3.1 Chronic Disease Prevention and Control

Alternative Perspective

Julia Fox-Rushby

Acknowledgements: This paper was prepared with a small grant from the Copenhagen Consensus Centre

Introduction

Jha and colleagues introduce the case for increased funding of five health interventions to control chronic disease in low and middle income countries: a 33% tax on tobacco; acute management of heart attacks with low cost drugs; prevention of heart attacks and stroke through salt reduction by a mix of voluntary manufacturing changes, behaviour change using mass media and other awareness raising campaigns; prevention of hepatitis B through immunisation; and secondary prevention of heart attacks and stroke through a combination of 3–
4 drugs in a ‘generic risk’ pill\(^1\). The benefit/cost ratios range, in order, from 40:1 to 4:1.

The determination of priorities begins with a focus on the current and expected future burden of disease, as measured by deaths, avoidable mortality, and cost of illness. The ‘very approximate’ (Jha et al 2012\(^{BIB-5}\)) discounted benefit-cost ratios are based on comparing a monetised value of a disability adjusted life year (DALY) with intervention cost.

Evidence on interventions draws largely from the second Disease Control Priorities Project (DCP 2) (Jamison et al 2006\(^{BIB-3\_1}\)), Copenhagen Consensus 2008 paper on disease control (Jamison et al 2008\(^{BIB-3\_1}\)) and selected other literature with a reflection that the investments proposed reflect views of other similar exercises. The five benefit-cost ratios are subject to sensitivity analyses of single and combined changes in the following assumptions; changing the discount rate from 3% to 5%, increasing all costs by 300%, and increasing the value of a DALY from $1000 to $5000.

The benefit-cost ratios are supplemented, to indicate a move to an ‘idealised’ version, by ‘accounting’ for the value of financial protection and non-financial costs (e.g. transaction, organisational and administrative effort to implement the intervention). The ‘accounting’ is a

\(^1\) E.g. use of aspirin, a statin and an antihypertensive drug (Jamison et al 2008\(^{BIB-3\_1}\))
categorisation that relies on: a literature review of various aspects of
health system capacity and; a review of the (limited) evidence on costs
and effects of the Chronic Care Model and its very limited adapted
application to low resource settings. This, at least partly, influences the
qualitative ratings based on the ‘speculative’ judgement of financial
protection and ‘non-financial’ costs by the authors. All interventions are
argued to offer high financial protection with only the impact of
‘capacity’ differentiating the proposed interventions; tobacco taxation is
considered to have low capacity requirements, a salt reduction
programme to have medium capacity requirements and the others to have
high capacity requirements.

The paper ends by calling for an increased role for donor
assistance in controlling chronic diseases despite a concern that this ‘may
not be politically feasible in the short or even medium term’. This role is
also charged to ‘conduct research which makes the marginal costs of
(interventions) affordable’ and includes both more research and
development of relevant health technologies as well as implementation
research to close the gap between knowledge and action.

There is a real challenge in drawing together a justified list of
priorities for funding in an area which is recognised as being both short
of evidence in terms of geographical coverage and range of interventions
evaluated (Suhkre et al 2012) and hampered by poor quality
studies (Mulligan et al 2006[^Mulligan2006]). The paper by Jha and colleagues is therefore a valiant effort to put forward the case for investment in an area of human life that has a worrying future health and economic impact.

This perspective paper considers whether the best interventions for investing in the improvement of chronic disease are presented in the challenge paper. It considers: the influence analysis of burden of illness analysis might have had and should have; the construction and testing of BENEFIT-COST ratios for the five interventions selected; and the approach taken to reflecting uncertainty. The paper ends by suggesting alternative interventions for the expert panel to consider.

### Questioning the influence of burden of illness

The paper appears to reflect the premise that the decision problem should be framed in terms of the burden of disease and, having accounted for the size of burden, focus on the set of cost-effective interventions to reduce the burden. Evidence presented points to mental health conditions having the highest economic burden using the cost-of-illness method and the second largest using the value of lost output method. However, no interventions are proposed for addressing this burden. By implication the authors may have applied a burden of disease approach inconsistently, adopted a very restricted definition of burden of disease or considered evidence on benefit-cost ratios for all mental health interventions to be less than 4:1. These possibilities are considered below.
It is not clear how estimates of burden in the challenge paper have been used in practice to narrow down towards the selected interventions. For example, a burden of illness approach based on mortality rates in Table 1 would suggest that ischemic and hypertensive heart disease should be the focus of all interventions. However, this is not the case as the selected of interventions aim at alleviating heart disease, stroke and cancer\(^2\). Use of avoidable mortality might explain the discrepancy but these data are not provided by disease and therefore the potential influence of this approach is unclear. Two further possibilities are that either the burden of disease approach has been applied inconsistently or it not been the lens through which cost-effective interventions are selected. However, if burden of disease is not the original frame it doesn’t explain why so much information on burden of disease presented without reference to the impact of health interventions.

Perhaps interventions to improve mental health are absent because the impact on mortality is comparatively low. There is a notable absence of cause of death attributed directly to mental health in Table 1 and a statement that “we focus chiefly here on changes in mortality ….. simply because it is far less likely to be misclassified than are the more

\(^2\) Given an assumption that mortality gains from tobacco tax are split equally between cancer and heart disease.
subjective measures of disability”. Valuation of health benefits in the
benefit-cost ratio therefore only appear to account for disability averted
when tied to cases of premature mortality. This suggests first that the
burden and impact of chronic disease is massively underestimated as
highly morbid low mortality chronic diseases will be missing from any
estimate of burden presented here. Indeed co-authors of the
challenge paper conclude elsewhere (Bloom et al, 2011) that cardiovascular
disease and mental health conditions are the
dominant contributors to the global economic burden of non-
communicable diseases. Secondly, it implies a further restriction
imposed by the particular burden of disease approach adopted in the
challenge paper – that cost-effective interventions aimed at alleviating
conditions with lower mortality rates are highly unlikely to be
recommended regardless of their cost-effectiveness. For a proposal
focussed on best buys for reducing chronic disease, this seems somewhat
limited and means that the investment proposals presented are unlikely to
reflect the best possible investment possibilities for reducing chronic
disease.

The possibility that the benefit-cost ratios for all mental health
interventions are less than 4:1 is a moot point and the authors provide no
evidence to support or refute this position. However, evidence from
DCP2 (Jamison et al 2006, p40), on which the challenge paper
itself draws, supports the case that interventions to reduce mental health
are valid contenders to the proposals offered in the challenge paper.

Evidence from DCP2 (Jamison et al 2006\textsuperscript{BIB-3_1}, p40) indicates
cost-effectiveness ratios for mental health interventions in the area of
alcohol abuse are around $600–800/DALY averted and that treatment for
depression by drugs with episodic or maintenance psychosocial
treatment) is roughly $900–3000/DALY averted. The detailed
DCP2 chapter by Hymen et al (2006)\textsuperscript{BIB-3_1} suggested that treatment of
depression with episodic treatment using older tricyclic antidepressants
ranged (by World Bank region) between $478–1,288/DALY averted. More recent evidence suggests that several mental health interventions
could be provided for under $1000/DALY averted in both sub-Saharan
Africa and South East Asia. These include a bundle aimed at alcohol
reduction (including tax increase, reduced access and tax enforcement),
episodic treatment of depression with newer antidepressants (selective
serotonin reuptake inhibitors) and treatment of epilepsy with older anti-
epileptics at 80% coverage (Chisholm et al 2012\textsuperscript{BIB-3_1}).

Evidence presented in Jamison et al (2006\textsuperscript{BIB-3_1}, p41) for the
five selected interventions suggests that interventions to improve mental
health compare well. For example, legislation with public education to
reduce salt content was shown to have a cost/DALY averted of around
$2,000 and secondary treatment of AMI and stroke with a polypill to be
around $700/DALY averted. It is likely therefore, that benefit-cost ratios
of 4:1 or greater for mental health interventions may exist and be on a par
with several of the interventions proposed. This is particularly likely
because the challenge paper converts disability adjusted life years
(DALYs) lost to a monetary value to estimate benefit-cost-ratios without
accounting for other non-money values.

While the absence of interventions for improving mental health
may be of concern, it is only an example and many other cost-effective
interventions could be missing. Of particular concern, given the lack of
clarity in the use of burden of disease estimates in selecting interventions
in this case, is that the proposals could be systematically biased against
recommending the most cost-effective interventions. Why are some
potentially cost-effective treatments of chronic diseases missing? Some
justification of interventions narrowly missing inclusion (e.g. in terms of
benefit-cost ratios or the other criteria) would have helped illuminate the
authors approach more clearly.

Whilst there is unease with the mechanics of using the burden of
illness approach adopted here, of much greater concern is why a burden
is illness approach is used to structure the decision problem. Counting the
size of the epidemiologic or economic problem may indicate problems
for which there are no solutions and could lead to distorted priorities as
more cost-beneficial interventions might never even be considered
Beginning with benefit-cost ratios first is more appropriate as it is a solution focussed approach. It allows a fuller range of potential interventions to be considered regardless of the focus of disease. It is possible that the most cost beneficial intervention would also address the disease of highest burden, but not necessarily.

It is important to recognise that the challenge paper authors were limited to recommending a maximum of five interventions. In this case it is not unreasonable to consider burden of disease estimates in order to benefit from more of the set budget of $75bn. However, to provide the best buy would require considering benefit-cost ratios before considering burden of disease. As the methods of combining information on disease burden and benefit-cost ratios are not clear, it is possible this was done, but this would be important to see.

Construction and sensitivity of the benefit-cost ratios:

‘Indicative’ benefit-cost ratios are presented in Table 7 of the challenge paper with details of calculation presented in the text and sensitivity analysis in the Appendix. Reflecting past research on immunisation for hepatitis B (Brenzel et al 2006; Sanderson 2005) I opted to replicate and reconsider one of the options, using the approach presented in the paper. Column 2 of Table 1 shows the replication. This indicates a 7:1 ratio which, through the rounding in Table 7 and further recalculation
to reflect the rounding was increased by the authors to 10:1 (Verguet, personal communication). The replication therefore satisfactorily reflects the assumptions of the challenge paper.

The assumptions specific to the hepatitis B vaccination option were:

<listing>

a. cost per vaccinated child was $3.6, reflecting a study of India’s national hepatitis B vaccination programme,

b. all benefits would occur 40 years after immunisation;

c. of the 600,000 annual deaths from hepatitis B reported by WHO, a quarter were considered avoidable by increasing global vaccination rates from 75% to 100%.

</listing>

While vaccine effectiveness was referred to as 75 and 95%, the increase from 75–100% coverage appears to implicitly assume 100% effectiveness, as all 150,000 deaths were considered avertable. All other assumptions (e.g. value of a DALY averted, discount rate, DALYs lost per death) were constant across investment options.

In reviewing the benefit-cost calculations three questions arose; Why were particular data and assumptions adopted?; How valuable were the sensitivity analyses in exploring these issues?; and, What is the potential impact of adopting different assumptions?
Little justification was provided for the hepatitis B vaccination-specific parameter values. As the sensitivity analyses only evaluated generic assumptions across all options, no sensitivity analysis considered the impact of option-specific assumptions. Therefore little consideration was given to the possibility that the benefit-cost ratios might change in relation to each other. If one (or more) intervention could move significantly closer to another, differences between options diminish and this could be of decisional importance. As it is relatively easy to choose alternative assumptions to effect change in these benefit-cost ratios, the reasoning for choosing alternative values is important. Therefore this quick reanalysis reflects sources the authors have cited, and applies health sector specific evidence to well versed economic arguments (i.e. rising marginal cost to achieve maximum coverage) to support four cumulative analyses:

<table>
<thead>
<tr>
<th>For achieving more favourable benefit-cost ratios</th>
<th>For achieving less favourable benefit-cost ratios</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Used mean cost from Brenzel et al (2006)</td>
<td>1. Doubled cost of achieving last 10%-point increase in coverage to</td>
</tr>
<tr>
<td>referenced in challenge paper (range $2.02-$2.37)</td>
<td></td>
</tr>
</tbody>
</table>

Johns and Baltussen (2004)\(^3\) showed that marginal costs rose by 70–100% roughly double for achieving the last 10% coverage of a hygiene outreach programme
and inflated to the publication year for Indian cost data used in base case. New cost was $2.7 per vaccinated child. Achieve 100% from $3.6 to $7.2 per child vaccinated for (the effective average cost increased to $5.04 from 75–100% coverage).

2. No amendment made for avoidable mortality as assumptions already appeared favourable (future burden likely to decline given increasing hep B vaccination rates and assumption of 100% efficacy)


3. Used a slightly older coverage rate of 64% vaccine coverage from Duclos et al (2009)\textsuperscript{[BIB-3_1]}. While out of date, the% will reflect the position for some countries.

3. Assumed increase of 3% in global coverage rates since 2010.

4. Assumed benefits occurred in 30 rather than 40 years.

4. Assumed benefits occurred in 50 rather than 40 years.

1 Results for the final cumulative step are given in Table 1. The more favourable assumptions move the benefit-cost ratio from 7:1 to 9:1 and 13:1. The less favourable assumptions move the benefit-cost ratio from 7:1 to 5:1 to 4:1, and finally to 3:1, which is on a par with the generic risk pill. Further investigation of the impact of alternative option-specific assumptions for the four other interventions may reveal a credible alternative positioning of benefit-cost ratios, both in absolute and relative terms.

A Treatment of uncertainty
The challenge paper refers to uncertainty\(^4\) in a number of ways: the size and shape of the future tobacco hazards; greater misclassification of morbidity compared with mortality statistics; methodological uncertainty about completeness of data, age weighting and discount rates; effectiveness of interventions to prevent elevated blood pressure, blood lipids, and diabetes; and adherence to the polypill. To reflect this, the benefit-cost estimates are referred to as ‘indicative’ and parameters to being a ‘ballpark idea’ (e.g. of the economic cost at the macro level). In each case further information on these issues would reduce uncertainty and provide more precise estimates.

The challenge paper judges that, given the “often broad ranges in CE ratios, and hence in benefit-cost ratios, it makes little sense to conclude with precise estimates or with attempts to quantify statistical uncertainty around the point estimates”. While there may be little possibility, given the uncertainties noted, of providing precise estimates, the conclusion that quantification of uncertainty should therefore be

\(^4\) This should be distinguished from variation for which further information could not increase precision as heterogeneity in patient (e.g. age, severity of disease, health outcomes) or health system (e.g. price) characteristics refers to real differences. Jha et al mention additionally variation in prices, scale of the intervention and epidemiological environment.
avoided is a little hasty. Indeed, its avoidance may result in inappropriate recommendations.

Briggs (1995) showed clearly that knowing the precision of an incremental cost-effectiveness ratio can affect the decision about which intervention to implement and indicated that choices may differ from that implied by point estimates alone. For example, in Figure 1 a decision maker with a willingness to pay of £10,000 per quality adjusted life year (QALY) might justifiably prefer intervention C above intervention A or B, because it is a more precise estimate of the incremental cost-effectiveness ratio even though the point estimate of the cost per QALY is higher. Since this work, much progress has been made in defining, measuring and interpreting uncertainty in the context of using economic evaluation to aid both investment adoption decisions as well as defining the need for further research. It has also led to much greater emphasis on the systematic search and review of evidence, as well as methods for eliciting expert opinion and analysis of evidence that influences the choice of parameter estimates in economic evaluations of health interventions (Griffin S and Claxton C 2011).

As uncertainty in both costs and effects can vary by intervention (e.g. Sassi et al, 2009) it is possible that the benefit-cost ratios presented in the challenge paper could be differentially affected by
uncertainty. While it is unusual for uncertainty to be reflected in benefit-cost cost ratios, the analysis of benefit by Jha and colleagues relies heavily on the value of DALYs averted and is not intrinsically different from the majority of economic evaluations presented in the health sector. Therefore analysis of uncertainty could be expected and decisions made without reference to it could badly mislead understanding of the likelihood of future costs and benefits.

Evidence to substantiate, refute and counter the priorities recommended

Two exercises designed to help encourage and guide investment decisions for controlling chronic disease have recently been published. The WHO produced three related reports (WHO 2011a, 2011b, 2011c) outlining the ‘best buys’ for controlling chronic disease and detailed the costs of scaling up the proposed interventions (to a level where 80% coverage is achieved within 15 years). A ‘best buy’ was considered to be an intervention that averts one DALY for less than the average annual income per capita but is also considered “cheap, feasible and culturally acceptable to implement”\(^5\).

\(^5\) This contrasts with ‘good buys’ which are other interventions that may cost more or generate less health gain but are still considered to provide good value for money.
As Jha et al state, all five interventions proposed are, at least partially, reflected in the listing of ‘best buys’. While this is important corroboration of the value of their investment proposal, there are two important caveats to accepting this as sufficient validation. First, further inspection of the ‘best buys’ indicates that several other interventions could have been selected, but the challenge paper is silent on both their non-selection and the reasons for their non-selection. The missing interventions include entire areas, such as controlling alcohol, as well as competing and complementary interventions for the risk factors addressed. Secondly, the reference point for the WHO reports was a

6 The need to select is, however, clear as the total cost of the package was expected to be $170bn with an average annual cost of $11.4 billion per year.

7 This included restricting access, enforce bans on advertising, raising taxes on alcohol, monitoring, advocacy/support. The authors explained (personal communication) that, while excess deaths in Russia can be linked clearly to binge drinking, the net effect in other populations is less clear. However, this decision also appears to be another impact of linking morbidity only to cases of mortality.

8 For diet, these include promoting public awareness about diet and physical activity, replacing trans fat with polyunsaturated fat. For tobacco it includes smoke-free indoor workplaces and public places,
focus on “four diseases; cardiovascular disease, cancer, diabetes and
chronic respiratory disease….(which are) largely caused by four shared
behavioural risk factors; tobacco use, harmful alcohol use, physical
inactivity, and unhealthy diet” (WHO 2011c, p10). Therefore,
confirmation is less convincing as a case for accepting that the best
investments have been presented in the challenge paper, as good
alternatives may exist outside of these disease areas.

A second exercise conducted by WHO focussed on the cost-
effectiveness of over 500 single or combined interventions for the
prevention and control of non-communicable diseases and injuries in
countries in sub-Saharan Africa and South East Asia that have high adult
and child mortality (Chisholm and Saxena 2012\textsuperscript{BIB-3.1}, Chisholm et al
2012\textsuperscript{BIB-3.1}, Ginsberg et al 2012\textsuperscript{BIB-3.1}, Ortegón, Lim, Chisholm and
Mendis 2012\textsuperscript{BIB-3.1}, Ortegon et al 2012\textsuperscript{BIB-3.1}, Baltussen and Smith
2012\textsuperscript{BIB-3.1}). This is interesting for a number of reasons: the analysis
extends beyond the disease areas of the challenge paper and the ‘best
buy’ analysis, including road traffic injuries, mental health, and sensory
health information and warning, bans on advertising, promotion and
sponsorship. Other possibilities to reduce CVD and cancer risks not
presented include; screening in primary care for CVD risk, counselling
and multi-drug therapy for individuals with >30 CVD risk, prevention
of cervical cancer through screening and lesion removal.
loss disorders; it provides a more accountable and direct comparison of a broader range of interventions; and, for the interventions that are not dominated\(^9\) (within disease clusters), a probabilistic cost-effectiveness analysis indicates some degree of the uncertainty. However, there are still limitations with using this analysis as a full critique or validation of investment options presented in the challenge paper. For example, the analysis is restricted to two WHO regions, one intervention proposed by Jha et al is excluded entirely (hepatitis B vaccination\(^10\)), and the drug based interventions proposed in the challenge paper are potentially grouped slightly differently\(^11\).

The second exercise, led by Chisholm, provides strong support for increasing tobacco tax as it is a particularly cost-effective intervention for both WHO regions (see Table 2). However, salt reduction and all salt

\(^9\) An intervention is ‘dominated’ if it is more costly and/or less effective than other (more efficient) interventions

\(^10\) Because treatment of liver disease was considered not to have strong evidence of effectiveness and aspects of prevention of hepatitis B and cirrhosis were ‘covered’ already in some of the alcohol interventions evaluated (Ginsberg et al 2012\(^{11}\)).

\(^11\) This isn’t entirely clear as the WHO based analysis does allow combinations of therapies.
based interventions were dominated by other options (within their
disease/risk factor cluster), as was treatment of AMI with aspirin, ace
inhibitor and beta blockers and all of the, drug therapy based,
secondary/tertiary prevention of myocardial infarction. This indicates that
other interventions could achieve greater DALY gain per $ spent.

| Insert table 3.1.3 here |

Chisholm et al (2012)^{BIB-3-1^}\note that, compared with all other
interventions for controlling chronic disease, “antibiotic treatment of
chronic otitis media (a persistent inflammation of the middle ear) is the
most cost-effective intervention in the two regions (<$Int100/DALY
saved), while extraction of cataracts and proactive screening for hearing
loss are among the biggest contributors to population health gain”. The
detailed results are provided in Table 3 and it can be seen that, even in
comparison with tax increases for tobacco, these interventions are more
cost-effective. However, with a population of 2 million needing cataract
surgery in Africa and 4.2 million in South East Asia (Baltussen and
Smith), the annual treatment is unlikely make a significant dent in the
hypothetical budget facing the Copenhagen Consensus Panel given that
the number of interventions selected are restricted to five. However, this
is unlikely to be the case for an intervention such as treatment based on
absolute risk of a cardiovascular event in next 10 years with statin,
diuretic, β blocker, and aspirin for cardiovascular risk of 5% (CVD-11).
In this case, the annual DALYs saved per million population is 3,163 at a cost of Int$ 0.33 per capita and both an average and incremental cost-effectiveness ratio of Int$104 per DALY averted.

**Conclusion**

Whether an additional investment of up to $75 billion should comprise the five interventions proposed by Jha and colleagues is questionable. The initial filtering through calculations of disease burden combined with a lack accounting for uncertainty and a sensitivity analysis that did not question the relative rankings of interventions suggests that the best buys are unlikely to be presented. Other evidence suggests that alternative interventions could indeed provide a better return on investment. Examples include cataract surgery, antibiotic treatment for otitis media and primary prevention of CVD. However, the cost-effectiveness analysis on which the latter suggestions are made do not account for the level of health system support needed. Jha et al do discuss this at length and it would have been interesting to see both a quantification of health system support needed for the proposed interventions in the challenge paper as well as understanding why this would not support the range of alternative interventions highlighted in the recent series of papers in the British Medical Journal.

**Bibliography**


Sanderson, C. et al., 2005. *Modelling the impact and incremental cost-effectiveness in Bangladesh and Peru of introducing vaccines against hepatitis B, Haemophilus influenzae type b, and rotavirus into routine infant immunisation programmes, and of modifications to current programmes with a particular focus on*
the measles and pertussis components. London: Department of

International Development.

Sassi, F. et al., 2009. Improving lifestyles, tackling obesity: the health
and economic impact of prevention strategies [Online]. OECD

Health working papers series no. 48. Available at:

http://dx.doi.org/10.1787/220087432153 [Accessed: 30 April

2012]

Suhrcke, M., Boluarte, T. and Niessen, L., 2012. A systematic review
of economic evaluations of interventions to tackle cardiovascular
disease in low- and middle-income countries. BMC Public

Health, 12:2.

Williams, A., 1999. Calculating the global burden of disease: time for a

strategic re-appraisal. Health Economics, 8, pp.1–8.


World Health Organisation, 2011a. From burden to ‘best buys’: reducing

economic impact of non-communicable disease in low and


World Health Organization. WHO 2011b. Global status report on non-


Organization
Table 3.1.1 Replication and extension of Jha et al estimate for hepatitis B vaccination

<table>
<thead>
<tr>
<th></th>
<th>Jha et al estimates</th>
<th>Less favourable assumptions</th>
<th>More favourable assumptions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Birth cohort</td>
<td>136,000,000</td>
<td>136,000,000</td>
<td>136,000,000</td>
</tr>
<tr>
<td>Average cost vaccination</td>
<td>3.6</td>
<td>4.6</td>
<td>2.7</td>
</tr>
<tr>
<td>Annual cost of vaccinating all children</td>
<td>489,600,000</td>
<td>625,600,000</td>
<td>367,200,000</td>
</tr>
<tr>
<td>Proportion vaccinated</td>
<td>0.75</td>
<td>0.64</td>
<td>0.75</td>
</tr>
<tr>
<td>New proportion to be vaccinated</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>1% linear cost</td>
<td>4,896,000</td>
<td>6,256,000</td>
<td>3,672,000</td>
</tr>
<tr>
<td>Extra% coverage re expected cost</td>
<td>122,400,000</td>
<td>225,216,000</td>
<td>91,800,000</td>
</tr>
<tr>
<td>Deaths from Hep B</td>
<td>600,000</td>
<td>1,400,000</td>
<td>600,000</td>
</tr>
<tr>
<td>Deaths assumed potentially savable from HBV given current and future vaccination coverage</td>
<td>150,000</td>
<td>176,400</td>
<td>150,000</td>
</tr>
<tr>
<td>DALYs lost per death</td>
<td>20</td>
<td>20</td>
<td>20</td>
</tr>
<tr>
<td>DALYs</td>
<td>3,000,000</td>
<td>3,528,000</td>
<td>3,000,000</td>
</tr>
<tr>
<td>Value of death/DALY averted</td>
<td>1,000</td>
<td>1,000</td>
<td>1,000</td>
</tr>
<tr>
<td>Value of death averted</td>
<td>150,000,000</td>
<td>176,400,000</td>
<td>150,000,000</td>
</tr>
<tr>
<td>Description</td>
<td>Value 1</td>
<td>Value 2</td>
<td>Value 3</td>
</tr>
<tr>
<td>----------------------------------------</td>
<td>----------</td>
<td>----------</td>
<td>----------</td>
</tr>
<tr>
<td>Value of DALY averted</td>
<td>3,000,000,000</td>
<td>3,528,000,000</td>
<td>3,000,000,000</td>
</tr>
<tr>
<td>Undiscounted B:C ratio (death)</td>
<td>1</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Undiscounted B:C ratio (DALYs)</td>
<td>25</td>
<td>16</td>
<td>33</td>
</tr>
<tr>
<td>discounted deaths (3%, 40yrs)</td>
<td>45,179</td>
<td>39,360</td>
<td>60,985</td>
</tr>
<tr>
<td>discounted DALYs</td>
<td>903,583</td>
<td>787,203</td>
<td>1,219,709</td>
</tr>
<tr>
<td>Discounted value deaths</td>
<td>45,179,132</td>
<td>39,360,160</td>
<td>60,985,449</td>
</tr>
<tr>
<td>Discounted value DALYs</td>
<td>903,582,636</td>
<td>787,203,205</td>
<td>1,219,708,979</td>
</tr>
<tr>
<td>Discounted benefit-cost ratio deaths</td>
<td>0</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Discounted benefit-cost ratio DALYs</td>
<td>7</td>
<td>3</td>
<td>13</td>
</tr>
</tbody>
</table>
Table 3.1.2 Costs and effects of a 50% increase in tobacco tax (from 40–60%)

<table>
<thead>
<tr>
<th>WHO Africa Region</th>
<th>WHO South East Asia Region</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Annual DALYs saved per million population</strong></td>
<td>687</td>
</tr>
<tr>
<td><strong>Annual cost per capita (Int $)</strong></td>
<td>0.31</td>
</tr>
<tr>
<td><strong>Average cost-effectiveness ratio (Int $)</strong></td>
<td>448</td>
</tr>
<tr>
<td><strong>Incremental cost-effectiveness ratio (Int $)</strong></td>
<td>448</td>
</tr>
<tr>
<td><strong>Sensitivity</strong></td>
<td>horizontal ellipse stretching from roughly Int$ 0.1–0.7 per capita and 200–1,200 DALYS averted per year per million population (i.e. most uncertainty with effectiveness)</td>
</tr>
</tbody>
</table>

Table 3.1.3 Costs and effects of two alternative interventions for investment

<table>
<thead>
<tr>
<th>Costs and effects of achieving 95% coverage of cataract, extracapsular cataract extraction with posterior chamber lens implant (CAT-6)</th>
<th>WHO Africa Region</th>
<th>WHO South East Asia Region</th>
</tr>
</thead>
<tbody>
<tr>
<td>Annual DALYs saved per million population</td>
<td>6,281</td>
<td>6,447</td>
</tr>
<tr>
<td>Annual cost per capita (Int $)</td>
<td>0.73</td>
<td>0.63</td>
</tr>
<tr>
<td>Average cost-effectiveness ratio (Int $)</td>
<td>116</td>
<td>97</td>
</tr>
<tr>
<td>Incremental cost-effectiveness ratio (Int $)</td>
<td>117</td>
<td>97</td>
</tr>
</tbody>
</table>

Sensitivity: Not possible to read from graph. Horizontal ellipse from (roughly 1,800–10,800 DALYs and Int $0.1–1.0 per capita.

Treatment based on absolute risk of a cardiovascular event in next 10 years with statin, diuretic, β blocker, and aspirin for cardiovascular risk of 5% (CVD-11)

<p>| Annual DALYs saved per million population | 3163 | 2984 |
| Annual cost per capita (Int $) | 0.33 | 0.41 |
| Average cost-effectiveness ratio (Int $) | 104 | 138 |
| Incremental cost-effectiveness ratio | 104 | 146 |</p>
<table>
<thead>
<tr>
<th>Sensitivity</th>
<th>Horizontal ellipse from (roughly) 800–5,200 DALYs lost per million population and (roughly) $0.2 to 0.5 per capita</th>
<th>Horizontal ellipse from (roughly) 1,000–5,000 DALYs lost per million population and (roughly) $0.2 to 0.5 per capita</th>
</tr>
</thead>
</table>

Sources: Baltussen and Smith (2012), Ortegon et al. (2012)
Figure 3.1.1 Variability in point estimates of incremental cost-effectiveness following sensitivity analysis

Source: Briggs (1995)