1	Colonoscopic surveillance for prevention
2	of colorectal cancer in patients with
3	ulcerative colitis, Crohn's disease or
4	adenomas
5	
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7	Part 2
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Appendix 7 – Health economic evaluation Cost-effectiveness analysis for inflammatory bowel disease

4 **1** Introduction

5 The Department of Health asked NICE to produce a short clinical guideline on

6 colonoscopic surveillance for the prevention of colorectal cancer in patients

7 with ulcerative colitis, Crohn's disease and polyps.

- 8 A cost-effectiveness analysis has been carried out to support the Guideline
- 9 Development Group (GDG) in making recommendations for adults with
- 10 inflammatory bowel disease considered to be at high risk of developing
- 11 colorectal cancer. This analysis has been conducted according to the
- 12 methods outlined in the NICE Guide to the methods of technology appraisal
- 13 2008 and the Guidelines Manual 2009. The methods used follow the NICE
- 14 reference case, which is the framework NICE request all cost-effectiveness
- 15 analyses to follow.
- 16 Given the quality of the data available this analysis should be considered an
- 17 exploration of the cost effectiveness of colonoscopic surveillance for
- 18 inflammatory bowel disease.

192Acknowledgements

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- 22 the development of this guideline by providing the uplifted cost data for stage-
- 23 specific colorectal cancer.

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1 4 Decision problem

- 2 Table 1 outlines the decision problem that is addressed in this guideline and is
- 3 based on the final scope.

4 **Table 1 Decision problem**

	Scope	Approach taken	
Population	People with inflammatory bowel disease (IBD: ulcerative colitis or Crohn's disease)	People considered to be at high risk of colorectal cancer with flat dysplastic lesions (low grade or high grade), age 30 to 85 years	
Interventions	Colonoscopic surveillance using colonoscopy, chromoscopy, computerised tomography colonoscopy, narrow band imaging, double-barium contrast enema	Colonoscopic surveillance using colonoscopy	
Comparators	No colonoscopic surveillance	No colonoscopic surveillance	
Outcome(s)	Costs, quality-adjusted life years (QALYs) and cost per QALY gained	Cost per QALY gained	

5

6 **4.1 Population**

7 Ulcerative colitis and Crohn's disease are collectively termed inflammatory

8 bowel disease (IBD). People with these conditions share the same risk of

9 developing colorectal cancer given a similar extent and duration of disease.

10 For the economic evaluation both conditions have therefore been grouped

11 together.

12 Based on the data available at the time of guideline development, a model

13 was developed assuming that surveillance intervals would depend on the

14 degree of dysplasia (because dysplasia is a precancerous marker for

15 colorectal cancer). The model simulated men and women aged 30–85 years

16 with flat dysplastic lesions (that is, non-resectable low- or high-grade

17 dysplasia) who had declined surgery. However, at the final meeting of the

- 18 GDG, it was decided that the surveillance intervals should depend on a
- 19 person's risk of developing colorectal cancer and the IBD surveillance

- 1 schedule was stratified accordingly. The GDG identified three groups for
- 2 surveillance: people at low risk, intermediate risk and high risk of developing
- 3 colorectal cancer

4 Because of the tight timelines between the final GDG meeting and

- 5 consultation, the cost effectiveness of surveillance based on the dysplasia
- 6 model was determined only for the high-risk group. People at high risk (as
- 7 defined by the GDG), include people with a previous history of primary
- 8 sclerosing cholangitis, ongoing moderate or severe active inflammation,
- 9 dysplasia or colonic strictures, or a family history of colorectal cancer in a first-
- 10 degree relative aged under 50 years. For more details please see the main
- 11 guideline.

12 The choice of 30 years as the starting age in the model was based on the

13 British Society of Gastroenterology (BSG) guidelines for IBD (British Society

14 of Gastroenterology 2004), which reported that both ulcerative colitis and

15 Crohn's disease affect young people and have a peak incidence between the

ages of 10 and 40 years in the UK. The GDG members agreed with this.

17 4.2 Interventions

To demonstrate that surveillance is beneficial for people with IBD, a reduction in mortality caused by colorectal cancer in people with IBD having surveillance would have to be shown in clinical studies. Because colonoscopic surveillance was found to reduce mortality from colorectal cancer for people with IBD, the intervention used in the model was colonoscopy. It was assumed that surveillance colonoscopy should be performed when colonic disease is in remission (as recommended in the updated BSG 2010 guidelines for IBD).

25 4.3 Comparators

Surveillance is not consistently offered across the NHS. Therefore 'no surveillance' was considered as the comparator for surveillance. The GDG pointed out that some people are offered surgery (colectomy) depending on their degree of dysplasia. Although managing dysplasia with surgery was not considered in this model because no evidence was reviewed, surgery upon, colorectal cancer detection has been factored into the mean lifetime cost of

- 1 colorectal cancer treatment (section 8.1.1.2). For simplicity, it was assumed
- 2 that all people who enter the model have confirmed dysplasia (either low or
- 3 high grade) at baseline colonoscopy and have declined surgery. The
- 4 surveillance schedule proposed by the GDG is consistent with the BSG 2010
- 5 guidelines:
- 6 Low risk surveillance every 5 years
- 7 Intermediate risk surveillance every 3 years
- 8 High risk surveillance every year.

9 4.4 Outcomes

10 In line with the NICE reference case a cost–utility analysis was used to

- 11 analyse the cost effectiveness of colonoscopic surveillance for people with
- 12 non-resectable dysplastic lesions who are considered to be at high risk of
- 13 developing colorectal cancer and require surveillance every year. This
- 14 required the calculation of resource use and quality-adjusted life years
- 15 (QALYs) to assess effectiveness.

5 Review of existing cost-effectiveness analyses

17 **5.1** Search for cost-effectiveness analyses

18 A search for cost-effectiveness studies did not identify any relevant papers 19 that examined colonoscopic surveillance for the prevention of colorectal cancer in people with IBD. However, during the search, three studies were 20 21 identified (Nguyen et al. 2009; Provenzale et al. 1995; Delco et al. 2000) that 22 examined colorectal cancer surveillance using colonoscopy for people with 23 ulcerative colitis. Two of the studies (Nguyen et al. and Provenzale et al.) 24 compared surveillance with surgery. All three studies explored approaches to 25 modelling strategies such as decision tree versus Markov models, and when applicable, informed the model structure. Given the absence of any 26 appropriate analysis that addressed the decision problem directly, a new cost-27 effectiveness model was developed based on the views of the GDG and 28

29 clinical data available at the time of guideline development.

1 5.2 Modelling approach

IBD is a chronic condition; a Markov model appeared to be most appropriate
to answer the decision problem.

The Markov model split the single state of dysplasia into two mutually
exclusive states of low-grade and high-grade dysplasia. Similarly, the single
colorectal cancer state was broken down into four mutually exclusive states of

7 Dukes' A, Dukes' B, Dukes' C and Dukes' D colorectal cancer.

8 The model started at age 30. It was assumed that the person had symptoms

9 of colitis for at least 10 years (that is, symptoms began on average at age 20),

10 had a screening colonoscopy that identified dysplasia, and subsequently

11 entered a surveillance programme. A cycle length of 3 months seemed most

12 appropriate, because surveillance for the high-risk group occurs every year

13 and this cycle length was long enough to allow the possible development of

14 asymptomatic and symptomatic cancer between colonoscopies.

15 The analysis was run over a 55-year time horizon, until age 85, and examined

16 the use of colonoscopy for surveillance compared with no surveillance for

17 people at high risk of developing colorectal cancer (section 4.1).

18 **5.3** *Natural history review*

19 A major component of the IBD model is the natural history of dysplasia,

20 because dysplasia is used as a precancerous marker of colorectal cancer risk.

21 Because of the lack of resources and time, a full systematic review of the data

22 on the natural history of dysplasia to calculate transition probabilities was not

23 possible. Therefore, a clinical study that reported the 30-year follow-up of a

24 UK colonoscopic surveillance programme for neoplasia in ulcerative colitis

25 (Rutter et al. 2006) was used to calculate the progression of low- and high-

26 grade dysplasia to colorectal cancer using a Bayesian dirichlet method. The

27 Bayesian approach was needed to calculate unobserved transitions. Further

28 details are provided in the transition probability section (section 6.2).

29 Data on the natural history of colorectal cancer were also obtained from a

30 published cost-effectiveness study by Tappenden et al. (2004) that

- 1 systematically reviewed cost-effectiveness studies for colorectal cancer
- 2 screening in the UK. Colorectal cancer transition probabilities (that is,
- 3 progression to symptomatic and/or asymptomatic colorectal cancer and
- 4 cancer-related mortality) were obtained from this study and followed the
- 5 Bayesian approach.

6 6 Model

7 6.1 Model structure

- 8 Figure 1 shows the basic outline of the surveillance model for the high-risk
- 9 group.

10 Figure 1 Colonoscopic surveillance model for people with IBD in the 11 high-risk group



12

- 13 LGD: low-grade dysplasia; HGD: high-grade dysplasia; DA: Dukes' A; DB: Dukes' B; DC:
- 14 Dukes' C; DD: Dukes' D; CRC: colorectal cancer
- 15

166.1.1Surveillance and natural history

- 17 Colonoscopic surveillance is recommended every year in the high-risk group
- 18 (every fourth cycle in the model) and it was assumed that colonoscopy was
- 19 carried out at the beginning of the scheduled cycle. In the model, the

1 development of colorectal cancer could be sequential, that is, progress from 2 low-grade to high-grade dysplasia to cancer; or from low-grade dysplasia 3 directly to colorectal cancer because some people do not progress through a 4 detectable phase of high-grade dysplasia. People with high-grade dysplasia 5 could also progress directly to colorectal cancer and were assumed not to 6 regress to low-grade dysplasia. Progression to colorectal cancer could occur 7 either asymptomatically or symptomatically between the scheduled 8 surveillance colonoscopies. Over time, if people had no evidence of 9 progression they would remain in the same state. Any other cause of mortality 10 was also considered in all states in the model.

11 6.1.2 Colorectal cancer

12 Cancer states were stratified by tumour stage at diagnosis using Dukes' 13 staging. If a person developed Dukes' A colorectal cancer, they could either 14 continue to progress to a higher Dukes' stage or stay in the same state. 15 According to the literature, colorectal cancer mortality occurs only at Dukes' 16 stages B, C and D, so it was applied only to each of these states in the model. 17 After cancer was diagnosed, the person was assumed to enter a cancer 18 management programme; that is, the person receives chemotherapy, surgery 19 and/or radiotherapy. All the cancer states were allocated both costs and utility 20 values.

21 6.1.3 Complications

22 The model assumed there were no complications from colonoscopy during the

23 55 years of surveillance. Although perforation and bleeding are serious risks

of colonoscopy, they occur infrequently and were assumed to be negligible.

Likewise, the cost-effectiveness study by Nguyen et al. (2009) that included

colectomy as a comparator to enhanced surveillance assumed that acute

27 complications from colonoscopy and colectomy were negligible.

28 6.1.4 Compliance

29 It was assumed that everyone participating in the surveillance programme

- 30 adhered to the colonoscopic surveillance protocol. This seemed reasonable,
- 31 because people are more likely to adhere to a programme when they are

informed that they have a high risk of developing colorectal cancer. The study
 by Rutter et al. (2006) reported a long-term compliance rate for surveillance of
 94.3%.

4 6.2 Transition probabilities

5 Two sets of transition probabilities were included in the model, for the natural 6 history of dysplasia and for colorectal cancer.

7 The probabilities derived from the observational study by Rutter et al. (2006)

8 were chosen because the study followed a UK population for 30 years of

9 colonoscopic surveillance. The study reported the first and maximal neoplasia

10 as required by the cost-effectiveness model. The cancer outcomes were also

11 reported as Dukes' staging and the study was included in the clinical-

12 effectiveness data for this guideline. Therefore, it was considered appropriate

13 to use this study as the basis to calculate transition probabilities for the natural

14 history of dysplasia. It was assumed that having a colonoscopy does not alter

15 the risk of colorectal cancer because for people with non-resectable dysplastic

16 lesions, colonoscopy would be used as a diagnostic tool rather than as an

17 interventional procedure.

18 The transition probabilities for the natural history of colorectal cancer were

19 taken from Tappenden et al. (2004) and were used in conjunction with the

20 transition probabilities for neoplasia calculated by Rutter et al. (2006) using a

21 Bayesian dirichlet method. This method permits the probabilities to be

22 calculated for unobserved transitions.

23 Age-related mortality rates were assumed for low- and high-grade dysplasia 24 and the asymptomatic cancer states. It seemed reasonable to assume that 25 people in the asymptomatic cancer states have the same probability of dying as people in the general population at the same age because they are unlikely 26 27 to have an increased risk of death until their cancer progresses. Annual 28 colorectal cancer-related mortality was taken from Tappenden et al. (2004) 29 and was used for all symptomatic cancer states. Age-related mortality was 30 applied in addition to colorectal cancer mortality for all symptomatic cancer 31 states.

- 1 Data from published interim life tables for the UK (Office of National Statistics,
- 2 2009) were used to produce age-related mortality probabilities. Because these
- 3 probabilities vary with time they were subtracted from the probabilities of
- 4 staying in the same health state. This ensured that all probabilities summed to
- 5 one.
- 6 To convert the 30-year observational data from Rutter et al. (2006) into a
- 7 yearly cycle length, the following formula was used where p is the yearly
- 8 probability (Briggs et al. 2003):

9 yearly probability = $1 - e^{((\ln 1 - P) \cdot (1/30))}$

10 The transition matrix for natural history is presented in table 2.

11 Table 2 Natural history transition matrix (yearly)

12

	LGD	HGD	DA	DB	DC	DD	mCRC	mOther	
LGD	#	0.0095	0.0050	0.0000	0.0000	0.0000	0.0000	Age	
HGD	0.0000	#	0.0037	0.0000	0.0000	0.0000	0.0000	Age	
DA	0.0000	0.0000	#	0.5830	0.0228	0.0029	0.0000	Age	
DB	0.0000	0.0000	0.0000	#	0.6560	0.0000	0.0100	Age	
DC	0.0000	0.0000	0.0000	0.0000	#	0.8650	0.0600	Age	
DD	0.0000	0.0000	0.0000	0.0000	0.0000	#	0.3870	Age	
# 1 minus other states; LGD: low-grade dysplasia; HGD: high-grade dysplasia; DA: Dukes' A; DB: Dukes' B; DC: Dukes' C; DD: Dukes' D; mCRC: colorectal cancer mortality; mOther: other cause mortality									
I he gre	y shaded a	areas repre	esent annu	ual transiti	ons, availa	able from	appender	i et al.	

- 14 The method used to calculate unobserved events is also the preferred method
- 15 of incorporating uncertainty into a Markov model with several states, using the
- 16 dirichlet distribution in a Bayesian framework.
- 17 The dirichlet distribution is a multinomial equivalent of the beta distribution (a
- 18 probability distribution that is bounded by 0 and 1). This allows distributions to
- 19 be placed on a parameter while maintaining the axiom of probabilities
- 20 (summing to one).

- 1 The Bayesian approach allows calculation of a probability based on
- 2 understanding the probability distribution of an event and on any prior
- 3 information. These two parts are called the posterior and the prior.
- 4 In this case prior beliefs can be included for transitions for which there are no
- 5 observed data but which can occur. Therefore, for transitions where a
- 6 transition probability was needed, uninformative priors were used, which
- 7 allowed these transitions to be calculated. For more details on the method
- 8 please see Briggs et al. (2003).
- 9 The chosen priors are presented in table 3.

	LGD	HGD	DA	DB	DC	DD	mCRC	mOther	
LGD	0.12	0.12	0.12	0	0	0	0	Age	
HGD	0	0.12	0.12	0	0	0	0	Age	
DA	0	0	0.12	0.12	0.12	0.12	0	Age	
DB	0	0	0	0.12	0.12	0.12	0.12	Age	
DC	0	0	0	0	0.12	0.12	0.12	Age	
DD	0	0	0	0	0	0.12	0.12	Age	
LGD: low-grade dysplasia; HGD: high-grade dysplasia; DA: Dukes' A; DB: Dukes' B; DC: Dukes' C; DD: Dukes' D; mCRC: colorectal cancer mortality; mOther: other cause mortality									

Table 3 Priors for the natural history transition matrix

- 12 A value of 0.12 was chosen for the uninformative priors because of a
- 13 calculating error in Excel (the small numbers involved resulted in num! errors)
- 14 which meant smaller priors were not possible. This was resolved by
- 15 increasing the size of the observed data by multiplying them by 1000 to
- 16 maintain the relative difference between the priors and the observed data. All
- 17 the transitions that were expected to occur within the model were given the
- 18 same prior value (0.12) so that each data set (Rutter et al. 2006 and
- 19 Tappenden et al 2004) contributed equally to the model.
- 20 Calculating the probabilities from Rutter et al. (2006) and the dirichlet
- 21 framework, the following transition matrices for natural history (table 4) were
- used. These represent the 3-monthly (or quarter of a year) transitions used in
- the model.

	LGD	HGD	DA	DB	DC	DD	mCRC	mOther			
LGD	0.99466	0.00354	0.00180	0.00000	0.00000	0.00000	0.00000	Age			
HGD	0.00000	0.99759	0.00241	0.00000	0.00000	0.00000	0.00000	Age			
DA	0.00000	0.00000	0.85793	0.13559	0.00572	0.00075	0.00000	Age			
								-			
DB	0.00000	0.00000	0.00000	0.84623	0.15122	0.00003	0.00253	Age			
								-			
DC	0.00000	0.00000	0.00000	0.00000	0.79066	0.19443	0.01491	Age			
								-			
DD	0.00000	0.00000	0.00000	0.00000	0.00000	0.90778	0.09222	Age			
								-			
LGD: lo	w-grade d	vsplasia: H	GD: hiah-c	rade dvsp	asia: DA: I	Dukes' A: D	B: Dukes'	B: DC:			
Dukes'	Dukes' C: DD: Dukes' D: mCRC: colorectal cancer mortality: mOther: other cause mortality										

1 Table 4 Final natural history transition matrix (every 3 months)

2 7 Quality of life

3 NICE recommends that changes in HRQoL as a result of an intervention or

- 4 treatment should be directly reported by patients. These changes should be
- 5 based on preferences determined using a choice-based method in a
- 6 representative sample of the UK general public. Ideally a full systematic
- 7 review would be carried out to identify health-related quality of life (HRQoL)
- 8 studies and appropriate values to include in a health economic model.
- 9 However, because of the lack of resources and time a search was carried out
- 10 for quality of life studies. The cost-effectiveness studies that were used to
- 11 explore approaches to modelling strategies were also searched for QALY

12 data.

13 **7.1** *Literature search*

- 14 The search identified one paper, a study by Gregor et al. (1997) that
- 15 examined quality of life in patients with Crohn's disease. The study reported
- 16 utility values by disease severity calculated using the time-trade-off method.
- 17 Several studies reported values obtained from a disease-specific
- 18 questionnaire (the Inflammatory Bowel Disease Questionnaire). However,
- 19 these values could not be used for calculating QALYs because they did not
- 20 report the values on a 0–1 scale, which is the format for generic
- 21 questionnaires.

1 7.2 People's health states

NICE recommends the use of the EuroQol 5 dimensions (EQ-5D) or another generic tool that enables patients to describe their health state and how the public values their health state. Although Gregor et al. (1997) reported utility values using a generic tool; the study was not in complete accordance with NICE methods. The values obtained in the study were collected from people with Crohn's disease who were asked to value health states that described their disease severity, specifically mild, moderate and severe Crohn's disease.

9 7.3 Low- and high-grade dysplasia

10 The GDG agreed that the values obtained from Gregor et al. (1997) could be 11 used to represent the utility values for people with low- and high-grade 12 dysplasia. The utility value for mild Crohn's disease was used as a proxy for 13 low-grade dysplasia and the utility value for moderate Crohn's disease was 14 used as a proxy for high-grade dysplasia. This approach seemed acceptable 15 because the patient experts on the GDG felt that a person with low-grade 16 dysplasia has a lower quality of life than a person in the general population 17 and a person with high-grade dysplasia has a lower quality of life than a 18 person with low-grade dysplasia.

19 **7.4 Cancer**

Stage-specific utility values for symptomatic colorectal cancer were obtained from Ness et al. (1999) and were applied to each symptomatic Dukes' state. Asymptomatic cancers were assigned the same utility value as their diagnostic state because if cancer is asymptomatic it is unlikely to affect the quality of life of the person until it is detected (that is, until it becomes symptomatic).

26 **7.5** Age-related quality of life

For all the health states in the model the specific health state utility values were multiplied by the age-related utility values. Age-related utility values for the UK population were available from Kinder et al. (1999). This approach was taken because it was assumed that as a person ages their quality of life

- 1 steadily decreases and if the same person has a condition that affects their
- 2 life, this multiplies the effect.

3 7.6 Final quality of life values

4 Table 5 Final health-related quality of life estimates

Health state	Mean value	Standard error	Reference
All health states	Age dependent	N/A	Kinder et al. (1999)
LGD (mild Crohn's disease)	0.95	0.008014	Gregor et al. (1997)
HGD (moderate Crohn's			Gregor et al. (1997)
disease)	0.88	0.014416	
Dukes' A	0.74	0.031276	Ness et al. (1999)
Dukes' B	0.7	0.051192	Ness et al. (1999)
Dukes' C	0.5	0.061521	Ness et al. (1999)
Dukes' D	0.25	0.206870	Ness et al. (1999)
LGD: low-grade dysplasia; HGD: high	h-grade dysplasia	a	

5

6 Uncertainty about utility values that were not time dependent was captured

7 using a lognormal distribution.

8 8 Resource use

9 8.1 Literature search

- 10 The initial search identified three studies (Hanauer et al. 1998; Stark et al.
- 11 2006; Bodger et al. 2002) that examined resource use for people with IBD.
- 12 The study by Hanauer et al. was excluded because it reported the cost of
- 13 Crohn's disease from a US perspective. The study by Stark et al. was
- 14 excluded because it reported the cost of IBD from a German perspective.
- 15 Bodger et al. was the only UK study looking at the cost of Crohn's disease in
- 16 one hospital. However, the study did not include a breakdown of the costs,
- 17 which were reported in US dollars, as required by the model.
- 18 Only one study provided information on the lifetime costs of colorectal cancer
- 19 in the UK by Dukes' staging (Tappenden et al. 2004).

2 8.1.1 Specific costs for the model

3 The main cost inputs that required consideration included:

- colonoscopy (procedure and biopsy specimens)
 - cancer (diagnosis, treatment and follow-up).
- 6

5

7 8.1.1.1 Colonoscopy

8 The cost of colonoscopy was obtained from a GDG member and was

9 validated using NHS reference costs 2008/09.

10 8.1.1.2 Cancer

- 11 The estimated mean lifetime costs associated with the diagnosis, treatment
- 12 (chemotherapy, radiotherapy, surgery) and follow-up of colorectal cancer were
- 13 reported in the study by Tappenden et al. (2004). The 2004 costs were
- 14 updated to 2010 costs by the lead author of the study and are listed in table 6.
- 15 These were only applied to people transitioning into the colorectal cancer
- 16 health state.

17 **8.1.1.3 Distributions of estimates**

- 18 Briggs et al. (2003) recommends that the gamma distribution is the most
- 19 appropriate probability distribution for costs. To fit a gamma distribution the
- 20 standard error is required for each value. Costs taken from NHS reference
- 21 costs 2008/09 and published papers, which have a stated standard error,
- were used in the model. For the cancer pathology costs, standard errors were
- 23 calculated because only the mean values were available.

2 Table 6 Mean costs and standard errors used in the probabilistic

3 sensitivity analysis

Parameter	Mean cost (£)	Standard error (£)
Symptomatic Dukes' A	11,965.78	6,490.90
Symptomatic Dukes' B	16,224.50	3811.55
Symptomatic Dukes' C	21,033.60	2368.03
Symptomatic Dukes' D	24,096.80	3050.62
Cancer pathology	250.00	277.98
Histology/histopathology	25.72	21.10
Colonoscopy	516.78	178.92

4

5 9 Assumptions

6 9.1 Cycle length

7 A cycle length of 3 months was assumed to be the most appropriate, because

8 surveillance for the high-risk group occurs every year and a 3-month cycle

9 allowed possible development of asymptomatic and symptomatic cancer

10 between colonoscopies.

11 9.2 Histopathology

12 The GDG recommended a median of eight biopsy specimens per

13 colonoscopy, with a lower limit of five and an upper limit of ten. Uncertainty

14 was captured using a simple uniform distribution with the minimum and

15 maximum because no information on the distribution was available.

16 9.3 Age dependency

- 17 The age-dependent variables used in the model were other cause mortality
- 18 and age-related utilities. All other variables were independent of time. Other
- 19 cause mortality was age dependent because it was assumed that people with
- 20 IBD have the same mortality as the rest of the UK population.

1 9.4 Misdiagnosis

2 It was assumed that no misdiagnoses were made during colonoscopy. This

- 3 follows the assumption that there may have been some degree of
- 4 misdiagnosis in the study by Rutter et al. (2006). Therefore, to include it would
- 5 double count the number of misdiagnoses.

6 9.5 Complications

7 It was assumed that people on surveillance have no complications caused by8 colonoscopy, such as perforations or bleeding.

9 9.6 Compliance

10 It was assumed that everyone participating in the surveillance programme

11 adhered to the colonoscopic surveillance protocol.

12 9.7 Diagnosis and treatment of cancer

13 It was assumed that cancer is detected once it becomes symptomatic and

14 asymptomatic cancer is only detected by surveillance colonoscopy.

15 Cancer costs and benefits have been separated, with costs applied only when

16 a person enters the colorectal cancer health state and benefits applied for

17 each time period in that state. This was assumed in the cost-effectiveness

- 18 study by Tappenden et al. (2004) and was a limitation of that study. This
- 19 limitation could potentially lead to conflicting conclusions about the effect of
- 20 colorectal cancer. However, because modelling the entire colorectal cancer
- 21 pathway is not possible in this guideline, this was considered an acceptable
- 22 simplification.

2 10 Results

- 3 The overall deterministic results are presented in table 7. Uncertainty about
- 4 the results follows in section 11.1.

5 **Table 7 Deterministic analysis over a 55-year period**

	QALYs	Cost (£)	Incremental QALYs	Incremental costs (£)	ICER (£)			
No surveillance	16.42	2320.44						
Surveillance – high-								
risk group only	17.19	15,785.13	0.77	13,464.69	17,557.32			
QALY: quality-adjusted life year; ICER: incremental cost-effectiveness ratio								

6

- 7 The analysis suggested that surveillance for the high-risk group is cost
- 8 effective.
- 9

10 **11 Sensitivity analysis**

- 11 Two approaches to testing the robustness of the model results were taken: a
- 12 series of one-way deterministic sensitivity analyses and a probabilistic
- 13 sensitivity analysis.

14 **11.1 Deterministic sensitivity analyses**

- 15 A one-way sensitivity analysis describes the process of changing one
- 16 parameter in the model and analysing the results of the model to see if this
- 17 parameter influences any of the overall results.
- 18 Sources of uncertainty were the number of biopsy specimens per
- 19 colonoscopy, the utility values and the costs. These were investigated using a
- 20 one-way sensitivity analysis. For each of the variables either the lower or the
- 21 upper point estimate was used, keeping all other variables constant. The
- 22 resulting incremental cost-effectiveness ratios (ICERs) are reported for each
- 23 variable in table 8.

Parameter	Base case	Range values			Determi	nistic ICER (£)			
				Distribution					
		Lower	Upper		Lower	Upper			
Biopsy specimen per colonoscopy	8	5	10	Uniform	15,654.07	18,826.15			
Utility values									
LGD	0.95	0.94	0.97	Gamma	17,511.19	17,650.29			
HGD	0.88	0.86	0.91	Gamma	17,452.29	17,717.24			
Dukes' A	0.74	0.69	0.78	Gamma	19,911.93	16,039.92			
Dukes' B	0.7	0.63	0.77	Gamma	17,299.27	17,823.18			
Dukes' C	0.5	0.44	0.56	Gamma	17,392.96	17,724.80			
Dukes' D	0.25	0.16	0.36	Gamma	17,511.85	17,613.21			
Cost parameters									
Histopathology	£25.72	£7.33	£35.80	Beta	13,928.47	19,546.67			
Colonoscopy	£516.78	£392.91	£634.27	Beta	14,501.94	20,455.62			
Dukes' A	£11,965.78	£10,387.24	£19,143.46	Beta	17,303.59	18,711.88			
Dukes' B	£16,224.50	£14,009.49	£19,151.27	Beta	17,609.59	17,488.60			
Dukes' C	£21,033.60	£19,445.98	£22,640.46	Beta	17,640.86	17,473.07			
Dukes' D	£24,096.80	£22,032.30	£26,147.59	Beta	17,617.74	17,497.60			
ICER: incremental cost-effectiveness ratio	ICER: incremental cost-effectiveness ratio; LGD: low-grade dysplasia; HDG: high-grade dysplasia								

Table 8 Varying the point estimate showing different ICERs

- 1 The results from table 8 suggest that the variables with the greatest impact on
- 2 the ICER are the number of biopsy specimens per colonoscopy, the utility
- 3 value allocated to stage Dukes' A, and the costs of histopathology and
- 4 colonoscopy.

5 **11.2** *Probabilistic sensitivity analysis*

6 The major limitation of a one-way sensitivity analysis is that there is often

- 7 uncertainty about many parameters at the same time. The joint impact of
- 8 altering all parameters simultaneously was therefore estimated using
- 9 probabilistic sensitivity analysis. The analysis was run 1000 times and for
- 10 each simulation, different values were picked from the various distributions for
- 11 each variable in the model.

12 The overall analysis is presented in table 9.

13

14 **Table 9 Probabilistic sensitivity analysis over a 55-year period**

	QALYs	Costs (£)	Incremental QALYs	Incremental costs (£)	ICER (£)	Probability of being cost effective at £20,000 per QALY gained			
No surveillance	13.04	7368.92	_	_	_	-			
Surveillance – high- risk group only	14.64	16,316.82	1.61	8947.90	5571.44	99%			
OALY: quality-adjusted li	OALX: quality-adjusted life year: ICEP: incremental cost-offectiveness ratio								

- 16 The incremental cost-effectiveness ratio (ICER) from the probabilistic
- 17 sensitivity analysis was lower than the ICER from the deterministic sensitivity
- 18 analysis. This suggests that there may be a high degree of uncertainty
- 19 associated with some model parameters, which resulted in a large change in
- 20 the ICER. However, in spite of the uncertainty the probabilistic sensitivity
- 21 analysis suggests that there is a 99% probability that colonoscopic
- 22 surveillance for the high-risk group (among the three risk groups) with IBD is
- 23 cost effective at the usual threshold of £20,000 per QALY gained.

- 1 Figure 2 shows the results of the 1000 simulations of the probabilistic
- 2 sensitivity analysis represented on the cost-effectiveness plane.



3 Figure 2 Cost-effectiveness plane for the high-risk group (IBD)

4

5

6 11.2.1 Cost-effectiveness acceptability curves

7 Figure 3 shows the cost-effectiveness acceptability curve for the high-risk

8 surveillance strategy. At a threshold of £20,000 per QALY gained and higher,

9 it shows the probability of being cost effective as nearly 100% for the high-risk

10 group compared with a no surveillance strategy.

- 1 Figure 3 The cost-effectiveness acceptability curve for the high-risk
- 2 surveillance strategy
- 3



6 12 Discussion

7 12.1 Strengths of the model

8 This model is similar to models used in previously published cost-

- 9 effectiveness studies on ulcerative colitis. One advantage this model has over
- 10 the others is that cancer has been divided into mutually exclusive states
- 11 representing Dukes' staging. Therefore, it more accurately considers the
- 12 different outcomes according to the stage of cancer. This allows better
- 13 identification of whether annual colonoscopies detect early-stage cancer,
- 14 which reduces cancer-related mortality.
- 15 A probabilistic sensitivity analysis was conducted to explore uncertainties in
- 16 the data.

1 **12.2** *Limitations of the model*

2 **12.2.1** Transition probabilities

The clinical data used to derive the transition probabilities were from an
observational study of low quality (Rutter et al. 2006). No randomised
controlled trial data were available because of the ethical issues of denying
people surveillance if they have an increased risk of cancer.

7 12.2.2 Management of dysplasia: high-risk group

8 It is recognised that people diagnosed with dysplasia may opt for surgery

- 9 (such as colectomy) depending on their personal preference and their
- 10 clinician's judgement. However, to simplify the model structure the cohort was
- 11 made up of people that decline surgery. This may have overestimated the
- 12 number of people that stay in surveillance.

13 **12.2.3 Misdiagnosis**

It was assumed that colonoscopy was associated with 100% sensitivity and 15 100% specificity. The GDG discussed the current sensitivity and specificity of colonoscopy to be around 95%. In addition, clinical data were mainly obtained from observational studies in which misdiagnosis was accounted for in the published literature. However, further work could incorporate the sensitivity and specificity of the chosen surveillance method where appropriate.

20 12.2.4 Complications

21 The potential complications of colonoscopy were not considered because of a

- 22 lack of time and resources. The inclusion of this factor could increase the
- 23 ICERs and make surveillance less cost effective.

24 **12.2.5 Quality of life data**

- 25 Uncertainty remains about the appropriate method to account for quality of life
- 26 associated with dysplasia because it is asymptomatic, whereas other risk
- 27 factors such as inflammation are symptomatic. The patient experts and clinical
- 28 specialists in the GDG considered that the psychological burden of being
- 29diagnosed with dysplasia and the grade of dysplasia could be very high. The
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approach taken to address the uncertainty was to conduct both a one-way
 sensitivity analysis and a probabilistic sensitivity analysis, varying the utility
 values.

4 **12.2.6** Treatment pathway

A large proportion of people may opt for surgery during the course of their
surveillance. The number of people requiring annual surveillance based on
their dysplasia may therefore have been overestimated. In either case it is
likely that colonoscopic surveillance will remain cost effective.

9 12.2.7 Chromoscopy

10 Chromoscopy was recommended for use in routine surveillance for people 11 with IBD. According to the NHS reference costs 2008/09, chromoscopy costs 12 the same as conventional colonoscopy. The GDG felt that although the 13 procedure may cost the same, the time needed to train healthcare 14 professionals to use chromoscopy is longer than training them to use 15 colonoscopy. Unfortunately, staff training time is usually already incorporated 16 into the reference costs therefore this cost-effectiveness model was unable to 17 compare conventional colonoscopy with chromoscopy. The GDG also stated 18 that chromoscopy takes longer to perform than colonoscopy. However, the 19 difference was not found to be statistically significant. Finally, for a true 20 comparison, sensitivity and specificity would need to be incorporated to 21 differentiate between the two types of colonoscopy.

22 **12.2.8 Costing**

- 23 Costs based on NHS reference costs may not be representative of the true
- 24 costs of the procedure. However, these are published costs and they
- 25 represent the average NHS costs across the country.

26 **13 Conclusions**

- 27 The analysis indicates that colonoscopic surveillance is a cost-effective
- 28 programme for people considered at high risk of developing colorectal cancer
- among the three risk groups for IBD surveillance, with an ICER below £20,000

1 per QALY gained when deterministic and probabilistic analyses are

2 considered.

3 **14** Future work

4 Because of the lack of time between the final meeting of the GDG when the 5 surveillance schedule was created and consultation for the guideline, it was 6 not possible to construct a new cost-effectiveness model to assess 7 surveillance for all three risk groups because transition probabilities would 8 depend on several factors in any given risk group. There is the possibility that 9 surveillance may not be cost effective for all three groups simultaneously. For 10 the future it will be important to evaluate whether surveillance for all three risk 11 groups, including those with resectable lesions, will be cost effective for 12 people with IBD.

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5 16 Appendices

6 16.1 Inclusion and exclusion criteria

Figure 4 Flowchart of the number of cost-effectiveness studies included
 and excluded



1 **16.2** *Quality checklist for de novo cost effectiveness*

2 **IBD high-risk group**

Guideline topic: Colonoscopic surveillance for IBD Question no: by Y Rajput 2010			
Check list completed by K Jeong			
Section 1: Applicability	Yes/ partly/ no/unclear/ NA	Comments	
1.1 Is the study population appropriate for the guideline?	Partly	30-year-old men and women who have had colitis symptoms for 10 years and are considered to be at high risk of developing colorectal cancer. Low- and intermediate- risk groups were not modelled.	
1.2 Are the interventions appropriate for the guideline?	2 Are the interventions appropriate for Partly e guideline?		
1.3 Is the healthcare system in which the study was conducted sufficiently similar to the current UK NHS context?	Yes		
1.4 Are costs measured from the NHS and personal social services (PSS) perspective?	Yes		
1.5 Are all direct health effects on individuals included?	Partly	QALY data from the USA using standard gamble technique used.	
1.6 Are both costs and health effects discounted at an annual rate of 3.5%?	Yes		
1.7 Is the value of health effects expressed in terms of quality-adjusted life years (QALYs)?	Yes		
1.8 Are changes in health-related quality of life (HRQoL) reported directly from patients and/or carers?	Yes		
1.9 Is the valuation of changes in HRQoL (utilities) obtained from a representative sample of the general public?	No	IBD QALY data were taken from a Crohn's disease study using time trade off.	

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				Colorectal cancer QALY data from the USA using standard gamble technique	
				used.	
1.10 Overall judgement: directly applic	able/par	tially app	licable	e/not applicable	
There is no need to use section 2 of the	e check	list if the	study	is considered 'not	
applicable'. Partially applicable					
Other comments					
Section 2: Study limitations (the	Yes/ p	artly/	Cor	nments	
level of methodological quality)	no/ un	clear/			
This checklist should be used once it has	NA				
been decided that the study is sufficiently	Comm	nents			
quideline					
2.1 Does the model structure	Yes		Use	e of a younger	
adequately reflect the nature of the			pop	population than other	
health condition under evaluation?			chro	chronic conditions	
2.2 Is the time horizon sufficiently	Yes		55 v	/ears	
long to reflect all important					
differences in costs and outcomes?					
2.3 Are all important and relevant	Yes				
health outcomes included?					
2.4 Are the estimates of baseline	Yes		Obs	Observational study in the	
health outcomes from the best			UK	setting	
available source?					
2.5 Are the estimates of relative	elative Yes		Bes	t quality studies	
treatment effects from the best			ider	ntified from clinical	
available source?	le source?		revi	ew	
2.6 Are all important and relevant	2.6 Are all important and relevant Yes				
costs included?					
2.7 Are the estimates of resource use	7 Are the estimates of resource use Yes		NHS	S specific	
from the best available source?					
2.8 Are the unit costs of resources	2.8 Are the unit costs of resources Yes				
from the best available source?	No.				
2.9 Is an appropriate incremental	res				
analysis presented of can it be					
2 10 Are all important parameters	Voo				
2.10 Are all important parameters	res				
whose values are uncertain					
analysis?					
2 11 is there no notential conflict of	No				
interest?					
2 12 Overall assessment : minor limitations/notentially serious limitations/yorv					
serious limitations					
Potentially serious limitation, only one subaroup in the high-risk group was evaluated					
However the ICER for the high-risk arc	oup was	robust (a	as den	nonstrated in the	
probabilistic sensitivity analysis).	1	(- · · ·	
IBD: inflammatory bowel disease: ICER: incremental cost-effectiveness ratio: QALY:					

quality adjusted life year

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Appendix 8 – Health economic evaluation

2 **Cost-effectiveness analysis of colonoscopic**

3 surveillance: adenomas

4 **1** Introduction

5 The Department of Health asked NICE to produce a short clinical guideline on

6 colonoscopic surveillance for the prevention of colorectal cancer in patients

- 7 with ulcerative colitis, Crohn's disease and polyps.
- 8 A cost-effectiveness analysis has been carried out to support the Guideline
- 9 Development Group (GDG) in making recommendations for adults with
- 10 adenomas considered to be at high risk of developing colorectal cancer. This
- 11 analysis has been conducted according to the methods outlined in the NICE
- 12 Guide to the methods of technology appraisal 2008 and the Guidelines
- 13 Manual 2009. The methods used follow the NICE reference case, which is the
- 14 framework NICE requests all cost-effectiveness analyses to follow. In the
- 15 model, it is assumed that people at the endpoint of colonoscopic surveillance
- 16 would return to the UK population norm then enter the NHS Bowel Cancer
- 17 Screening Programme according to the current criteria.
- 18 Given the quality of the data available this analysis should be considered an
- 19 exploration of the cost effectiveness of colonoscopic surveillance for
- 20 adenomas.

21 2 Acknowledgements

- 22 On behalf of the GDG and NICE technical team, we would like to
- 23 acknowledge and thank Paul Tappenden and Hazel Pilgrim for their support
- 24 and help in the development of this guideline by providing the uplifted costing
- 25 data for stage-specific colorectal cancer.

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2 4 Decision problem

- 3 Table 1 outlines the decision problem that is addressed in this guideline and is
- 4 based on the final scope.

5 **Table 1 Decision problem**

	Scope	Approach taken
Population	People with polyps including adenomas in the colon and rectum	People aged 50 years who have adenomas removed in the colon and rectum at baseline colonoscopy
Interventions	Colonoscopic surveillance using colonoscopy, chromoscopy, computerised tomography colonoscopy, narrow band imaging, double-barium contrast enema	Colonoscopic surveillance using colonoscopy
Comparator	No colonoscopic surveillance	No colonoscopic surveillance
Outcome(s)	Costs, quality-adjusted life years (QALYs) and cost per QALY gained	Cost per QALY gained

6

7 4.1 Population

- 8 The estimated prevalence of colonic adenomas is 30–40% at age 60 years
- 9 (Williams et al. 1982) and the lifetime cumulative incidence of colorectal
- 10 cancer is 5.5% (Lieberman et al. 2000). Adenomas are diagnosed on average
- 10 years before colorectal cancer (Olsen et al. 1988). Therefore, the model
- 12 simulated men and women aged 50 years in order to identify precancerous
- 13 polyps. People entering the model had adenomas and were at high risk of
- 14 developing colorectal cancer. Any detected adenomas were removed at
- 15 baseline colonoscopy and during subsequent surveillance.

16 **4.2** *Interventions*

- 17 From the clinical review there was no direct evidence for or against routine
- 18 colonoscopic surveillance for the prevention and early detection of colorectal
- 19 cancer after removal of adenomas. Currently there is no national guidance
- 20based on the clinical and cost effectiveness of surveillance in the NHS. The
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- 1 model assessed the cost effectiveness of current practice in the NHS, which
- 2 broadly follows the British Society of Gastroenterology (BSG) guidelines –
- 3 people are offered colonoscopic surveillance after removal of adenomas
- 4 (Atkin and Saunders 2002).

5 4.3 Comparators

- 6 Colonoscopy is the gold standard for surveillance and screening for colorectal
- 7 cancer in the NHS. Therefore, colonoscopic surveillance using colonoscopy is
- 8 the main comparator in the surveillance model compared with no surveillance.
- 9 Colonoscopic surveillance after adenoma removal is consistent with the BSG
- 10 guidelines (Atkin and Saunders 2002; Cairns et al. 2010). The person's risk
- 11 state is determined after the baseline colonoscopy and is based on the
- 12 number and size of adenomas removed at the baseline colonoscopy. In the
- 13 model, surveillance in low-, intermediate- and high-risk groups is referred to
- 14 as surveillance in all risk groups. The BSG guidelines are not definitive for
- 15 surveillance of people at low risk of developing colorectal cancer, therefore
- 16 surveillance in the intermediate- and high-risk groups only is also considered.
- 17 An outline of the surveillance strategies considered in the model is given in
- 18 table 2.

Risk status	Schedule
Low risk :one or two adenomas smaller than 10 mm	No surveillance is recommended. However surveillance at 5 years, then no surveillance if subsequent colonoscopy results are negative can be considered and will be explored in the analysis.
Intermediate risk: three or four adenomas smaller than 10 mm or one or two adenomas if one is 10 mm or larger	Surveillance is offered every 3 years until there are two consecutive negative colonoscopies, then surveillance is stopped.
High risk: five or more adenomas smaller than 10 mm or three or more adenomas if one is 10 mm or larger	A colonoscopy is offered at or within 1 year to detect missed lesions:
	 if high-risk adenomas are detected, the person remains high risk
	• if results are negative, or low- or intermediate- risk adenomas are detected, the surveillance programme for people at intermediate risk is followed.

19 Table 2 Surveillance schedule after adenoma removal in the model

2 **4.4 Outcomes**

In line with the NICE reference case, a cost-utility analysis was used to
assess the cost effectiveness of colonoscopic surveillance using conventional
colonoscopy. Given the absence of an appropriate analysis, a Markov model
was developed to fit the decision problem. This required the calculation of
resource use and quality-adjusted life years (QALYs) to assess effectiveness.

8 5 Review of existing cost-effectiveness
 9 analyses

10 **5.1** Search for cost-effectiveness analyses

A search for cost-effectiveness, quality of life and resource papers was carried out. These papers were then subject to a systematic search. Papers were initially excluded, for example, on the basis of the title, subject, intervention or condition. Of the remaining papers, abstracts were then searched to see if they contained relevant information. These papers were then categorised into: cost effectiveness – colonoscopic surveillance, cost effectiveness – natural history, quality of life and resource use.

18 **5.2**

19

Review of cost-effectiveness studies – colonoscopic surveillance

Of 289 studies identified for both polyps and inflammatory bowel disease, 234 were excluded based on the title and an abstract review. The applicability of 55 studies was assessed using a checklist. Of 55 studies of potential interest, 54 were excluded based on NICE methods and the NICE reference case

- 24 using modified GRADE methods. Only one analysis was relevant to
- 25 surveillance for adenomas (Tappenden et al. 2004). A modified GRADE table
- that summarises this analysis is presented in section 16.5.
- 27 The study by Tappenden et al. (2004) was considered of high quality and
- 28 provided valuable information on the modelling approach. However, the study

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1

- 1 had limited applicability because the population and comparators were
- 2 different to the decision problem and so a new Markov model was developed.

3 5.3 Modelling approach

4 Colonic polyps and recurrent adenomas are chronic conditions that require 5 lifetime surveillance to prevent colorectal cancer (Atkin and Saunders 2002). The transformation of adenomas to invasive colorectal cancer is slow and can 6 7 take 10–15 years (South West Cancer Intelligence Service 1995). Therefore, 8 a Markov model was developed over a lifetime horizon (50 years). This was 9 associated with risks of developing colorectal cancer over time and the 10 importance of detecting the transformation of adenoma to cancer. The three 11 diagnostic states in the model, low, intermediate and high risk, differ only in 12 terms of the surveillance offered. Movement between diagnostic states is only 13 possible through surveillance or symptomatic presentation of colorectal 14 cancer. The health states represented the progression of the condition from 15 adenoma free after adenoma removal, to new non-advanced adenomas after adenoma removal, to asymptomatic and symptomatic colorectal cancer (using 16 17 Dukes' A, B, C and D classification; Dukes 1932) to death. The GDG 18 acknowledged that the future risk of developing colorectal cancer or advanced 19 adenomas after removal of adenomas depends on the number and size of 20 adenomas removed at baseline colonoscopy, as indicated in the BSG 21 guidelines (Atkin and Saunders 2002; Cairns et al. 2010) (see figure 1).

22

- 2 Figure 1 Surveillance after adenoma removal (Atkin and Saunders 2002)
- 3

1



SURVEILLANCE FOLLOWING ADENOMA REMOVAL

5 For simplicity, people in the surveillance programme were assumed to adhere 6 to the schedule, but in reality this is unlikely. For the purpose of the guideline,

7 when comparing a surveillance programme with no surveillance, the

8 sensitivity and specificity of colonoscopy were assumed to be 100% in

9 adenoma detection. This was agreed with the GDG. In reality, the actual rates

10 would be lower; however, the clinical data may take this into account in the

11 estimates of progression. It was also assumed that each colonoscopy is

12 complete, which increases uncertainty in the results.

13 5.4 Natural history

14 It is widely accepted that most colorectal cancers arise from pre-existing

- adenomas, based on epidemiological, clinical, post-mortem, and molecular
- 16 biology evidence. The size of adenomas is correlated with malignant potential
- 17 (Muto et al. 1975). It is unlikely for a small adenoma (≤5 mm) to progress to
- invasive cancer in less than 5 years (Eide 1986). Winawer et al. (1993b)
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1 reported that people with a history of adenoma removal are more likely to 2 develop subsequent adenomas. The majority of recurrent adenomas are 3 found to be predominantly small (≤ 5 mm) at follow-up (Winawer et al. 1993b). 4 Outcomes of clinical treatment can be determined by using the natural history 5 of adenomas leading to colorectal cancer. The clinical results of treatment can 6 be extrapolated to a lifetime horizon to account for the long-term benefits of 7 treatment. Because of a lack of resources and time a full systematic review of 8 the natural history data to calculate transition probabilities was not possible. 9 Therefore, all cost-effectiveness studies were reviewed to estimate the 10 progression of polyps to colorectal cancer. One analysis was found that 11 reported the cost effectiveness and cost-utility of colorectal cancer screening 12 options in England (Tappenden et al. 2004). Tappenden et al. obtained 13 estimates from two sources: the National Polyp Study (Winewar et al. 1993a) 14 and calibrating their model against 60,000 random iterations, of which around 15 400 potential solutions were identified that appeared to fit the published 16 incidence and mortality data. These data on the natural history of undetected 17 colorectal cancer, polyp incidence and growth rates, the rate at which high-18 risk adenomas develop into cancer, and stage-specific colorectal cancer-19 related mortality represented the best available source and were used in the 20 model.

21

22 **6 Model**

23 6.1 Model structure

- 24 The structure of the colonoscopic surveillance model for people with
- adenomas is given in figure 2.

26

27



- 17 NAA: non-advanced adenoma; AAi: advanced adenoma, intermediate risk; AAh: advanced
- 18 adenoma, high risk; DA: Dukes' A; DB: Dukes' B; DC: Dukes' C; DD: Dukes' D; CRC:
- 19 colorectal cancer.
- 20
- 21

1 Limited evidence exists about the actual risk of developing advanced 2 adenomas and colorectal cancer after removal of adenomas. Figure 3 gives 3 an outline of the natural history model. Given the slow transformation of 4 adenomas to colorectal cancer and the fact that most recurrent adenomas are 5 small (≤5 mm) (Winawer et al.1993b), the risk state at baseline colonoscopy is 6 inversely correlated with the size of the group in the initial starting cohort. For 7 example, the higher the risk of developing colorectal cancer, the smaller the 8 group in the initial starting cohort. The proportions chosen were based on 9 Lieberman et al. (2008) with 64% of the cohort starting in the low-risk, 20% in 10 the intermediate-risk and 16% in the high-risk group. This reflects the fact that 11 small adenomas are most commonly found during colonoscopy. The main 12 components of the model were the surveillance and natural history strategies.

13 **6.1.1** Surveillance and natural history

14 The effectiveness of colonoscopic surveillance was modelled as an 15 intervention under near-perfect conditions to determine whether colonoscopic 16 surveillance using colonoscopy for the early detection of adenomas and 17 colorectal cancer was clinically and cost effective compared with no 18 surveillance. Health states included being polyp free and having recurrent 19 adenomas to incorporate the natural history of recurring adenomas after 20 adenoma removal. The strategies analysed were surveillance of all risk 21 groups and surveillance of the intermediate- and high-risk groups only. 22 Movement between the three diagnostic states was only possible through

surveillance or symptomatic presentation of colorectal cancer. According to
surveillance criteria, people could drop out of surveillance and be assumed to
return to UK population norms.

26 6.1.2 Colorectal cancer

Symptomatic and asymptomatic colorectal cancers were included in the
surveillance model. It was assumed that colorectal cancer is diagnosed by
symptomatic presentation or surveillance. People who stop surveillance were
assumed to have the same mortality risk as the general population. The GDG
agreed with this assumption. The cost of treating colorectal cancer was not
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- 1 varied according to the method of diagnosis, that is, by surveillance or
- 2 symptoms. Costs were varied by stage of cancer Dukes' A, B, C or D.
- 3 Therefore, if cancers are detected early, the average cost is reduced.

4 6.1.3 Complications

5 No complications or adverse events were assumed in the model. This

6 assumption was agreed with the GDG.

7 6.1.4 After removal of adenomas (tunnel states)

8 In the model, tunnel states were used to represent two health states after
9 removal of adenomas, depending on whether the person had previous
10 adenomas, to determine the surveillance strategy:

- adenoma free after removal of non-advanced adenoma at year 1 and
 year 2 onwards
- adenoma free after removal of advanced adenoma at year 1 and year 2
 onwards.
- 15 It was assumed that all adenomas are removed endoscopically during
- 16 surveillance. It was also agreed with the GDG to assume that all colorectal
- 17 cancers arise from pre-existing adenomas.
- 18 The main consideration in this model is that the long-term outcomes from
- 19 repeated colonoscopic surveillance depend on two factors: timing of adenoma
- 20 removal (prevention of colorectal cancer) and timing of cancer detection
- 21 (detection of early colorectal cancer). This affects the proportion of people
- 22 who can be treated with surgery only (Dukes' A colorectal cancer) and
- 23 subsequent long-term survival. Therefore, the treatment benefit distinguishes
- 24 between cancer that is detected early and is asymptomatic, and symptomatic
- 25 cancer, which is reflected in the costs and health benefits (QALYs). It was
- 26 assumed that people diagnosed with colorectal cancer (asymptomatic or
- 27 symptomatic) received identical stage-specific treatments. This allowed a
- 28 comparison of colonoscopic surveillance with no surveillance under identical

conditions. It was also assumed that people could progress to death from all
 health states.

3 6.2 Transition probabilities

4 The transition probabilities were taken from Tappenden et al. (2004). Although 5 these data were from the best available source, transferring them to another model was not ideal and there was potential uncertainty. In addition, there 6 7 were no data for people at intermediate risk of developing colorectal cancer. 8 Therefore, the data for the high-risk group were extrapolated to this group. To 9 reflect the fact that the probability of the intermediate-risk group progressing 10 from adenoma free is likely to be less than for the high-risk group, a variable 11 factor was added to reduce the rate of progression. A base-case value of 0.85 12 was chosen at random, but it was varied in the probabilistic sensitivity 13 analysis. 14 Data from published interim life tables for the UK (Office of National Statistics 15 2009) were used to calculate age-related mortality probabilities. It was

- 16 assumed that for people in the asymptomatic colorectal cancer health states
- 17 the probability of dying is the same as the age-related probability of dying.
- 18 This appears to be reasonable because people without symptoms are unlikely
- 19 to have an increased risk of death until their cancer progresses. This ensured
- 20 that all probabilities added up to one (table 3).

	AF	NAA	AA	DA	DB	DC	DD	mCRC	mOthers
AF(NAAR) year 1	#	0.18	0	0	0	0	0	0	Age
AF (NAAR) year 2+	#	0.05	0	0	0	0	0	0	Age
AF (AAR) year 1	#	0.25	0	0	0	0	0	0	Age
AF (AAR) year 2+	#	0.06	0	0	0	0	0	0	Age
NAA	0	#	0.021	0	0	0	0	0	Age
AA	0	0	#	0.0326	0	0	0	0	Age
DA	0	0	0	#	0.5829	0	0	0.0	Age
DB	0	0	0	0	#	0.6555	0	0.010	Age
DC	0	0	0	0	0	#	0.864 8	0.0602	Age
DD	0	0	0	0	0	0	#	0.3867	Age
mCRC	0	0	0	0	0	0	0	1	0
mOthers	0	0	0	0	0	0	0	0	1

1 Table 3 Natural history yearly transition matrix

AF: adenoma free; NAAR: non-advanced adenoma removed; AAR: advanced adenoma removed; NAA: nonadvanced adenoma; AA: advanced adenoma; DA: Dukes' A colorectal cancer (CRC); DB: Dukes' B CRC; DC: Dukes' C CRC; DD: Dukes' D CRC; mCRC: death caused by CRC; mOthers: death from other causes, # 1 minus other states; Age: age dependent.

2

3 7 Quality of life

4 NICE recommends that changes in HRQoL as a result of an intervention or

5 treatment should be directly reported by patients. These changes should be

- 6 based on preferences determined using a choice-based method in a
- 7 representative sample of the UK general public. Ideally a full systematic
- 8 review would be carried out to identify HRQoL studies and appropriate values
- 9 to include in a health economic model. However, because of the lack of
- 10 resources and time, a search was carried out for quality of life studies. The
- 11 quality of life data included in the cost-effectiveness analyses identified in
- 12 section 5 are reviewed.

13 7.1 Literature search

14 A literature search found studies relating to quality of life in people with polyps

15 or adenomas. Quality of life evidence for people with colorectal cancer was

limited.

1 7.1.1 Review of the literature

The main study identified was Ness et al. (1999), which assessed utility values associated with the stage of cancer and treatment. It is crucial to capture utility values that include pre-cancerous stages and any possible positive and/or negative impact of the test results on the person's wellbeing. However, there was no evidence identified from the search demonstrating a decrease in utility values associated with colonoscopic surveillance.

8 7.2 People's health states

9 NICE recommends the use of the EuroQol 5 dimensions (EQ-5D) or another 10 generic tool that enables patients to describe their health states and how the 11 public values their health states. In addition, no one set of values can be used 12 for the entire model. There are also potential issues when different values are 13 used from different sources, which may lead to inconsistencies. For example, 14 time trade off and standard gamble techniques have a tendency to produce 15 different estimates for the same health states. To minimise potential 16 inconsistency, studies were chosen that follow the NICE methods and also 17 share similar populations and methods of determining and valuing health 18 states.

Ness et al. (1999) assessed utility values associated with the stage of cancer and treatment in the USA. People were asked to assess utility values for stage-dependent outcome states using the standard gamble technique. These states were not valued by the UK public. The GDG considered the very limited evidence on the colorectal cancer stage-specific utilities, and agreed that the use of utility values from Ness et al. was appropriate in the model.

25 **7.3 Cancer-free and adenoma-free health states**

Utility values associated with the cancer-free and the adenoma-free health states were assumed to be the same as the 'no known adenomas' heath state with a utility value of 0.91 (Ness et al. 2000; Tappenden et al. 2004). This was considered to be a reasonable assumption because adenomas are likely to be asymptomatic.

1 7.4 Cancer

Evidence about people's quality of life, especially in stage-specific colorectal cancer, was very limited. No published studies were found that considered the quality of life impact of colonoscopic surveillance, diagnosis and subsequent treatment of colorectal cancer. Ness et al. (1999) interviewed 90 people who had previously had colorectal adenomas removed to assess utility values associated with stage-specific colorectal cancer using a standard gamble technique.

Asymptomatic cancer and undiagnosed cancer were assigned the same utility
value as their diagnostic state (0.91) because if cancer is asymptomatic it is
unlikely to affect the quality of life of the person until it is detected (that is, until
it becomes symptomatic).

13 **7.5 Colonoscopy**

The patient experts in the GDG felt that the utility value for the cancer-free health state would be less than 0.91 because of the significant temporary disability caused by intensive bowel preparation and the recovery period after the procedure. Therefore discomfort associated with the procedure was explored in the sensitivity analysis using a disutility value of 0.0025 (Saini et al. 2010; Syngal et al. 1998).

1 7.6 Final quality of life values

Health state	Mean value	Standard error	Reference
Cancer-free state	0.91	0.015306	Ness et al. (2000)
Asymptomatic cancer	0.91	0.051020	Ness et al. (2000)
Dukes' A CRC	0.74	0.022959	Ness et al. (1999)
Dukes' B CRC	0.70	0.035714	Ness et al. (1999)
Dukes' C CRC	0.50	0.030612	Ness et al. (1999)
Dukes' D CRC	0.25	0.030612	Ness et al. (1999)
CRC: colorectal cancer	•	-	-

2 Table 4 Final health-related quality of life estimates

3

4 8 Resource use

5 8.1 Literature search

6 From the initial search, only one study was identified that examined resource

7 use in the NHS (Tappenden et al. 2004) that was applicable to the model.

8 Stage-specific colorectal cancer treatment costs were uplifted to incorporate

9 the relevant NICE guidance published since 2004 (personal communication

10 with Paul Tappenden and Hazel Pilgrim, 8 April 2010).

11 8.1.1 Specific costs for the model

- 12 The main cost inputs that required consideration include:
- 13 colonoscopy and pathology
 - lifetime treatment costs for stage-specific colorectal cancer.
- 15

14

16 8.1.1.1 Endoscopy (colonoscopy)

- 17 The cost of endoscopy was obtained from NHS reference costs 2008/09
- 18 (£517; NHS cost code FZ26A endoscopic or intermediate large intestine
- 19 procedures 19 years and over).

20

2 8.1.1.2 Pathology for adenomas

3 The cost of pathology for adenomas was obtained from NHS reference costs

4 2008/09 (NHS cost code DAP824 – histology or histopathology.

5 8.1.1.3 Stage-specific treatment costs for colorectal cancer

6 Recently uplifted stage-specific treatment costs for colorectal cancer were

7 based on Tappenden et al. (2004) (personal communication with Paul

8 Tappenden and Hazel Pilgrim, 8 April 2010). These broadly include

9 chemotherapy, surgery or radiotherapy (if appropriate), follow-up, and

10 palliative care.

11 8.1.1.4 Distributions of estimates

12 The gamma distribution is recommended as the appropriate probability

- 13 distribution for costs (Briggs et al. 2003). To fit a gamma distribution the
- 14 standard error is required for each value. The standard errors for the costs
- 15 obtained from Tappenden and Pilgrim were calculated using the mean costs,
- 16 97.5% and 2.5% credibility intervals (Tappenden and Pilgrim 2010). There is
- 17 no agreed method on how to calculate standard errors for reference costs.
- 18 Only the mean and quartile values (except the median) are available.
- 19 Therefore the method used was the solver function in Excel to find the
- 20 variables for the gamma function that produce the relevant estimates of the
- 21 upper and lower quartiles.

22 9 Assumptions

- 23 The GDG agreed that the model would only examine factors relating to
- 24 colorectal cancer development; other epidemiological factors would be
- 25 considered only when a risk of developing colorectal cancer can be
- 26 demonstrated.

1 9.1 Age of cohort and cycle length

The GDG agreed on a cohort age of 50 years and a cycle length of 3 months, which allows transition to other states in between surveillance visits. The GDG agreed that no further surveillance would be carried out after 85 years because of the slow transformation of adenomas to colorectal cancer over 10–15 years (Winawer 1993a), and the potential risks and benefits of colonoscopic surveillance. Therefore the model was run over 50 years with a surveillance duration of 35 years.

9 9.2 Age dependency

10 Apart from other cause mortality all other variables were independent of time.

- 11 This was because of a lack of information on the relationship between time
- 12 and a number of important variables such as the rate of cancer progression.
- 13 Other cause mortality was age dependent because it was assumed that
- 14 people with adenomas have the same mortality as the rest of the UK
- 15 population. This seemed appropriate because there is no other reported
- 16 difference in life expectancy other than increased rate of recurrent adenomas
- 17 and increased colorectal cancer rate in people with adenomas.

18 **9.3** *Misdiagnosis*

19 The GDG acknowledged that the underlying data from observational studies 20 already included a degree of misdiagnosis. Therefore it was assumed in the 21 model that there was no misdiagnosis.

22 9.4 Complications

- 23 For simplicity, no complications relating to colonoscopy or adenoma removal
- 24 were assumed in the model. The GDG discussed potential risks associated
- 25 with colonoscopy and adenoma removal, including bowel perforation and
- 26 bleeding. The GDG noted that the number of colonoscopy-related
- 27 complications reported was small but these events could be fatal.

1 9.5 Compliance

In the model, the cohort was assumed to adhere to the colonoscopy schedule.
The GDG discussed the higher compliance rate in people who were informed
of an increased risk of developing colorectal cancer and this assumption was
considered to be reasonable.

6 9.6 Stopping surveillance

7 The GDG agreed that the low-risk group would not have further surveillance 8 after one negative colonoscopy. The intermediate-risk group would have a 9 follow-up at 3 years, then would stop surveillance after two consecutive 10 negative results. The high-risk group would have a follow-up colonoscopy at 11 1 year, which would determine the surveillance strategy: if the colonoscopy is 12 negative, or low- or intermediate adenomas are found, they would follow the 13 frequency of surveillance for the intermediate-risk group; if high-risk 14 adenomas are found, they would have colonoscopic surveillance at 1 year. 15 This surveillance schedule follows the current BSG guidelines (Atkin and 16 Saunders 2002; Cairns et al. 2010). Health benefits (QALY gains) of people 17 who meet the criteria for stopping surveillance were accounted for in the surveillance models. 18

19 9.7 Diagnosis and treatment of cancer

20 Colonoscopy, removal of adenomas and pathology were included for the 21 surveillance and treatment of adenomas detected during surveillance. 22 Surgery, chemotherapy and radiotherapy were included for the treatment for 23 colorectal cancer. Appropriate NICE guidance for the treatment of colorectal 24 cancer was also taken into account. Therefore, the impact of colonoscopic 25 surveillance on the cost effectiveness is the relative benefit of prevention or 26 early detection of colorectal cancer. Costs incurred at each stage of colorectal 27 cancer and detrimental to quality of life were also included.

- 28 Cancer costs and benefits were separated, with costs applied only when a
- 29 person entered the colorectal cancer health state and benefits applied for
- 30 each time period in the state. This was assumed in Tappenden et al. (2004)

and was a limitation of that study. This limitation could lead to conflicting
 conclusions over the effect of colorectal cancer. However, because modelling
 the entire colorectal cancer pathway was not possible, this was considered an
 acceptable simplification.

5 9.8 Adenoma recurrence rate during surveillance

6 in the model it was assumed that he probability of people in the high-risk 7 group who have had adenomas removed developing further adenomas is 8 higher than for people with no previous history of adenomas. In the 9 surveillance model two tunnel states represent post-adenoma removal (see 10 section 6.1.4). Tappenden et al. (2004) gave the key uncertainties in their 11 analysis, including the probability of progressing through undiagnosed cancer 12 states, the probability of clinical presentation by cancer stage, polyp incidence and growth rates, the rate at which high-risk adenomas develop into cancer, 13

14 and stage-specific colorectal cancer mortality rate.

15 9.9 Transition probabilities

16 Estimated transition probabilities were assumed to be constant with the

17 exception of age-specific adenoma incidence (Tappenden et al. 2004) and

18 age-specific mortality rates, which were taken from government sources.

19 Because of limited evidence the GDG agreed that all transitions from one

20 health state to the next in the model are progressive.

21 **9.10** Utility values for cancer-free health states

Because a person with adenomas who is cancer free is likely to be
asymptomatic, the utility value estimate for this health state was assumed to
be the same as that for the general population (Ness et al. 2000; Tappenden
et al. 2004). The GDG considered this was necessary because most people
with adenomas are asymptomatic.

27 9.11 Colorectal cancer

28 Probabilities of cancer progression were assumed to be equivalent in both the

29 distal and proximal colon. This appears to be a reasonable assumption

- 1 because the population included in the model have no familial or previous
- 2 history of colorectal cancer.

3 9.12 Final costs

- 4 Stage-specific colorectal cancer treatment costs, obtained from NHS
- 5 reference costs, were uplifted (personal communication with Tappenden and
- 6 Pilgrim, 8 April 2010). The final values and a breakdown are presented in
- 7 table 5.

8 Table 5 Mean costs and standard errors used in the base-case analysis 9 and probabilistic sensitivity analysis

		Standard	
Costs	Mean (£)	error (£)	Reference
Diagnostic/therapeutic colonoscopy			NHS reference costs
	517.00	178.92	2008/09 (2010)
Pathology for adenoma			NHS reference costs
	26.00	21.50	2008/09 (2010)
Pathology for cancer	250.00	277.98	Tappenden et al. (2004)
Lifetime cost			
Dukes' A			Tappenden and Pilgrim
	11,965.78	6490.90	(2010)
Dukes' B			Tappenden and Pilgrim
	16,224.50	3811.55	(2010)
Dukes' C			Tappenden and Pilgrim,
	21,033.60	2368.03	(2010)
Dukes' D			Tappenden and Pilgrim
	24,096.80	2050.62	(2010)

10 **10 Sensitivity analysis**

11 **10.1 Deterministic sensitivity analysis**

12 Deterministic sensitivity analysis was carried out on a range of variables,

- 13 including all costs and utility values. The key areas of uncertainty (see section
- 14 9.8) were explored by examining two sets of transition matrices: higher values
- 15 from the literature and another set of lower values. The full matrices are given
- 16 in table 6. Costs were reduced and increased by 50% to examine this effect. A
- 17 person's quality of life was explored in relation to the potential (dis)utility
- 18 associated with full bowel preparation and the recovery period (Sandi et al.

19 2010).

- 1 Table 6 Transition probabilities through model calibration (Tappenden et
- 2 al. 2004)

Annual transition probability		Parameter estimate used in base-case	Uniform distribution used in calibration		
State from	State to	analysis	Minimum	Maximum	
LR	HR	0.02	0.005	0.0400	
HR	DA	0.033	0.0100	0.0600	
DA	DB	0.5830	0.3000	0.9000	
DB	DC	0.6560	0.3000	0.9000	
DC	DD	0.8650	0.3000	0.9000	
PSDA	-	0.0700	0.0200	0.1500	
PSDB	-	0.3200	0.1000	0.3500	
PSDC	-	0.4900	0.5000	0.9000	
PSDD	_	0.8540	0.5000	0.9000	
DA	mCRC	0.000	0.000	0.0050	
DB	mCRC	0.0100	0.0050	0.0300	
DC	mCRC	0.0600	0.0200	0.1500	
DD	mCRC	0.3870	0.3500	0.4500	

LR: low risk; HR: high risk; DA: Dukes' A colorectal cancer (CRC); DB: Dukes' B CRC; DC: Dukes' C CRC; DD: Dukes' D CRC; mCRC: death caused by CRC; mOthers: death from other causes; PSDA: probability of presenting with symptomatic Dukes' A CRC; PSDB: probability of presenting with symptomatic Dukes' B CRC; PSDC: probability of presenting with symptomatic Dukes' C CRC; PSDD: probability of presenting with symptomatic Dukes' D CRC; DC: Dukes' D CRC

3

4 10.2 Structural sensitivity analysis

5 The following structural assumptions and variables were explored.

6 **10.2.1 Age of the cohort**

- 7 The base case assumes an average age of 50 years for the cohort because
- 8 most published cost-effectiveness analyses use 45 years based on limited
- 9 prevalence data. Average cohort ages of 35, 40 and 45 (varying the duration
- 10 of surveillance) were explored.

1 10.2.2 Stopping surveillance at different ages

- 2 The cutoff age for stopping surveillance was altered from 85 to 65, 70 and 75,
- 3 because remaining life expectancy is likely to be less than the average time
- 4 required for adenomas to develop into colorectal cancer.

5 10.3 Probabilistic sensitivity analysis

- 6 All transition probabilities in the natural history were varied using beta
- 7 distributions. Because no standard errors were available, a sample size of 100
- 8 was assumed. This value and the mean were used to calculate the relevant
- 9 factors.

10 **10.3.1 Utility values**

- 11 Beta distributions of the differences between the estimates were used to
- 12 ensure that the results remained consistent. Table 7 outlines the utility values
- 13 varied according to their difference.

		Standard	Distribution
State	Mean	error	
Cancer free	0.91	0.02	Beta
Undiagnosed asymptomatic colorectal cancer	0.91	0.02	Beta
Dukes' A	0.74	0.02	Beta
Dukes' B	0.70	0.04	Beta
Dukes' C	0.50	0.03	Beta
Dukes' D	0.25	0.05	Beta

14 Table 7 Probabilistic sensitivity analysis calculations for quality of life

15

16 **10.3.2 Costs**

- 17 Table 8 outlines the costs and standard errors that were modelled using a
- 18 gamma distribution.

1 Table 8 Probabilistic sensitivity analysis: gamma or normal distribution

2 of costs

Costs	Mean (£)	Standard error (£)
Colonoscopy	517.00	178.92
Pathology for adenoma	26.00	21.50
Pathology for cancer	250.00	277.98
Lifetime treatment cost		
Dukes' A	11,965.78	6490.90
Dukes' B	16,224.50	3811.55
Dukes' C	21,033.60	2368.03
Dukes' D	24,096.80	3050.62

3

4 11 Results

5 11.1 Deterministic results

- 6 Table 9 presents results of the deterministic base case. Colonoscopic
- 7 surveillance following the BSG guidelines and the inclusion of the low-risk
- 8 group are both associated with ICERs below £20,000 per QALY gained
- 9 compared with no surveillance. These results appear to have face validity
- 10 because the total cost of surveillance according to the BSG guidelines 2010
- 11 was estimated to be £1100, which is consistent with the value given in table 9.

12 Table 9 Deterministic analysis over a 50-year period

	QALYs (utilities)	Costs (£)	Incremental QALYs	Incremental costs (£)	ICER (£)	
No	16.11	641.06	_	_	_	
surveillance						
Colonoscopic surveillance in intermediate- and high-risk groups	16.16	841.54	0.05	200.49	4235.75	
Colonoscopic						
surveillance in						
all risk groups	16.26	1177.03	0.15	535.970	3669.70	
QALY: quality-adjusted life year; ICER: incremental cost-effectiveness ratio						

13

14 The cost-effectiveness plane for the base-case analysis is shown in figure 4.

1 Figure 4 Cost-effectiveness plane



3

2

These results indicate that surveillance of the intermediate- and high-risk groups (denoted as BSG in Figure 4) is extendly dominated by suveillance of all risk groups. This is because surveillance of all risk groups is associated with a lower ICER. These results suggest that surveillance of all risk groups (denoted as BSG + Low risk in Figure 4) is the most cost-effective strategy.

9 11.1.1 Transition matrices

Table 10 presents the results if the upper estimates for transition probabilitiesare used.

1 Table 10 Deterministic results with upper estimates for transition

2 probabilities

50-year time horizon	QALYs (utilities)	Costs (£)	Incremental QALYs	Incremental costs (£)	ICER (£)		
No surveillance	15.74	1532.85					
Intermediate- and high-risk groups	15.90	1468.76	0.16	-64.09	-388.66		
All risk groups	16.24	1229.93	0.50	-302.92	Extended dominance		
QALY: quality-adjusted life year; ICER: incremental cost-effectiveness ratio.							

3

- 4 Table 11 presents the results if the lower estimates for transition probabilities
- 5 are used.

6 Table 11 Deterministic results with lower estimates for transition

7 probabilities

50-year time horizon	QALYs (utilities)	Costs (£)	Incremental QALYs	Incremental costs (£)	ICER (£)	
No surveillance	16.259	38.91				
Intermediate- and high-risk groups	16.261	420.20	0.002	381.28	191,602	
All risk groups	16.265	1151.88	0.006	1112.97	181,288.36	
QALY: quality-adjusted life year; ICER: incremental cost-effectiveness ratio.						

8

- 9 The natural history transitions have a significant impact on the estimates of
- 10 cost effectiveness. However, the deterministic results of cost effectiveness
- 11 were consistent when colonoscopic surveillance in intermediate- and high-risk
- 12 groups was a cost-effective strategy compared with no surveillance.

13 **11.1.2** Potential disutility associated with colonoscopy

- 14 The GDG agreed that potential discomfort and recovery from sedation
- 15 associated with colonoscopy would have an effect on the QALYs gained. A
- 16 potential disutility of 0.0025 was used in the base-case analysis to explore the
- 17 impact of disutility on the ICERs (see table 12).

Strategy	QALYs (utilities)	Costs (£)	Incremental QALYs	Incremental costs (£)	ICER (£)	
No surveillance	16.07	641.06	-	-		
Intermediate- and high-risk groups	16.12	841.54	0.05	200.49	4242.84	
All risk groups		1177.0				
	16.22	3	0.15	535.97	3675.82	
QALY: quality-adjusted life year; ICER: incremental cost-effectiveness ratio.						

1 Table 12 Disutility of 0.0025 associated with colonoscopy

2

3 The GDG discussed the potential psychological and physical impacts of

4 colonoscopy, including anxiety and discomfort. It was agreed that despite the

5 inconvenience related to full bowel preparation and the recovery time after

6 each procedure, the long-term benefit of colonoscopic surveillance outweighs

7 the short-term discomfort. The estimated ICERs for each strategy showed

8 little variation. Therefore surveillance following the BSG guidelines and

9 including the low-risk group remained more cost effective.

10 **11.2** Structural sensitivity analysis

11 **11.2.1** Age of the cohort

12 The age of the cohort was varied from 50 to 35, 40 and 45 years and

13 surveillance was stopped at 85 years for each strategy. The model was run for

14 50 years to see the costs and health benefits of surveillance over a lifetime for

each strategy. Table 13 outlines the results. The overall trends of ICER

16 estimates show that colonoscopic surveillance in all risk groups is a cost-

17 effective strategy compared with no surveillance at £20,000 and £30,000 per

18 QALY gained. The results indicate that the younger the cohort, the more cost-

19 effectiveness the strategy. This is an important consideration when examining

20 other published cost-effectiveness analyses because most examine a cohort

21 age of 50 years. However, the transitions from adenomas to colorectal cancer

22 were assumed to be constant. Therefore there is some uncertainty about the

23 results for cohorts younger than the base case.

Age of cohort			Costs	Incremental	Incrementa	
(years)	Strategy	QALYS	(£)	QALYS	l costs (£)	ICER (£)
35	No surveillance	19.41	1095.10			
	Intermediate- and					
	high-risk groups	19.51	1172.34	0.10	77.24	772.44
	All risk group	20.71	1229.36	0.32	134.26	419.44
40	No surveillance	18.54	943.39			
	Intermediate- and					
	high-risk groups	18.63	1061.51	0.08	118.11	1416.22
	All risk group	18.81	1218.69	0.26	275.30	1040.05
45	No surveillance	17.43	791.33			
	Intermediate- and					
	high-risk groups	17.50	951.16	0.06	159.83	2458.97
	All risk group	17.63	1202.24	0.20	410.91	2016.25
QALY: qu	ality-adjusted life year;	ICER: incr	emental cost	-effectiveness rati	0.	

1 Table 13 ICER estimates when varying age of cohort

2 **11.2.2 Stopping surveillance at different ages**

- 3 Table 14 outlines the results of stopping surveillance at different ages over a
- 4 lifetime horizon (50 years).

Stopping				Incrementel	Incrementel				
age (years)	Strategy	QALYs	Costs (£)	QALYs	costs (£)	ICER (£)			
65	No surveillance	16.11	641.06						
	Intermediate - and high- risk groups	16.16	841.54	0.047	200.49	4235.38			
	All risk groups	16.25	1127.48	0.142	486.42	3414.14			
70	No surveillance	16.11	641.06						
	Intermediate - and high- risk groups	16.16	841.54	0.047	200.49	4235.75			
	All risk groups	16.26	1155.91	0.145	514.85	3543.54			
75	No surveillance	16.11	641.06						
	Intermediate - and high- risk groups	16.16	841.54	0.052	200.49	4235.45			
	All risk groups	16.26	1169.41	0.1506	528.36	3620.90			
80	No surveillance	16.11	641.06						
	Intermediate - and high- risk groups	16 16	941 54	0.05	200.40	4225 75			
	All risk	10.16	041.34	0.05	200.49	4230.70			
	groups	16.26	1175.22	0.15	534.17	3657.73			
	QALY: quality-a	QALY: quality-adjusted life year; ICER: incremental cost-effectiveness ratio.							

1 Table 14 Stopping surveillance at different ages

3 The results show that stopping surveillance at 60, 65 or 75 years has little

4 impact on ICERs and the results are consistent with the base-case results.

5 Surveillance in all risk groups is therefore a cost-effective strategy.

6 11.3 Probabilistic sensitivity analysis

7 Probabilistic sensitivity analysis enables the uncertainty associated with

8 parameters to be reflected in the results of the model. In non-linear decision

9 models, probabilistic sensitivity analysis provides the best estimates of mean

- 1 costs and health consequences in terms of QALYs gained. Table 15 outlines
- 2 the results. The costs are slightly higher and given the low incremental QALYs
- 3 do cause the ICERs to increase compared with the deterministic results, but
- 4 not significantly.

5	Table 15	Probabilistic	sensitivity	analysis	over a S	50-year	period
---	----------	---------------	-------------	----------	----------	---------	--------

	QALYs	Costs (£)	Incremental QALYs	Incremental costs (£)	ICER (£)	Probability of being cost effective at £20,000 per QALY gained (%)
No surveillance	16.12	562.91	_	_	_	_
Intermediate- and high-risk groups	16.17	786.25	0.04	223.33	5298.03	78
All risk groups	16.25	1167.77	0.13	604.85	4626.57	81
QALY: quality-adjusted life year; ICER: incremental cost-effectiveness ratio.						

7 11.3.1 Cost-effectiveness plane

8 Figures 5 and 6 show the results of the probabilistic sensitivity analysis plotted

- 9 on a graph of incremental costs and QALYs. It appears that effectiveness and
- 10 cost are negatively correlated. That is, the more progressive the condition, the
- 11 more people develop colorectal cancer, and therefore, the greater the
- 12 potential savings from reduced surveillance. It is also apparent that
- 13 surveillance in all risk groups is associated with greater variation in values, but
- 14 also potentially greater gains.

- 1 Figure 5 Cost-effectiveness (CE) plane intermediate-risk (IR) and high-
- 2 risk (HR) group surveillance





2 Figure 6 Cost-effectiveness (CE) plane – all risk groups

3

1

5 **11.3.2 Cost-effectiveness acceptability curves**

Figure 7 presents the cost-effectiveness acceptability curves for the different
surveillance strategies. At a threshold of £20,000 per QALY gained,
colonoscopic surveillance in the intermediate- and high-risk groups is
associated with a probability of being cost effective of over 78% compared
with no surveillance. In all risk groups the probability of being cost effective is

11 81%. At £30,000 per QALY gained these figures increase to 87% and 88%

12 respectively.

13

- 1 Figure 7 The cost-effectiveness acceptability curve for different
- 2 surveillance strategies
- 3



- 5
- 6 The results support findings from the base-case analysis that surveillance of
- 7 all risk groups is the preferred option because it is associated with the lowest
- 8 ICERs and the least uncertainty.

9 **11.3.3** Cost-effectiveness acceptability frontiers

- 10 Figure 8 presents the cost-effectiveness acceptability frontiers for the different
- 11 surveillance strategies.

- **1** Figure 8 Cost-effectiveness acceptability frontiers for different
- 2 surveillance strategies



- 4 These results indicate that at £20,000 per QALY gained and £30,000 per
- 5 QALY gained the optimum strategy is the all risk groups surveillance strategy.

6 12 Discussion

7 12.1 Strengths of the model

- 8 The main strength of the model is its comprehensiveness, using the most up-
- 9 to-date evidence available in the public domain. Extensive sensitivity analyses
- 10 were performed to explore any uncertainty in the data and the model. The
- 11 model included projected health benefits and related resource use following
- 12 the BSG guidelines, taking into account different recurrence rates of
- 13 adenomas in the NHS.

1 **12.2** *Limitations of the model*

2 12.2.1 Natural history data

3 Because of a lack of time, a systematic review was not carried out examining

- 4 the natural history of the progression of adenomas into colorectal cancer.
- 5 However, the GDG agreed to use assumptions consistent with a published
- 6 analysis by Tappenden et al. (2004). Although the analysis by Tappenden et
- 7 al. would not have taken into account newly published evidence, it was
- 8 confirmed in the recently updated BSG guidelines (Cairns et al. 2010) that
- 9 there is no new evidence associated with polyps and adenoma surveillance.
- 10 The model focused on colonoscopic surveillance and so different treatment
- 11 options and chemoprevention for stage-specific colorectal cancer were not
- 12 distinguished in the model because of a lack of time and resources. Ideally
- 13 these options would have been included in the model to show different health
- 14 benefits and subsequent resource use.

15 **12.2.2 Clinical data**

- 16 Limitations include the lack of directly observed progression and regression
- 17 data for the development of adenomas. The transition probabilities in the
- 18 model were obtained from Tappenden et al. (2004). Transferring these data to
- 19 another model was not ideal and there was potential uncertainty.
- 20 In the model it was assumed that all colorectal cancers arise from pre-existing
- 21 adenomas. However, direct evidence suggests that new colorectal cancers
- can also arise. This assumption led to bias in favour of surveillance comparedwith no surveillance.

24 **12.2.3 Misdiagnosis**

- 25 For adenoma detection, 100% sensitivity and specificity were assumed. The
- 26 GDG discussed the current sensitivity and specificity of colonoscopy to be
- around 95%. In addition, clinical data were mainly obtained from observational
- studies in which misdiagnosis was accounted for in the published literature.

- 1 Further work could incorporate the sensitivity and specificity of the chosen
- 2 surveillance method where appropriate.

3 12.2.4 Complications

The probabilities of perforation during colonoscopy with and without adenoma
removal were reported to be 0.17% and 0.08% respectively (Tappenden et al.
2004). Because of a lack of time and resources these complications were not
considered in the model.

8 12.2.5 Quality of life data

9 Uncertainty remains about the appropriate method to account for quality of life

- 10 for people with polyps and colorectal cancer. The patient experts and clinical
- 11 specialists in the GDG considered that the psychological burden of being
- 12 diagnosed with adenomas at high risk of progressing to colorectal cancer
- 13 could be very high. The GDG also highlighted the discomfort and
- 14 inconvenience associated with full bowel preparation before colonoscopy and
- 15 the recovery period after each procedure. However, the GDG acknowledged
- 16 that referral for colonoscopic surveillance was broadly reassuring and not
- 17 associated with adverse psychological consequences in the long term (Miles
- 18 et al. 2009). More work will be required on the short- and long-term benefits of
- 19 colonoscopic surveillance in preventing colorectal cancer.

20 **12.2.6 Surveillance using colonoscopy**

21 The updated BSG guidelines (Cairns et al. 2010) highlighted the user-

- 22 dependency of colonoscopy and the importance of careful and thorough
- 23 colonoscopy in preventing colorectal cancer with a 'fail-safe system' in place
- 24 for recall of patients at high risk.

25 **12.2.7 Costing**

- 26 NHS reference costs are published costs and represent the average NHS
- 27 costs across the country. However, the GDG highlighted that these costs
- 28 could potentially underestimate the true cost of the procedure. This was
- 29 explored by increasing the costs in the deterministic sensitivity analysis. It
- 30should be noted that the incremental costs are the most important figures, not
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- 1 the absolute costs. A true micro-costing exercise in a UK setting would have
- 2 been the preferred option.

3 12.2.8 Systematic reviews

- Ideally systematic reviews would have been carried out for all model inputs so
 that the most robust evidence was selected. However, the GDG agreed that
- 6 the approach was acceptable given the limited time and resources for
- 7 guideline development.

8 12.2.9 Full care pathway modelling

9 The current analysis simplifies the actual treatment by modelling identical

- 10 treatment pathways for stage-specific colorectal cancer. It was necessary to
- 11 explore the cost effectiveness of colonoscopic surveillance for the detection
- 12 and prevention of colorectal cancer in the given timeframe. The model does
- 13 not take into account the possibility of a person progressing between
- 14 treatments, loss to follow-up or colorectal cancer arising from other causes. If
- 15 improved clinical-effectiveness data were to be collected, these should be
- 16 included in a more comprehensive model in the future to allow a more detailed
- 17 comparison to be made.

18 **13** Conclusions

- 19 This analysis indicates that colonoscopic surveillance in all risk groups is the
- 20 most cost-effective strategy for people with adenomas at high risk of
- 21 developing colorectal cancer. ICER estimates below £20,000 and £30,000 per
- 22 QALY gained are apparent when deterministic and probabilistic analyses are
- 23 considered. However, the GDG acknowledged that there was uncertainty
- 24 about the clinical benefits of colonoscopic surveillance in the low-risk group.
- 25 The GDG discussed the potential risks of perforation and bleeding associated
- 26 with colonoscopy and adenoma removal in the low-risk group, which could
- 27 outweigh potential benefits (Ransohoff et al. 1991). In the absence of
- 28 evidence for increased detection of adenomas and colorectal cancer leading
- 29 to reduced mortality in the low-risk group, the GDG agreed that colonoscopic
- 30 surveillance in this group would not be recommended as routine practice in
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- 1 the NHS. The GDG highlighted, however, that clinical judgement should be
- 2 used for people with small adenomas (≤5 mm): their age, co-morbidities,
- 3 potential risks of bleeding and perforation should be considered.

4 **14** Future work

5 A better understanding of the natural history of colonic polyps and the 6 progression of adenomas to colorectal cancer is a priority so that the full 7 course of the disease, from diagnosis to the stage-specific treatments for 8 colorectal cancer, can be modelled in the future. Therefore, the potential for 9 discrete event simulation should be considered to make modelling less time 10 consuming.

- 11 Carrying out audits of current surveillance for people with adenomas will
- 12 provide valuable data for identifying gaps in the evidence, training and
- 13 development needs in clinical practice, as well as the provision of patient
- 14 information. Audit should include colonoscopy adherence, complications
- 15 associated with colonoscopy, a breakdown of possible causes of
- 16 complications, and the outcomes and additional techniques used when the
- 17 results of colonoscopy are inconclusive and/or incomplete. Audit will also
- 18 provide information about areas for training needs.
- 19 Ongoing research on the long-term safety of a no surveillance strategy for
- 20 people at low risk of developing colorectal cancer is expected to report
- 21 outcomes in the next 2 years (Cairns et al. 2010). This will provide invaluable
- 22 evidence for future guidance development.
- 23 The NHS Bowel Cancer Screening Programme was fully rolled out in 2009
- 24 and so reports and outcomes will be available soon. Careful consideration and
- 25 further study of the relationship between the population eligible for the
- screening programme and the colonoscopic surveillance population are
- 27 needed. This will ensure that the most appropriate and timely interventions
- are in place for reducing mortality associated with colorectal cancer and
- 29 improving relevant health benefits in the NHS.

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1

2 16 Appendices

3 16.1 Inclusion and exclusion criteria

Studies identified (IBD and Polyps) N=289

Studies of potential interest N=55

Studies included (0 IBD; 1 polyps) N=1 Excluded based on title/abstract review N=234

Excluded studies based on NICE reference case N=54

4

5 16.2 Review of Tappenden et al. (2004)

6 The objective of the report was to conduct a detailed assessment of research

7 evidence and to develop a mathematical model to estimate the costs, benefits

- 8 and capacity implications of alternative screening options for colorectal cancer
- 9 in England. As part of the report, the authors considered subsequent
- 10 colonoscopic surveillance in people with high-risk polyps at index
- 11 colonoscopy, which broadly follows the current BSG guidelines.





1

- 2 In this model people are allocated to a risk state based on a baseline
- 3 colonoscopy: low risk, intermediate risk or high risk. People can then progress
- 4 or regress in each diagnostic state and will stay there until surveillance re-
- 5 classifies them or until they develop cancer. If there is no surveillance then
- 6 colorectal cancer is only diagnosed when the person becomes symptomatic.
- 7 Asymptomatic cancer can be detected by surveillance. Death from other
- 8 causes is based on age-related mortality. This model does not include
- 9 misdiagnosis from surveillance, but allows an initial misdiagnosis at baseline
- 10 colonoscopy, because the natural history data take misdiagnosis into account.
- 11 The overall quality of the report was very high and all assumptions and
- 12 variables were justified. The possible limitations of the report are that the
- 13 surveillance strategies examined include faecal occult blood testing, flexible
- 14 sigmoidoscolonoscopy, and colonoscopy in a general population. The
- 15 population for this analysis was people with polyps who are at high risk of
- 16 developing colorectal cancer.
- 17

16.3 Quality checklist for Tappenden et al. (2004) study

2

1

Study name	Colorectal cancer screening options appraisal: cost effectiveness, cost–utility and resource impact of alternative options for colorectal concer (2004)				
	Tappenden P, E	ggington S, Nixon R et al.			
Study question	Grade (yes/ no/ not clear/ N/A)	Comments			
	Study design				
1. Was the research question stated?	Yes				
2. Was the economic importance of the research question stated?	Yes				
3. Was/were the viewpoint(s) of the analysis clearly stated and justified?	Yes				
4. Was a rationale reported for the choice of the alternative programmes or interventions compared?	Yes				
5. Were the alternatives being compared clearly described?	Yes				
6. Was the form of economic evaluation stated?	Yes				
7. Was the choice of form of economic evaluation justified in relation to the questions addressed?	Yes				
Data collection					
8. Was/were the source(s) of effectiveness estimates used stated?	Yes	From systematic review and additional published studies			
9. Were details of the design and results of the effectiveness study given (if based on a single study)?	Yes				
10. Were details of the methods of synthesis or meta-analysis of estimates given (if based on an overview of a number of effectiveness studies)?	Yes	Because of a lack of RCT evidence no meta-analysis was conducted, but the means of obtaining probabilities were stated.			

11. Were the primary outcome measure(s) for the economic evaluation clearly stated?	Yes	
12. Were the methods used to value health states and other benefits stated?	Yes	In the absence of utility values in stage-specific colorectal cancer using EQ-5D as the preferred method in line with the NICE reference case, utility estimates were used from published sources that used standard gamble
13. Were the details of the subjects from whom valuations were obtained given?	Yes	
14. Were productivity changes (if included) reported separately?	N/A	
15. Was the relevance of productivity changes to the study question discussed?	N/A	
16. Were quantities of resources reported separately from their unit cost?	No	Use of NHS reference costs implies that there is no requirement to separately calculate unit costs as all costs are included in estimates.
17. Were the methods for the estimation of quantities and unit costs described?	Yes	NHS reference cost codes quoted where possible. Uplifted treatment cost data for stage- specific colorectal cancer were obtained from personal communications.
18. Were currency and price data recorded?	Yes	
19. Were details of price adjustments for inflation or currency conversion given?	Yes	
20. Were details of any model used given?	Yes	
21. Was there a justification for the choice of model used and the key parameters on which it was based?	Yes	
Analysis	and interpretation	of results
22. Was the time horizon of cost and benefits stated?	Yes	
23. Was the discount rate stated?	Yes	

24. Was the choice of rate justified?	Yes	All costs and health outcomes are discounted at 3.5% per year as recommended by NICE.
25. Was an explanation given if costs or benefits were not discounted?	N/A	
26. Were the details of statistical test(s) and confidence intervals given for stochastic data?	Yes	
27. Was the approach to sensitivity analysis described?	Yes	
28. Was the choice of variables for sensitivity analysis justified?	Yes	
29. Were the ranges over which the parameters were varied stated?	Yes	
30. Were relevant alternatives compared? (That is, were appropriate comparisons made when conducting the incremental analysis?)	Yes	
31. Was an incremental analysis reported?	Yes	
32. Were major outcomes presented in a disaggregated as well as an aggregated form?	Yes	
33. Was the answer to the study question given?	Yes	
34. Did conclusions follow from the data reported?	Yes	
35. Were conclusions accompanied by the appropriate caveats?	Yes	
36. Were generalisability issues addressed?	Yes	
Adapted from Drummond and Jeffers Dissemination (2008).	son (1996). Cited in (Centre for Reviews and

1

2

1 **16.4 Quality checklist for new cost-effectiveness analysis**

Guideline topic: colonoscopic surveillance in polyps by Y Rajput					
Cost-effectiveness modelling for colonoscopic surveillance in people with adenomas by K Jeong 2010					
Section 1: Applicability		Yes/ par no/uncle NA	tly/ ear/	Comments	
1.1 Is the study population appropriate the guideline?	e for	Yes		50-year old men and women who have adenomas removed at baseline colonoscopy with a high risk of developing colorectal cancer	
1.2 Are the interventions appropriate f the guideline?	or	Yes		All clinically effective interventions/strategies included within the scope	
1.3 Is the healthcare system in which study was conducted sufficiently simila the current UK NHS context?	the ar to	Yes			
1.4 Are costs measured from the NHS personal social services (PSS) perspective?	S and	Yes			
1.5 Are all direct health effects on individuals included?		Partly		QALY data from USA using standard gamble technique, there is very limited evidence available on the colorectal cancer stage-specific utility data	
1.6 Are both costs and health effects discounted at an annual rate of 3.5%?)	Yes			
1.7 Is the value of health effects expre in terms of quality-adjusted life years (QALYs)?	Yes				
1.8 Are changes in health-related qua life (HRQoL) reported directly from pa and/or carers?	Yes				
1.9 Is the valuation of changes in HRC (utilities) obtained from a representative sample of the general public?	No		QALY data from USA using standard gamble technique used		
1.10 Overall judgement: Directly appli There is no need to use section 2 of the applicable' Directly applicable Other comments	icable/p e check	artially ap list if the s	plicat tudy	ble/not applicable is considered 'not	
Section 2: Study limitations (the level of methodological quality) This checklist should be used once it has	Yes/pa no/uno NA	artly/ clear/	Con	nments	

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been decided that the study is sufficiently applicable to the context of the clinical quideline	Comments	
2.1 Does the model structure adequately reflect the nature of the health condition under evaluation?	Yes	
2.2 Is the time horizon sufficiently long to reflect all important differences in costs and outcomes?	Yes	45-year time horizon, uncertainty verified using different starting age of cohort (50, 55, 60, 65 years)
2.3 Are all important and relevant health outcomes included?	Yes	
2.4 Are the estimates of baseline health outcomes from the best available source?	Yes	
2.5 Are the estimates of relative treatment effects from the best available source?	Yes	Best quality studies identified from clinical review
2.6 Are all important and relevant costs included?	Yes	
2.7 Are the estimates of resource use from the best available source?	Yes	NHS specific
2.8 Are the unit costs of resources from the best available source?	Yes	
2.9 Is an appropriate incremental analysis presented or can it be calculated from the data?	Yes	
2.10 Are all important parameters whose values are uncertain subjected to appropriate sensitivity analysis?	Yes	
2.11 Is there no potential conflict of interest?	No	
2.12 Overall assessment : Minor limit serious limitations Minor limitations	ations/potentially	serious limitations/very

16.5	Modified GRADE for health economic literature
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	Ref ID	Country	Population	Comparators	Outcome measure	Study design	Cost-effectiveness results (base case)	Applicability
Tappenden et al. (2004)	Identified through lateral search	UK	Cohort at age 30	Biennial FOBT 50– 69 years; biennial FOBT 60– 69 years; FSIG once at 55 years; FSIG once at 60 years; FSIG once at 60 years, followed by biennial FOBT 61–70 years	QALY	DES	Screening using FOBT and/or FSIG is potentially a cost-saving strategy for the early detection of colorectal cancer. However, the practical feasibility of alternative screening programmes is inevitably limited by current pressures on endoscopy services.	Applicable