A comparative case study of bowel cancer screening in the UK and Australia: Evidence lost in translation?

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Conclusions—Insufficient funding has forced programme administrators to make trade-offs that may undermine the potential net population benefits achieved in randomized controlled trials. Such policy compromise contravenes the principle of evidence-based practice which is dependent on adequate funding being made available.

INTRODUCTION

Bowel cancer, also known as colorectal cancer, is the second largest cause of cancer death in men and women combined in all four UK countries and in Australia. The UK National Screening Committee (NSC) and the Australian Health Technology Advisory Committee concluded that screening for bowel cancer using faecal occult blood tests (FOBTs) satisfied the criteria for population screening programmes. The UK and Australian pilot programmes reviewed the evidence for the benefits, harms and cost-effectiveness of bowel cancer screening in their own jurisdictions. Subsequently England, Scotland, Wales, Northern Ireland and Australia have implemented different versions of FOBT bowel cancer screening programmes.

A 2008 review of bowel cancer screening initiatives across 17 countries found FOBTs were the most commonly used screening modality. There are good reasons for governments to allocate funding for FOBT screening programmes. If the 25% relative mortality reduction demonstrated in the intention-to-treat data from the most recent meta-analysis was achieved, almost 4,000 lives in the UK and over 1,000 lives in Australia could be saved each year. Bowel cancer screening can also lead to lower incidence of bowel cancer in the screened population, as the removal of pre-cancerous adenomas prevents the disease occurring. Conversely, the number and severity of potential harms from bowel cancer screening are relatively small, as serious adverse events are rare, and anxiety associated with it is believed to be relatively short-term and minor.

In order to achieve the same or greater magnitude of benefits demonstrated in randomized controlled trials (RCTs), FOBT programmes should be based on the best available evidence. This paper focuses on seven key characteristics we believe are most salient for a population cancer-screening programme and explores the policy trade-offs in these characteristics that result from inadequate funding. Trade-offs are defined as compromises made in one area to obtain benefits in another.

METHODOLOGY

Comparative case study methodology was used to explore in detail the different versions of FOBT screening programmes. This approach emphasizes the importance of describing and interpreting events and their context, in order to illuminate more general issues. For us, the wider issue was the relationship between evidence-informed policy decisions and the implementation of these policy decisions in practice.
**Methods**

**Document collection**—Major documents relating to the five programmes were examined, beginning with background documents to the establishment of the UK and Australian Pilot Programmes. The aim was to establish as complete a set of programme documents as possible. Other sources included commissioned reports, guidelines, press releases, election campaign documents, websites and federal government budget papers. Interview respondents were also helpful in identifying additional relevant documents and some provided copies of background correspondence.

**Interviews**—A total of 42 key respondents were interviewed between August 2007 and October 2009 (34 from Australia and 8 from the UK). Interviews were semi-structured and lasted between 30 and 90 minutes. Questions varied according to the participant’s role in bowel cancer screening policy. All interviews were digitally recorded and transcribed by a professional transcription service.

The Australian interviews were conducted first and purposive sampling was used to achieve a maximum diversity sample, across clinical, methodological and policy expertise and involvement across the policy development timeline. Members of the two major government review committees\(^4,6\) were initially interviewed and our sample was extended based on further document analysis and suggestions from respondents. Interviewees included representatives from cancer organizations, academics, clinicians, former and current state and federal bureaucrats and political advisors.

The UK interviewees were also selected using purposive sampling, with interviews conducted in England and Scotland over a two-week period. Interviewees included those involved in establishing and analysing the Pilots, as well as those responsible for running the English and Scottish programmes. It was a small but highly knowledgeable group.

**Email correspondence**—Email correspondence was used to obtain information on the Welsh and Northern Irish programmes, updates on the implementation of the English and Scottish programmes, and information on the implementation within the six Australian states and two territories. This added an additional six UK and eight Australian respondents to the 42 key respondents interviewed via telephone or face-to-face, making a total of 56 informants. For confidentiality reasons, respondents are referred to only by their identification number, preceded by UK (United Kingdom) or an A (Australia) given in square brackets after quotations or references attributable to them.

**Data analysis**—Information obtained from the documents, interviews and email correspondence was used to compile a table of the seven most salient programme characteristics, as determined by discussion among the authors. Based on our reading of the literature and interview responses from key informants, we proposed a set of ‘optimal’ features for each characteristic. We then compared what had actually happened in practice in each of the five countries (Table 1) with these optimal features (Table 2). Interview data provided insights into the rationale behind some programme implementation choices.

**RESULTS**

Our main findings are presented in Tables 1 and 2, followed by commentary on the types of trade-offs involved for each of the seven characteristics. Details of programme implementation within Australia are provided in Appendix A.
1) Screening frequency

Australia is the only country to not offer screening every two years, despite biennial screening being part of the Australian Pilot programme’s suggested framework and the Australian National Health and Medical Research Council (NHMRC) guidelines recommending at least biennial FOBT screening. Although screening less often reduces the cost of the programme, it is unlikely to improve its cost-effectiveness, as the expected benefits of the programme are also reduced. The trade-off here is between the up-front costs versus the longer-term cost-effectiveness of the programme. It is unlikely that the mortality reductions attained in the RCTs are achievable in the Australian programme which only offers screening to individuals at 50, 55 and 65.

2) Population coverage

The more people included in the target population, the greater the cost of a screening programme. England’s total target population is five times larger than Scotland’s and eighteen times larger than Australia’s. Bowel cancer screening is the first organized screening programme offered to men and women, so the target population is much higher than for programmes offered only to women in a similar age range. For example, in Scotland the target population for breast cancer screening (women aged 50–70) is 672,200, compared with that for bowel cancer screening (men and women aged 50–74) which is 1,471,800. Scotland’s choice to offer screening to 50–74 year olds was based on the Council of the European Union’s recommendation, but suited Scotland’s desire for age-based equity: “... and everyone has to have equitable access to health services and there shouldn’t be different tiers of delivery of health services” [UK03]. Because 74 year olds today are healthier and have a longer life expectancy than when the trials were conducted in the late 1970s to early 1990s, demand may increase in the future. As one respondent noted “we’re now beginning to find people over the age of 74 knocking on the door and saying what about us?” [UK04].

The decision to initially limit screening to 60–69 year olds in England, Scotland and Northern Ireland was pragmatic, reflecting the shortage of colonoscopy capacity in some areas [UK05, UK11, UK13], but was in line with the then recommendations of the UK NSC to routinely screen those in their 60s while allowing older people to request screening. The lack of screening for those in their 50s was not an ideal arrangement, as the risk of bowel cancer incidence and mortality does increase steadily from age 50[28–29] and trial evidence included participants in this age range. Following approval by the UK NSC in April 2011, National Health Service (NHS) England has announced plans for once-only flexible sigmoidoscopy screening at age 55, in addition to FOBT screening.

Given the limited resources however, the decision to focus FOBT screening on the above 60s, where the majority of cancerous and pre-cancerous lesions will be found, makes sense. In Australia, the decision to limit screening to 55 and 65 year olds (later expanded to include 50 year olds) was justified on the basis of the need to manage the capacity of health services, although Australia has a much higher colonoscopy capacity per capita than the UK. As a result, many of the people who may benefit most from screening, particularly those aged 66–75 are not screened at all in the current Australian programme.

3) Quality of test

The choice of an FOBT represents a trade-off between the more costly but more accurate immunochemical test (iFOBT), versus the less expensive but less accurate guaiac test (gFOBT). Only Australia uses a quantitative automated iFOBT in their programme, which has been found to have superior clinical benefits over gFOBTs used in the UK.
The choice of test also affects the screening algorithm, which is a trade-off between accuracy and convenience of testing for the screening participant. The iFOBT gives an actionable result (test is positive or negative) and avoids the need for repeated retesting. In England, participants may be required to complete a gFOBT up to three times and the programme loses many participants to further testing due to the extended screening episodes [UK03]. The programmes in Wales and Northern Ireland have adopted the Scottish algorithm, which involves using gFOBT as the first test, and then following up people who have a ‘weakly positive’ test (that is, 1–4 of 6 windows positive) with a qualitative iFOBT. This ‘two tier’ approach is not without its problems. Although this strategy may reduce the number of colonoscopies (by reducing the number of false positive tests), the overall programme sensitivity is still limited by the sensitivity of the initial gFOBT.34

Ideally, a screening programme should be capable of finding all the pathology that exists in the target population. In practice, this is usually not feasible. As van Rossum et al. have noted: “Policy makers will determine the optimal cutoff value [the point at which the test is considered to be positive or negative] on the basis of a largely arbitrary balance between the acceptability of missing cancer and the possibility and acceptability of assigning essential resources”.35 Scotland is currently conducting a trial of a quantitative automated iFOBT to replace the current algorithm, but even if funded, the positivity rate will still need to be set at around 2% as their health system will not cope with additional colonoscopies required to deal with a larger pathology yield [UK04].

4) Programme model

The English, Welsh and Northern Irish programmes follow a single funder exclusive designated provider model. It separates the provision of screening colonoscopies from symptomatic or surveillance colonoscopies, and so follows the model for breast cancer screening. The developers of the UK breast cancer screening programme deliberately differentiated between screening and non-screening cases, arguing that this was preferable, because even though initially the women being screened for breast cancer may have better facilities, it was important to first establish those high standards [UK07].

Scotland intentionally took a different approach because they did not want to create a ‘two-tier’ system where people receiving colonoscopy as part of the screening programme were guaranteed high quality colonoscopy, but those with symptoms or on a surveillance programme were not [UK03]. Each of the fourteen Scottish health boards are responsible for ensuring a positive FOBT is treated in the same way as an urgent GP referral with fast-tracked investigation and treatment [UK03]. In all four UK countries, most screening follow-up is delivered in the public sector, without GP involvement.

In Australia, the bowel cancer screening programme was established as a ‘usual care’ model, with the federal government funding the screening pathway only up to the point of a positive FOBT. The state and territory governments, which are responsible for funding of hospital-based services, were left to cover the costs of colonoscopy facilities for follow-up of positive FOBTs performed in the public sector.

5) Quality of follow-up of positive test

In the designated provider jurisdictions in the UK, a central body is responsible for organising pre-colonoscopy assessment appointments by specialized staff for those with positive FOBTs. In Scotland pre-colonoscopy assessment is arranged by local NHS boards and the quality may vary [UK04]. Similarly, in Australia there is no nationally coordinated process for pre-colonoscopy assessment, which is organized through GPs who may lack screening expertise or the time to devote to discussion of the pros and cons of further testing.
There is the potential for screening colonoscopies to impact on waiting times for people needing surveillance colonoscopies or those presenting with symptoms. Countries with limited colonoscopy capacity may find this difficult to manage. In Australia, over 500,000 colonoscopies are performed annually. As at January 2010, less than 29,000 colonoscopies were undertaken through the National Bowel Cancer Screening Program (NBCSP),

so colonoscopy capacity is not the major issue. The choice of the ‘usual care’ model, however, has led to wide variation in the quality of follow-up between the States (see Appendix A).

Over 75% of colonoscopies performed in Australia are carried out in the private sector.

Although this reduces the financial impact of a screening programme on public sector facilities, the federal government still pays the majority of colonoscopy costs through the fee for service-based Medicare Benefits Schedule. Participants are reimbursed through their private health insurance (if covered) but are liable for any ‘gap’ payment if there is a difference between the private health fund reimbursement and the private colonoscopist’s fee (which is very likely). Reliance on the private sector may discriminate against those without private health insurance (55% of all Australians) as colonoscopy waiting times are often longer in the public sector.

6) Quality of colonoscopy

The quality of the colonoscopy itself is determined by the training and experience of the colonoscopist and the standard of facilities (personnel and equipment) available. England has established high standards in certification and accreditation procedures, and the other UK countries have largely followed its lead. A recent report from the Australian NBSCP Quality Working Group has recommended the introduction of a national scheme for certification and re-certification of competence in colonoscopy – based on many features of the English model – but applying to all colonoscopy, not just screening. The ‘usual care’ model adopted in Scotland and most of Australia, can lead to difficulties in assessing the quality of screening colonoscopy, which may vary across regions.

7) Quality of data collection

The quality of data collection is closely related to the programme model. England, Wales and Northern Ireland, where clinical outcomes data is linked to a central register, are much better positioned than Scotland and Australia which rely on this information being forwarded to a central register. In Scotland, the NHS health boards collect clinical data on different IT systems and are responsible for feeding that data back to the central Information Services Division (ISD) within NHS Scotland. Although the health boards are required to deliver the data, they do always provide all the necessary data in the appropriate form.

In Australia, the situation is even less coordinated. The programmes in all states except Queensland rely on individual clinicians – GPs, colonoscopists and histopathologists – to return data to the central programme register run by Medicare Australia, and the rates of return, particularly by histopathologists, have been poor. England, despite having an efficient data collection system in place, and a programme underway since mid-2006, has not yet published any programme outcomes, although they are expected to be produced annually in the near future. England has made meaningful comparison of the relative performance of each programme impossible.
DISCUSSION

None of the programmes incorporated all of our proposed optimal features, with different programmes using different combinations of them. Insufficient funding has forced programme administrators to compromise on ideal implementation characteristics. England, with a much larger population, has opted for a less expensive, less accurate test, while Australia uses a more expensive, more accurate test, but only offers testing to three select age groups: 50, 55 and 65. It is clear that a trade-off between accuracy of test (and cost) and population coverage has been considered necessary in both settings. In contrast, Scotland is the only country to offer screening to the ideal age range of 50–74, as it has prioritized a philosophical commitment to age-based equity [UK03] over offering a more accurate (and expensive) test to a more restricted age range. These kind of trade-offs are necessary in all five countries because governments have baulked at the upfront costs of offering an optimal bowel cancer screening programme.

For example, the Australian government has made a continuing commitment to fund biennial breast and cervical cancer screening programmes for all women in the recommended age ranges (50–69 for breast cancer; 18–69 for cervical cancer). Bowel cancer kills more people each year than each of these cancers combined, and FOBT screening has consistently been reported to be highly cost-effective. Yet the Australian Government appears to be weighting short-term costs more heavily than the potential future cost savings and improved health outcomes.

Limitations of this study include the fact that the seven most salient programme characteristics and some of our proposed optimal features are based on our detailed examination of the different programmes and our considered opinion about what appears to work best. This has been necessary because of the paucity of clinical outcomes data available to support all of our assertions.

Further research could explore the potential efficiency gains of having one central funding source for all cancer screening, and redistributing funding according to relative cost-effectiveness.

CONCLUSION

This review raises the broader question of the role of evidence in policy and practice, by demonstrating the clear link between funding and evidence-based practice. All five bowel cancer screening programmes are being delivered in different, but potentially inefficient, ways. The irony is that by delivering programmes that do not optimize operational efficiency, governments may not be achieving the net health benefits of FOBT screening that justified the initial decision to fund them in the first place.

Acknowledgments

The authors wish to thank the key informants who generously gave their time to provide valuable insights into the implementation of these bowel cancer screening programmes, and the programme correspondents who so helpfully provided additional unpublished data.

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Ethics committee approval statement: Ethics approval for this research was granted by the Human Research Ethics Committee of the University of Sydney (Approval no. 05-2007/9971).
References


### Table 1

FOBT bowel cancer screening programmes as at January 2011

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Feature</th>
<th>England</th>
<th>Scotland</th>
<th>Wales</th>
<th>Northern Ireland</th>
<th>Australia</th>
</tr>
</thead>
<tbody>
<tr>
<td>1) Frequency</td>
<td>N/A</td>
<td>2 yearly</td>
<td>2 yearly</td>
<td>2 yearly</td>
<td>2 yearly</td>
<td>Once-only</td>
</tr>
<tr>
<td></td>
<td>2) Population coverage</td>
<td>60–74&lt;sup&gt;a&lt;/sup&gt;</td>
<td>50–74</td>
<td>60–71&lt;sup&gt;b&lt;/sup&gt;</td>
<td>60–69</td>
<td>50&lt;sup&gt;c&lt;/sup&gt;, 55, 65</td>
</tr>
<tr>
<td></td>
<td>Invited age range</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Number offered screening (% of population aged 50–74)</td>
<td>7,486,600 (54.64%)</td>
<td>1,471,800 (100%)</td>
<td>413,400 (47.55%)</td>
<td>168,800 (38.43%)</td>
<td>786,039 (13.96%)</td>
</tr>
<tr>
<td>Participation rates</td>
<td>Around 53% as at Dec 2010.</td>
<td></td>
<td>53% as at Dec 2010.</td>
<td>58% as at Dec 2010.</td>
<td>N/A. 60% is target.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>3) Quality of test</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Type of test</td>
<td>gFOBT: Hema Screen™ (Immunostics Inc. New Jersey USA).</td>
<td>gFOBT: Hema Screen™ Qualitative iFOBT: Hema-screen SPECIFIC (both manufactured by Immunostics Inc. New Jersey USA).</td>
<td>Same as Scotland.</td>
<td>Same as Scotland.</td>
<td>Quantitative iFOBT: Magstream HemSp with Magstream HT autoanalyser (Fujirebio Inc, Tokyo).</td>
</tr>
<tr>
<td></td>
<td>Screening algorithm</td>
<td>gFOBT&lt;sup&gt;d&lt;/sup&gt; if 'unclear'&lt;sup&gt;e&lt;/sup&gt;, repeat gFOBT if second test negative, repeat to make sure. If positive, colonoscopy.</td>
<td>gFOBT&lt;sup&gt;d&lt;/sup&gt; if 'weakly positive'&lt;sup&gt;e&lt;/sup&gt;, iFOBT&lt;sup&gt;f&lt;/sup&gt; if positive, colonoscopy.</td>
<td>same as Scotland.</td>
<td>same as Scotland.</td>
<td>iFOBT if positive, colonoscopy.</td>
</tr>
<tr>
<td></td>
<td>Test positivity rate</td>
<td>Around 1.8% as at December 2010</td>
<td>2.1%</td>
<td>2.6%</td>
<td>N/A</td>
<td>6.6%</td>
</tr>
<tr>
<td></td>
<td>Positive predictive value (PPV)</td>
<td>5.3% for cancer 32.1% for all neoplasia&lt;sup&gt;g&lt;/sup&gt; 10.6% for cancer 28.9% for high risk adenoma&lt;sup&gt;h&lt;/sup&gt;</td>
<td>8.6% for cancer 11.7% for cancer and high risk adenoma&lt;sup&gt;g&lt;/sup&gt;,&lt;sup&gt;h&lt;/sup&gt;</td>
<td>Similar to Scotland (same tests but will vary with prevalence)</td>
<td>Similar to Scotland (same tests but will vary with prevalence)</td>
<td>5.1% for cancer 20.1% for cancer and advanced adenomas&lt;sup&gt;g&lt;/sup&gt; Data for Queensland for Phase 1: 23.9% for cancer and advanced adenomas.&lt;sup&gt;g&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>Invitation register</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Funding</td>
<td>Single funder. Funded up to point of cancer diagnosis or return to screening.</td>
<td>Same as England.</td>
<td>Same as England.</td>
<td>Same as England.</td>
<td>Single funder up to point of positive test, and for register, data collection and reporting. Multiple funders for colonoscopy provision – federal</td>
</tr>
<tr>
<td>Characteristic</td>
<td>Feature</td>
<td>England</td>
<td>Scotland</td>
<td>Wales</td>
<td>Northern Ireland</td>
<td>Australia</td>
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<td>---------------</td>
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<td>-----------</td>
</tr>
<tr>
<td>Provider</td>
<td>Exclusive/designated provider model</td>
<td>Usual care</td>
<td>Same as England</td>
<td>Same as England</td>
<td>Varies between states and territories. See Table 3 in Appendix A.</td>
<td></td>
</tr>
<tr>
<td>5) Quality of follow up of positive test</td>
<td>Who by?</td>
<td>Face to face pre-assessment by local specialist screening nurse clinics.</td>
<td>Either face to face or telephone pre-assessment by NHS local health board nurses.</td>
<td>Telephone pre-assessment by local specialist screening practitioners.</td>
<td>Face to face pre-assessment by local specialist screening nurse clinics.</td>
<td>GP organizes referral to specialist.</td>
</tr>
<tr>
<td>Goal for waiting times</td>
<td>Within 28 days.</td>
<td>Within 31 days.</td>
<td>Same as England.</td>
<td>Same as England.</td>
<td>Within 30 days from GP referral to specialist.</td>
<td></td>
</tr>
<tr>
<td>6) Quality of colonoscopy</td>
<td>Certification of colonoscopist</td>
<td>Screening Assessment and Accreditation System, involving questionnaire and direct observation over two consecutive cases. Colonoscopists must be attached to a screening centre, have performed a minimum of 150 examinations in the previous year and have a documented completion rate of at least 90%.</td>
<td>Clinical standards developed by NHS Quality Improvement Scotland. Colonoscopist has demonstrated at least 90% completion in continuous audit and has undergone Joint Advisory Group (JAG) on Gastrointestinal Endoscopy-approved course. Desirable QIS criterion is for colonoscopist to have undertaken screening colonoscopy accreditation.</td>
<td>Similar certification procedures as England. Must be approved by Bowel Screening Wales before they can work in the screening programme.</td>
<td>Colonoscopists must undertake an approved competency-based training programme before they are able to do screening colonoscopies.</td>
<td>Recognition of training by the Conjoint Committee for the Recognition of Training in Gastrointestinal Endoscopy. Queensland has its own additional requirements for specialists wishing to become authorized providers, with colonoscopists required to apply for certification and meet minimum set criteria as determined by the QBCSP State Coordination Unit.</td>
</tr>
<tr>
<td></td>
<td>Accreditation of colonoscopy facility</td>
<td>National external accreditation by JAG using the web-based Global Bowel Rating Scale (GRS).</td>
<td>Screening colonoscopy is undertaken in a unit participating in the GRS.</td>
<td>Screening centres are inspected by a team from Bowel Screening Wales and must satisfy BSW criteria.</td>
<td>All screening units must be externally accredited by JAG.</td>
<td>No uniform licensing arrangements exist across jurisdictions.</td>
</tr>
<tr>
<td>7) Quality of data collection</td>
<td>Return of data</td>
<td>All 58 screening centres and 5 hubs are linked to the one national database.</td>
<td>14 local health boards responsible for return of data to central database. Have experienced problems with getting suitable data from local health boards.</td>
<td>Data from all screening centres (11 currently, another expected soon, and one in England for border referrals) collected centrally through the Bowel Screening Information Management System (BSIMS).</td>
<td>Data from all 5 screening colonoscopy centres is collected centrally through the BSIMS (developed by Wales, but also used by Northern Ireland).</td>
<td>GPs, colonoscopists and histopathologists are requested to return data to the central Medicare register. Have experienced problems with getting data from individual practitioners.</td>
</tr>
<tr>
<td>Characteristic</td>
<td>Feature</td>
<td>England</td>
<td>Scotland</td>
<td>Wales</td>
<td>Northern Ireland</td>
<td>Australia</td>
</tr>
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<td>---------------------</td>
<td>-------------------------------------------------------------------------</td>
<td>-------------------------------------------------------------------------</td>
<td>-------------------------------------------------------------------------</td>
<td>------------------------------------------------</td>
<td>------------------</td>
<td>-----------------------------------------------------------</td>
</tr>
<tr>
<td>Reporting of data</td>
<td>No published reports as yet, but plan to produce annual reports.</td>
<td>Annual reports of key performance indicators prepared by Information Services Division and available on website.</td>
<td>Monthly internal reporting only at this stage.</td>
<td>Same as England.</td>
<td>The Australian Institute of Health and Welfare has produced four reports since April 2008.</td>
<td></td>
</tr>
</tbody>
</table>

\[\text{\textsuperscript{a}}\text{Extended from 60–69 in January 2010. Plan to cover 60–74 by 2014.}\]

\[\text{\textsuperscript{b}}\text{Extended from 60–69 in late 2010. Plan to extend age range to 70–74 by 2015.}\]

\[\text{\textsuperscript{c}}\text{Added to programme in 2008.}\]

\[\text{\textsuperscript{d}}\text{Non-hydrated guaiac tests used with no dietary restrictions (despite evidence that certain foods have been shown to increase the likelihood of a false positive test).}\]

\[\text{\textsuperscript{e}}\text{'Unclear' and 'weakly positive' are defined as 1–4 positive windows out of 6.}\]

\[\text{\textsuperscript{f}}\text{Scotland is piloting the quantitative iFOBT OC Sensor (Eikin Chemical Co., Ltd, Japan) as the only test.}\]

\[\text{\textsuperscript{g}}\text{England, Scotland and Australia use different terminology to report PPV for lesions other than cancer, so direct comparison is not possible. Scotland reports PPV for “the current screening test” which refers to the whole test algorithm (ie it includes the guaiac and qualitative immunochromatic tests).}\]

\[\text{\textsuperscript{h}}\text{Updated figures for August 2006–August 2009. Personal communication, Dr Thomas Lee.}\]

\[\text{\textsuperscript{i}}\text{Only screening-identified assessment or surveillance colonoscopies performed at these centres.}\]

\[\text{\textsuperscript{j}}\text{Screening and non-screening colonoscopies performed at these centres.}\]
Table 2
Trade-offs involved in FOBT cancer screening programmes with inadequate funding

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Features</th>
<th>More funding/resources</th>
<th>Less funding/resources</th>
<th>Optimal features</th>
</tr>
</thead>
<tbody>
<tr>
<td>Frequency</td>
<td>N/A</td>
<td>More often</td>
<td>Less often</td>
<td>At least biennial(^d)</td>
</tr>
<tr>
<td>Population coverage</td>
<td>Invited age range</td>
<td>More people in the</td>
<td>Less people in the</td>
<td>50 to 74 or 75 year olds(^c)</td>
</tr>
<tr>
<td></td>
<td>appropriate age range</td>
<td>appropriate age range</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Participation rates</td>
<td>Maximizes for all people, including population sub-groups(^a) to ensure equity of access</td>
<td>Maximize for all people,</td>
<td>Aim for 60% but may be a one-size fits all programme</td>
<td>Minimum of 60% of the unscreened population(^b), maximising equity of access for population sub-groups(^c)</td>
</tr>
<tr>
<td>Speed of implementation</td>
<td>Faster</td>
<td>Faster</td>
<td>Slower</td>
<td>As fast as possible, without decreasing quality and increasing waiting times(^d)</td>
</tr>
<tr>
<td>Quality of test</td>
<td>Type of test</td>
<td>Most accurate</td>
<td>Most affordable</td>
<td>Quantitative immunochemical FOBT(^16)–(^22)</td>
</tr>
<tr>
<td>Test positivity rate</td>
<td>Higher</td>
<td>Higher</td>
<td>Lower</td>
<td>Between 5–8% to minimize missed cancers and advanced adenomas(^d)</td>
</tr>
<tr>
<td>Positive predictive value</td>
<td>Higher</td>
<td>Higher</td>
<td>Lower</td>
<td>The combined PPVs for cancer and advanced adenomas(^d)</td>
</tr>
<tr>
<td>Screening algorithm</td>
<td>Minimizes re-testing and inconvenience</td>
<td>Minimizes re-testing and inconvenience</td>
<td>May involve up to three FOBTs</td>
<td>FOBT with actionable result after one test (positive or negative), and no dietary or medication restrictions(^d)</td>
</tr>
<tr>
<td>Programme model</td>
<td>Invitation register</td>
<td>Able to exclude in advance those not suitable for FOBT screening</td>
<td>General invitation to all in the target age group</td>
<td>Utilizes existing medical information in targeting invitees(^d)</td>
</tr>
<tr>
<td>Funding</td>
<td>Single funder</td>
<td>Single funder</td>
<td>Multiple funders</td>
<td>Single funder to provide adequate levels of funding to maximize benefits and minimize harms, while avoiding cost-shifting(^d)</td>
</tr>
<tr>
<td>Provider</td>
<td>Designated provider</td>
<td>Usual care(^d)</td>
<td></td>
<td>Designated providers to set minimum quality standards and mandate return of data to register(^d)</td>
</tr>
<tr>
<td>Quality of follow-up of</td>
<td>Who by?</td>
<td>Staff with bowel cancer screening knowledge and expertise</td>
<td>Usual care</td>
<td>Staff with specific expertise to provide tailored advice on benefits and harms and procedural matters, increase adherence to follow-up and ensure safety</td>
</tr>
<tr>
<td>positive test</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Characteristic</td>
<td>Features</td>
<td>More funding/resources</td>
<td>Less funding/resources</td>
<td>Optimal features</td>
</tr>
<tr>
<td>---------------------</td>
<td>-----------------------------------------------</td>
<td>-------------------------</td>
<td>------------------------</td>
<td>----------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Waiting times</td>
<td>Shorter</td>
<td>Longer</td>
<td></td>
<td>by assessing pre-colonoscopy risk&lt;sup&gt;d&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Within 28 days from positive test to colonoscopy&lt;sup&gt;d&lt;/sup&gt;</td>
</tr>
<tr>
<td>Quality of colonoscopy</td>
<td>Certification of colonoscopists</td>
<td>National certification with minimal standards for training</td>
<td>No minimum standards for training</td>
<td>Specialized training to ensure consistency across regions and maximize quality and safety&lt;sup&gt;d&lt;/sup&gt;</td>
</tr>
<tr>
<td>Accreditation of colonoscopy facility</td>
<td>National accreditation with minimal standards for facilities</td>
<td>No minimum standards for facilities</td>
<td></td>
<td>National accreditation standards to ensure consistency across regions and maximize quality and safety&lt;sup&gt;d&lt;/sup&gt;</td>
</tr>
<tr>
<td>Quality of data collection</td>
<td>Return of data</td>
<td>Specific IT systems created for the bowel cancer screening programmes</td>
<td>Usual IT systems</td>
<td>Specialized IT systems to maximize quantity and quality of data returned to register&lt;sup&gt;d&lt;/sup&gt;</td>
</tr>
<tr>
<td>Reporting of data</td>
<td>Regular internal and external reporting</td>
<td>Some internal but less external reporting</td>
<td></td>
<td>At least quarterly internal and annual external reporting to allow for continuing programme evaluation&lt;sup&gt;d&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

<sup>a</sup>Irrespective of age, ethnicity, income or geographical location.

<sup>b</sup>The idea of setting participation ‘targets’ is contentious as it may lead to people being encouraged to be screened, rather than them making their own genuinely informed choice about whether to be screened or not. Nevertheless, participation rates are used as a de facto measure of a screening program’s success. A participation rate close to 60% was achieved in the Nottingham RCT<sup>23</sup> and this has become an unofficial ‘acceptable’ or target rate. The NHS Quality Improvement Scotland (QIS) standard for bowel screening uptake is 60%.<sup>13</sup> This figure was also used by English researchers to model potential population-based bowel cancer screening programmes<sup>24</sup> and representatives from all four UK screening programmes were keen to establish participation rates at around this level or higher [UK02, UK05, UK11, UK13], although there is evidence that high participation in bowel cancer screening programmes is not necessary to achieve cost-effectiveness.<sup>25</sup> In Australia, where there is a large amount of opportunistic bowel cancer screening outside the programme (mainly via colonoscopy but also through some independent FOBT programmes such as the one run by Rotary) a participation rate of 60% may be more unrealistic.

<sup>c</sup>Participation in both the UK<sup>5</sup> and Australian<sup>6</sup> pilots varied according to population sub-groups, and this was identified as an issue that needed to be addressed.

<sup>d</sup>In our considered opinion. These ideal features are based on what seems to work best and/or appears to have the best clinical outcomes, in the absence of definitive evidence.

<sup>e</sup>The usual care model, while less expensive than a designated provider model, may also be chosen on equity, rather than cost grounds. It ensures all people requiring colonoscopy, whether screen-detected or not, have access to the same standard of care.

<sup>f</sup>This is the ideal maximum waiting time specified in the English, Welsh and Northern Irish programmes. It is not achieved in all areas of these countries.
## APPENDIX A

### Programme models within Australia

<table>
<thead>
<tr>
<th>Jurisdiction</th>
<th>Model</th>
<th>Additional state/territory funding?</th>
<th>Features</th>
</tr>
</thead>
</table>
| New South Wales [A40] | Usual care           | No                                 | • Participants referred back to GP to organize follow-up of positive FOBT.  
• Participants can be referred to either the public sector or private sector for their assessment colonoscopy.  
• NSW Health is paying for a new Endoscopy Information System, using the same software as Queensland, which will allow improved reporting to the national register. |
| Queensland[^1] [A35] | Designated provider | Yes – $10.4 million (~£6.7 million) over 3 years in 2006 to develop its own programme (the Queensland Bowel Cancer Screening Program). | • Offers an integrated screening pathway for public patients which commences at FOBT invitation and concludes at the point of histopathological diagnosis or re-screening.  
• 28 designated facilities that offer high quality and timely assessment colonoscopies.  
• Facilities selected for their capacity to provide additional colonoscopy services.  
• Some public patients may be offered colonoscopy in a private facility in areas where public waiting lists are long.  
• Both facilities and proceduralists required to demonstrate compliance with state and international quality requirements (not national) as part of the authorization process.  
• Facilities are paid $1,240 (~£800) per colonoscopy – includes all costs (proceduralist, sedationist, and histopathology).  
• Established a State Coordination Unit, a network of 12 Gastroenterology Nurse Coordinators and 11 Health Promotion Officers.  
• Captured 100% of data at designated centres.  
• Piloting an alternative delivery model in some remote Indigenous communities where kits are distributed through Indigenous health workers/health centres.  
• From July 2009, implementation of the Endoscopy Services Information System Solution (ESISS) will significantly enhance the integrity and timelines of clinical data. |
| South Australia [A39] | Mixed                | Yes – they pay the greater share of costs for pathway coordinators (the rest paid for by the Commonwealth). | • Five nurse pathway coordinators located at five public hospitals who conduct pre-colonoscopy assessments.  
• Flinders Medical Centre offers one dedicated NBCSP list per week. |
<p>| Tasmania [A38]      | Usual care           | No                                 | • As for NSW.                                                                                                                                 |
| Victoria [A37]      | Mixed                | Yes – $14 million (~£9 million) over four years allocated in 2007. | • Usual care but with some designated providers (19 as at February 2011) for public patients. |</p>
<table>
<thead>
<tr>
<th>Jurisdiction</th>
<th>Model</th>
<th>Additional state/territory funding?</th>
<th>Features</th>
</tr>
</thead>
<tbody>
<tr>
<td>Western Australia [A36]</td>
<td>Usual care</td>
<td>No</td>
<td>• Designated providers are selected for their ability to provide colonoscopy capacity to support the programme.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Participants can still go to non-designated public and private providers.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• As for NSW.</td>
</tr>
<tr>
<td>Australian Capital Territory (ACT) [A41]</td>
<td>Mixed</td>
<td>No</td>
<td>• One hospital (The Canberra Hospital) offers a clinic for NBCSP participants, where they are given priority booking for colonoscopy.</td>
</tr>
<tr>
<td>Northern Territory (NT) [A42]</td>
<td>Opportunistic screening⁴</td>
<td>No</td>
<td>• Kits not sent directly to participants.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• When people aged 45 or over present to a health clinic, they are offered an adult health check which includes an FOBT.</td>
</tr>
</tbody>
</table>

*Usual care is not considered appropriate for the NT because it has a high proportion of Indigenous people (36.1%) ³⁹, the vast majority (81%) of whom live in remote or very remote areas. ⁴⁰ This raises both practical and cultural barriers. The practical issues include the absence of postal addresses in remote communities, where all mail is sent to the health clinic or community council. The kits themselves are often addressed incorrectly, as Aboriginal people may have several names over their lifetime. Moreover, most patients in the NT have several co-morbidities such as diabetes, renal failure and chronic obstructive pulmonary disease, and require more targeted invitational approaches. The cultural issues include the fact that faeces is considered shameful in Aboriginal culture, so clinical staff need to be involved in explaining the collection process, helping with the collection and returning the kits to the pathology lab [A42].