1 hour for preparation). To these costs we added $24 \%$ for benefits (e.g., dental, longterm disability, etc.), $40 \%$ for non-productive paid hours (e.g., statutory holidays, vacations, sick time, educational leave, etc.) and $50 \%$ for overhead costs (e.g., use of the facility and support staff). Based on these assumptions, the estimated costs per behavioural counselling intervention would be $\$ 487$ (Table 7, row $n$ ). We have

[^144]assumed that each session would be attended by an average of 5 individuals (Table 7, row $l$ ).

- Costs per infection avoided - The direct medical costs per infection avoided are taken from a US study (Table 7, rows $x-d d$ ). ${ }^{656}$ These costs, provided in 2010 US dollars, were adjusted to 2017 CAD. When costs were provided separately for males and females, we estimated the combined average costs based on the proportion of infections by sex expected in BC (Table 2 and 3 ) (see Table 6).

|  | 2010 US\$ |  |  | 2016 Can\$ |  |  | 2017 Can\$ |  |  | \% M/F | Est | Range |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| STI Sex | Est | Range |  | Est | Range |  | Est | Rang |  |  |  |  |
| Chlamydia |  |  |  |  |  |  |  |  |  |  |  |  |
| Male | \$30 | \$15 | \$45 | \$40 | \$20 | \$59 | \$29 | \$14 | \$43 |  |  | \$114 \$343 |
| Female | \$364 | \$182 | \$546 | \$481 | \$241 | \$722 | \$346 | \$173 | \$519 | 63\% |  |  |
| Gonorrhea |  |  |  |  |  |  |  |  |  |  |  |  |
| Male | \$79 | \$40 | \$119 | \$104 | \$53 | \$157 | \$75 | \$38 | \$113 | 64\% | \$169 | \$85 \$254 |
| Female | \$354 | \$177 | \$531 | \$468 | \$234 | \$702 | \$337 | \$168 | \$505 | 36\% |  |  |
| HBV | \$2,667 | \$2,172 | \$2,924 | \$3,525 | \$2,871 | \$3,865 | \$2,536 | \$2,065 | \$2,780 |  |  |  |
| HIV | \$304,500 | \$229,300 | \$379,700 | \$402,494 | \$303,093 | \$501,895 | \$289,543 | \$218,037 \$ | \$361,049 |  |  |  |
| HPV |  |  |  |  |  |  |  |  |  |  |  |  |
| Male | \$45 | \$23 | \$78 | \$59 | \$30 | \$103 | \$43 | \$22 | \$74 | 50\% |  | \$57 \$194 |
| Female | \$191 | \$96 | \$329 | \$252 | \$127 | \$435 | \$182 | \$91 | \$313 | 50\% |  | \$57 \$194 |
| HSV-2 |  |  |  |  |  |  |  |  |  |  |  |  |
| Male | \$761 | \$381 | \$1,142 | \$1,006 | \$504 | \$1,510 | \$724 | \$362 | \$1,086 | 31\% |  |  |
| Female | \$621 | \$311 | \$932 | \$821 | \$411 | \$1,232 |  |  |  | 69\% |  | \$16 \$948 |
| Syphilis | \$709 | \$355 | \$1,064 | \$937 | \$469 | \$1,406 | \$674 | \$338 | \$1,012 |  |  |  |

- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the CE associated with behavioural counselling interventions for the prevention of sexually transmitted diseases is $\$ 10,267$ per QALY (Table 7, row $k k$ ).

[^145]Table 7: CE of Behavioural Counselling Interventions for the Prevention of Sexually Transmitted Infections in a Birth Cohort of 40,000

| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Years of life between the ages of 15 and 59 in birth cohort | 1,758,398 | Tables 2 and 3 |
| b | Proportion of years sexually active | 81\% | Table 5 |
|  | Costs of intervention |  |  |
| c | Frequency of screening to determine sexual activity (every x years) | 4 | Assumed |
| d | Total number of screens | 439,600 | = a/c |
| e | Cost of 10-minute office visit | \$34.85 | Ref Doc |
| f | Value of patient time and travel for office visit | \$59.38 | Ref Doc |
| g | Portion of 10-minute office visit for screen | 50\% | Ref Doc |
| h | Cost of screening | \$20,711,730 | $=d^{*}(e+f) * g$ |
| i | Screen positive for sexual activity | 356,076 | $=d^{*} \mathrm{~b}$ |
| j | Adherence with behavioural counselling | 29\% | Table 4, row q |
| k | Attendance at a behavioural counselling intervention | 103,262 | = ${ }^{*}$ j |
| 1 | Individuals per behavioural counselling intervention | 5 | Assumed |
| m | Total number of behavioural counselling interventions | 20,652 | = $\mathrm{k} / \mathrm{m}$ |
| n | Cost per behavioural counselling intervention | \$487 | V |
| 0 | Value of patient time and travel for behavioural counselling intervention | \$89.07 | $\checkmark$ |
| p | Cost of behavioural counselling interventions | \$19,255,251 | $=\left(m^{*} \mathrm{n}\right)+(\mathrm{k} * \mathrm{o})$ |
|  | Cost avoided |  |  |
| q | Estimated \# of chlamydia infections avoided | 946 | Table 4, row r |
| r | Estimated \# of HIV infections avoided | 12 | Table 4, row s |
| s | Estimated \# of gonorrhea infections avoided | 183 | Table 4, row t |
| t | Estimated \# of Hep B-Acute infections avoided | 0.2 | Table 4, row u |
| $u$ | Estimated \# of syphilis infections avoided | 38 | Table 4, row v |
| v | Estimated \# of HPV infections avoided | 21,428 | Table 4, row w |
| w | Estimated \# of HSV-2 infections avoided | 233 | Table 4, row x |
| x | Cost of chlamydia infection avoided | \$229 | $\checkmark$ |
| y | Cost of HIV infection avoided | \$289,543 | $\checkmark$ |
| z | Cost of gonorrhea infection avoided | \$169 | $\checkmark$ |
| aa | Cost of Hep B-Acute infection avoided | \$2,536 | $\checkmark$ |
| bb | Cost of syphilis infection avoided | \$674 | $\checkmark$ |
| cc | Cost of HPV infection avoided | \$112 | $\checkmark$ |
| dd | Cost of HSV-2 infection avoided | \$632 | $\checkmark$ |
|  | CE calculation |  |  |
| ee | Cost of intervention over lifetime of birth cohort | \$39,966,981 | $=\mathrm{h}+\mathrm{p}$ |
| ff | Costs avoided | \$6,239,820 | $\begin{gathered} =q^{*} x+r^{*} y+s^{*} z+t^{*} a a \\ +u^{*} b b+v^{*} c c+w^{*} d d \end{gathered}$ |
| gg | QALYs saved | 3,285 | Table 4, row ff |
| hh | Cost of intervention over lifetime of birth cohort (1.5\% discount) | \$29,128,113 | Calculated |
| ii | Costs avoided (1.5\% discount) | \$4,547,608 | Calculated |
| jj | QALYs saved (1.5\% discount) | 2,394 | Calculated |
| kk | CE (\$/QALY saved) | \$10,267 | $=(\mathrm{hh}-\mathrm{ii}) / \mathrm{jj}$ |

$V=$ Estimates from the literature
We also modified several major assumptions and recalculated the cost per QALY as follows:

- Assume the effectiveness of high intensity behavioural counselling interventions in reducing the incidence of STIs is reduced from $62 \%$ to $40 \%$ in adolescents and from $30 \%$ to $13 \%$ in adults (Table 4, rows $o \& p$ ): CE = $221,687 /$ QALY.
- Assume the effectiveness of high intensity behavioural counselling interventions in reducing the incidence of STIs is increased from $62 \%$ to $74 \%$ in adolescents and from $30 \%$ to $44 \%$ in adults (Table 4, rows $o$ \& $p$ ): $\mathrm{CE}=\$ 6,921 / \mathrm{QALY}$.
- Assume screening to determine sexual activity is less frequent, carried out once every 5 years rather than once every 4 years (Table 7, rows $c$ ): $\mathrm{CE}=\$ 7,833 / \mathrm{QALY}$.
- Assume screening to determine sexual activity is more frequent, carried out once every 3 years rather than once every 4 years (Table 7, rows $c$ ): $C E=\$ 14,322 /$ QALY.
- Assume the average number of individuals attending each behavioural counselling intervention is increased from 5 to 10 (Table 7, rows $l$ ): $\mathrm{CE}=\$ 8,736 / \mathrm{QALY}$.
- Assume the average number of individuals attending each behavioural counselling intervention is reduced from 5 to 1 (Table 7, rows $l$ ): $\mathrm{CE}=\$ 22,513 / \mathrm{QALY}$.
- Assume the average direct cost per HIV infection is reduced from $\$ 289,543$ to $\$ 218,037$ (Table 7, rows $y$ ): $\mathrm{CE}=\$ 10,524 / \mathrm{QALY}$.
- Assume the average direct cost per HIV infection is increased from $\$ 289,543$ to $\$ 361,049$ (Table 7, rows y): $\mathrm{CE}=\$ 10,010 / \mathrm{QALY}$.
- Assume the average direct cost per HPV infection is reduced from $\$ 112$ to $\$ 57$ (Table 7, rows $c c$ ): $\mathrm{CE}=\$ 10,625 / \mathrm{QALY}$.
- Assume the average direct cost per HPV infection is increased from \$112 to \$194 (Table 7, rows $c c$ ): $\mathrm{CE}=\$ 9,732 / \mathrm{QALY}$.


## Summary

| Table 8: Behavioural Counselling Interventions for the Prevention of Sexually Transmitted Infections in a Birth Cohort of 40,000 Summary |  |  |  |
| :---: | :---: | :---: | :---: |
|  |  |  |  |
|  |  | Range |  |
| CPB (Potential QALYs Gained) |  |  |  |
| Gap between 0\% and Best in the World (29\%) |  |  |  |
| 1.5\% Discount Rate | 2,394 | 1,243 | 3,278 |
| 3\% Discount Rate | 1,790 | 929 | 2,451 |
| 0\% Discount Rate | 3,285 | 1,706 | 4,49 |
| CE (\$/QALY) including patient time costs |  |  |  |
| 1.5\% Discount Rate | \$10,267 | \$6,921 | \$22,513 |
| 3\% Discount Rate | \$10,267 | \$6,921 | \$22,513 |
| 0\% Discount Rate | \$10,267 | \$6,921 | \$22,513 |
| CE (\$/QALY) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | \$3,494 | \$1,974 | \$15,740 |
| 3\% Discount Rate | \$3,494 | \$1,974 | \$15,740 |
| 0\% Discount Rate | \$3,494 | \$1,974 | \$15,740 |

## Smoking Cessation Advice and Help to Quit

## United States Preventive Services Task Force Recommendations (2009)

Tobacco use, cigarette smoking in particular, is the leading preventable cause of death in the United States. Tobacco use results in more than 400000 deaths annually from cardiovascular disease, respiratory disease, and cancer. Smoking during pregnancy results in the deaths of about 1000 infants annually and is associated with an increased risk for premature birth and intrauterine growth retardation. Environmental tobacco smoke contributes to death in an estimated 38000 people annually.
The USPSTF strongly recommends that clinicians screen all adults for tobacco use and provide tobacco cessation interventions for those who use tobacco products. (A Recommendation).
The USPSTF strongly recommends that clinicians screen all pregnant women for tobacco use and provide augmented pregnancy-tailored counseling to those who smoke. (A Recommendation) ${ }^{657}$

## Canadian Task Force on Preventive Health Care Recommendations (1994)

A large body of evidence has accumulated regarding the health effects of smoking. Tobacco use has been consistently linked with a variety of serious pulmonary, cardiovascular and neoplastic diseases. Evaluation of this evidence is beyond the scope of this chapter but detailed reviews and estimates of relative risk for the many tobacco associated diseases have been published elsewhere. Likewise, reviews of the evidence regarding the health consequences of ETS are published elsewhere. In 1992 the U.S. Environmental Protection Agency (EPA) named ETS a Group A carcinogen (shown to cause cancer in humans) at typical environmental levels.
There is good evidence to support counselling for smoking cessation in the periodic health examination of individuals who smoke (A Recommendation). Nicotine replacement therapy can be effective as an adjunct (A Recommendation).
There is fair evidence to support physicians also referring patients to other programs after offering cessation advice (B Recommendation).
There is insufficient evidence to evaluate counselling to reduce ETS exposure ( $C$ Recommendation) but it may be useful to combine such counselling with cessation advice, again based on the burden of suffering, the potential benefits of the intervention and the effectiveness of cessation advice. ${ }^{658}$

## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with behavioural counselling and interventions for the prevention of tobacco use in a British Columbia birth cohort of 40,000.

In estimating CPB , we made the following assumptions:

[^146]- The proportion of the BC population that are light smokers (less than 10 cigarettes per day), moderate smokers (10-19 cigarettes per day) and heavy smokers ( 20 or more cigarettes per day) by age group is based on 2014 CCHS data. ${ }^{659}$ No data is available for ages $80+$ so we assumed a $50 \%$ decline in smoking rate between the ages of 79 and 84 and further $50 \%$ decline between the ages of 85 and 89 . Between the ages of 18 and 89 , the proportion of life years lived with light smoking is $8.0 \%$ ( 200,747 of $2,524,990$ life years), moderate smoking is $3.9 \%$ ( 98,886 of $2,524,990$ life years) and heavy smoking is $2.4 \%$ ( 59,461 of $2,524,990$ life years) (see Table 1).

| Table 1: Years of Life Lived and Current Smoking <br> Between the Ages of 18 and 89 <br> in a British Columbia Birth Cohort of 40,000 |  |  |  |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Age Group | Mean Survival Rate | Individuals in Birth Cohort | \% of BC Population Current Smokers |  |  | BC Population Current Smokers |  |  |  | Life Years Lived | Years Lived as Current Smokers |  |  |
|  |  |  | Light | Mod | Heavy | Light | Mod | Heavy | Total |  | Light | Mod | Heavy |
| 18-19 | 0.994 | 39,744 | 10.3\% | 0.4\% | 0.4\% | 4,092 | 143 | 143 | 4,378 | 79,488 | 8,183 | 286 | 287 |
| 20-24 | 0.992 | 39,682 | 20.5\% | 1.9\% | 0.4\% | 8,131 | 767 | 176 | 9,074 | 198,408 | 40,654 | 3,835 | 879 |
| 25-29 | 0.989 | 39,570 | 14.9\% | 5.2\% | 2.3\% | 5,905 | 2,074 | 907 | 8,885 | 197,850 | 29,523 | 10,368 | 4,533 |
| 30-34 | 0.986 | 39,458 | 16.6\% | 5.2\% | 1.3\% | 6,552 | 2,048 | 518 | 9,118 | 197,290 | 32,759 | 10,242 | 2,589 |
| 35-39 | 0.983 | 39,310 | 8.9\% | 6.7\% | 1.2\% | 3,513 | 2,645 | 489 | 6,647 | 196,550 | 17,566 | 13,224 | 2,444 |
| 40-44 | 0.978 | 39,105 | 6.8\% | 5.0\% | 3.5\% | 2,672 | 1,939 | 1,385 | 5,996 | 195,526 | 13,360 | 9,693 | 6,927 |
| 45-49 | 0.970 | 38,814 | 4.4\% | 2.9\% | 3.2\% | 1,726 | 1,119 | 1,247 | 4,092 | 194,070 | 8,632 | 5,593 | 6,235 |
| 50-54 | 0.960 | 38,390 | 7.6\% | 4.1\% | 4.6\% | 2,918 | 1,560 | 1,766 | 6,244 | 191,948 | 14,590 | 7,799 | 8,832 |
| 55-59 | 0.944 | 37,757 | 3.9\% | 7.9\% | 4.3\% | 1,468 | 2,987 | 1,635 | 6,089 | 188,786 | 7,341 | 14,933 | 8,173 |
| 60-64 | 0.920 | 36,800 | 3.9\% | 4.7\% | 3.5\% | 1,427 | 1,746 | 1,289 | 4,462 | 183,998 | 7,137 | 8,728 | 6,446 |
| 65-69 | 0.883 | 35,332 | 4.7\% | 3.5\% | 3.0\% | 1,654 | 1,235 | 1,061 | 3,950 | 176,658 | 8,269 | 6,176 | 5,304 |
| 70-74 | 0.827 | 33,072 | 3.7\% | 3.6\% | 2.1\% | 1,208 | 1,207 | 701 | 3,116 | 165,362 | 6,038 | 6,033 | 3,507 |
| 75-79 | 0.741 | 29,628 | 2.9\% | 0.9\% | 1.4\% | 857 | 253 | 423 | 1,532 | 148,142 | 4,283 | 1,264 | 2,115 |
| 80-84 | 0.614 | 24,551 | 1.4\% | 0.4\% | 0.7\% | 355 | 105 | 175 | 635 | 122,756 | 1,775 | 524 | 876 |
| 85-89 | 0.441 | 17,632 | 0.7\% | 0.2\% | 0.4\% | 127 | 38 | 63 | 228 | 88,158 | 637 | 188 | 315 |
| Total |  |  | 8.0\% | 3.9\% | 2.4\% |  |  |  |  | 2,524,990 | 200,747 | 98,886 | 59,461 |

- A significant proportion of smokers quit on their own. ${ }^{600}$ According to the Treating Tobacco Use and Dependence: 2008 Update document, individuals who quit on their own have a success (abstinence rate) of $10.9 \%$. This increases to $28.0 \%$ ( $95 \% \mathrm{CI}$ of $23.0 \%-33.6 \%$ ) with 2-3 brief counselling interventions with a primary care provider and the use of medications. ${ }^{661}$ We used the rate of $10.9 \%$ to populate row $w$ in Table 2 and the $28.0 \%$ to populate row $x$.
- Other assumptions used in assessing the clinically preventable burden are detailed in the Reference Document.

Based on these assumptions, the CPB associated with behavioural counselling and interventions for the prevention of tobacco use is 5,944 QALYs (Table 2, row ac). The CPB of 5,944 represents the gap between no coverage and the 'best in the world' coverage estimated at $51 \%$.

[^147]Table 2: CPB of Behavioural Counselling and Interventions to Prevent Tobacco Use in a BC Birth Cohort of 40,000

| Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
|  | Estimated current status |  |  |
| a | \# of life years lived between the ages of 18-89 in birth cohort | 2,524,990 | Table 1 |
| b | \% of life years at light smoking (<10 cigarettes / day) | 8.0\% | Table 1 |
| c | \# of life years at light smoking | 200,747 | = (a*b) |
| d | \% of life years at moderate smoking (10-19 cigarettes / day) | 3.9\% | Table 1 |
| e | \# of life years at moderate smoking | 98,886 | = (a*d) |
| f | \% of life years at heavy smoking ( $\geq 20$ cigarettes / day) | 2.4\% | Table 1 |
| g | \# of life years at heavy smoking | 59,461 | $=(\mathrm{a} * \mathrm{f})$ |
|  | Life years lost due to Smoking |  |  |
| h | \% of life years lost due to light smoking | 10.2\% | Ref Doc |
| i | \# of life years lost due to light smoking | 20,478 | = (c* h ) |
| j | \% of life years lost due to moderate smoking | 18.4\% | Ref Doc |
| k | \# of life years lost due to moderate smoking | 18,188 | $=(\mathrm{e} * \mathrm{j})$ |
| 1 | \% of life years lost due to heavy smoking | 28.0\% | Ref Doc |
| m | \# of life years lost due to heavy smoking | 16,634 | $=(\mathrm{g} *$ I) |
| n | Life years lost due to smoking | 55,300 | $=\mathrm{i}+\mathrm{k}+\mathrm{m}$ |
|  | QALYs lost due to Smoking |  |  |
| o | \% of QoL lost due to light smoking | 3.7\% | Ref Doc |
| p | \# of QALYs lost due to light smoking | 6,590 | $=(\mathrm{c}-\mathrm{i}) * \mathrm{o}$ |
| q | \% of QoL lost due to moderate smoking | 3.9\% | Ref Doc |
| r | \# of QALYs lost due to moderate smoking | 3,140 | $=(e-k) * q$ |
| s | \% of QoL lost due to heavy smoking | 7.3\% | Ref Doc |
| t | \# of QALYs lost due to heavy smoking | 3,131 | $=(\mathrm{g}-\mathrm{m}) * \mathrm{~s}$ |
| u | QALYs lost due to smoking | 12,862 | $=p+r+t$ |
| v | Total QALYs lost due to smoking | 68,162 | $=\mathrm{n}+\mathrm{u}$ |
|  | Benefits if 51\% of smokers received counselling and an intervention |  |  |
| w | Quit rate without intervention | 10.9\% | $\checkmark$ |
| x | Quit rate with intervention | 28.0\% | $\checkmark$ |
| y | QALYs gained without intervention | 7,430 | $=v^{*} \mathrm{w}$ |
| z | QALYs gained with intervention with $100 \%$ adherence | 19,085 | = ${ }^{*}$ * |
| aa | Net QALYs gained with 100\% adherence | 11,656 | = $\mathrm{z}-\mathrm{y}$ |
| ab | Estimated adherence with screening and intervention | 51\% | Ref Doc |
| ac | Potential QALYs gained, Screening \& Intervention from 0\% to 51\% | 5,944 | = aa * ab |

$v=$ Estimates from the literature
We also modified a number of major assumptions and recalculated the CPB as follows:

- Assume the disutility of light smoking is reduced from $3.7 \%$ to $2.1 \%$ (Table 2, row $o$ ), the disutility of moderate smoking is reduced from $3.9 \%$ to $2.2 \%$ (Table 2 , row $q$ ) and the disutility of heavy smoking is reduced from $7.3 \%$ to $5.0 \%$ (Table 2, row $s$ ): CPB $=5,499$ QALYs.
- Assume the disutility of light smoking is increased from $3.7 \%$ to $5.3 \%$ (Table 2, row $o$ ), the disutility of moderate smoking is increased from $3.9 \%$ to $5.5 \%$ (Table 2, row $q$ ) and the disutility of heavy smoking is increased from $7.3 \%$ to $9.7 \%$ (Table 2, row $s): \mathrm{CPB}=6,408$ QALYs.
- Assume that the quit rate with intervention (2-3 sessions + medication) is reduced from $28.0 \%$ to $23.0 \%$ (Table 2, row $x$ ): $\mathrm{CPB}=4,206$ QALYs.
- Assume that the quit rate with intervention (2-3 sessions + medication) is increased from $28.0 \%$ to $33.6 \%$ (Table 2, row $x$ ): $\mathrm{CPB}=7,891$ QALYs.


## Modelling Cost-Effectiveness

In this section, we will calculate the CE associated with behavioural counselling and interventions for the prevention of tobacco use in a British Columbia birth cohort of 40,000.

In estimating CE, we made the following assumptions:

- For modelling purposes, we assumed that of the smokers who would successfully quit as a result of the intervention, $50 \%$ would quit at age $30,25 \%$ at age 40 and $25 \%$ at age 50 .
- Average cost of smoking cessation aids per quit attempt - in 2011, BC PharmaCare estimated the costs for pharmacological aids to smoking cessation based on a 12 week supply including mark-up and dispensing fees. ${ }^{662}$ Varenicline (Champix®) was estimated to cost $\$ 336$, buproprion (Zyban®) $\$ 209$, nicotine patch $\$ 273$ and nicotine gum \$122-\$289. In deriving the average cost we assumed that 57\% of all smokers would use either varenicline or buproprion and $43 \%$ of all smokers would use either the nicotine patch or nicotine gum. The mid-point for the cost estimate of nicotine gum was used. Based on these assumptions, the average cost of smoking cessation aids per quit attempt in BC was $\$ 257.87$ (in 2011 CAD ) or \$272.41 (in 2017 CAD).
- Portion of counseled who use a smoking cessation aid - Because the effectiveness of the intervention is based on 2-3 brief counselling sessions and the use of medication, we have assumed the $100 \%$ of those counselled would use a smoking cessation aid.
- In estimating the costs avoided due to the intervention, we assumed annual costs avoided of $\$ 785$ per light smoker, $\$ 1,386$ per moderate smoker and $\$ 2,050$ per heavy smoker (see Reference Document). These costs avoided, however, are not fully realized until 20 years following smoking cessation. ${ }^{63,664}$ This gradual increase in costs avoided was incorporated into the model.
- The later in life smoking cessation occurs, the fewer the benefits. Based on data provided by Jha and colleagues, ${ }^{665}$ we have assumed that $91.3 \%$ of potential benefits would occur if smoking cessation occurred at age $30,82.6 \%$ at age 40 and $56.5 \%$ at age 50 .
- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the CE associated with behavioural counselling and interventions for the prevention of tobacco use is $-\$ 1,863$ / QALY (Table 3, row $y$ ).

[^148]Table 3: CE of Behavioural Counselling and Interventions to Prevent Tobacco Use in a BC Birth
Cohort of 40,000

| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | \# of life years lived between the ages of 18-89 in birth cohort | 2,524,990 | Table 1 |
| b | \# of life years lived as smokers between the ages of 18-89 in birth cohort | 359,095 | Table 2, row c + Table 2, row e + Table 2, row g |
|  | Estimated cost of screening |  |  |
| c | Number of annual screens to assess willingness to quit | 359,095 | = b |
| d | Proportion of office visit required | 50\% | See Ref Doc |
| e | Cost of 10-minute office visit | \$34.85 | See Ref Doc |
| f | Patient time costs / office visit | \$59.38 | See Ref Doc |
| g | Estimated cost of screening | \$16,918,757 | $=(e+f) * d * c$ |
|  | Estimated cost of intervention |  |  |
| h | Average \# of smokers in birth cohort ages 20-29 | 8,979 | Table 1 |
| i | Estimated adherence with screening and intervention | 51\% | Table 2, row ab |
| j | \# of brief counselling interventions | 3 | $\checkmark$ |
| k | Cost of smoking cessation aids | \$272.41 | $\checkmark$ |
| I | Estimated cost of intervention | \$5,037,004 | $=\left((\mathrm{h} * \mathrm{i}) * \mathrm{j}^{\text {j }}\right.$ ) $(\mathrm{e}+\mathrm{f}+\mathrm{k})$ |
| m | Average \# of smokers in birth cohort ages 30-39 | 7,882 | Table 1 |
| n | Estimated cost of intervention | \$4,421,696 | $=\left(\left(m^{*}\right)^{*}\right)^{\prime}{ }^{*}(\mathrm{e}+\mathrm{f}+\mathrm{k})$ |
| 0 | Average \# of smokers in birth cohort ages 40-49 | 5,044 | Table 1 |
| p | Estimated cost of intervention | \$2,829,413 | $=\left(\left(0{ }^{*}\right)^{*}{ }^{\text {j }}\right.$ ) ${ }^{*}(\mathrm{e}+\mathrm{f}+\mathrm{k})$ |
| q | Total cost of interventions | \$12,288,114 | = $1+n+p$ |
| r | Estimated costs avoided due to intervention | \$49,085,691 | Calculated |
|  | CE Calculation |  |  |
| s | Cost of intervention over lifetime of birth cohort | \$29,206,871 | $=\mathrm{g}+\mathrm{q}$ |
| t | Costs avoided due to intervention over lifetime of birth cohort | \$49,085,691 | = r |
| $u$ | QALYs saved | 5,944 | Table 2, row ac |
| v | Cost of intervention over lifetime of birth cohort (1.5\% discount) | \$21,019,352 | Calculated |
| w | Costs avoided due to intervention over lifetime of birth cohort (1.5\% discount) | \$27,143,609 | Calculated |
| x | QALYs saved (1.5\% discount) | 3,287 | Calculated |
| y | CE (\$/QALY saved) | -\$1,863 | = (v-w) / x |

$\checkmark=$ Estimates from the literature

We also modified a number of major assumptions and recalculated the cost per QALY as follows:

- Assume the disutility of light smoking is reduced from $3.7 \%$ to $2.1 \%$ (Table 2, row $o$ ), the disutility of moderate smoking is reduced from $3.9 \%$ to $2.2 \%$ (Table 2 , row $q$ ) and the disutility of heavy smoking is reduced from $7.3 \%$ to $5.0 \%$ (Table 2, row $s$ ): $C E=-\$ 2,014$.
- Assume the disutility of light smoking is increased from $3.7 \%$ to $5.3 \%$ (Table 2, row $o$ ), the disutility of moderate smoking is increased from $3.9 \%$ to $5.5 \%$ (Table 2, row $q$ ) and the disutility of heavy smoking is increased from $7.3 \%$ to $9.7 \%$ (Table 2, row $s): \mathrm{CE}=-\$ 1,728$.
- Assume that the quit rate with intervention (2-3 sessions + medication) is reduced from $28.0 \%$ to $23.0 \%$ (Table 2, row $x$ ): $\mathrm{CE}=\$ 779$
- Assume that the quit rate with intervention (2-3 sessions + medication) is increase from $28.0 \%$ to $33.6 \%$ (Table 2, row $x$ ): $\mathrm{CE}=-\$ 3,441$.
- Assume the proportion of an office visit required for screening is reduced from $50 \%$ to $33 \%$ (Table 3, row $d$ ): $\mathrm{CE}=-\$ 3,122$.
- Assume the proportion of an office visit required for screening is increased from $50 \%$ to $67 \%$ (Table 3, row $d$ ): $\mathrm{CE}=-\$ 604$.


## Summary

| Summary |  |  |  |
| :---: | :---: | :---: | :---: |
|  | $\begin{aligned} & \hline \text { Base } \\ & \text { Case } \\ & \hline \end{aligned}$ | Range |  |
| CPB (Potential QALYs Gained) |  |  |  |
| Gap between No Service and 'Best in the World' (51\%) |  |  |  |
| 1.5\% Discount Rate | 3,287 | 2,326 | 4,364 |
| 3\% Discount Rate | 1,833 | 1,297 | 2,433 |
| 0\% Discount Rate | 5,944 | 4,206 | 7,891 |
| Gap between BC Current (19\%) and 'Best in the World' (51\%) |  |  |  |
| 1.5\% Discount Rate | 1,225 | 867 | 1,626 |
| 3\% Discount Rate | 683 | 483 | 906 |
| 0\% Discount Rate | 2,214 | 1,567 | 2,940 |
| CE (\$/QALY) including patient time costs |  |  |  |
| 1.5\% Discount Rate | -\$1,863 | -\$3,441 | \$779 |
| 3\% Discount Rate | -\$226 | -\$1,867 | \$3,731 |
| 0\% Discount Rate | -\$3,344 | -\$4,556 | -\$1,314 |
| CE (\$/QALY) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | -\$4,633 | -\$5,527 | -\$3,135 |
| 3\% Discount Rate | -\$3,449 | -\$4,635 | -\$1,462 |
| 0\% Discount Rate | -\$5,472 | -\$6,160 | -\$4,322 |

## Alcohol Misuse Screening and Brief Intervention

## United States Preventive Services Task Force Recommendations (2013)

The USPSTF uses the term "alcohol misuse" to define a spectrum of behaviors, including risky or hazardous alcohol use (for example, harmful alcohol use and alcohol abuse or dependence). Risky or hazardous alcohol use means drinking more than the recommended daily, weekly, or per-occasion amounts resulting in increased risk for health consequences. For example, the National Institute on Alcohol Abuse and Alcoholism (NIAAA) and the U.S. Department of Agriculture define "risky use" as consuming more than 4 drinks on any day or 14 drinks per week for men, or more than 3 drinks on any day or 7 drinks per week for women (as well as any level of consumption under certain circumstances). "Harmful alcohol use" (defined by the International Statistical Classification of Diseases and Related Health Problems, Tenth Revision) is a pattern of drinking that causes damage to physical or mental health.
"Alcohol abuse" (defined by the Diagnostic and Statistical Manual of Mental Disorders, Fourth Edition) is drinking that leads an individual to recurrently fail in major home, work, or school responsibilities; use alcohol in physically hazardous situations (such as while operating heavy machinery); or have alcohol-related legal or social problems. "Alcohol dependence" (or alcoholism) (defined by the Diagnostic and Statistical Manual of Mental Disorders, Fourth Edition) includes physical cravings and withdrawal symptoms, frequent consumption of alcohol in larger amounts than intended over longer periods, and a need for markedly increased amounts of alcohol to achieve intoxication.

An estimated $30 \%$ of the U.S. population is affected by alcohol misuse, and most of these persons engage in risky use. More than 85000 deaths per year are attributable to alcohol misuse; it is the estimated third leading cause of preventable deaths in the United States.

The U.S. Preventive Services Task Force recommends screening and behavioral counseling interventions to reduce alcohol misuse by adults, including pregnant women, in primary care settings ( $B$ Recommendation).

The USPSTF concludes that the evidence is insufficient to recommend for or against screening and behavioral counseling interventions to prevent or reduce alcohol misuse by adolescents in primary care settings (I Statement). ${ }^{666}$

## Canadian Task Force on Preventive Health Care Recommendations (1994)

In 1989 the Canadian Task Force on the Periodic Health Examination concluded that there was fair evidence that routine case-finding for problem drinking, and that brief counselling intervention in patients identified thereby was effective in reducing alcohol consumption and related consequences. The studies which yielded this evidence have since been confirmed by seven new randomized controlled trials in study populations that included both men and women aged 18-60 years. Standardized interviewing strategies and questionnaires are more sensitive than clinical judgement and can be used routinely with all adults to raise the index of clinical suspicion of problem drinking. When problem drinkers are identified, either simple advice or brief counselling is effective in reducing alcohol consumption and diminishing the negative consequences of drinking. The intervention of simple advice or brief counselling is appropriate for the patient with mild to moderate as opposed to severe alcohol
${ }^{666}$ Moyer VA. Screening and behavioral counseling interventions in primary care to reduce alcohol misuse: U.S. preventive services task force recommendation statement. Annals of Internal Medicine. 2013; 159(3): 210-8.
dependency. Problem drinking or mild to moderate, rather than severe dependency is the focus of this report.
Routine active case-finding of problem drinking by physicians is highly recommended on the basis of the high prevalence of this problem in medical practices, its association with adverse consequences before the stage of dependency is reached, and its amenability to a counselling intervention by physicians. Detection by biomarkers is not recommended, although these may be used to confirm clinical suspicions raised by use of the CAGE query, MAST or AUDIT questionnaires, and may be useful for monitoring the patient's progress. Either simple advice or the brief counselling intervention may be used with equal effectiveness in reducing alcohol consumption in problem drinkers. The counselling intervention is probably most effective in the context of an established and effective doctor-patient relationship. ${ }^{667}$

## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with behavioural counselling and interventions for the prevention of alcohol misuse in a British Columbia birth cohort of 40,000.

In estimating CPB, we made the following assumptions:

- The proportion of the BC population with low alcohol use (less than 1.5 drinks a day for females and 3 drinks a day for males), hazardous alcohol use ( 1.5 to 3 drinks a day for females and 3 to 4.5 drinks per day for males) and harmful alcohol use (more than 3 drinks a day for females and 4.5 drinks a day for males) by age group is based on 2014 CCHS data. ${ }^{668}$ Alcohol consumption rates are adjusted for underreporting. ${ }^{669,670,671}$ Individuals who consume alcohol are grouped into these three categories based on their weekly consumption patterns.
- A significant proportion of individuals with low alcohol consumption levels consume their alcohol via binge drinking. A female binge drinker is defined as a female who consumes at least four drinks (containing 13.6 g of ethanol) on one occasion at least once per month during the past 12 months. A male binge drinker is defined as a male who consumes at least five drinks on one occasion at least once per month during the past 12 months.
- For modelling purposes, alcohol misuse is defined as any individuals with hazardous or harmful alcohol consumption levels and binge drinkers within the low consumption category.
- In a BC birth cohort of 40,000 , an estimated $39.1 \%$ of life years lived (between the ages of 18 and 79 ( 905,864 of $2,314,076$ ) are lived with alcohol misuse (see Table 1).

[^149]| Table 1: Years of Life Lived and Current Alcohol Use Between the Ages of 18 and 79 in a British Columbia Birth Cohort of 40,000 |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Age Group | Mean Survival Rate | Individuals in Birth Cohort | $\begin{gathered} \text { \% of BC Population Current } \\ \text { Drinkers } \\ \text { Low Hazardous Harmful } \end{gathered}$ |  |  | BC Population Current DrinkersLow-Non- Low-LowBinge Binge Hazardous Harmful |  |  |  |  | Life Years Lived | Years Low-NonBinge | Lived as <br> Low- <br> Binge | urrent Drink <br> Hazardous | ers <br> Harmful |
| 18-19 | 0.994 | 39,744 | 49.2\% | 5.5\% | 5.4\% | 19,555 | 9,247 | 10,308 | 2,192 | 2,127 | 79,488 | 18,494 | 20,615 | 4,385 | 4,254 |
| 20-24 | 0.992 | 39,682 | 49.2\% | 5.5\% | 5.3\% | 19,523 | 9,232 | 10,291 | 2,188 | 2,123 | 198,408 | 46,160 | 51,455 | 10,940 | 10,613 |
| 25-29 | 0.989 | 39,570 | 49.1\% | 5.4\% | 5.2\% | 19,442 | 9,194 | 10,248 | 2,153 | 2,069 | 197,850 | 45,968 | 51,240 | 10,765 | 10,347 |
| 30-34 | 0.986 | 39,458 | 57.5\% | 6.0\% | 5.0\% | 22,693 | 10,731 | 11,962 | 2,383 | 1,966 | 197,290 | 53,655 | 59,809 | 11,916 | 9,831 |
| 35-39 | 0.983 | 39,310 | 57.5\% | 6.0\% | 5.0\% | 22,616 | 10,695 | 11,921 | 2,377 | 1,964 | 196,550 | 53,473 | 59,607 | 11,886 | 9,820 |
| 40-44 | 0.978 | 39,105 | 57.5\% | 6.0\% | 5.0\% | 22,491 | 10,635 | 11,855 | 2,362 | 1,949 | 195,526 | 53,177 | 59,276 | 11,810 | 9,745 |
| 45-49 | 0.970 | 38,814 | 57.1\% | 6.8\% | 4.6\% | 22,147 | 10,473 | 11,674 | 2,652 | 1,777 | 194,070 | 52,365 | 58,372 | 13,262 | 8,885 |
| 50-54 | 0.960 | 38,390 | 57.1\% | 6.8\% | 4.6\% | 21,904 | 10,358 | 11,546 | 2,623 | 1,757 | 191,948 | 51,791 | 57,731 | 13,116 | 8,785 |
| 55-59 | 0.944 | 37,757 | 57.1\% | 6.8\% | 4.6\% | 21,545 | 10,188 | 11,357 | 2,580 | 1,729 | 188,786 | 50,941 | 56,784 | 12,901 | 8,644 |
| 60-64 | 0.920 | 36,800 | 54.0\% | 7.4\% | 3.5\% | 19,886 | 9,404 | 10,483 | 2,706 | 1,293 | 183,998 | 47,019 | 52,413 | 13,529 | 6,465 |
| 65-69 | 0.883 | 35,332 | 54.0\% | 7.4\% | 3.5\% | 19,092 | 9,028 | 10,064 | 2,598 | 1,239 | 176,658 | 45,142 | 50,320 | 12,992 | 6,197 |
| 70-74 | 0.827 | 33,072 | 43.1\% | 8.3\% | 3.1\% | 14,262 | 6,744 | 7,518 | 2,751 | 1,040 | 165,362 | 33,722 | 37,590 | 13,757 | 5,199 |
| 75-79 | 0.741 | 29,628 | 43.0\% | 8.4\% | 3.1\% | 12,742 | 6,025 | 6,717 | 2,481 | 924 | 148,142 | 30,127 | 33,583 | 12,403 | 4,622 |
| Total |  |  | 53.2\% | 6.6\% | 4.5\% |  |  |  |  |  | 2,314,076 | 582,035 | 648,794 | 153,663 | 103,407 |
|  |  |  |  |  |  |  |  |  |  |  |  |  | 28.0\% | 6.6\% | 4.5\% |

- Alcohol misuse results in life years lost due to both chronic and acute (binge drinking) conditions. Solberg and colleagues estimated that life years lost due to acute conditions are 2.14 times that of chronic conditions. ${ }^{672}$ That is, for every death due to chronic alcohol conditions, there would be 2.14 deaths due to acute alcohol conditions (Table 2, row $j$ ).
- The meta-analysis for the USPSTF found an improvement of $10.9 \%$ ( $95 \%$ CI of $8.3 \%$ to $13.4 \%$ ) in the proportion of adults achieving recommended drinking limits associated with brief counselling interventions (Table 2, row $s$ ). ${ }^{673}$
- Other costs and assumptions used in assessing CPB are detailed in the Reference Document.

Based on these assumptions, the CPB associated with behavioural counselling for the prevention of alcohol misuse is 2,175 QALYs (Table 2, row $v$ ). The CPB of 2,175 represents the gap between no coverage and the 'best in the world' coverage estimated at $30 \%$.

[^150]Table 2: CPB of Behavioural Counselling to Prevent Alcohol Misuse in a BC Birth Cohort of 40,000

| Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
|  | Estimated current status |  |  |
| a | \# of life years lived between the ages of 18-79 in birth cohort | 2,314,076 | Table 1 |
| b | \% of life years at low alcohol use - binge | 28.0\% | Table 1 |
| c | \# of life years at low alcohol use - binge | 648,794 | = (a*b) |
| d | \% of life years at hazardous alcohol use | 6.6\% | Table 1 |
| e | \# of life years at hazardous alcohol use | 153,663 | = (a*d) |
| f | \% of life years at harmful alcohol use | 4.5\% | Table 1 |
| g | \# of life years at harmful alcohol use | 103,407 | $=(\mathrm{a} * \mathrm{f}$ ) |
|  | Life years lost due to Alcohol Misuse |  |  |
| h | \% of life years lost due to harmful alcohol use | 4.8\% | Ref Doc |
| i | \# of life years lost due to chronic harmful alcohol use | 4,955 | = ( g * h ) |
| j | Ratio of life years lost to acute vs. chronic alcohol misuse | 2.14 | $\checkmark$ |
| k | \# of life years lost due to acute alcohol misuse | 10,605 | $=\mathrm{i}^{*} \mathrm{j}$ |
| I | Life years lost due to alcohol misuse | 15,559 | $=\mathrm{i}+\mathrm{k}$ |
|  | QALYs lost due to Alcohol Misuse |  |  |
| m | \% of QoL lost due to hazardous alcohol use | 14.5\% | Ref Doc |
| n | \# of QALYs lost due to hazardous alcohol use | 22,288 | = ${ }^{*} \mathrm{~m}$ |
| 0 | \% of QoL lost due to harmful alcohol use | 27.7\% | Ref Doc |
| p | \# of QALYs lost due to harmful alcohol use | 28,656 | = $\mathrm{F}^{*}$ o |
| q | QALYs lost due to alcohol misuse | 50,945 | $=\mathrm{n}+\mathrm{p}$ |
| $r$ | Total QALYs lost due to alcohol misuse | 66,504 | $=1+q$ |
|  | Benefits if $\mathbf{3 0 \%}$ of individuals who misuse alcohol received counselling |  |  |
| s | \% of adults achieving recommended drinking levels with intervention | 10.9\% | $\checkmark$ |
| t | QALYs gained with intervention with $100 \%$ adherence | 7,249 | = ${ }^{*}$ s |
| u | Estimated adherence with screening and intervention | 30\% | Ref Doc |
| v | Potential QALYs gained, Screening \& Intervention from 0\% to 51\% | 2,175 | = ${ }^{*}$ u |

V = Estimates from the literature

We also modified several major assumptions and recalculated the CPB as follows:

- Assume that the QoL reduction associated with hazardous alcohol consumption is reduced from $14.5 \%$ to $9.7 \%$ and the QoL reduction associated with harmful alcohol consumption is reduced from $27.7 \%$ to $18.9 \%$ (Table 2, rows $m \& o$ ): $\mathrm{CPB}=1,633$.
- Assume that the QoL reduction associated with hazardous alcohol consumption is increased from $14.5 \%$ to $20.9 \%$ and the QoL reduction associated with harmful alcohol consumption is reduced from $27.7 \%$ to $38.6 \%$ (Table 2, rows $m \& o$ ): $\mathrm{CPB}=$ 2,861.
- Assume that the effectiveness of counselling at changing behaviour is reduced from $10.9 \%$ to $8.3 \%$ (Table 2, row $s$ ): $\mathrm{CPB}=1,656$.
- Assume that the effectiveness of counselling at changing behaviour is increased from $10.9 \%$ to $13.4 \%$ (Table 2, row $s$ ): $\mathrm{CPB}=2,673$.


## Modelling Cost-Effectiveness

In this section, we will calculate the CE associated with behavioural counselling for the prevention of alcohol misuse in a British Columbia birth cohort of 40,000.

In estimating CE, we made the following assumptions:

- For modelling purposes, we assumed that $50 \%$ of the prevention of alcohol misuse due to the intervention would occur at age $30,25 \%$ at age 40 and $25 \%$ at age 50 .
- BC guidelines for alcohol screening and brief interventions recommend screening annually ${ }^{674}$ while economic evaluations have assumed that screening would occur at least once a year to at least once every 10 years. ${ }^{675,677,677}$ For modelling purposes we assumed screening would occur annually in the base case and modified this to once every 5 years in the sensitivity analysis.
- The 2013 USPSTF review found no evidence to determine the optimal interval for screening but did note that brief multi-contact (each contact is 6 to 15 minutes) interventions are most effective, requiring up to 120 minutes of total counseling contact. ${ }^{678}$ For modelling purposes we assumed 9 contacts of 10 -minutes in the base case analysis (Table 3, row $j$ ) and modified this from 6 to 12 contacts of 10 -minutes in the sensitivity analysis.
- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the CE associated with behavioural counselling for the prevention of alcohol misuse is $\$ 23,607$ / QALY (Table 3, row $x$ ).

[^151]Table 3: CE of Behavioural Counselling to Prevent Alcohol Misuse in a BC Birth Cohort of $\mathbf{4 0 , 0 0 0}$

| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | \# of life years lived between the ages of 18-79 in birth cohort | 2,314,076 | Table 1 |
| b | Screening rate | 35\% | Ref Doc |
|  | Estimated cost of screening |  |  |
| c | Number of annual screens to assess alcohol consumption habits | 809,927 | = ${ }^{*} \mathrm{~b}$ |
| d | Proportion of office visit required | 50\% | See Ref Doc |
| e | Cost of 10-minute office visit | \$34.85 | See Ref Doc |
| f | Patient time costs / office visit | \$59.38 | See Ref Doc |
| g | Estimated cost of screening | \$38,159,692 | $=(e+f) *{ }^{*}{ }^{*}$ |
|  | Estimated cost of intervention |  |  |
| h | \# of drinkers who misuse alcohol at age 30 | 16,311 | Table 1 |
| i | Estimated adherence with intervention | 30\% | Table 2, row u |
| j | \# of brief counselling interventions | 9 | $\checkmark$ |
| k | Estimated cost of intervention | \$4,149,934 | $=\left(\mathrm{h}^{*}{ }^{*} \mathrm{j}\right) *(\mathrm{e}+\mathrm{f})$ |
| I | \# of drinkers who misuse alcohol at age 40 | 16,166 | Table 1 |
| m | Estimated cost of intervention | \$4,113,041 | $=\left(1{ }^{*}{ }^{*}{ }^{\text {j }}\right)^{*}(\mathrm{e}+\mathrm{f})$ |
| n | \# of drinkers who misuse alcohol at age 50 | 15,926 | Table 1 |
| o | Estimated cost of intervention | \$4,052,034 | $=\left(0^{*}{ }^{*}{ }^{\mathrm{j}}\right.$ ) ${ }^{*}(\mathrm{e}+\mathrm{f})$ |
| p | Total cost of interventions | \$12,315,009 | = $1+\mathrm{n}+\mathrm{p}$ |
| q | Estimated costs avoided due to intervention | \$14,351,678 | Calculated |
|  | CE Calculation |  |  |
| $r$ | Cost of intervention over lifetime of birth cohort | \$50,474,700 | $=\mathrm{g}+\mathrm{p}$ |
| s | Costs avoided due to intervention over lifetime of birth cohort | \$14,351,678 | = q |
| t | QALYs saved | 2,175 | Table 2, row v |
| $u$ | Cost of intervention over lifetime of birth cohort (1.5\% discount) | \$36,325,203 | Calculated |
| v | Costs avoided due to intervention over lifetime of birth cohort (1.5\% discount) | \$7,936,250 | Calculated |
| w | QALYs saved (1.5\% discount) | 1,203 | Calculated |
| x | CE (\$/QALY saved) | \$23,607 | = (u-v) / w |

V = Estimates from the literature
We also modified several major assumptions and recalculated the CE as follows:

- Assume that the QoL reduction associated with hazardous alcohol consumption is decreased from $14.5 \%$ to $9.7 \%$ and the QoL reduction associated with harmful alcohol consumption is decreased from $27.7 \%$ to $18.9 \%$ (Table 2, rows $m \& o$ ): $\mathrm{CE}=$ \$31,444.
- Assume that the QoL reduction associated with hazardous alcohol consumption is increased from $14.5 \%$ to $20.9 \%$ and the QoL reduction associated with harmful alcohol consumption is increased from $27.7 \%$ to $38.6 \%$ (Table 2, rows $m \& o$ ): $\mathrm{CE}=$ \$17,941.
- Assume that the effectiveness of counselling at changing behaviour is reduced from $10.9 \%$ to $8.3 \%$ (Table 2, row $s$ ): $\mathrm{CE}=\$ 33,069$.
- Assume that the effectiveness of counselling at changing behaviour is increased from $10.9 \%$ to $13.4 \%$ (Table 2, row $s$ ): $\mathrm{CE}=\$ 19,972$.
- Assume that screening is carried out less frequently, once every five years rather than annually (Table 3, row $c$ ): $\mathrm{CE}=\$ 5,338$.
- Assume that the portion of an office visit used for screening is reduced from $50 \%$ to $33 \%$ (Table 3, row $d$ ): $\mathrm{CE}=\$ 15,843$.
- Assume that the portion of an office visit used for screening is increased from $50 \%$ to $67 \%$ (Table 3, row $d$ ): $\mathrm{CE}=\$ 31,372$.
- Assume that the number of brief counselling interventions is reduced from 9 to 6 (Table 3, row $j$ ): $\mathrm{CE}=\$ 21,150$.
- Assume that the number of brief counselling interventions is increased from 9 to 12 (Table 3, row j): CE = \$26,064.


## Summary

| Misuse in a BC Birth Cohort of 40,000 |  |  |  |
| :---: | :---: | :---: | :---: |
| Summary |  |  |  |
|  | $\begin{aligned} & \hline \text { Base } \\ & \text { Case } \\ & \hline \end{aligned}$ | Range |  |
| CPB (Potential QALYs Gained) |  |  |  |
| Gap between No Service and 'Best in the World' (30\%) |  |  |  |
| 1.5\% Discount Rate | 1,203 | 903 | 1,582 |
| 3\% Discount Rate | 671 | 503 | 882 |
| \%\% Discount Rate | 2,175 | 1,633 | 2,861 |
| CE (\$/QALY) including patient time costs |  |  |  |
| 1.5\% Discount Rate | \$23,607 | \$5,338 | \$33,069 |
| 3\% Discount Rate | \$33,475 | \$9,237 | \$46,029 |
| 0\% Discount Rate | \$16,611 | \$2,573 | \$23,881 |
| CE (\$/QALY) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | \$4,572 | -\$2,185 | \$8,072 |
| 3\% Discount Rate | \$8,222 | -\$742 | \$12,864 |
| 0\% Discount Rate | \$1,985 | -\$3,207 | \$4,674 |

## Screening for and Management of Obesity

## Canadian Task Force on Preventive Health Care (2015)

We recommend measuring height and weight and calculating BMI at appropriate primary care visits. (Strong recommendation; very low-quality evidence)

We recommend that practitioners not offer formal, structured interventions aimed at preventing weight gain in normal-weight adults. (Weak recommendation; very lowquality evidence)

For adults who are obese (BMI 30-39.9) and are at high risk of diabetes, we recommend that practitioners offer or refer to structured behavioural interventions aimed at weight loss. (Strong recommendation; moderate-quality evidence)

For adults who are overweight or obese, we recommend that practitioners offer or refer to structured behavioural interventions aimed at weight loss. (Weak recommendation; moderate-quality evidence)

For adults who are overweight or obese, we recommend that practitioners not routinely offer pharmacologic interventions (orlistat or metformin) aimed at weight loss. (Weak recommendation; moderate-quality evidence) ${ }^{679}$

## United States Preventive Services Task Force Recommendations (2012)

The USPSTF recommends screening all adults for obesity. Clinicians should offer or refer patients with a body mass index (BMI) of $30 \mathrm{~kg} / \mathrm{m}^{2}$ or higher to intensive, multicomponent behavioral interventions. This is a $B$ recommendation.

Intensive, multicomponent behavioral interventions for obese adults include the following components:

- Behavioral management activities, such as setting weight-loss goals
- Improving diet or nutrition and increasing physical activity
- Addressing barriers to change
- Self-monitoring
- Strategizing how to maintain lifestyle changes

The USPSTF found that the most effective interventions were comprehensive and of high intensity ( 12 to 26 sessions in a year).

Behavioral intervention participants lost an average of $6 \%$ of their baseline weight (4 to 7 kg [ 8.8 to 15.4 lb ]) in the first year with 12 to 26 treatment sessions compared with little or no weight loss in the control group participants. A weight loss of $5 \%$ is considered clinically important by the U.S. Food and Drug Administration (FDA). ${ }^{680}$

[^152]
## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with screening for and management of obesity in adults aged 18 or older in a British Columbia birth cohort of 40,000 .

In modelling CPB, we made the following assumptions:

- Based on 2014 prevalence rates of obesity (based on self-reported height and weight) by age group and sex in BC, ${ }^{681}$ a total of 344,743 life years lived between the ages of 18 and 79 in a birth cohort of 40,000 individuals are in the obese class I or II category (Tables $1 \& 2$, Table 3, row $a$ ).

| Age | Mean Survival | Individuals in Birth | Years of Life in Birth | Prevalen | ce of Ex | xcess We | eight | \# of Year | s with E | cess W | ight |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Group | Rate | Cohort | Cohort | Overweight | Class I | Class II | Class III | Overweight | Class I | Class II | Class III |
| 18-19 | 0.993 | 19,867 | 39,733 | 19.3\% | 4.8\% | 0.3\% | 0.2\% | 7,653 | 1,903 | 118 | 61 |
| 20-24 | 0.991 | 19,813 | 99,065 | 31.2\% | 7.7\% | 0.7\% | 0.2\% | 30,913 | 7,629 | 660 | 211 |
| 25-29 | 0.987 | 19,734 | 98,672 | 36.6\% | 9.3\% | 2.4\% | 0.8\% | 36,082 | 9,191 | 2,372 | 746 |
| 30-34 | 0.983 | 19,658 | 98,289 | 42.7\% | 14.4\% | 4.6\% | 0.0\% | 41,927 | 14,137 | 4,493 | 0 |
| 35-39 | 0.978 | 19,560 | 97,798 | 27.8\% | 21.0\% | 3.6\% | 0.1\% | 27,234 | 20,573 | 3,500 | 118 |
| 40-44 | 0.971 | 19,427 | 97,134 | 37.4\% | 20.2\% | 3.5\% | 0.1\% | 36,284 | 19,656 | 3,396 | 56 |
| 45-49 | 0.962 | 19,241 | 96,203 | 45.4\% | 10.4\% | 5.5\% | 0.2\% | 43,678 | 9,991 | 5,304 | 195 |
| 50-54 | 0.949 | 18,971 | 94,855 | 37.1\% | 25.8\% | 1.3\% | 0.3\% | 35,186 | 24,473 | 1,232 | 290 |
| 55-59 | 0.929 | 18,570 | 92,852 | 47.3\% | 11.4\% | 2.0\% | 1.6\% | 43,958 | 10,565 | 1,855 | 1,476 |
| 60-64 | 0.898 | 17,967 | 89,835 | 41.2\% | 15.8\% | 3.1\% | 1.7\% | 36,989 | 14,225 | 2,822 | 1,567 |
| 65-69 | 0.853 | 17,052 | 85,261 | 44.9\% | 16.2\% | 4.2\% | 0.2\% | 38,256 | 13,818 | 3,565 | 158 |
| 70-74 | 0.783 | 15,668 | 78,342 | 47.7\% | 17.4\% | 3.6\% | 0.4\% | 37,342 | 13,633 | 2,802 | 308 |
| 75-79 | 0.681 | 13,616 | 68,078 | 34.3\% | 8.0\% | 3.0\% | 0.7\% | 23,374 | 5,439 | 2,072 | 478 |
| Total Ag | es 18-79 |  | 1,136,117 | 38.6\% | 14.5\% | 3.0\% | 0.5\% | 438,876 | 165,233 | 34,191 | 5,665 |


| Age | Mean Survival | Individuals in Birth | Years of Life in Birth | Prevalen | ce of Ex | xcess We | eight | \# of Years | with Ex | ess We |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Group | Rate | Cohort | Cohort | Overweight | Class I | Class II | Class III | Overweight | Class I | Class II | Class III |
| 18-19 | 0.995 | 19,891 | 39,781 | 10.2\% | 3.5\% | 0.0\% | 0.0\% | 4,050 | 1,403 | 0 | 0 |
| 20-24 | 0.993 | 19,865 | 99,323 | 17.7\% | 3.5\% | 1.0\% | 0.0\% | 17,582 | 3,488 | 957 | 0 |
| 25-29 | 0.992 | 19,833 | 99,163 | 15.2\% | 4.0\% | 4.2\% | 0.2\% | 15,082 | 3,928 | 4,117 | 150 |
| 30-34 | 0.990 | 19,795 | 98,975 | 20.2\% | 5.7\% | 3.7\% | 1.9\% | 19,963 | 5,645 | 3,675 | 1,918 |
| 35-39 | 0.987 | 19,741 | 98,706 | 21.7\% | 11.0\% | 5.5\% | 2.0\% | 21,463 | 10,849 | 5,436 | 2,021 |
| 40-44 | 0.983 | 19,662 | 98,311 | 23.9\% | 10.7\% | 1.2\% | 4.0\% | 23,531 | 10,500 | 1,215 | 3,947 |
| 45-49 | 0.977 | 19,546 | 97,730 | 29.4\% | 6.2\% | 0.5\% | 0.9\% | 28,771 | 6,083 | 516 | 919 |
| 50-54 | 0.969 | 19,375 | 96,873 | 30.3\% | 15.4\% | 2.2\% | 1.3\% | 29,385 | 14,871 | 2,166 | 1,264 |
| 55-59 | 0.956 | 19,118 | 95,591 | 28.1\% | 8.2\% | 3.1\% | 2.1\% | 26,884 | 7,853 | 2,944 | 2,008 |
| 60-64 | 0.936 | 18,726 | 93,630 | 27.3\% | 14.4\% | 6.0\% | 3.0\% | 25,572 | 13,491 | 5,630 | 2,777 |
| 65-69 | 0.906 | 18,113 | 90,567 | 34.5\% | 11.6\% | 5.0\% | 1.2\% | 31,222 | 10,482 | 4,517 | 1,059 |
| 70-74 | 0.857 | 17,144 | 85,720 | 24.6\% | 9.4\% | 5.9\% | 1.9\% | 21,068 | 8,054 | 5,070 | 1,625 |
| 75-79 | 0.780 | 15,608 | 78,041 | 28.0\% | 14.3\% | 1.6\% | 0.9\% | 21,847 | 11,153 | 1,265 | 702 |
| Total Ages 18-79 |  |  | 1,172,411 | 24.4\% | 9.2\% | 3.2\% | 1.6\% | 286,419 | 107,802 | 37,508 | 18,390 |

- Research for the USPSTF found that behavioral intervention participants lost an average of $6 \%$ or $3 \mathrm{~kg}(6.6 \mathrm{lb})$ of their baseline weight $(95 \% \mathrm{CI}$ of 4 to 7 kg [8.8 to $15.4 \mathrm{lb}]$ ) in the first year with 12 to 26 treatment sessions, compared with little or no

[^153]weight loss in the control group participants. ${ }^{682}$ Research for the CTFPHC found similar results with an average weight loss of $3.02 \mathrm{~kg}(95 \% \mathrm{CI}$ of 2.52 to 3.52$) .{ }^{683} \mathrm{In}$ addition, waist circumference was reduced by an average of $2.78 \mathrm{~cm}(95 \% \mathrm{CI}$ of 2.22 to 3.34 ) and BMI was reduced by $1.11 \mathrm{~kg} / \mathrm{m}^{2}(95 \% \mathrm{CI}$ of 0.84 to 1.39$)$. On average, one out of every five participants ( $95 \%$ CI of 4 to 7 ) lost at least $5 \%$ of their body weight (Table 3, row $c$ ) and one out of nine ( $95 \%$ CI of 7 to 12) lost more than $10 \%$ of their body weight. A weight loss of $5 \%$ is considered clinically important.

- Other assumptions used in assessing the clinically preventable burden are detailed in the Reference Document.

Based on these assumptions, the CPB associated with screening for and management of obesity is 2,287 QALYs (Table 3, row $i$ ).

Table 3: CPB of Screening for and Management of Obesity in Adults in a Birth Cohort
of 40,000

| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Years of life lived with Class I or II obesity | 344,733 | Tables 1 and 2 |
| b | Adherence with an intensive, multicomponent behavioral intervention | 33\% | Ref Doc |
| C | Number needed to treat to achieve a clinically important reduction in weight ( $\geq 5 \%$ of body weight) | 5 | $\checkmark$ |
| d | Reduced years of life lived with Class I or II obesity due to intervention | 22,752 | $=(\mathrm{a} * \mathrm{~b}) / \mathrm{c}$ |
|  | Benefits Associated with Screening and Management |  |  |
| e | Reduction in quality of life - Class I / II obesity vs. overweight | 6.96\% | Ref Doc |
| f | QALYs gained | 1,584 | = ${ }^{*}$ e |
| g | Reduction in years of life lived - Class I / II obesity vs. overweight | 3.09\% | Ref Doc |
| h | QALYs gained | 703 | $=d^{*} \mathrm{~g}$ |
| i | Potential QALYs gained, management increasing from 0\% to 33\% | 2,287 | = $\mathrm{f}+\mathrm{h}$ |

$V=$ Estimates from the literature
We also modified a major assumption and recalculated the CPB as follows:

- Assume that one out of every four participants lost at least $5 \%$ of their body weight after completing an intensive, multicomponent behavioral intervention, rather than one out of every five participants (Table 3, row $c$ ): $\mathrm{CPB}=2,858$ QALYs.
- Assume that one out of every seven participants lost at least $5 \%$ of their body weight after completing an intensive, multicomponent behavioral intervention, rather than one out of every five participants (Table 3, row $c$ ): $\mathrm{CPB}=1,633$ QALYs.


## Modelling Cost-Effectiveness

In this section, we will calculate the CE associated with screening for and management of obesity in adults aged 18 or older in a British Columbia birth cohort of 40,000 .

In modelling CE, we made the following assumptions:

[^154]- Frequency of screening - We assumed that a general practitioner would measure a patient's height and weight in order to calculate BMI and discuss physical activity and healthy eating once every two years (Table 4 , row $g$ ).
- Cost of an intensive, multicomponent behavioral intervention - The per person costs of such interventions in the literature vary substantially, ranging from $\$ 269$ to $\$ 3,267$ (converted to 2017 CAD). ${ }^{684,685,686,687}$ The difference in costs is largely attributable to the ratio of facilitators to clients. The intervention costing $\$ 3,267$ per person involved case managers teaching a 16 -week curriculum on a one-to-one basis. ${ }^{688}$ The intervention costing $\$ 269$ per person was set up for 16 group sessions of up to 18 persons. ${ }^{689}$ We used the mean cost of three of the four interventions (excluding the $\$ 3,267$ per person intervention) for an estimated cost of $\$ 607$ per person per intervention (Table 4, row $m$ ).
- Patient time costs for intensive, multicomponent behavioral intervention - We assumed three hours of patient time would be required (including travel to and from the session) for an average of 18 sessions, the mid-point between 12 and 24 sessions (Table 4, rows $q$ ).
- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the CE associated with screening for and management of obesity is $\$ 12,160$ per QALY (Table 4, row $f f$ ).

[^155]Table 4: CE of Screening for and Management of Obesity in Adults in a Birth
Cohort of 40,000

| $\begin{aligned} & \text { Row } \\ & \text { Label } \end{aligned}$ | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Individuals in birth cohort at age 40 | 39,089 | Tables 1 \& 2 |
| b | Total life years between age 18 and 70 | 1,998,347 | Tables 1 \& 2 |
| c | Proportion of years with Class I/ II obesity without intervention | 14.9\% | Tables 1 \& 2 |
| d | Years with Class I/ II obesity without intervention | 344,733 | Tables 1 \& 2 |
| e | Adherence with screening in primary care | 73\% | Ref Doc |
| f | Adherence with an intensive, multicomponent behavioral intervention | 33\% | Ref Doc |
|  | Costs of intervention |  |  |
| g | Frequency of measuring height and weight and asking about physical activity and diet between age 18 and 70 (every x years) | 2 | Assumed |
| h | Total number of screens | 729,397 | $=(\mathrm{b}$ *e) / g |
| i | Cost of 10-minute office visit | \$34.85 | Ref Doc |
| j | Value of patient time and travel for office visit | \$59.38 | Ref Doc |
| k | Portion of 10-minute office visit for screen | 50\% | Ref Doc |
| I | Cost of screening | \$34,365,530 | $=h^{*}(\mathrm{l}+\mathrm{j}) * \mathrm{k}$ |
| m | Costs per person of an intensive, multicomponent behavioral intervention | \$607 | $\checkmark$ |
| n | Individuals eligible for an intensive, multicomponent behavioral intervention | 5,837 | = ${ }^{*}$ c |
| o | Individuals enrolled in an intensive, multicomponent behavioral intervention | 1,926 | $=\mathrm{n}^{*} \mathrm{f}$ |
| p | Costs of an intensive, multicomponent behavioral intervention | \$1,169,244 | = ${ }^{*} \mathrm{~m}$ |
| q | \# of treatments per intensive, multicomponent behavioral intervention | 18 | $\checkmark$ |
| r | Value of patient time and travel for per intervention treatment | \$89.07 | $\checkmark$ |
| $s$ | Value of patient time and travel for intervention | \$3,088,306 | $=0 * q * r$ |
|  | Cost avoided |  |  |
| t | Number needed to treat to achieve a clinically important reduction in weight ( $\geq 5 \%$ of body weight) | 5 | $\checkmark$ |
| u | Individuals achieving a clinically important reduction in weight ( $\geq 5 \%$ of body weight) | 385 | $=0 / \mathrm{t}$ |
| v | Years with Class I / II obesity avoided with intervention | 22,752 | $=(\mathrm{u} / \mathrm{n}) * \mathrm{~d}$ |
| w | Excess direct costs per year attributable to obesity | \$805 | Ref Doc |
| x | Excess direct costs per year attributable to overweight | \$227 | Ref Doc |
| w | Costs avoided | \$13,150,883 | $=(\mathrm{w}-\mathrm{x})$ * v |
|  | CE calculation |  |  |
| z | Cost of intervention over lifetime of birth cohort | \$38,623,081 | $=1+p+s$ |
| aa | Costs avoided | \$13,150,883 | = w |
| bb | QALYs saved | 2,287 | Table 3, row i |
| cc | Cost of intervention over lifetime of birth cohort (1.5\% discount) | \$26,777,542 | Calculated |
| dd | Costs avoided (1.5\% discount) | \$9,117,562 | Calculated |
| ee | QALY saved (1.5\% discount) | 1,452 | Calculated |
| ff | CE (\$/QALY saved) | \$12,160 | = (cc-dd)/ee |

$V=$ Estimates from the literature
We also modified several major assumptions and recalculated the cost per QALY as follows:

- Assume that one out of every four participants lost at least 5\% of their body weight after completing an intensive, multicomponent behavioral intervention rather than one out of every five participants (Table 3, row $c$ ): $\mathrm{CE}=\$ 8,472$ per QALY.
- Assume that one out of every seven participants lost at least $5 \%$ of their body weight after completing an intensive, multicomponent behavioral intervention rather than one out of every five participants (Table 3, row $c$ ): $\mathrm{CE}=\$ 19,535$ per QALY.
- Assume that the frequency of measuring height and weight and asking about physical activity and diet would occur every year rather than once every two years (Table 4, row $g$ ): $\mathrm{CE}=\$ 28,565$ per QALY.
- Assume that the frequency of measuring height and weight and asking about physical activity and diet would occur every three years rather than once every two years (Table 4, row $g$ ): $\mathrm{CE}=\$ 6,691$ per QALY.
- Assume the proportion of an office visit required for screening/referral is reduced from $50 \%$ to $33 \%$ (Table 4, row $k$ ): $\mathrm{CE}=\$ 6,582$ per QALY.
- Assume the proportion of an office visit required for screening/referral is increased from $50 \%$ to $67 \%$ (Table 4, row $k$ ): CE $=\$ 17,738$ per QALY.
- Assume that the costs per person of an intensive, multicomponent behavioral intervention are reduced from $\$ 607$ to $\$ 269$ (Table 4, row $m$ ): $\mathrm{CE}=\$ 11,849$ per QALY.
- Assume that the costs per person of an intensive, multicomponent behavioral intervention are increased from $\$ 607$ to $\$ 3,267$ (Table 4, row $m$ ): $\mathrm{CE}=\$ 14,606$ per QALY.


## Summary

| Adults in a | Cohort <br> mary | $40,000$ |  |
| :---: | :---: | :---: | :---: |
|  | Base Case | Range |  |
| CPB (Potential QALYs Gained) |  |  |  |
| Gap between 0\% and Best in the World (33\%) |  |  |  |
| 1.5\% Discount Rate | 1,452 | 1,037 | 1,815 |
| 3\% Discount Rate | 959 | 685 | 1,199 |
| 0\% Discount Rate | 2,287 | 1,633 | 2,858 |
| $\overline{\mathrm{CE}}$ (\$/QALY) including patient time costs |  |  |  |
| 1.5\% Discount Rate | \$12,160 | \$6,582 | \$28,565 |
| 3\% Discount Rate | \$13,219 | \$7,155 | \$31,053 |
| 0\% Discount Rate | \$11,140 | \$6,030 | \$26,169 |
| CE (\$/QALY) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | \$348 | -\$1,715 | \$6,415 |
| 3\% Discount Rate | \$378 | -\$1,865 | \$6,974 |
| 0\% Discount Rate | \$318 | -\$1,571 | \$5,877 |

## Falls in Community-Dwelling Elderly

## United States Preventive Service Task Force Recommendations (2012)

Falls are the leading cause of injury in adults aged 65 years or older. Between 30\% and $40 \%$ of community dwelling adults aged 65 years or older fall at least once per year.
The USPSTF recommends exercise or physical therapy and vitamin $D$ supplementation to prevent falls in community-dwelling adults aged 65 years or older who are at increased risk for falls. (Grade B recommendation)
The USPSTF does not recommend automatically performing an in-depth multifactorial risk assessment in conjunction with comprehensive management of identified risks to prevent falls in community-dwelling adults aged 65 years or older because the likelihood of benefit is small. In determining whether this service is appropriate in individual cases, patients and clinicians should consider the balance of benefits and harms on the basis of the circumstances of prior falls, comorbid medical conditions, and patient values. (Grade C recommendation) ${ }^{690}$

More specifically, the USPSTF suggests annual screening for risk using "a pragmatic, expert-supported approach to identifying high risk persons (based on) a history of falls and mobility problems and the results of a timed Get-Up-and-Go test. The test is performed by observing the time it takes a person to rise from an armchair, walk 3 meters ( 10 feet), turn, walk back, and sit down again." Exercise should consist of at least 150 minutes of moderate intensity activity per week while Vitamin D supplementation of 800 IU per day should occur for at least one year. ${ }^{691}$

Note that the 2003 recommendations from the CTFPHC apply only to individuals living in long-term care facilities, rather than the general population of community-dwelling elderly. ${ }^{692}$

## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with preventing falls in the communitydwelling elderly.

In estimating CPB, we made the following assumptions:

- We first estimated the number of life years lived in a BC cohort of 40,000 from age 65 to death as well as the average life expectancy for this cohort (see Table 1). The 765,288 life years lived was used to populate row $a$ of Table 2 while the average life expectancy of 12.5 years was used to populate row $c$ of Table 2 .

[^156]| Table 1: Deaths and Years of Life Lived Between the Ages of 65 and Death in a British Columbia Birth Cohort of 40,000 |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: |
| Age Group | Mean Survival Rate | $\begin{aligned} & \text { Individuals } \\ & \text { in Birth } \\ & \text { Cohort } \\ & \hline \end{aligned}$ | Life Years Lived | Life <br> Expectancy |
| 60-64 | 0.920 | 36,800 |  |  |
| 65-69 | 0.883 | 35,332 | 176,658 | 19.2 |
| 70-74 | 0.827 | 33,072 | 165,362 | 15.3 |
| 75-79 | 0.741 | 29,628 | 148,142 | 11.8 |
| 80-84 | 0.614 | 24,551 | 122,756 | 8.7 |
| 85-89 | 0.441 | 17,632 | 88,158 | 6.1 |
| 90+ | 0.321 | 12,842 | 64,212 | 4.8 |
| Total |  |  | 765,288 | 12.5 |

- An estimated $94.3 \%$ of life years in this cohort are lived in the community (Table 1, row $b$ ). ${ }^{693}$
- Fall-related hospitalizations occur at a rate of 14.19 per 1,000 elderly in BC (Table 1 , row $d$ ). ${ }^{694}$
- An estimated $30 \%$ of individuals die within one year after a fall-related hospitalization (Table 1, row f). ${ }^{695}$
- Individuals who survive a fall-related hospitalization have a $20 \%$ reduced life expectancy (Table 1, row h). ${ }^{696}$
- Individuals who survive a fall-related hospitalization have a .20 reduction in quality of life in year 1 following the hospitalization (Table 1 , row $k$ ) and 0.06 reduction per year thereafter (Table 1 , row $m$ ). ${ }^{697}$
- Interventions involving exercise or physical therapy in reducing falls in communitydwelling elderly have an effectiveness rate of $13 \%$ (RR of $0.87: 95 \% \mathrm{CI}$ of 0.81 to 0.94) (Table 1 , row $p$ ). ${ }^{698}$
- Current delivery of screening and counselling regarding exercise interventions is assumed to be $18 \%$ (Table 1, row $r$ ) (see Reference Document).
- Adherence with exercise intervention is assumed to be $30 \%$ (Table 1 , row $s$ ).
- Other assumptions used in assessing the clinically preventable burden are detailed in the Reference Document.

[^157]The role of vitamin D in fracture prevention is contentious. ${ }^{699,700,701}$ The 2012 USPSTF review noted above, for example, has suggested that vitamin D supplementation reduced the risk of falling by $17 \%$ (RR of 0.83 [ $95 \% \mathrm{CI}$ of 0.77 to 0.89$]$ ). ${ }^{702}$ The Cochrane review, on the other hand, found no reduction in the risk of falling associated with vitamin D supplementation ( $(\mathrm{RR}$ of 0.96 [ $95 \% \mathrm{CI}$ of 0.89 to 1.03$]$ ) although the reviewers did acknowledge that vitamin D supplementation may lower this risk in "people with lower vitamin D levels before treatment. ${ }^{י 703}$ Both groups agree, however, that group and home based exercise as well as home safety interventions reduce the rate of falls and the risk of falls.

Since the 2012 USPSTF review and recommendations regarding the prevention of falls in the community-dwelling elderly, the USPSTF has released (in May 2013) an updated assessment of the use of vitamin D and calcium supplementation to prevent fractures in adults. ${ }^{704,705}$ The updated recommendations include the following:

The USPSTF concludes that the current evidence is insufficient to assess the balance of the benefits and harms of combined vitamin D and calcium supplementation for the primary prevention of fractures in premenopausal women or in men. (Grade I recommendation)

The USPSTF concludes that the current evidence is insufficient to assess the balance of the benefits and harms of daily supplementation with greater than 400 IU of vitamin $D_{3}$ and greater than 1,000 mg of calcium for the primary prevention of fractures in noninstitutionalized postmenopausal women. (Grade I recommendation)

The USPSTF recommends against daily supplementation with 400 IU or less of vitamin $D_{3}$ and 1,000 mg or less of calcium for the primary prevention of fractures in noninstitutionalized postmenopausal women. (Grade D recommendation).

We have therefore focused on the role of exercise in the prevention of falls in the communitydwelling elderly.

Based on these assumptions, the CPB associated with screening and interventions to reduce falls in community-dwelling elderly is 429 (see Table 2, row $t$ ). The CPB of 429 represents the gap between no coverage and the 'best in the world' coverage estimated at $18 \%$ for screening for risk and $30 \%$ for adherence with recommended exercise regimen.

[^158]Table 2: CPB of Screening and Intervention to Reduce Falls in a Birth Cohort of $\mathbf{4 0 , 0 0 0}$ (B.C.)

| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Years lived ages 65+ | 765,288 | Table 1 |
| b | Adjusted for community-dwelling elderly | 0.943 | $\checkmark$ |
| C | Average life expectancy | 12.5 | Table 1 |
| d | Fall-related hospitalizations /1,000 | 14.19 | $\checkmark$ |
| e | Fall-related hospitalizations | 10,240 | $=(a * b) / 1000 * d$ |
| f | Deaths in year following hospital admission | 0.30 | $\checkmark$ |
| g | Fall-related hospitalization LYs lost due to deaths | 38,473 | =e*f* |
| h | Reduced life expectancy for survivors of fall-related hospitalization | 0.20 | $\checkmark$ |
| 1 | Fall-related hospitalization LYs lost in survivors | 17,954 | $=e^{*}(1-\mathrm{f}) *{ }^{*} \mathrm{~h}$ |
| j | Fall-related hospitalization LYs lived in survivors | 71,817 | $={ }^{*}(1-\mathrm{f}) * \mathrm{c}-\mathrm{i}$ |
| k | Reduction in QoL associated with surviving a fall-related hospitalization - Year 1 | 0.20 | $\checkmark$ |
| I | QALYs lost associated with surviving a fall-related hospitalization - Year 1 | 1,434 | $={ }^{*}(1-\mathrm{f}) * \mathrm{k}$ |
| m | Reduction in QoL associated with surviving a fall-related hospitalization - subsequent years | 0.06 | $\checkmark$ |
| n | QALYs lost associated with surviving a fall-related hospitalization - subsequent years | 3,232 | $=(j-(1-f)-i) * m$ |
| 0 | Total QALYs lost | 61,093 | $=g+i+k+n$ |
| p | Effectiveness of exercise at reducing falls | 13.0\% | $\checkmark$ |
| q | QALYs gained based on 100\% adherence | 7,942 | $=0$ * p |
| $r$ | Delivery of screening and counseling | 18.0\% | Ref Doc |
| S | Adherence with exercise | 30.0\% | Assumed |
| t | QALYs gained, CPB | 429 | $=q^{*} r^{*} \mathrm{~s}$ |

$\checkmark=$ Estimates from the literature

We also modified a number of major assumptions and recalculated the CPB as follows:

- Assume that the proportion of the elderly who die within one year following their falls-related hospitalization is decreased from $30 \%$ to $25 \%$ (Table 2, row f): $\mathrm{CPB}=$ 395.
- Assume that the proportion of the elderly who die within one year following their falls-related hospitalization is increased from $30 \%$ to $35 \%$ (Table 2, row f): $\mathrm{CPB}=$ 463.
- Assume the effectiveness of exercise interventions is decreased from $13 \%$ to $6 \%$ (Table 2, row $p$ ): $\mathrm{CPB}=198$.
- Assume the effectiveness of exercise interventions is increased from $13 \%$ to $19 \%$ (Table 2, row $p$ ): $\mathrm{CPB}=627$.


## Modelling Cost-Effectiveness

In this section, we will calculate the CPB associated with preventing falls in the communitydwelling elderly.

In estimating CE, we made the following assumptions:

- Cost per hour of exercise - This is easily the most significant cost and thus drives the estimate of CE (Table 3, row $m$ ). We have estimated the cost of $\$ 5.00$ per hour (e.g., the approximate cost of admission to a community exercise facility), but have also included a sensitivity analysis from $\$ 0$ (e.g., walking) to $\$ 15$ (e.g., the cost per hour for a commercially-based group exercise program). ${ }^{706}$
- Falls-related hospitalization - The cost of a falls-related hospitalization is taken from the Canadian Institute of Health Information Patient Cost Estimator. ${ }^{707}$ We used the average cost in British Columbia associated with a hospitalization for a primary procedure of case-mix group 727 Fixation/repair hip/femur of $\$ 11,897$ (Table 3, row $o)$.
- Other costs and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the CE associated with screening and interventions to reduce falls in community-dwelling elderly are estimated at $\$ 35,213 /$ QALY (see Table 3, row z).

[^159]Table 3: CE of Screening and Intervention to Reduce Falls in a Birth Cohort of
40,000 (B.C.)

| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Years lived ages 65+ as community dwelling elderly | 721,667 | Table 2, row a * Table 2, row b |
|  | Costs of screening |  |  |
| b | Cost of 10-minute office visit | \$34.85 | Ref Doc |
| c | Value of patient time and travel for office visit | \$59.38 | Ref Doc |
| d | Portion of 10-minute office visit for screen | 50\% | Ref Doc |
| e | Delivery of screening and counseling | 18\% | Table 2, row r |
| f | Cost of screening over lifetime of birth cohort | \$6,120,238 | $=(a * e) *(b+c) * d$ |
|  | Costs of interventions |  |  |
| g | Proportion of elderly with falls in previous year | 0.30 | $\checkmark$ |
| h | Portion of 10-minute office visit for referral to exercise program | 50\% | Ref Doc |
| i | Cost of referrals | \$1,836,071 | $=(a * f) * e *((b+c) *$ <br> d) |
| j | Adherence with exercise recommendation | 30\% | Table 2, row s |
| k | Life years lived with exercise in at risk individuals | 11,691 | = ${ }^{*} \mathrm{e}^{*} \mathrm{~g}$ * j |
| 1 | Hours of exercise (3 times per week for 1 hour) | 1,823,796 | = ${ }^{*} 52 * 3$ |
| m | Cost per hour of exercise | \$5.00 | $\checkmark$ |
| n | Cost of intervention (exercise) | \$9,118,979 | $=1 * \mathrm{~m}$ |
|  | Costs avoided |  |  |
| $\bigcirc$ | Reduction in fall-related hospitalizations | 166 | $=(\mathrm{k} / \mathrm{a}) * \text { Table 2, row }$ |
| p | Cost of a fall-related hospitalization | \$11,897 | $\checkmark$ |
| q | Cost avoided | \$1,973,656 | = * p |
|  | CE calculation |  |  |
| $r$ | Cost of initial screen | \$6,120,238 | = f |
| S | Costs of referral and intervention | \$10,955,050 | $=\mathrm{i}+\mathrm{n}$ |
| t | Costs avoided | \$1,973,656 | = q |
| u | QALYs saved | 429 | Table 2, row t |
| v | Cost of initial screen (1.5\% discount rate) | \$5,226,698 | Calculated |
| w | Costs of referral and intervention (1.5\% discount rate) | \$9,355,639 | Calculated |
| X | Costs avoided (1.5\% discount rate) | \$1,685,507 | Calculated |
| y | QALYs saved (1.5\% discount rate) | 366 | Calculated |
| z | CE (\$/QALY saved) | \$35,213 | $=(v+w-x) / y$ |

$V=$ Estimates from the literature
We also modified a number of major assumptions and recalculated the CE as follows:

- Assume that the proportion of the elderly who die within one year following their falls-related hospitalization is decreased from $30 \%$ to $25 \%$ (Table 2 , row $f$ ): $\mathrm{CE}=$ \$38,213 / QALY.
- Assume that the proportion of the elderly who die within one year following their falls-related hospitalization is increased from $30 \%$ to $35 \%$ (Table 2, row f): $\mathrm{CE}=$ \$32,649 / QALY.
- Assume the effectiveness of exercise interventions is decreased from $13 \%$ to $6 \%$ (Table 2, row $p$ ): $\mathrm{CE}=\$ 76,294 / \mathrm{QALY}$.
- Assume the effectiveness of exercise interventions is increased from $13 \%$ to $19 \%$ (Table 2, row p): CE = 24,093 / QALY.
- Assume the cost of an hour of exercise is decreased from $\$ 5$ to $\$ 0$ (Table 3, row $m$ ): CE = \$13,950 / QALY.
- Assume the cost of an hour of exercise is increased from $\$ 5$ to $\$ 15$ (Table 3, row $m$ ): CE $=\$ 77,738 /$ QALY.


## Summary

$\left.\begin{array}{cc}\text { Table 4: Screening and Intervention to Reduce Falls in the } \\ \text { Community-Dwelling Elderly } \\ \text { Summary }\end{array}\right]$

## Preventive Medication / Devices

## Routine Aspirin Use for the Prevention of Cardiovascular Disease and Colorectal Cancer

## Background

In 2007, the USPSTF recommended "against the routine use of aspirin... to prevent colorectal cancer in individuals at average risk for colorectal cancer" with a D recommendation. ${ }^{708}$ In 2009, the USPSTF recommended "the use of aspirin for men age 45 to 79 years when the potential benefit due to a reduction in myocardial infarctions outweighs the potential harm due to an increase in gastrointestinal hemorrhage". The USPSTF also recommended "the use of aspirin for women age 55 to 79 years when the potential benefit of a reduction in ischemic strokes outweighs the potential harm of an increase in gastrointestinal hemorrhage". Both of these 2009 recommendations were A recommendations. ${ }^{709}$

In a 2014 update of the BC LPS, members of the Lifetime Prevention Schedule Expert Committee (LPSEC) reviewed key research that had been published since the 2009 USPSTF recommendations ${ }^{710,711,712}$ calling into question the clinical effectiveness of low-dose aspirin in primary prevention. ${ }^{713,714,715} \mathrm{~A}$ major concern of this new research was that the evidence used for the 2009 USPSTF recommendations appeared to overestimate the benefits of the use of aspirin in primary prevention (e.g. a reduction in cardiovascular disease) and to underestimate the harms (e.g. gastrointestinal bleeding and hemorrhagic stroke). Based on this updated evidence on clinical effectiveness, the LPSEC found that the routine use of lowdose aspirin in primary prevention no longer passed the initial test for inclusion on the BC LPS, namely that the maneuver is not clinically effective (i.e. benefits do not significantly outweigh harms). ${ }^{716}$

In the process of updating both their 2007 and 2009 recommendation on the routine use of aspirin to prevent colorectal cancer and cardiovascular diseases, the USPSTF commissioned

[^160]three systematic evidence reviews ${ }^{717,718,719}$ and one decision analysis using simulation modelling. ${ }^{720}$

The systematic review by Guirguis-Blake and colleagues noted that very-low dose aspirin use ( $\leq 100 \mathrm{mg}$ daily) for primary prevention reduced the risk of nonfatal myocardial infarction by $17 \%$ (RR of $0.83,95 \%$ CI of $0.74-0.94$ ) and nonfatal stroke by $14 \%$ (RR of $0.86,95 \% \mathrm{CI}$ of $0.76-0.98$ ) but they found no reduction in all-cause or cardiovascular mortality. ${ }^{721}$

The systematic review by Chubak and co-authors noted that using aspirin (in dosages ranging from 50 to 500 mg daily) for primary prevention reduced the incidence of colorectal cancer by $40 \%$ (RR of $0.60,95 \%$ CI of $0.47-0.76$ ) but only in secondary studies which followed individuals for at least 10 years. In addition, the use of aspirin for approximately 5 years reduced the risk of death from CRC about 20 years later by $33 \%$ (RR of $0.67,95 \% \mathrm{CI}$ of 0.52 - 0.86). Aspirin's effect on total cancer mortality and incidence was not clearly established. ${ }^{722}$

The systematic review by Whitlock et al. found that very-low dose aspirin use $(\leq 100 \mathrm{mg}$ daily or every other day) increased the risk of major gastrointestinal bleeding by $58 \%$ (RR of 1.58 , $95 \%$ CI of $1.29-1.95$ ) and the risk of haemorrhagic stroke by $27 \%$ (RR of $1.27,95 \% \mathrm{CI}$ of $0.96-1.68) .{ }^{723}$

To help disentangle the "uncertain relationship between the benefits and harms of long-term aspirin use", the USPSTF commissioned the decision analysis by Dehmer and colleagues. ${ }^{724}$ The decision analysis found that the results of net gains (as measured by QALYs) were quite sensitive to all assumptions about the relative risks of both benefits and harms, including baseline risks for GI bleeding. In addition, the results are highly sensitive to assumptions made about the potential disutility associated with regular aspirin use. Their base-case scenario assumed no disutility associated with regular aspirin use.

The collation of this evidence resulted in the following recommendation by the USPSTF.

[^161]
## United States Preventive Services Task Force Recommendations (2016) ${ }^{725}$

The USPSTF recommends initiating low dose aspirin use for the primary prevention of CVD and CRC in adults aged 50 to 59 years who have a $10 \%$ or greater 10-year CVD risk, are not at increased risk for bleeding, have a life expectancy of at least 10 years, and are willing to take low-dose aspirin daily for at least 10 years. ( $B$ recommendation)

The decision to initiate low-dose aspirin use for the primary prevention of CVD and CRC in adults aged 60 to 69 years who have a $10 \%$ or greater 10-year CVD risk should be an individual one. Persons who are not at increased risk for bleeding, have a life expectancy of at least 10 years, and are willing to take low-dose aspirin daily for at least 10 years are more likely to benefit. Persons who place a higher value on the potential benefits than the potential harms may choose to initiate low-dose aspirin. (C recommendation)

Risk factors for gastrointestinal (GI) bleeding with aspirin use include higher dose and longer duration of use, history of GI ulcers or upper GI pain, bleeding disorders, renal failure, severe liver disease, and thrombocytopenia. Other factors that increase risk for GI or intracranial bleeding with low-dose aspirin use include concurrent anticoagulation or nonsteroidal anti-inflammatory drug (NSAID) use, uncontrolled hypertension, male sex, and older age.

## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with initiating low dose aspirin use for the primary prevention of CVD and CRC in adults between the ages of 50 and 59 years who have a $10 \%$ or greater 10 -year CVD risk, are not at increased risk for bleeding, have a life expectancy of at least 10 years, and are willing to take low-dose aspirin daily for at least 10 years.

In estimating CPB, we made the following assumptions:

- Based on BC life tables for 2010 to 2012, there are a total of 380,576 life years lived between the ages of 50 and 59 in a BC birth cohort of 40,000 (see Table 1).
- Based on BC life tables for 2010 to 2012, a total of 1,072 deaths would be expected between the ages of $50-59$, a further 2,460 deaths between the ages of $60-69$ and 5,808 deaths between the ages of 70-79 in a BC birth cohort of 40,000 (see Table 1).
- Based on BC vital statistics data, 601 of 5,076 (11.8\%) deaths in 45-64 year olds in 2011 were due to cardiovascular disease (ICD-10 codes I00-I51) and 191 of 5,076 (3.8\%) deaths were due to cerebrovascular disease (ICD-10 codes I60-I69). ${ }^{726}$ This data was used to estimate that approximately 190 of the 1,611 (11.8\%) deaths between the ages of 55-64 in the birth cohort would be due to cardiovascular disease and $61(3.8 \%)$ due to cerebrovascular disease (see Table 1).

[^162]- Based on BC Cancer Agency data, there were 3,021 ${ }^{727}$ new cases of colorectal cancers (CRC) in BC in 2012 and $1,099^{728}$ deaths due to CRC that same year. An estimated $19.9 \%^{729}$ of deaths (or 219 in BC in 2012) from CRC are in individuals between the ages of 60-69. Since the effectiveness of aspirin on reducing the incidence of CRC only appears after approximately ten years, the age range of 65-74 is being used in the modelling when considering CRC incidence. Similarly, the age range of $75-84$ is being used in the modelling when considering CRC mortality due to the 20 -year lag time observed for this outcome in the research. ${ }^{730}$ An estimated $26.9 \%^{731}$ of deaths (or 296 in BC in 2012) from CRC are in individuals between the ages of 70-79.
- Based on BC vital statistics data, there were 31,776 deaths in BC in 2011. ${ }^{732}$ An estimated $12.5 \%$ of these deaths (or 3,972 ) are in individuals between the ages of 6069 and $22.2 \%$ (or 7,065 ) in individuals between the ages of $70-79 .{ }^{733}$ The 219 deaths from CRC between the ages of 60-69 therefore represents approximately $5.3 \%$ of all deaths in this age cohort. In the birth cohort of $40,000,5.3 \%$ of deaths between the ages of 60-69 represents 130 deaths due to CRC (see Table 1). The 296 deaths from CRC represents approximately $4.2 \%$ of all deaths in this age cohort. In the birth cohort of $40,000,4.2 \%$ of deaths between the ages of 70-79 represents 244 deaths due to CRC (see Table 1).

[^163]| Table 1: Deaths and Selected Causes of Death <br> Between the Ages of 50 and 84 <br> in a British Columbia Birth Cohort of 40,000 |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Age <br> Group | Mean Survival Rate |  | Individuals in Birth Cohort |  |  | Life Years Lived | Deaths in Birth Cohort |  | Cardiovascular Disease |  | Deaths due to Cerebrovascular Disease |  | Colorectal Cancer |  |
|  | Males | Females | Males | Females | Total |  | \% | \# | \% | \# | \% | \# | \% | \# |
| 45-49 | 0.963 | 0.977 | 19,263 | 19,546 | 38,809 |  |  |  |  |  |  |  |  |  |
| 50-54 | 0.950 | 0.969 | 19,003 | 19,375 | 38,378 | 191,890 | 1.1\% | 431 |  |  |  |  |  |  |
| 55-59 | 0.931 | 0.956 | 18,619 | 19,118 | 37,737 | 188,686 | 1.7\% | 641 | 11.8\% | 76 | 3.8\% | 24 |  |  |
| 60-64 | 0.902 | 0.936 | 18,041 | 18,726 | 36,767 | 183,834 | 2.6\% | 970 | 11.8\% | 115 | 3.8\% | 37 | 5.3\% | 51 |
| 65-69 | 0.858 | 0.906 | 17,164 | 18,113 | 35,277 | 176,387 | 4.2\% | 1,489 |  |  |  |  | 5.3\% | 79 |
| 70-74 | 0.792 | 0.857 | 15,837 | 17,144 | 32,981 | 164,903 | 7.0\% | 2,297 |  |  |  |  | 4.2\% | 96 |
| 75-79 | 0.693 | 0.780 | 13,861 | 15,608 | 29,469 | 147,346 | 11.9\% | 3,511 |  |  |  |  | 4.2\% | 147 |
| 80-84 | 0.553 | 0.661 | 11,053 | 13,228 | 24,281 | 121,405 | 21.4\% | 5,188 |  |  |  |  | 4.2\% | 218 |

- We are not aware of any information which indicates the proportion of adults aged 50 to 59 years in BC who have had a cardiovascular or bleeding risk assessment. Nor are we aware of BC-specific data on the proportion of adults at intermediate or higher risk of CVD and low bleeding risk who are taking aspirin over the longer term for primary prevention purposes. Research suggests that $73.3 \%$ of Canadians between the ages of 40 and 59 are at low risk (defined as a mean 10-year risk of a CVD event of less than $10 \%$ ), $10.3 \%$ are at intermediate risk (mean 10-year risk of a CVD event of $10 \%-19 \%$ ) and $16.4 \%$ are at high risk (mean 10-year risk of a CVD event of $\geq 20 \%)^{734}$ (see Table 2).

Table 2: Estimated Number of Canadian Adults Ages 20-79
By CVD Risk Status, 2007 to 2011

| Age <br> Group | Population | Estimated \# by CVD Risk Status |  |  | Estimated \% by CVD Risk Status |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Low | Int. | High | Low | Int. | High |
| 20-39 | 8,983,467 | 8,893,999 | 4,335 | 85,133 | 99.0\% | 0.05\% | 0.95\% |
| 40-59 | 9,863,690 | 7,231,730 | 1,014,437 | 1,617,523 | 73.3\% | 10.3\% | 16.4\% |
| 60-79 | 5,186,843 | 1,011,071 | 1,148,828 | 3,026,944 | 19.5\% | 22.1\% | 58.4\% |
| Total | 24,034,000 | 17,136,800 | 2,167,600 | 4,729,600 | 71.3\% | 9.0\% | 19.7\% |

- We assumed that the average age at which a cardiovascular or cerebrovascular event was prevented due to the use of aspirin would be 60 (Table 3 , rows $q \& x$ ). For the prevention of a CRC event, this would be 70.4 (Table 3, row $a e$ ). For the prevention of a death due to CRC, this would be 80 (Table 3, row $a j$ ). Based on BC life tables for 2010 to 2012, the average life expectancy of a 60 year old is 25.1 years (Table 3, rows $y \& z$ ), that of a 70.4 year old is 16.5 years (Table 3, rows af \& ag) and that of an 80 year old is 9.9 years (Table 3, row $a k$ ). ${ }^{735}$
- Very-low dose aspirin use ( $\leq 100 \mathrm{mg}$ daily) for primary prevention reduces the risk of nonfatal myocardial infarction by $17 \%$ (RR of $0.83,95 \%$ CI of $0.74-0.94$ ) (Table 3, row $a o$ ) and nonfatal stroke by $14 \%$ (RR of $0.86,95 \%$ CI of $0.76-0.98$ ) (Table 3, row $a q$ ), but does not reduce all-cause or cardiovascular mortality. ${ }^{736}$
- Use of aspirin (in dosages ranging from 50 to 500 mg daily) for primary prevention reduces the incidence of colorectal cancer by $40 \%$ (RR of $0.60,95 \%$ CI of $0.47-$

[^164]0.76 ) (Table 3, row as) but only in secondary studies which followed individuals for at least 10 years. ${ }^{77}$

- The use of aspirin for approximately 5 years reduces the risk of death from CRC about 20 years later by $33 \%$ (RR of $0.67,95 \% \mathrm{CI}$ of $0.52-0.86$ ) (Table 3, row au). ${ }^{738}$
- The rate of a major bleeding event in a 50-69 year old not taking aspirin is 1.99 per 1,000 person-years ( $95 \%$ CI 1.82 to 2.18) (Table 3, row $a z$ ). The rate of a major bleeding event in a 50-69 year old who is taking aspirin increases to 3.21 per 1,000 person-years ( $95 \%$ CI 2.93 to 3.53 ) (Table 3, row $b a$ ). Sixty-five percent of bleeding events are episodes of gastrointestinal bleeding (Table 3, row $b c$ ) while $35 \%$ are episodes of intracranial hemorrhage (Table 3, row $b d$ ). ${ }^{739}$
- In a study of 936 patients with acute upper gastrointestinal bleeding (AUGIB) in the UK, 42 ( $4.5 \%$ ) had died by day 28 following the bleeding episode (Table 3, row $b g$ ). The mean QoL score at 28 days for surviving patients was 0.735 compared to 0.86 for the general UK population, a disutility of $14.5 \%$ (Table 3, row bo). We have assumed that this disutility lasts for a one-year period. ${ }^{740}$
- An estimated $40 \%$ of patients die within 28 days after a haemorrhagic stroke (Table 3 , row $b h$ ). ${ }^{741}$
- Other assumptions used in assessing the clinically preventable burden are detailed in the Reference Document.
Based on these assumptions, the CPB associated with screening for and initiating use of low-dose aspirin for the primary prevention of CVD and CRC in adults aged 50 to 59 years who have a $10 \%$ or greater 10 -year CVD risk, are not at increased risk for bleeding, have a life expectancy of at least 10 years, and are willing to take low-dose aspirin daily for at least 10 years is 1,098 QALYs (Table 3, row $b s$ ). This is based on the assumption of moving from no aspirin use in this intermediate to high risk cohort to $24 \%$ of this cohort initiating and sustaining aspirin use.

[^165]Table 3: CPB of Screening for and Initiating Use of Aspirin in Adults Between the Ages of 50 and 59 Years with an Intermediate or Higher Risk of CVD in a Birth Cohort of 40,000

| Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
|  | Estimated current status |  |  |
| a | \# of life years lived between the ages of 55-64 in birth cohort | 372,520 | Table 1 |
| b | \% of life years at low risk of CVD | 73.3\% | Table 2 |
| c | \% of life years at intermediate risk of CVD | 10.3\% | Table 2 |
| d | \% of life years at high risk of CVD | 16.4\% | Table 2 |
| e | \# of life years at low risk | 273,119 | $=(\mathrm{a} * \mathrm{~b})$ |
| f | \# of life years at intermediate risk | 38,312 | $=(\mathrm{a} * \mathrm{c})$ |
| g | \# of life years at high risk | 61,089 | = (a*d) |
| h | Total deaths in birth cohort between the ages of 55-64 | 1,611 | Table 1 |
| i | Cardiovascular deaths in birth cohort between the ages of 55-64 | 190 | Table 1 |
| j | Cerebrovascular deaths in birth cohort between the ages of 55-64 | 61 | Table 1 |
| k | Total deaths in birth cohort between the ages of 65-74 | 3,786 | Table 1 |
| I | Colorectal cancer deaths in birth cohort between the ages of 65-74 | 175 | Table 1 |
| m | Total deaths in birth cohort between the ages of 75-84 | 8,700 | Table 1 |
| n | Colorectal cancer deaths in birth cohort between the ages of 75-84 | 365 | Table 1 |
| 0 | \# of nonfatal cardiovascular events per fatal event | 5.09 | See Ref Doc |
| p | \# of nonfatal cardiovascular events | 968 | = ( ${ }^{*}$ o) |
| q | Average age of individual with a cardiovascular event | 60 | $\checkmark$ |
| $r$ | Life years lived with a nonfatal cardiovascular event | 18.8 | $\checkmark$ |
| S | Life years lost due to a nonfatal cardiovascular event | 6.3 | See Ref Doc |
| t | QoL reduction living with a nonfatal cardiovascular event (for 1 month) | 0.125 | See Ref Doc |
| u | QALYs lost due to nonfatal cardiovascular events | 6,286 | $=\left(p^{*} \mathrm{~s}\right)+\left(\mathrm{p}^{*} \mathrm{r}^{*} \mathrm{t}\right) / 12$ |
| v | Ratio of nonfatal cerebrovascular events per fatal event | 4.58 | See Ref Doc |
| w | \# of nonfatal cerebrovascular events | 280 | $=(\mathrm{j} * \mathrm{u})$ |
| x | Average age of individual with a cerebrovascular event | 60 | $\checkmark$ |
| y | Life years lived with a nonfatal cerebrovascular event | 19.7 | $\checkmark$ |
| z | Life years lost due to a nonfatal cerebrovascular event | 5.5 | See Ref Doc |
| aa | QoL reduction living with a nonfatal cerebrovascular event | 0.264 | See Ref Doc |
| ab | QALYs lost due to nonfatal cerebrovascular events | 3,001 | = ( w*z) + ( w ${ }^{\text {c }}$ * aa ) |
| ac | Ratio of nonfatal colorectal cancer events per fatal event | 4.32 | See Ref Doc |
| ad | \# of nonfatal colorectal cancer events, ages 65-74 | 758 | = ( ${ }^{*}$ aa) |
| ae | Average age of individual with colorectal cancer | 70.4 | See Ref Doc |
| af | Life years lived with colorectal cancer | 6.6 | See Ref Doc |
| ag | Life years lost due to nonfatal colorectal cancer | 9.9 | See Ref Doc |
| ah | QoL reduction living with a nonfatal colorectal cancer event | 0.065 | See Ref Doc |
| ai | QALYs lost due to nonfatal colorectal cancer events | 7,825 | = (ad * ag) + (ad * af * ah) |
| aj | Average age of individual dying from colorectal cancer | 80 | $\checkmark$ |
| ak | Life expectancy of a 80 year old in BC | 9.9 | $\checkmark$ |
| al | QALYs lost due to deaths from colorectal cancer | 3,617 | = (n * ak) |


| Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
|  | Benefits if 24\% of intermediate \& high risk individuals were on aspirin |  |  |
| am | \% of life years at intermediate or high risk on aspirin | 24\% | See Ref Doc |
| an | \# of life years at intermediate or high risk on aspirin | 23,856 | $=(\mathrm{f}+\mathrm{g}) * \mathrm{am}$ |
| ao | \% reduction in risk of cardiovascular disease associated with aspirin use | 17\% | $\checkmark$ |
| ap | QALYs gained due to cardiovascular disease events avoided with 24\% aspirin usage | 256 | $=\left(u^{*} \mathrm{am} * \mathrm{ao}\right)$ |
| aq | \% reduction in cerebrovascular events associated with aspirin use | 14\% | $\checkmark$ |
| ar | QALYs gained due to cerebrovascular disease events avoided with $\mathbf{2 4 \%}$ aspirin usage | 101 | $=(a b * a m * a q)$ |
| as | \% reduction in colorectal cancer events associated with aspirin use, ages 60-69 | 40\% | $\checkmark$ |
| at | QALYs gained due to a reduction in nonfatal colorectal cancer events associated with $24 \%$ aspirin use | 751 | $=(\mathrm{ai} * \mathrm{am} * \mathrm{as})$ |
| au | \% reduction in colorectal cancer deaths associated with aspirin use, ages 70-79 | 33\% | V |
| av | QALYs gained due to a reduction in colorectal cancer deaths associated with 24\% aspirin use | 286 | $=(\mathrm{al}$ * am * au) |
| aw | Total QALYs gained if 24\% of intermediate \& high risk individuals were on aspirin | 1,395 | $=(a n+a q+a t+a v)$ |
|  | Harms if $\mathbf{2 4 \%}$ of intermediate \& high risk individuals were on aspirin |  |  |
| ax | Disutility per year associated with taking pills for cardiovascular prevention | -0.0032 | See Ref Doc |
| ay | Disutility associated with taking pills for cardiovascular prevention | -76 | = (an * ax) |
| az | Risk of major bleeding event in age group $50-69$ per 1,000 person-years, no aspirin | 1.99 | $\checkmark$ |
| ba | Risk of major bleeding event in age group 50-69 per 1,000 person-years, with aspirin | 3.21 | $\checkmark$ |
| bb | Major bleeding events in cohort due to aspirin | 29 | $\begin{gathered} =((\mathrm{ak} / 1000) * \mathrm{ba})- \\ ((\mathrm{ak} / 1000) * \mathrm{az}) \end{gathered}$ |
| bc | Proportion of major bleeding events - gastrointestinal bleeding | 0.65 | V |
| bd | Proportion of major bleeding events - haemorrhagic stroke | 0.35 | $\checkmark$ |
| be | Gastrointestinal bleeding events attributable to aspirin use | 19 | $=\left(\mathrm{bb}{ }^{*} \mathrm{bc}\right)$ |
| bf | Haemorrhagic strokes attributable to aspirin use | 10 | $=(\mathrm{bb} * \mathrm{bd})$ |
| bg | Death rate following a gastrointestinal bleeding event | 0.045 | $\checkmark$ |
| bh | Death rate following a haemorrhagic stroke | 0.40 | $\checkmark$ |
| bi | Deaths due to a gastrointestinal bleeding event | 0.9 | $=(\mathrm{be} * \mathrm{bg})$ |
| bj | Deaths due to a haemorrhagic stroke | 4.1 | $=(\mathrm{bf}$ * bh) |
| bk | Average age of individual with a major bleeding event | 60 | $\checkmark$ |
| bl | Life years lived following a non-fatal gastrointestinal bleeding event | 29.6 | $\checkmark$ |
| bm | Life years lived following a non-fatal haemorrhagic stroke | 24.1 | = (bl - bn) |
| bn | Life years lost following a non-fatal haemorrhagic stroke | 5.5 | See Ref Doc |
| bo | QoL reduction living with a gastrointestinal bleed (1 year only) | -0.145 | $\checkmark$ |
| bp | QALYs lost due to gastrointestinal bleeding | -28 | =(-bi*bl)+((be-bi)*bo) |
| bq | QALYs lost due to haemorrhagic stroke | -193 | $\begin{gathered} =(-\mathrm{bj} * \mathrm{bl})-((\mathrm{bf}-\mathrm{bj}) * \mathrm{bn})-((\mathrm{bf} \\ \mathrm{bj}) * b m * a \mathrm{a}) \end{gathered}$ |
| br | Total QALYs lost if 100\% of intermediate \& high risk individuals were on aspirin | -297 | $=a y+b p+b q$ |
| bs | Net QALYs gained, Screening \& Intervention from 0\% to 24\% | 1,098 | $=(\mathrm{aw}+\mathrm{br})$ |

V = Estimates from the literature
For our sensitivity analysis, we modified a number of major assumptions and recalculated the CPB as follows:

- Assume that decreased risk of cardiovascular disease events associated with aspirin use is reduced from $17 \%$ to $6 \%$ (Table 3, row ao), the decreased risk of cerebrovascular disease events is reduced from $14 \%$ to $2 \%$ (Table 3, row $a q$ ), the decreased risk of incident CRC is reduced from $40 \%$ to $24 \%$ (Table 3, row as) and the decreased risk of mortality due to CRC is reduced from $33 \%$ to $14 \%$ (Table 3 , row $a u$ ): $\mathrm{CPB}=380$.
- Assume that decreased risk of cardiovascular disease events associated with aspirin use is increased $17 \%$ to $26 \%$ (Table 3, row ao), the decreased risk of cerebrovascular disease events is increased from $14 \%$ to $24 \%$ (Table 3, row $a q$ ), the decreased risk of incident CRC is increased from $40 \%$ to $53 \%$ (Table 3, row as) and the decreased risk
of mortality due to CRC is increased from $33 \%$ to $48 \%$ (Table 3, row $a u$ ): $\mathrm{CPB}=$ 1,680.
- Assume that the disutility per year associated with taking pills for cardiovascular prevention is reduced from -0.0033 to 0.000 (Table 3, row $a x$ ): $\mathrm{CPB}=1,174$.
- Assume that the disutility per year associated with taking pills for cardiovascular prevention is increased from -0.0033 to -0.0044 (Table 3, row $a x$ ): $\mathrm{CPB}=1,069$.
- Assume that the rate of a major bleeding event in a 50-69 year old not taking aspirin is reduced from 1.99 to 1.82 per 1,000 person-years (Table 3, row $a z$ ) while the rate of a major bleeding event in a $50-69$ year old who is taking aspirin is reduced from 3.21 to 2.93 per 1,000 person-years (Table 3, row $b a$ ): $\mathrm{CPB}=1,118$.
- Assume that the rate of a major bleeding event in a 50-69 year old not taking aspirin is increased from 1.99 to 2.18 per 1,000 person-years (Table 3, row $a z$ ) while the rate of a major bleeding event in a $50-69$ year old who is taking aspirin is increased from 3.21 to 3.53 per 1,000 person-years (Table 3, row $b a$ ): $\mathrm{CPB}=1,074$.


## Modelling Cost-Effectiveness

In this section, we will calculate the CE associated with initiating low dose aspirin use for the primary prevention of CVD and CRC in adults between the ages of 50 and 59 years who have a $10 \%$ or greater 10 -year CVD risk, are not at increased risk for bleeding, have a life expectancy of at least 10 years, and are willing to take low-dose aspirin daily for at least 10 years.

In estimating CE, we made the following assumptions:

- Screening for CVD risk - The USPSTF notes that it used the ACC/AHA Pooled Cohort Equations to calculate the 10-year risk of CVD events in their analysis and identified key risk factors for GI bleeding: higher doses and longer duration of aspirin use, GI ulcers or upper GI pain, bleeding disorders, renal failure, severe liver disease, thrombocytopenia, concurrent anticoagulation or nonsteroidal anti-inflammatory drug use, uncontrolled hypertension, male sex and older age. ${ }^{742}$
- The need to concurrently screen for CVD risk, bleeding risk and willingness to take low-dose aspirin daily for at least 10 years has recently led to the development of a clinical decision support tool called the Aspirin Guide. ${ }^{743,744}$
- We have assumed that the CVD screening and bleeding risk assessment would take place three times between the ages of 50 and 59 (beginning, mid-point and end of this age range). This would involve screening individuals to determine their risk status and whether or not aspirin would be recommended as well as for follow-up purposes for individuals taking aspirin for primary prevention purposes (Table 3, row $e$ ).
- Completion of a CVD risk assessment includes a physician visit and a full lipid profile (total cholesterol [TC]; high density lipoprotein cholesterol [HDL-C]; lowdensity lipoprotein cholesterol [LDL-C], non-HDL-C; and triglycerides [TG]). The

[^166]full lipid profile costs $\$ 21.31$ (Table 3, row $l$ ). ${ }^{745}$ Note that a CVD risk assessment is required when considering both statins (see previous modelling section) and aspirin for the primary prevention of CVD.

- We assumed that a 10-minute office visit would be required for the initial screening. If the results indicate a low risk of CVD, then the follow-up would consist of a phone call to the patient. If the results indicate an intermediate or high risk of CVD, then a follow-up visit would be required to discuss the results and the possibility of taking aspirin.
- Cost of aspirin therapy - The cost of $100-81 \mathrm{mg}$ aspirin tablets at London Drugs is $\$ 14.99 .{ }^{746}$ We assumed an annual cost of $\$ 54.70$ (Table 3, row $t$ ).
- We assumed an annual follow-up visit with a clinician for patients taking aspirin for preventative purposes (Table 3, row $v$ ).
- Other costs incurred or avoided and assumptions used in assessing cost-effectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the CE associated with screening for and initiating use of lowdose aspirin for the primary prevention of CVD and CRC in adults between the ages of 50 and 69 years who have a $10 \%$ or greater 10-year CVD risk, are not at increased risk for bleeding, have a life expectancy of at least 10 years, and are willing to take low-dose aspirin daily for at least 10 years is $\$ 2,302$ / QALY (Table 3, row bi).

[^167]| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | \# of individuals alive at age 59 in birth cohort | 37,737 | Table 1 |
| b | \# of life years lived between the ages of 55-64 in birth cohort | 372,520 | Table 3 |
| c | $\%$ of life years at intermediate or high risk | 26.7\% | Table 3 |
| d | \# of life years at intermediate or high risk | 99,401 | = (b*c) |
| e | Lifetime number of screens | 3.0 | Assumed |
| f | Adherence with offers to receive screening | 33\% | See Ref Doc |
| g | Total \# of screens in birth cohort | 37,360 | = (a*e*f) |
|  | Estimated cost of screening |  |  |
| h | Number of office visits associated with screening - low risk | 1 | Expert Opinion |
| i | Number of office visits associated with screening - medium or high risk | 2 | Expert Opinion |
| j | Cost of 10-minute office visit | \$34.85 | $\checkmark$ |
| k | Cost of a follow-up phone call | \$15.00 | $\checkmark$ |
| I | Cost to measure cholesterol | \$21.31 | $\checkmark$ |
| m | Health care costs of screening - low risk | \$1,949,142 | $=(1-\mathrm{c}) * \mathrm{~g} \mathrm{~h}^{*}(\mathrm{j}+\mathrm{k}+\mathrm{l})$ |
| n | Health care costs of screening - medium and high risk | \$907,264 | $=\left(\left(c^{*} \mathrm{~g}\right) * \mathrm{i}\right) *\left(\mathrm{j}+\mathrm{l}^{*} 0.5\right)$ |
| 0 | Patient time required / office visit (hours) | 2 | $\checkmark$ |
| p | Value of patient time (per hour) | \$29.69 | $\checkmark$ |
| q | Value of patient time and travel for screening | \$2,810,376 | $=\left(\left(\left(c^{*}{ }^{*} \mathrm{i}\right)+((1-c) * g * h)\right){ }^{*}{ }^{*} \mathrm{p}\right.$ |
|  | Estimated cost of intervention |  |  |
| r | Adherence with long-term aspirin therapy in intermediate \& high risk cohort | 24.0\% | See Ref Doc |
| s | Years on aspirin therapy | 23,856 | $=(\mathrm{d} * \mathrm{r})$ |
| t | Cost of aspirin therapy / year | \$54.70 | $\checkmark$ |
| u | Cost of aspirin therapy | \$1,304,933 | $=(\mathrm{s} * \mathrm{t}$ ) |
| v | Follow-up office visits / year on aspirin therapy | 1.0 | Expert Opinion |
| w | Health care costs of intervention | \$831,388 | $=s^{*} v^{*} \mathrm{j}$ |
| x | Value of patient time and travel for intervention | \$1,416,579 | = ${ }^{*}{ }^{*}{ }^{*}{ }^{*} \mathrm{p}$ |
|  | Estimated costs avoided due to intervention |  |  |
| y | \# of nonfatal cardiovascular events avoided | 39.5 | = Table 3, row p * Table 3, row ao *r |
| z | \# of nonfatal cerebrovascular events avoided | 9.4 | = Table 3, row w * Table 3, row aq * r |
| aa | \# of nonfatal colorectal cancer events avoided | 72.7 | = Table 3, row ad * Table 3, row as *r |
| ab | \# of fatal colorectal cancer events avoided | 28.9 | = Table 3, row n * Table 3, row au * r |
| ac | First year costs avoided per nonfatal cardiovascular event avoided | \$33,934 | See Ref Doc |
| ad | First year costs avoided per nonfatal cerebrovascular event avoided | \$21,139 | See Ref Doc |
| ae | First year costs avoided per nonfatal colorectal cancer event avoided | \$40,080 | See Ref Doc |
| af | Costs avoided per fatal colorectal cancer event avoided | \$49,197 | See Ref Doc |
| ag | First year costs avoided | \$5,878,221 | $=\left(y^{*} a c\right)+\left(z^{*} a d\right)+\left(a a^{*} a e^{\prime}+\left(a b^{*} a f\right)\right.$ |
| ah | Post-first-year annual costs avoided for nonfatal cardiovascular events avoided | \$2,278 | See Ref Doc |
| ai | Duration of post-first year annual costs | 12.1 | See Ref Doc |
| aj | Post-first-year annual costs avoided for nonfatal cerebrovascular events avoided | \$6,246 | See Ref Doc |
| ak | Duration of post-first year annual costs | 9.3 | See Ref Doc |
| al | Post-first-year annual costs avoided for nonfatal colorectal cancer events avoided | \$3,687 | See Ref Doc |
| am | Duration of post-first year annual costs | 6.6 | See Ref Doc |
| an | Post-first-year costs avoided for nonfatal cardiovascular events avoided | \$1,088,300 | = (y * ah *ai) |
| ao | Post-first-year costs avoided for nonfatal cerebrovascular events avoided | \$547,297 | $=(\mathrm{z} * \mathrm{aj} * \mathrm{ak})$ |
| ap | Post-first-year costs avoided for nonfatal colorectal cancer events avoided | \$1,770,154 | = (aa * al *am) |
| aq | Costs avoided due to intervention | \$9,283,971 | $=a g+a n+a o+a p$ |
|  | Estimated costs incurred due to intervention |  |  |
| ar | \# of gastrointestinal bleeds incurred | 18.9 | = Table 3, row be |
| as | \# of nonfatal haemorrhagic strokes incurred | 6.1 | = Table 3, row bf - Table 3, row bj |
| at | \# of fatal haemorrhagic strokes incurred | 4.1 | = Table 3, row bj |
| au | Costs per nonfatal gastrointestinal bleed | \$6,425 | See Ref Doc |
| av | Cost per fatal haemorrhagic stroke | \$9,583 | See Ref Doc |
| aw | First year costs per nonfatal cerebrovascular event | \$21,139 | See Ref Doc |
| ax | Post-first-year costs for nonfatal cerebrovascular events | \$6,246 | See Ref Doc |
| ay | Duration of post-first year annual costs | 9.3 | See Ref Doc |
| az | Costs incurred due to intervention | \$515,625 |  |
|  | CE Calculation |  |  |
| ba | Cost of intervention over lifetime of birth cohort | \$9,219,683 | $=m+n+q+u+w+x$ |
| bb | Costs avoided due to intervention over lifetime of birth cohort | \$9,283,971 | = aq |
| bc | Costs incurred due to intervention over lifetime of birth cohort | \$515,625 | = az |
| bd | Net QALYs saved | 1,098 | Table 3, row bs |
| be | Cost of intervention over lifetime of birth cohort (1.5\% discount) | \$8,045,187 | Calculated |
| bf | Costs avoided due to intervention over lifetime of birth cohort (1.5\% discount) | \$6,864,254 | Calculated |
| bg | Costs incurred due to intervention over lifetime of birth cohort (1.5\% discount) | \$449,939 | Calculated |
| bh | Net QALYs saved (1.5\% discount) | 708 | Calculated |
| bi | CE (\$/QALY saved) | \$2,302 | $=(\mathrm{be}+\mathrm{bg}-\mathrm{bf}) / \mathrm{bh}$ |

For our sensitivity analysis, we modified a number of major assumptions and recalculated the CE as follows:

- Assume that decreased risk of cardiovascular disease events associated with aspirin use is reduced from $17 \%$ to $6 \%$ (Table 3, row ao), the decreased risk of cerebrovascular disease events is reduced from $14 \%$ to $2 \%$ (Table 3, row aq), the decreased risk of incident CRC is reduced from $40 \%$ to $24 \%$ (Table 3, row as) and the decreased risk of mortality due to CRC is reduced from $33 \%$ to $14 \%$ (Table 3, row $a u$ ): $\mathrm{CE}=\$ 24,255$.
- Assume that decreased risk of cardiovascular disease events associated with aspirin use is increased $17 \%$ to $26 \%$ (Table 3, row ao), the decreased risk of cerebrovascular disease events is increased from $14 \%$ to $24 \%$ (Table 3, row $a q$ ), the decreased risk of incident CRC is increased from $40 \%$ to $53 \%$ (Table 3, row as) and the decreased risk of mortality due to CRC is increased from $33 \%$ to $48 \%$ (Table 3, row au): $\mathrm{CE}=-$ \$1,189.
- Assume that the disutility per year associated with taking pills for cardiovascular prevention is reduced from -0.0033 to 0.000 (Table 3, row $a x$ ): $\mathrm{CE}=\$ 2,105$.
- Assume that the disutility per year associated with taking pills for cardiovascular prevention is increased from -0.0033 to -0.0044 (Table 3, row $a x$ ): $\mathrm{CE}=\$ 2,387$.
- Assume that the rate of a major bleeding event in a 50-69 year old not taking aspirin is reduced from 1.99 to 1.82 per 1,000 person-years (Table 3, row $a z$ ) while the rate of a major bleeding event in a 50-69 year old who is taking aspirin is reduced from 3.21 to 2.93 per 1,000 person-years (Table 3, row $b a$ ): $\mathrm{CE}=\$ 2.191$.
- Assume that the rate of a major bleeding event in a 50-69 year old not taking aspirin is increased from 1.99 to 2.18 per 1,000 person-years (Table 3, row $a z$ ) while the rate of a major bleeding event in a $50-69$ year old who is taking aspirin is increased from 3.21 to 3.53 per 1,000 person-years (Table 3, row $b a$ ): $\mathrm{CE}=\$ 2,441$.


## Summary



## Folic Acid Supplementation in Reproductive-age Women for the Prevention of Neural

 Tube Defects (NTDs)United States Preventive Services Task Force Recommendations (2017) ${ }^{747}$
The USPSTF recommends that all women who are planning or capable of pregnancy take a daily supplement containing 0.4 to $0.8 \mathrm{mg}(400-800 \mu \mathrm{~g})$ of folic acid (Grade A recommendation).

The critical period of supplementation starts at least 1 month before conception and continues through the first 2 to 3 months.

## Modelling the Clinically Preventable Burden

In this section, we will calculate the CPB associated with advising all women of reproductive age to take a daily supplement containing 0.4 to $0.8 \mathrm{mg}(400-800 \mu \mathrm{~g})$ of folic acid.

In estimating CPB , we made the following assumptions:

## What are Neural Tube Defects?

- "NTDs are major birth defects of the brain and spine that occur early in pregnancy as a result of improper closure of the embryonic neural tube, which can lead to death or varying degrees of disability. The two most common NTDs are anencephaly and spina bifida." 748
- Anencephaly is a serious birth defect in which a baby is born without parts of the brain and skull.
- "Spina bifida is a congenital malformation in which the spinal column is split (bifid) as a result of failed closure of the embryonic neural tube, during the fourth week post-fertilization. ${ }^{י 749}$
- NTDs are caused by a variety of genetic and non-genetic factors, although the contributing role of each is not fully known. Between $10 \%$ and $60 \%$ of NTDs have a genetic component. Lack of folic acid is perhaps the best known risk factor but there are a number of potential behavioural and environmental risk factors, such as alcohol use, smoking, poor nutrition, valproic acid use and indoor air pollution.
Consequently, some women who take folic acid supplements in the periconceptional period still experience NTD-affected pregnancies. ${ }^{750}$
- The WHO has wrestled with determining what proportion of NTDs are preventable given optimal ( $<906 \mathrm{nmol} / \mathrm{L}$ ) red blood cell folate concentrations in the population. If

[^168]these levels are uniformly achieved, the rate of NTDs could fall somewhere within the range of 4 to 9 per 10,000 live births. ${ }^{751,752}$

## Prevalence of Neural Tube Defects

- Between 1993 and 2002, a total of 2,446 NTDs were among live births, still births and terminations of pregnancies in seven Canadian Provinces. ${ }^{753}$ Of the 2,446 neural tube defects identified in seven Canadian provinces between 1993 and 2002, 1,466 $(60 \%)$ were terminations of pregnancy, $112(5 \%)$ were stillbirth and $868(35 \%)$ were live birth. The majority of NTDs were either spina bifida (53\%) or anencephaly (34\%) (see Table 1). ${ }^{754}$

Table 1: NTDS by Diagnostic Category and Pregnancy Outcome
In Seven Canadian Provinces, 1993 to 2002.
Pregnancy Outcome


| Induced <br> Abortion Stillbirth Live Birth <br> Total\% of <br> Total |
| :--- |

Spina bifida
Anencephaly
Encephalocele
Unspecified NTD Iniencephaly All NTDs
\% of Total

|  |  |  |  |  |  |
| ---: | ---: | ---: | ---: | ---: | ---: |
| 595 | 35 | 656 | 1,286 | $53 \%$ |  |
| 668 | 67 | 95 | 830 | $34 \%$ |  |
| 160 | 8 | 115 | 283 | $12 \%$ |  |
| 24 | 0 | 0 | 24 | $1 \%$ |  |
| 19 | 2 | 2 | 23 | $1 \%$ |  |
|  | $\mathbf{1 , 4 6 6}$ | $\mathbf{1 1 2}$ | $\mathbf{8 6 8}$ | $\mathbf{2 , 4 4 6}$ |  |
| $60 \%$ | $5 \%$ | $35 \%$ |  |  |  |

- Based on data from these seven provinces between January 1, 1993 and September 30, 1997, the prevalence of NTDs among live births, still births and terminations of pregnancies was 15.8 per 10,000 live births. ${ }^{755} \mathrm{BC}$ 's rate, at 9.6 per 10,000 , was the lowest of the seven provinces (see Table 2).

Table 2: Prevalance of NTDS / 10,000 Births
In Seven Canadian Provinces
January 1, 1993 to September 30, 1997

| Province | Rate |
| :---: | :---: |
| N/L | 45.6 |
| NS | 27.2 |
| PEI | 20.8 |
| PQ | 17.7 |
| MB | 15.4 |
| AB | 11.2 |
| BC | 9.6 |
| Combined | 15.8 |

[^169]
## Evidence of the Effectiveness of Folic Acid Supplementation in Reducing the Prevalence of NTDs

- In Hungary in the mid-1980s, 7,540 women planning to conceive were randomly assigned to receive a prenatal vitamin supplement (including 0.8 mg of folic acid) or a trace element supplement, starting one month prior to conception and for three months after conception. In the evaluation of 4,704 pregnancies and 4,122 live births, 28 congenital malformations were observed in the experimental group vs. 47 in the control group. Six of the congenital malformations in the control group were neuraltube defects (NTDs) vs. none in the experimental group. ${ }^{756}$ Given the results of this trial, RCTs are no longer considered ethically possible because of the clear benefits of folic acid supplementation. ${ }^{757}$
- Other cohort and case control studies completed between 1976 and 1998 consistently found evidence of a protective effect associated with folic acid supplementation. ${ }^{758}$
- Case control studies since 1998 have not consistently demonstrated a protective association with folic acid supplementation, but these studies tend to be weakened by misclassification and recall bias. ${ }^{759}$


## Fortification of Grain Products with Synthetic Folic Acids

- The evidence of the effectiveness of folic acid supplementation in reducing the prevalence of NTDs noted above led to a 1992 recommendation by the US Public Health Service that all women of childbearing age consume $400 \mu \mathrm{~g}$ ( 0.4 mg ) of folic acid daily, followed by the US Food and Drug Administration authorization to add synthetic folic acid to grain products in March of 1996 with mandatory compliance by January of 1998. ${ }^{760}$
- In Canada, the milling industry began fortification early in 1997 to meet US requirements for imported flour. On November 11, 1998, fortification of all types of white flour, enriched pasta and cornmeal became mandatory in Canada. ${ }^{761,762}$
- The prevalence of NTDs among live births, still births and terminations of pregnancies declined from 10.7 cases per 10,000 live births before the implementation of food fortification in the US (1995 to 1996) to 7.0 cases per 10,000 live births after fortification. ${ }^{763}$
- In Canada, the prevalence of neural tube defects among live births, still births and terminations of pregnancies decreased from 15.8 to 8.6 per 10,000 live births between January 1, 1993 and December 31, 2002 (see Table 3). ${ }^{764}$ The time period was divided into three 'fortification' periods. The pre-fortification period ran from

[^170]January 1, 1993 to September 30, 1997 to coincide with the beginning of flour fortification in Canada. The partial fortification period ran from October 1, 1997 to March 31, 2000 based on evidence from Ontario that red-cell folate levels in the population started to increase in April 1997 and reached a plateau in February 1999. ${ }^{765}$ The full fortification period ran from April 1, 2000 to December 31, 2002. The biggest reduction between the pre-fortification and full fortification periods was observed in Newfoundland and Labrador (from 45.6 to 7.6 per 10,000) while the smallest reduction was observed in BC (from 9.6 to 7.5 per 10,000). BC already had the lowest prevalence of NTDs (at 9.6 per 10,000 ) in the country before fortification (see Table 3).

Table 3: Prevalance of NTDS / 10,000 Births
In Seven Canadian Provinces
According to Fortification Period


- The prevalence of neural tube defects among live births, still births and terminations of pregnancies declined from 11.3 cases per 10,000 live births before the implementation of food fortification in Ontario (1994 to 1997) to 5.8 cases per 10,000 live births after fortification (1998 to 2000). ${ }^{766}$ Ontario's data was not included in Tables 1 to 3 because the review by De Wals et al. focussed on seven provinces rather than all of Canada.


## Modelling in a BC Birth Cohort of $\mathbf{4 0 , 0 0 0}$

- Based on BC life tables for 2010 to 2012, an estimated 19,672 females would survive through to age 44 in a BC birth cohort of 40,000 (see Table 4). Note that the birth cohort includes both males and females. Our analysis focusses on just the females of reproductive age in this cohort. Based on age specific fertility rates, ${ }^{767}$ an estimated 28,110 live births would occur between the ages of 15 and 44 in this cohort of females (see Table 4).
- For modelling purposes, we have assumed that the pre-fortification rate of NTDs in BC would be approximately 11 / 10,000 live births, followed by a rate of 7.5 / 10,000 live births post-fortification (see Table 3). We have chosen the higher rate of 10.8 (rounded to 11) seen during the partial fortification period in BC (see Table 3) rather than the 9.6 seen during prefortification as a conservative approach (recognizing that the lower 9.6 seen during prefortification in BC may be an anomaly as the rate was reduced from prefortification to partial fortification in all provinces except BC).

[^171]Furthermore, we have assumed that this could be further reduced to 5.8 / 10,000 live births based on Ontario's full fortification rate noted above. ${ }^{768}$ In the sensitivity analysis, we modelled the effect of reducing this rate to $4.0 / 10,000$, the lowest range considered achievable by the WHO given optimal red blood cell folate concentrations in the population. ${ }^{769}$

- We have also assumed that $39 \%$ ( 830 of 2,116 ) of pregnancies with NTD would be anencephaly and $61 \%(1,286$ of 2,116 ) spina bifida (see Table 1). Furthermore, $11.4 \%$ of pregnancies with anencephaly and $51 \%$ of pregnancies with spina bifida would result in a live birth (see Table 1). Based on these assumptions, an estimated 9.6 live births with spina bifida would have occurred in the birth cohort prefortification. The estimated post-fortification status would be 6.5 live births with spina bifida with the potential to be further reduced to 5.1 live births with spina bifida if Ontario's rate of $5.8 / 10,000$ were achieved (see Table 4). Likewise, an estimated 0.9 live births with anencephaly would occur post-fortification with the potential to reduce this to 0.7 live births with anencephaly if Ontario's rate of $5.8 / 10,000$ were achieved (see Table 4).

|  |  |  |  |  | Estimated Prefortification Status |  |  |  |  | Estimated Current Status |  |  |  |  | Estimated Potential Status |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Age |  | Females | Life | \# of |  |  |  | Live Birt | $h$ with |  |  |  | Live Birt | $h$ with |  |  |  | Live Bir | with |
| Group | Survival <br> Females | in Birth Cohort | Years <br> Lived | Live Births | Est. \# of <br> NTDs | Anencephaly | Spina <br> Bifida | Anencephaly | Spina <br> Bifida | $\begin{gathered} \text { Est. \# of } \\ \text { NTDs } \\ \hline \end{gathered}$ | Anencephaly | Spina <br> Bifida | Anencephaly | Spina <br> Bifida | Est. \# of <br> NTDs | Anencephaly | Spina <br> Bifida | Anencephaly | Spina <br> Bifida |
| 15-19 | 0.995 | 19,900 | 99,499 | 759 | 0.8 | 0.3 | 0.5 | 0.0 | 0.3 | 0.6 | 0.2 | 0.3 | 0.0 | 0.2 | 0.4 | 0.2 | 0.3 | 0.0 | 0.1 |
| 20-24 | 0.993 | 19,868 | 99,339 | 3,241 | 3.6 | 1.4 | 2.2 | 0.2 | 1.1 | 2.4 | 1.0 | 1.5 | 0.1 | 0.8 | 1.9 | 0.7 | 1.1 | 0.1 | 0.6 |
| 25-29 | 0.992 | 19,836 | 99,179 | 7,489 | 8.2 | 3.2 | 5.0 | 0.4 | 2.6 | 5.6 | 2.2 | 3.4 | 0.3 | 1.7 | 4.3 | 1.7 | 2.6 | 0.2 | 1.3 |
| 30-34 | 0.990 | 19,799 | 98,997 | 9,894 | 10.9 | 4.3 | 6.6 | 0.5 | 3.4 | 7.4 | 2.9 | 4.5 | 0.3 | 2.3 | 5.7 | 2.3 | 3.5 | 0.3 | 1.8 |
| 35-39 | 0.987 | 19,748 | 98,738 | 5,575 | 6.1 | 2.4 | 3.7 | 0.3 | 1.9 | 4.2 | 1.6 | 2.5 | 0.2 | 1.3 | 3.2 | 1.3 | 2.0 | 0.1 | 1.0 |
| 40-44 | 0.984 | 19,672 | 98,358 | 1,153 | 1.3 | 0.5 | 0.8 | 0.1 | 0.4 | 0.9 | 0.3 | 0.5 | 0.0 | 0.3 | 0.7 | 0.3 | 0.4 | 0.0 | 0.2 |
| Total |  |  | 594,110 | 28,110 | 30.9 | 12.1 | 18.8 | 1.4 | 9.6 | 21.1 | 8.3 | 12.8 | 0.9 | 6.5 | 16.3 | 6.4 | 9.9 | 0.7 | 5.1 |

- A 2015 Cochrane Review found that there is high quality evidence that daily folic acid supplementation (alone or in combination with other vitamins and minerals) prevents NTDs when compared with no intervention/placebo or vitamins and minerals without folic acid ( RR of $0.31,95 \% \mathrm{CI}$ of 0.17 to 0.58 ). The review also found no evidence of an increase in cleft palate, cleft lip, congenital cardiovascular defects, miscarriages or any other birth defects associated with daily folic acid supplementation. ${ }^{770}$
- The 2017 USPSTF review found no significant evidence of potential harms associated with folic acid supplementation. ${ }^{771}$

[^172]- "Spina bifida results from the incomplete closure of the tissue and bone surrounding the spinal cord. Children born with spina bifida can have mild to severe disabilities depending on the location of the lesion along the spinal cord." ${ }^{772}$
- The mortality rate is substantially higher for individuals with moderate to severe spina bifida than for less severe cases. Oakeshott and colleagues have followed a cohort of individuals with spina bifida for 50 years and found that just $12 \%$ with moderate to severe spina bifida survived to age 50 , while $54 \%$ of those with less severe spina bifida survived to age $50 .{ }^{773,774}$
- We used this survival data to compare life expectancy in the general population vs. a population with a sacral lesion (least severe) or a lumbar lesion (moderate to severe) (see Table 5). If we use $100 \%$ to represent the normal life-span of the general population, a person with a sacral lesion will have a life expectancy of $60.6 \%$ (or a loss of $39.4 \%$ of a normal life expectancy, Table 6 , row $m$ ) and a person with a lumbar lesion will have a life expectancy of $25.1 \%$ (or a loss of $74.9 \%$ of a normal life expectancy, Table 6, row $n$ ).

| Table 5: Survival and Year of Life in a Birth Cohort of 40,000 The General Population Compared to Individuals with Spina Bifida |  |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Age Group | Mea <br> Male | Survival Female | Rate <br> Total | pulation <br> Individuals <br> in Birth <br> Cohort | Years of Life in Birth | Lower Mean Survival Rate | Lesion (les <br> Individuals in Birth Cohort | dividuals w severe) Years of Life in Birth | h Spina Bi <br> Higher <br> Mean <br> Survival <br> Rate | fida <br> Lesion (mor <br> Individuals <br> in Birth <br> Cohort | severe) <br> Years of Life in Birth |
| 0-4 | 0.996 | 0.996 | 0.996 | 39,846 | 199,230 | 0.818 | 32,727 | 163,636 | 0.649 | 25,965 | 129,825 |
| 5-9 | 0.995 | 0.996 | 0.996 | 39,823 | 199,115 | 0.764 | 30,545 | 152,727 | 0.526 | 21,053 | 105,263 |
| 「 10-14 | 0.995 | 0.995 | 0.995 | 39,809 | 199,043 | 0.745 | 29,818 | 149,091 | 0.491 | 19,649 | 98,246 |
| 15-19 | 0.994 | 0.995 | 0.994 | 39,773 | 198,864 | 0.691 | 27,636 | 138,182 | 0.456 | 18,246 | 91,228 |
| 20-24 | 0.991 | 0.993 | 0.992 | 39,683 | 198,417 | 0.673 | 26,909 | 134,545 | 0.368 | 14,737 | 73,684 |
| 25-29 | 0.987 | 0.992 | 0.989 | 39,572 | 197,859 | 0.655 | 26,182 | 130,909 | 0.333 | 13,333 | 66,667 |
| 30-34 | 0.983 | 0.990 | 0.986 | 39,451 | 197,253 | 0.618 | 24,727 | 123,636 | 0.298 | 11,930 | 59,649 |
| 35-39 | 0.977 | 0.987 | 0.982 | 39,293 | 196,463 | 0.600 | 24,000 | 120,000 | 0.211 | 8,421 | 42,105 |
| 40-44 | 0.971 | 0.983 | 0.977 | 39,075 | 195,375 | 0.545 | 21,818 | 109,091 | 0.175 | 7,018 | 35,088 |
| 45-49 | 0.961 | 0.977 | 0.969 | 38,765 | 193,826 | 0.545 | 21,818 | 109,091 | 0.123 | 4,912 | 24,561 |
| 50-54 | 0.947 | 0.969 | 0.958 | 38,310 | 191,551 | 0.534 | 21,363 | 106,816 | 0.111 | 4,457 | 22,286 |
| 55-59 | 0.926 | 0.955 | 0.941 | 37,627 | 188,136 | 0.517 | 20,680 | 103,401 | 0.094 | 3,774 | 18,872 |
| 60-64 | 0.894 | 0.935 | 0.915 | 36,591 | 182,955 | 0.491 | 19,644 | 98,220 | 0.068 | 2,738 | 13,690 |
| 65-69 | 0.847 | 0.904 | 0.875 | 35,009 | 175,045 | 0.452 | 18,062 | 90,310 | 0.029 | 1,156 | 5,780 |
| 70-74 | 0.776 | 0.854 | 0.815 | 32,600 | 162,999 | 0.391 | 15,653 | 78,265 |  | 0 | 0 |
| 75-79 | 0.673 | 0.777 | 0.725 | 28,992 | 144,961 | 0.301 | 12,045 | 60,226 |  | 0 | 0 |
| 80+ | 0.531 | 0.659 | 0.595 | 23,809 | 119,047 | 0.172 | 6,862 | 34,312 |  | 0 | 0 |
| Total |  |  |  |  | 3,140,140 |  |  | 1,902,458 |  |  | 786,945 |
| \% Compared to General Population |  |  |  |  |  |  |  | 60.6\% |  |  | 25.1\% |

[^173]- The research by Oakeshott and colleagues was based on 117 consecutive infants born with spina bifida between 1963 and 1971 in the UK who have been followed until 2013. Of these 117 infants, 40 ( $34 \%$ ) died before the age of $5 .{ }^{775}$ The 1 -year survival of infants born with spina bifida in the US has improved from $87.1 \%$ during 1983 to 1987 to $93.6 \%$ during 1998 to $2002 .{ }^{776}$ To take into account the possibility of better long-term survival of infants currently born with spina bifida, we increased the calculated life expectancy of infants with both a sacral (Table 6, row $m$ ) and lumbar lesion (Table 6, row $n$ ) by $25 \%$ in the sensitivity analysis.
- Based on a consecutive cohort of 117 children with spina bifida in the UK, the distribution of children were $33.9 \%$ (Table 6, row $g$ ) with a sacral lesion, $28.6 \%$ (Table 6, row $h$ ) with a lower lumbar lesion and $37.5 \%$ (Table 6, row $i$ ) with a higher lumbar lesion. ${ }^{777}$
- Based on a study of 98 children with spina bifida in Arkansas, the average loss in QoL associated with spina bifida was $41 \%$, ranging from $34 \%$ ( $6 \%$ to $62 \%$ ) for the sacral lesion (Table 6, row $j$ ), $42 \%$ ( $22 \%$ to $62 \%$ ) for the lower lumbar lesion (Table 6 , row $k$ ) and $52 \%$ ( $25 \%$ to $78 \%$ ) for the upper lumbar lesion (Table 6, row $l$ ). We used plus or minus one standard deviation provided by Tilford et al. in the sensitivity analysis. ${ }^{778}$ There was also a modest $5 \%$ reduction in the QoL of caregivers. This reduction, however, was only significantly different from control caregivers for the group of parents caring for the most severe children ( $10 \%$ reduction in QoL). A subsequent, more in depth analysis of these caregivers identified less sleep and less frequent engagement in leisure and social activities as key differences compared with a sample of control caregivers. ${ }^{79}$
- Verhoef and colleagues used the SF-36 to compare the QoL in 164 young adults (ages 16 to 25) with spina bifida in Holland. Compared to the average Dutch population ages $16-25$, young adults with spina bifida experienced a significant decrement in physical functioning ( $51 \%$ ), role limitations due to physical health problems ( $22 \%$ ), bodily pain ( $9 \%$ ) and general health ( $17 \%$ ). No significant differences were observed in vitality, social functioning and role limitations due to emotional health problems or mental health. ${ }^{780}$
- The life expectancy of an infant born in BC of 82.2 years (Table 6 , row $o$ ) is based on life tables for 2010 to 2012 for BC.
- De Wals and colleagues found that there were 656 live births with spina bifida in seven Canadian provinces between 1993 and 2002. At the same time, 1,466 pregnancies with a diagnosed NTD resulted in an induced abortion (see Table 1). ${ }^{781}$

[^174]We have assumed that for every live birth with spina bifida avoided, an estimated 2.23 abortions ( 1,466 / 656 ) would be avoided.

- Other assumptions used in assessing the clinically preventable burden are detailed in the Reference Document.

Based on these assumptions, the CPB associated with advising all women who are planning or capable of pregnancy to take a daily supplement containing 0.4 to $0.8 \mathrm{mg}(400-800 \mu \mathrm{~g})$ of folic acid is 95 QALYs (see Table 6, row $a c$ ). The 95 QALYs is based on moving from the current NTD rate in BC of 7.5 per 10,000 births to 5.8 per 10,000 births, the post fortification rate observed in Ontario.

Table 6: CPB Associated with Advising Women Ages 15 to 44 to Take a Daily Supplement Containing 0.4 to 0.8 mg of Folic Acid in a Birth Cohort of $\mathbf{4 0 , 0 0 0}$

| Row <br> Label | Variable | Base Case | Data Source |
| :---: | :---: | :---: | :---: |
| a | Average \# of females ages 15-44 in birth cohort | 19,767 | Table 4 |
| b | Life years lived between the ages of 15 and 44 | 594,110 | Table 4 |
| c | Live births between the ages of 15 and 44 | 28,110 | Table 4 |
| d | Estimated live births with spina bifida prefortification | 9.6 | Table 4 |
| e | Estimated live births with spina bifida currently | 6.5 | Table 4 |
| f | Estimated potential live births with spina bifida post fortification | 5.1 | Table 4 |
| g | Proportion of children with spina bifida with a sacral lesion (least severe) | 33.9\% | $\checkmark$ |
| h | Proportion of children with spina bifida with a lower lumbar lesion | 28.6\% | V |
| i | Proportion of children with spina bifida with a higher lumbar lesion (most severe) | 37.5\% | $\checkmark$ |
| j | Loss in QoL with a sacral lesion | 34.0\% | $\checkmark$ |
| k | Loss in QoL with a lower lumbar lesion | 42.0\% | $\checkmark$ |
| I | Loss in QoL with a upper lumbar lesion | 52.0\% | $\checkmark$ |
| m | Reduction in life expectancy with a sacral lesion | 39.4\% | $\checkmark$ |
| n | Reduction in life expectancy with a lumbar lesion | 74.9\% | $\checkmark$ |
| 0 | Average life expectancy in BC at birth (in years) | 82.2 | $\checkmark$ |
| p | Births with sacral lesion spina bifida avoided (9.6 to 5.1) | 1.5 | $=(\mathrm{d}-\mathrm{f}) * \mathrm{~g}$ |
| q | Births with lumbar lesion spina bifida avoided (9.6 to 5.1) | 3.0 | $=(\mathrm{d}-\mathrm{f})-\mathrm{p}$ |
| r | Life years gained due to sacral lesion spina bifida avoided | 49.8 | $=m^{*} o^{*} \mathrm{p}$ |
| s | Life years gained due to lumbar lesion spina bifida avoided | 184.4 | $=\mathrm{n}^{*} \mathrm{o}^{*} \mathrm{q}$ |
| t | QALYs gained due to sacral lesion spina bifida avoided | 26.0 | $=p^{*}(1-m) * o{ }^{*}$ |
| u | QALYs gained due to lumbar lesion spina bifida avoided | 29.0 | $\begin{gathered} =q^{*}(1-n) * o^{*}(k \\ +1) / 2 \end{gathered}$ |
| v | Total QALYs gained due to spina bifida avoided (9.6 to 5.1) | 289 | $=r+s+t+u$ |
| w | Births with sacral lesion spina bifida avoided (6.5 to 5.1) | 0.5 | $=(e-f) * g$ |
| x | Births with lumbar lesion spina bifida avoided (6.5 to 5.1) | 1.0 | $=(e-f)-w$ |
| y | Life years gained due to sacral lesion spina bifida avoided | 16.3 | $=m^{*} o^{*} w$ |
| z | Life years gained due to lumbar lesion spina bifida avoided | 60.3 | $=\mathrm{n}^{*} \mathrm{o}^{*} \mathrm{x}$ |
| aa | QALYs gained due to sacral lesion spina bifida avoided | 8.5 | $=w^{*}(1-m) * o * j$ |
| ab | QALYs gained due to lumbar lesion spina bifida avoided | 9.5 | $\begin{gathered} =x *(1-n) * o *(k+ \\ \mathrm{I}) / 2 \end{gathered}$ |
| ac | Total QALYs gained due to spina bifida avoided (6.5 to 5.1) | 95 | $=y+z+a a+a b$ |

$V=$ Estimates from the literature
For our sensitivity analysis, we modified a number of major assumptions and recalculated the CPB as follows:

- Assume that the loss in QoL associated with a sacral lesion is reduced from $34 \%$ to 6\% (Table 6, row $j$ ), the loss in QoL associated with a lower lumbar lesion is reduced from $42 \%$ to $22 \%$ (Table 6, row $k$ ) and the loss in QoL associated with an upper lumbar lesion is reduced from $52 \%$ to $25 \%$ (Table 6, row $l$ ): $\mathrm{CPB}=83$.
- Assume that the loss in QoL associated with a sacral lesion is increased from $34 \%$ to $62 \%$ (Table 6, row $j$ ), the loss in QoL associated with a lower lumbar lesion is
increased from $42 \%$ to $62 \%$ (Table 6, row $k$ ) and the loss in QoL associated with an upper lumbar lesion is increased from $52 \%$ to $78 \%$ (Table 6, row $l$ ): CPB $=106$.
- Assume that the reduction in life expectancy with either a sacral and lumbar lesion is increased by $25 \%$, giving people with spina bifida a longer lifespan. (Table 6, rows $m$ $\& n):$ CPB $=105$.
- Reduce the incidence of NTDs from 5.8 to $4.0 / 10,000$ live births: $\mathrm{CPB}=194$.


## Modelling Cost-Effectiveness

In this section, we will calculate the CE associated with advising all women of reproductive age to take a daily supplement containing 0.4 to $0.8 \mathrm{mg}(400-800 \mu \mathrm{~g})$ of folic acid.

In estimating $C E$, we made the following assumptions:

- Approximately half of all pregnancies are unplanned. Therefore clinicians should advise all women who are capable of pregnancy to take daily folic acid supplements. ${ }^{782}$
- In a survey of 499 women, the majority ( $95 \%$ ) indicated that they prefer to receive information about preconception health from their primary care physician. Only 39\% of these women, however, could recall their physician ever discussing this topic. ${ }^{783}$
- Mazza and colleagues in Australia found that low levels of engagement between primary care providers and women regarding preconception care are due to a number of perceived barriers, including "time constraints, the lack of women presenting at the preconception stage, the numerous competing preventive priorities within the general practice setting, issues relating to the cost of and access to preconception care, and the lack of resources for assisting in the delivery of preconception care guidelines., ${ }^{784}$
- Does a clinician's advice increase the uptake of daily folic acid supplements during the periconceptional period? In a study of 1,173 women with a median age of 32 in the UK, $51 \%$ reported receiving advice on issues such as smoking, alcohol use, healthy diet and folic acid intake from a health professional prior to becoming pregnant. Women who received this advice were significantly more likely to take folic acid supplements ( $76 \%$ ) than women who did not receive this advice $(37 \%){ }^{785}$
- For modelling purposes, we assumed that $70 \%$ (ranging from $60 \%$ to $80 \%$ in the sensitivity analysis) (Table 7, row $b$ ) of clinicians would advise women ages 15 to 44 to take a daily supplement containing 0.4 to 0.8 mg of folic acid and that $76 \%$ (ranging from $66 \%$ to $86 \%$ ) (Table 7, row $e$ ) of women would follow this advice.
- For modelling purposes, we assumed this advice would need to be given every three years (Table 7, row $c$ ) and modified this from every one to five years in the sensitivity analysis.

[^175]- Cost of folic acid supplements - The cost of folic acid supplements averages $\$ 0.043$ per tablet at London Drugs. ${ }^{786} \mathrm{We}$ assumed an annual cost of $\$ 15.70$ (Table 7, row g).
- Costs avoided - Average incremental medical expenditures comparing patients with spina bifida and those without are $\$ 41,460$ (in 2003 USD) in the first year of life, $\$ 14,070$ per year from ages $1-17, \$ 13,339$ per year from ages $18-44$ and $\$ 10,134$ per year from ages 45-64. ${ }^{787}$
- Based on a study of the same 98 children and their caregivers, the caregivers worked an average of 7.5 to 11.3 hours less per week (depending on their children's disability severity) than matched control caregivers. ${ }^{788}$
- Grosse and co-authors estimated the lifetime costs associated with spina bifida to be $\$ 791,900$ (in 2014 USD). This includes $\$ 513,500$ in medical costs, $\$ 63,500$ in special education and developmental service costs and $\$ 214,900$ in parental time costs. ${ }^{789}$ We converted the medical costs to equivalent 2017 Canadian costs; $\$ 454,745$ in medical costs (Table 7, row $r$ ), $\$ 79,203$ in special education and developmental service costs (Table 7, row $s$ ) and $\$ 268,043$ in parental time costs (Table 7, row $t$ ). ${ }^{790}$
- Parental time costs are excluded from the base model (Table 7, row $t$ ) but included in the sensitivity analysis. The literature on 'spillover effects' (e.g. when the illness of a child or family member has an economic or quality of life impact on the broader family or caregiver(s) is nascent and further work is required before these effects can be relied upon with confidence. ${ }^{791,792}$
- For every live birth with spina bifida avoided, an estimated 2.23 abortions would be avoided (Table 7, row $v$ ). The cost of an abortion is estimated at $\$ 609$ (Table 7, row w). ${ }^{793}$
- Anencephaly is uniformly fatal. However, an estimated $11.4 \%$ of pregnancies with anencephaly result in live births (Table 1). These infants survive an average of 2.11 days. ${ }^{794}$ According to the Canadian Institute for Health Information's Patient Cost Estimator, the average cost per day in BC in 2014 for CMG 599 (Neonate 2500+ grams, ages 0-28 days, other major problem) was $\$ 2,085$. ${ }^{795}$ We therefore calculated

[^176]an avoided cost of \$4,399 (2.11 * \$2,085) per anencephaly live birth avoided (Table 7 , row $p$ ).

- Other costs incurred, or avoided, and assumptions used in assessing costeffectiveness are detailed in the Reference Document.
- Discount rate of $1.5 \%$, varied from $0 \%$ to $3 \%$ in the sensitivity analysis.

Based on these assumptions, the CE associated with advising all women of reproductive age to take a daily supplement containing 0.4 to $0.8 \mathrm{mg}(400-800 \mu \mathrm{~g})$ of folic acid is $\$ 195,379$ / QALY (Table 7, row $a d$ ).

| Containing 0.4 to 0.8 mg of Folic Acid in a Birth Cohort of 40,000 |  |  |  |
| :---: | :---: | :---: | :---: |
| Row <br> Label | Variable | Base Case | Data Source |
| a | Life years lived between the ages of 15 and 44 | 594,110 | Table 6, row b |
| b | Clinician adherence in offering advice re: folic acid supplementation | 70\% | Assumed |
| c | Frequency of offering advice re: folic acid supplementation (every x years) | 3 | Assumed |
| d | Life years covered by advice re: folic acid supplementation | 415,877 | =a*b |
| e | Proportion of women taking folic acid supplementation after receiving advice | 76\% | $\checkmark$ |
| $f$ | Life years covered by folic acid supplementation | 316,067 | $=d^{*} \mathrm{e}$ |
| g | Annual cost of folic acid supplementation | \$15.70 | $\checkmark$ |
| h | Cost of folic acid supplementation | \$4,962,244 | = f * g |
| i | Cost of 10-minute office visit | \$34.85 | $\checkmark$ |
| j | Portion of 10-minute office visit for offering advice | 50\% | Assumed |
| k | Costs of office visits | \$2,415,552 | $=(\mathrm{d} / \mathrm{c}) * \mathrm{i} * \mathrm{j}$ |
| 1 | Patient time required per office visit (hours) | 2 | Assumed |
| m | Value of patient time (per hour) | \$29.69 | $\checkmark$ |
| n | Value of patient time and travel for intervention | \$4,115,796 | $=(\mathrm{d} / \mathrm{c}) *{ }^{*} \mathrm{~m}^{*} \mathrm{j}$ |
| 0 | Estimated cost of the intervention | \$11,493,593 | $=\mathrm{h}+\mathrm{k}+\mathrm{n}$ |
| p | Medical care costs avoided per anencephaly live birth avoided | -\$4,399 | $\checkmark$ |
| q | Cases of anencephaly live births avoided with intervention | 0.21 | Table 4 |
| r | Medical care costs avoided per case of spina bifida avoided | -\$454,745 | $\checkmark$ |
| S | Special education and developmental service costs avoided per case of spina bifida avoided | -\$79,203 | $\checkmark$ |
| t | Parental time costs avoided per case of spina bifida avoided | \$0 | $\checkmark$ |
| $u$ | Cases of spina bifida avoided with intervention | 1.48 | Table 6, row w + x |
| v | Abortions avoided per spina bifida live birth | 2.23 | $\checkmark$ |
| w | Costs avoided per abortion avoided | -\$609 | $\checkmark$ |
|  | CE Calculation |  |  |
| x | Cost of intervention over lifetime of birth cohort | \$11,493,593 | $=0$ |
| y | Costs avoided over lifetime of birth cohort | -\$793,981 | $\begin{gathered} =((\mathrm{r}+\mathrm{s}+\mathrm{t}) * \mathrm{u})+\left(\mathrm{u}^{*}\right. \\ \left.\mathrm{v}^{*} \mathrm{w}\right)+(\mathrm{p} * \mathrm{q}) \\ \hline \end{gathered}$ |
| z | QALYs saved | 95 | Table 6, row ac |
| aa | Cost of intervention over lifetime of birth cohort (1.5\% discount) | \$11,493,593 | Calculated |
| ab | Costs avoided over lifetime of birth cohort (1.5\% discount) | -\$697,164 | Calculated |
| ac | QALYs saved (1.5\% discount) | 55 | Calculated |
| ad | CE (\$/QALY saved) | \$195,379 | $=(a a+a b) / \mathrm{ac}$ |

$V=$ Estimates from the literature

For our sensitivity analysis, we modified a number of major assumptions and recalculated the CE as follows:

- Assume that the loss in QoL associated with a sacral lesion is reduced from $34 \%$ to 6\% (Table 6, row $j$ ), the loss in QoL associated with a lower lumbar lesion is reduced from $42 \%$ to $22 \%$ (Table 6, row $k$ ) and the loss in QoL associated with an upper lumbar lesion is reduced from $52 \%$ to $25 \%$ (Table 6, row $l$ ): $\mathrm{CE}=\$ 223,110$.
- Assume that the loss in QoL associated with a sacral lesion is increased from $34 \%$ to $62 \%$ (Table 6, row $j$ ), the loss in QoL associated with a lower lumbar lesion is increased from $42 \%$ to $62 \%$ (Table 76 row $k$ ) and the loss in QoL associated with an upper lumbar lesion is increased from $52 \%$ to $78 \%$ (Table 6, row $l$ ): $\mathrm{CE}=\$ 173,945$.
- Assume that the reduction in life expectancy with either a sacral and lumbar lesion is increased by $25 \%$ (Table 6, rows $m \& n$ ): $\mathrm{CE}=\$ 175,564$.
- Reduce the incidence of NTDs from 5.8 to $4.0 / 10,000$ live births: $\mathrm{CE}=\$ 88,410$.
- Assume that clinician adherence in offering advice re: folic acid supplementation is reduced from $70 \%$ to $60 \%$ (Table 7 , row $b$ ): $\mathrm{CE}=\$ 165,666$.
- Assume that clinician adherence in offering advice re: folic acid supplementation is increased from $70 \%$ to $80 \%$ (Table 7, row b): $\mathrm{CE}=\$ 225,093$.
- Assume that the frequency of offering advice re: folic acid supplementation is increased from every 3 years to every year (Table 7, row $c$ ): $\mathrm{CE}=\$ 431,720$.
- Assume that the frequency of offering advice re: folic acid supplementation is decreased from every 3 years to every 5 years (Table 7, row $c$ ): $\mathrm{CE}=\$ 148,101$.
- Assume the proportion of women taking folic acid supplementation after receiving advice is decreased from $76 \%$ to $66 \%$ (Table 7, row $e$ ): $\mathrm{CE}=\$ 183,563$.
- Assume the proportion of women taking folic acid supplementation after receiving advice is increased from $76 \%$ to $86 \%$ (Table 7, row $e$ ): $\mathrm{CE}=\$ 207,195$.
- Assume that the portion of 10 -minute office visit required for offering advice is reduced from $50 \%$ to $33 \%$ (Table 7, row $j$ ): $\mathrm{CE}=\$ 155,193$.
- Assume that the portion of 10 -minute office visit required for offering advice is increased from $50 \%$ to $66 \%$ (Table 7, row $j$ ): $\mathrm{CE}=\$ 233,202$.
- Include parental time costs avoided per case of spina bifida avoided (Table 7, row $t$ ): $C E=\$ 189,069$


## Summary

| Supplement Containing 0.4 to 0.8 mg of Folic Acid in a Birth Cohort of 40,000 <br> Summary |  |  |  |
| :---: | :---: | :---: | :---: |
|  | Base <br> Case |  |  |
| CPB (Potential QALYs Gained) |  |  |  |
| 1.5\% Discount Rate | 55 | 48 | 114 |
| 3\% Discount Rate | 35 | 31 | 72 |
| 0\% Discount Rate | 95 | 83 | 195 |
| CE (\$/QALY) including patient* time costs |  |  |  |
| 1.5\% Discount Rate | \$195,379 | \$88,410 | \$431,770 |
| 3\% Discount Rate | \$310,525 | \$141,800 | \$683,392 |
| 0\% Discount Rate | \$113,155 | \$50,643 | \$251,301 |
| CE (\$/QALY) excluding patient time costs |  |  |  |
| 1.5\% Discount Rate | \$120,897 | \$52,233 | \$208,324 |
| 3\% Discount Rate | \$193,042 | \$84,736 | \$330,943 |
| 0\% Discount Rate | \$69,628 | \$29,501 | \$120,720 |
| * Patient time costs do not normally include caregiver time costs (spillover effects). In this model, however, we have included caregiver time costs but only in the sensitivity analysis and not in the base case analysis. |  |  |  |

While the approach modelled above involving regular clinic-based reminders for women ages 15 to 44 to take a daily supplement containing folic acid is not cost-effective, folic acid supplementation is still highly recommended before conception and throughout pregnancy. The BC Perinatal Health Program's Maternity Care Pathway, for example, recommends "supplementation with folic acid before conception and throughout pregnancy. Folic acid supplementation as per patient risk ( $0.4 \mathrm{mg}-5 \mathrm{mg}$ per day pre pregnancy)." 796

[^177]
# The Lifetime Prevention Schedule 

## Establishing Priorities among Effective Clinical Prevention Services in British

 ColumbiaSummary and Technical Report October 2019 Update

An update of Clinically Preventable Burden and Cost-Effectiveness Estimates for All Services Reviewed to Date

Participating partner organizations:


BCGuidelines.ca
"By BC Physicians, for BC Physicians"


BC Centre for Disease Control An agency of the Provincial Health Services Authority


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    ${ }^{145}$ Growth monitoring consists of measurement of height or length, weight and BMI calculation or weight for length according to age.
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    ${ }^{147}$ Structured interventions are behavioural modification programs that involve several sessions that take place over weeks to months, follow a comprehensive-approach delivered by a specialized inter-disciplinary team, involve group sessions, and incorporate family and parent involvement. Behaviourally-based interventions may focus on diet, increasing exercise, making lifestyle changes, or any combination of these. These can be delivered by a primary health care team in the office or through a referral to a formal program within or outside of primary care, such as hospital-based, school-based or community programs.
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