Articles

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Ovarian cancer screening and mortality in the UK Collaborative Trial of Ovarian Cancer Screening (UKCTOCS): a randomised controlled trial

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Summary

Background Ovarian cancer has a poor prognosis, with just 40% of patients surviving 5 years. We designed this trial to establish the effect of early detection by screening on ovarian cancer mortality.

Methods In this randomised controlled trial, we recruited postmenopausal women aged 50–74 years from 13 centres in National Health Service Trusts in England, Wales, and Northern Ireland. Exclusion criteria were previous bilateral oophorectomy or ovarian malignancy, increased risk of familial ovarian cancer, and active non-ovarian malignancy. The trial management system confirmed eligibility and randomly allocated participants in blocks of 32 using computer-generated random numbers to annual multimodal screening (MMS) with serum CA125 interpreted with use of the risk of ovarian cancer algorithm, annual transvaginal ultrasound screening (USS), or no screening, in a 1:1:2 ratio. The primary outcome was death due to ovarian cancer by Dec 31, 2014, comparing MMS and USS separately with no screening, ascertained by an outcomes committee masked to randomisation group. All analyses were by modified intention to screen, excluding the small number of women we discovered after randomisation to have a bilateral oophorectomy, have ovarian cancer, or had exited the registry before recruitment. Investigators and participants were aware of screening type. This trial is registered with ClinicalTrials.gov, number NCT00058032.

Findings Between June 1, 2001, and Oct 21, 2005, we randomly allocated 202638 women: 50640 ($25 \cdot 0\%$) to MMS, 50639 ($25 \cdot 0\%$) to USS, and 101359 ($50 \cdot 0\%$) to no screening. 202546 ($>99 \cdot 9\%$) women were eligible for analysis: 50624 ($>99 \cdot 9\%$) women in the MMS group, 50623 ($>99 \cdot 9\%$) in the USS group, and 101299 ($>99 \cdot 9\%$) in the no screening group. Screening ended on Dec 31, 2011, and included 345570 MMS and 327775 USS annual screening episodes. At a median follow-up of 11·1 years (IQR 10·0–12·0), we diagnosed ovarian cancer in 1282 (0·6%) women: 338 (0·7%) in the MMS group, 314 (0·6%) in the USS group, and 630 (0·6%) in the no screening group. Of these women, 148 (0·29%) women in the MMS group, 154 (0·30%) in the USS group, and 347 (0·34%) in the no screening group had died of ovarian cancer. The primary analysis using a Cox proportional hazards model gave a mortality reduction over years 0–14 of 15% (95% CI –3 to 30; p=0·10) with MMS and 11% (–7 to 27; p=0·21) with USS. The Royston-Parmar flexible parametric model showed that in the MMS group, of 2% (–27 to 26) in years 0–7 and 23% (1–46) in years 7–14, and in the USS group, of 2% (–27 to 26) in years 0–7 and 21% (–2 to 42) in years 7–14. A prespecified analysis of death from ovarian cancer of MMS versus no screening with exclusion of prevalent cases showed significantly different death rates (p=0·021), with an overall average mortality reduction of 8% (–27 to 43) in years 0–7 and 28% (–3 to 49) in years 7–14 in favour of MMS.

Interpretation Although the mortality reduction was not significant in the primary analysis, we noted a significant mortality reduction with MMS when prevalent cases were excluded. We noted encouraging evidence of a mortality reduction in years 7–14, but further follow-up is needed before firm conclusions can be reached on the efficacy and cost-effectiveness of ovarian cancer screening.

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Introduction

The poor prognosis for ovarian cancer¹ motivated us to start a programme of screening research 30 years ago.² We have since reported CA125 as a predictor of ovarian cancer risk,^{3,4} high specificity² and preliminary evidence of a survival benefit⁵ of multimodal screening using CA125 interpreted with a cutoff with transvaginal ultrasound as a second-line test, development of a risk of ovarian cancer algorithm



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Research in context

Evidence before this study

During the 1990s, findings from large prospective studies of screening showed that both CA125 and ultrasound-based ovarian cancer screening could identify preclinical cases of ovarian cancer. These studies included some by our own group with use of a multimodal strategy incorporating CA125 with ultrasound as a secondary test. In the year before the start of this trial, a systematic review of 25 ovarian cancer screening studies commissioned by the National Health Service Health Technology Assessment Programme reported that although ultrasound and multimodal screening can detect ovarian cancer in asymptomatic women, the effect of screening on ovarian cancer was unproven. The authors concluded that screening should not be introduced into clinical practice until further information was available from randomised trials designed to assess the effect of ovarian cancer screening on mortality and its adverse effects and cost-effectiveness. After that publication, our group reported the findings of a pilot randomised controlled trial in 22000 postmenopausal women showing the feasibility of ovarian cancer screening with use of a multimodal screening strategy and provided preliminary evidence of a survival benefit in this population. During this trial, two large randomised controlled trials were done: the Shizuoka Cohort Study of Ovarian Cancer Screening and the ovarian component of the Prostate Lung Colorectal Ovarian (PLCO) Cancer Screening Trial. The Shizuoka Cohort Study of Ovarian Cancer Screening only reported on ovarian cancer detection and not on deaths. Findings from the ovarian component of the PLCO Cancer Screening Trial showed no difference in ovarian cancer deaths between the screening and control groups. We searched MEDLINE between Jan 1, 2001, and Nov 31, 2015, using the protocol described by Mosch and colleagues, with the following

search terms: "ovarian neoplasms", "Fallopian tube neoplasms", "ovar*", "fallopian tub* OR adnex*", "tumo*", "malignan*", "carcinoma* OR adenocarcinoma* OR neoplasm* OR mass*", "mass screening", "early detection of cancer", "randomized controlled trial", "controlled clinical trial", "randomized", "placebo", "clinical trials", "randomly", and "trial". This search yielded 234 publications, which, when limited to randomised controlled trials of human female adults published in the English language resulted in 64 articles, 11 of which were duplicated. The remaining 53 articles consisted of 28 pertaining to the PLCO Cancer Screening Trial, 11 from our own group, and one from Kobayashi and colleagues in 2008 (the Shizuoka Cohort Study of Ovarian Cancer Screening). We identified three key randomised controlled trials in ovarian cancer screening. Of these, only the PLCO Cancer Screening Trial thus far has reported mortality data.

Added value of this study

To our knowledge, this trial is the first randomised controlled trial of ovarian cancer screening to produce findings that show that in postmenopausal women from the general population, annual screening with use of the multimodal strategy is safe and could reduce deaths due to ovarian cancer. These findings are derived from one of the largest randomised trials ever done and renew hope that death rates from the most lethal of all gynaecological malignancies can be reduced through early detection.

Implications of all the available evidence

Our findings suggest that a multimodal approach to screening might detect ovarian cancer sufficiently early to reduce mortality. To establish the magnitude of this reduction in deaths, a longer duration of follow-up is needed. Meanwhile, efforts to refine ovarian cancer screening strategies should continue.

(ROCA) for interpretation of longitudinal CA125,⁶⁷ use of morphological criteria in second-line transvaginal ultrasound,⁸ and use of ROCA in a pilot randomised controlled trial.⁹ During this period, advances in treatment have only produced a slight reduction in disease mortality.^{10,11}

This research led us in 2001 to undertake a definitive randomised controlled trial, the UK Collaborative Trial of Ovarian Cancer Screening (UKCTOCS), of more than 200000 women.¹² The primary aim was to assess the effect of screening on disease mortality. We have previously reported on performance characteristics of screening,^{13,14} harms related to false-positive surgery,^{13,14} and the psychological morbidity associated with screening.¹⁵ This report is a key landmark for the programme, providing the first ovarian cancer mortality data from UKCTOCS.

Methods

Study design and participants

In this randomised controlled trial, we recruited women through 13 regional centres (RCs) in National Health Service (NHS) Trusts in England, Wales, and Northern Ireland, with the Queen Mary University of London as the coordinating centre (CC) between 2001 and 2004 and then University College London from 2004 onwards. We invited women identified through the age-sex registers of 27 participating primary care trusts within the RC catchment areas. We commissioned specialised software from the NHS to randomly select women aged 50-74 years and then flag them on primary care trusts' registers and allow electronic transfer of their personal and general practice details to the CC in lots of 5000 to 10000 every quarter. We then sent women personal invitations and logged replies on the trial management system. Women attended a recruitment clinic at the RC where they viewed an information video, completed a recruitment questionnaire, and provided written consent and a baseline serum sample. We scanned recruitment questionnaires at the CC into a bespoke trial management system.12 Eligibility criteria were 50-74 years of age and postmenopausal status. Exclusion criteria were self-reported previous bilateral oophorectomy or ovarian malignancy, increased risk of familial ovarian cancer, or active non-ovarian malignancy.

The trial design has been previously published¹²⁻¹⁴ and the protocol is available online. Ethical approval was by the UK North West Multicentre Research Ethics Committees (North West MREC 00/8/34).

Randomisation and masking

The trial management system confirmed eligibility and then randomly allocated women to annual screening using a multimodal screening (MMS) or ultrasound screening (USS) strategy or no screening in a 1:1:2 ratio. The randomisation was accomplished using the Visual Basic randomisation statement and the Rnd function. The trial management system allocated a set of 32 random numbers to each RC, of which eight were allocated to MMS, eight to USS, and the remaining 16 to no screening. We randomly allocated each successive volunteer within the RC to one of the numbers and subsequently randomly allocated them into a group. Investigators and participants were aware of screening type, but the outcomes committee was masked.

Procedures

Annual screening in the MMS group used serum CA125 concentration testing, with the pattern over time interpreted with use of the risk of ovarian cancer calculation, which identifies significant rises in CA125 concentration above baseline.67,9 Next, ROCA triaged women to normal (annual screening), intermediate (repeat CA125 concentration testing in 3 months), and elevated (repeat CA125 concentration testing and transvaginal USS as a second-line test in 6 weeks) risk. Annual screening in the USS group used transvaginal USS as the primary test, which was classified as normal (annual screening), unsatisfactory (repeat in 3 months), or abnormal (scan with a senior ultrasonographer within 6 weeks).13 In both groups, women with persistent abnormalities had clinical assessment and additional investigations within the NHS by a trial clinician.^{13,14} We deemed women who had surgery or a biopsy for suspected ovarian cancer after clinical assessment screen positive.13,14

Screening was implemented centrally by the CC using a web-based trial management system, which ensured that protocol deviations were kept to a minimum.¹² A quality assurance programme for transvaginal USS and accreditation for scanning of postmenopausal ovaries was overseen by the ultrasound committee.¹⁶ We linked women using their NHS number in England and Wales to the Health and Social Care Information Centre for cancer and death registrations and, in Northern Ireland, to the Central Services Agency and Northern Ireland Cancer Registry. For women from English RCs, we also obtained data for cancer diagnosis from the National Cancer Intelligence Network, and between April, 2001, and March, 2010, Hospital Episodes Statistics administrative records. Other sources were two postal follow-up questionnaires (3-5 years after randomisation and April, 2014) and direct communication from trial participants, their families, and physicians.

We interrogated all available data sources to identify women diagnosed after randomisation with any of 19 International Classification of Diseases (ICD)-10 codes (appendix p 3).14 We retrieved copies of medical notes for all except those with ICD-10 C80 (malignant neoplasm of uncertain origin) who also had another specific non-ovarian or non-peritoneal cancer registration (appendix p 25). All were reviewed by an outcomes review committee (two pathologists and two gynaecological oncologists) who were masked to the randomisation group. We used an algorithm to assign the final diagnosis. We based death due to ovarian cancer on disease progression (appearance of new lesions or increases in size of previously documented lesions with imaging, clinical worsening, or rising biomarker concentrations). We ascertained ovarian and adnexal surgery outside of the trial after randomisation through both self-reporting and Hospital Episodes Statistics records.

We also ascertained contamination in the no screening group. This contamination was based on self-reporting in the 2014 follow-up questionnaire in which women in the no screening group were asked whether they had had ovarian cancer screening after recruitment.

The original trial protocol specified six annual screens and follow-up for 7 years from randomisation.¹² In 2008, an analysis of overall and cause-specific standardised mortality in the no screening group showed a lower than expected mortality rate. We therefore extended screening in the USS and MMS groups to Dec 31, 2011, resulting in women being offered 7–11 screens depending on the year of randomisation. Follow-up was extended to Dec 31, 2014.

Outcomes

The primary outcome was ovarian cancer death by Dec 31, 2014, assessed with use of data from the sources described above. Ovarian cancer refers to malignant neoplasms of the ovary (ICD-10 C56), which include primary non-epithelial ovarian cancer, borderline epithelial ovarian cancer, and invasive epithelial ovarian cancer; malignant neoplasms of the fallopian tube (ICD-10 C570); and undesignated malignancies of the ovaries, fallopian tube, or peritoneum.¹⁷ It does not include primary peritoneal cancer, which was diagnosed on the basis of WHO 2003 criteria.¹⁸

A secondary outcome was death due to ovarian and primary peritoneal cancer (ICD-10 C48.1 and C48.2). Most peritoneal cancers are likely to be classified as tubal and ovarian cancer once wider acceptance of the WHO 2014 revision has occurred.¹⁹ Another secondary outcome was compliance with screening, which was the proportion of women who attended all tests that consisted of a screening episode out of the total who were eligible for that screening episode. A further secondary outcome was complications related to screening and false-positive Correspondence to: Prof lan J Jacobs, University of New South Wales, Sydney, NSW 2052, Australia **i jacobs@unsw.edu.au** Prof Usha Menon, Department of Women's Cancer, Institute for Women's Health, University College London, London W1T 7DN, UK **u.menon@ucl.ac.uk**

For the **protocol** see http://www. instituteforwomenshealth.ucl. ac.uk/womens-cancer/gcrc/ ukctocs/files/ukctocs_protocol71

See Online for appendix

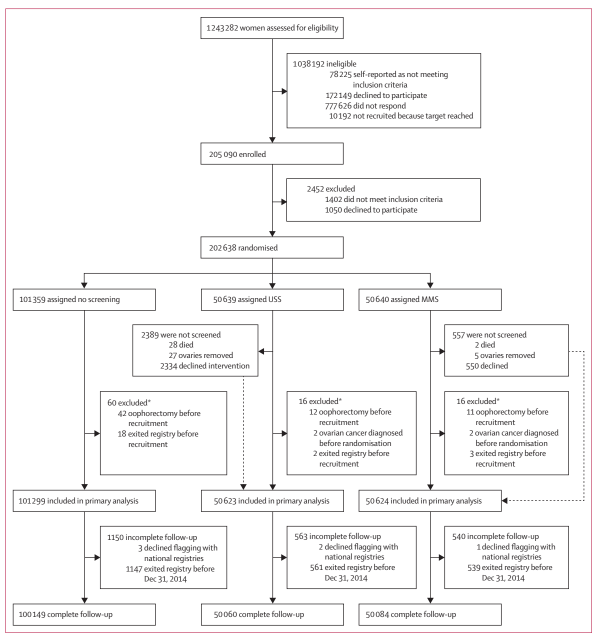


Figure 1: Trial profile

MMS=multimodal screening. USS=ultrasound screening. *Events occurred before recruitment, but discovered after randomisation.

surgery. Centres reported screening-related complications to a designated safety officer. We deemed screenpositive surgery resulting in benign pathology or normal adnexa false positive. Medical notes were centrally reviewed for complications, which were classified as major (resulting in sequelae) or minor by a designated gynaecological oncologist. Other secondary outcomes were to assess and compare the performance characteristics of the two screening strategies, assess the psychological effects of screening, and establish the resource implications of screening.

Statistical analysis

In 2000, we estimated that a sample size of 200 000 women at a two-sided 5% significance level for a difference in relative ovarian cancer mortality of 30% would give 90% power for the no screening versus combined screening groups and 80% power for the no screening versus MMS or no screening versus USS comparisons. After trial extension to Dec 31, 2014, our recalculated power remained 80% at a two-sided 5% significance level to detect a reduction of 30% in no screening versus MMS or no screening versus USS comparisons.

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The primary mortality analysis was an MMS versus no screening and USS versus no screening analysis of the primary outcome with use of a Cox proportional hazards model. The primary analysis initially consisted of comparison of the combined MMS and USS with the no screening group and individual comparisons of MMS with no screening and USS with no screening. The assumption was that sensitivity of the two screening strategies would be similar. During the trial, the data monitoring and ethics committee concluded that this assumption was not the case. On the basis of differences in performance characteristics of the two strategies at the initial screen,13 we updated our primary statistical analysis to MMS versus no screening and USS versus no screening comparisons. We defined survival time from date of randomisation to date of death due to ovarian cancer or censorship, or sooner if the volunteer died of another cause or was lost to follow-up. To allow for the fact that we were comparing two intervention arms against a control group, we made a Dunnett correction for multiple testing against a control to the critical α (α =0.0258). Mortality reduction estimates are 1 – hazard ratio (HR) estimates.

Early detection is likely to have less effect on women with ovarian cancer before screening starts (prevalent cases) than on those who develop it after the start of screening. The typical CA125 concentration profile noted before diagnosis in ovarian cancer cases is a baseline level, a change point, and then a rising CA125 concentration (appendix pp 10, 25).¹⁴ We did a prespecified MMS versus no screening subgroup analysis excluding prevalent cases for which the estimated change point was before randomisation, showing that the cancer was present before screening began (appendix pp 10, 24). For the prespecified MMS versus no screening and USS versus no screening secondary outcome analysis of ovarian and peritoneal cancer deaths, we used a Cox proportional hazards model, Royston-Parmar (RP) model, and post-hoc weighted log-rank (WLR) test.

In the original statistical plan, we did not make provision for the delayed effect on mortality that has been reported in other screening trials.²⁰⁻²³ The delayed effect is due to the inherent delay from randomisation to diagnosis and then death. Other cancer screening trials, including the Prostate Lung Colorectal Ovarian (PLCO) Cancer Screening Trial²⁴⁻²⁷ and the National Lung Screening Trial,28 used a WLR test for the primary analysis to address this delay.^{29,30} In view of the precedent of the PLCO Cancer Screening Trial screening trial, we did a sole post-hoc WLR analysis for the primary outcome, applying the WLR test with the same choice of weights proportional to pooled ovarian cancer mortality (appendix p 11) as that used in the primary PLCO Cancer Screening Trial analysis.²⁴ We applied the prespecified RP method³¹ that can model proportional and non-proportional hazards due to delayed effects^{22,23} to investigate the hazard functions for each group. We estimated changing HRs across time32 and calculated

average HRs for a split midway between the first (0–7) and second (7–14) 7 years to summarise delayed effects.

We calculated the preliminary number needed to screen to prevent one death from ovarian cancer as the reciprocal of the absolute difference in cumulative mortality from ovarian cancer between the MMS and no screening groups. We calculated prespecified ovarian cancer survival in the no screening group from date of diagnosis to explore external validity by comparison with UK ovarian cancer survival statistics. In addition to non-parametric Kaplan-Meier plots, we investigated the underlying ovarian cancer incidence (hazard) rates for each group with RP models.

We did sensitivity analyses using both a prespecified Cox proportional hazards model and the post-hoc WLR to establish the robustness of the primary analysis. We did a data source analysis restricting the outcome to events that had a national cancer registration or death certification to address the possibility of under-reporting in the no screening group. We also did potential within-centre correlation analyses by stratifying the model by centre and using cluster-robust SEs. Finally, we did competing-risk

	MMS (n=50624)	USS (n=50 623)	No screening (n=101299)*	Overall (n=202 546)
Age at randomisation (years)	60·61 (56·03–66·15)	60·61 (55·99–66·15)	60·58 (55·97–66·15)	60·59 (55·99–66·15)
Time since last period at randomisation (years)	11·36 (5·26–18·49)	11·34 (5·24–18·46)	11·26 (5·22–18·46)	11·3 (5·23-18·47)
Duration of HRT use in those who were on HRT at randomisation (years)	8.09 (4.56–11.99)	8·15 (4·55–12·11)	8·17 (4·5–12·09)	8·15 (4·53–12·07)
Duration of OCP use in those who had used it (years)	5 (2–10)	5 (2–10)	5 (2–10)	5 (2–10)
Pregnancies <6 months	0 (0–1)	0 (0–1)	0 (0–1)	0 (0–1)
Children (pregnancies >6 months)	2 (2-3)	2 (2-3)	2 (2–3)	2 (2–3)
Height (cm)	162·6 (157·5–165·1)	162·6 (157·5–165·1)	162·6 (157·5–165·1)	162·6 (157·5–165·1)
Weight (kg)	67·6 (60·3–76·2)	67·6 (60·3–76·2)	67·6 (60·3–76·2)	67·6 (60·3–76·2)
Ethnic origin				
White	48845 (96.5%)	48749 (96·3%)	97598 (96.3%)	195192 (96·4%)
Black	670 (1.3%)	717 (1.4%)	1377 (1.4%)	2764 (1.4%)
Asian	442 (0.9%)	477 (0.9%)	936 (0.9%)	1855 (0.9%)
Other	428 (0.8%)	424 (0.8%)	839 (0.8%)	1691 (0.8%)
Missing	239 (0.5%)	256 (0.5%)	549 (0.5%)	1044 (0.5%)
Hysterectomy	9680 (19·1%)	9496 (18.8%)	18990 (18·7%)	38166 (18.8%)
Ever use of OCP	30098 (59.5%)	30308 (59.9%)	60284 (59.5%)	120 690 (59.6%)
Use of HRT at recruitment	9457 (18.7%)	9383 (18·5%)	19150 (18.9%)	37990 (18.8%)
Personal history of cancer†	2973 (5·9%)	2974 (5·9%)	6105 (6.0%)	12052 (6.0%)
Personal history of breast cancer	1848 (3.7%)	1891 (3.7%)	3912 (3.9%)	7651 (3.8%)
Maternal history of ovarian cancer	802 (1.6%)	778 (1·5%)	1579 (1.6%)	3159 (1.6%)
Maternal history of breast cancer	3159 (6.2%)	3206 (6·3%)	6619 (6·5%)	12984 (6.4%)

Data are n (%) or median (IQR). MMS=multimodal screening. USS=ultrasound screening. HRT=hormone replacement therapy. OCP=oral contraceptive pill. *One woman asked for all her details to be removed. †Includes those with personal history of breast cancer.

Table 1: Baseline characteristics

analysis that treats other deaths as a competing risk rather than a censoring event. We also treated bilateral salpingooophorectomy done in women both within and outside of the trial as a competing (risk) event.³³

As detailed previously,^{13,14} we changed the risk of ovarian cancer calculation cutoffs in the MMS group in April, 2005, to maintain annual triage rates of 15% of women to intermediate-risk and 2% to increased-risk categories. Safety data related to screening was monitored annually by the data monitoring and ethics committee, who also assessed performance characteristics of the screening strategies. We did an interim analysis of the primary outcome when about half of the anticipated number of ovarian cancer deaths had occurred in the no screening group. We had no stopping guidelines for futility, and the critical significance level guideline for stopping for benefit was small (significance level of 0.001).

All analyses were by modified intention to screen. We analysed all randomly allocated women except for those who we later came to know had a bilateral oophorectomy,

	Total*	Screen positives	Cancers not detected by screening				
			Screen negatives <1 year from last test of screening episode†	Screen negatives >1 year after last test of screening episode	After screening phase‡	Never attended screening	
MMS (50 624 women, 548 533 women-years)							
Primary ovarian cancer	338 (100%)	199 (59%)	38 (11%)	41 (12%)	57 (17%)	3 (1%)	
Primary non-epithelial neoplasm of ovary (ICD C56)	11 (100%)	7 (64%)	2 (18%)	2 (18%)	0	0	
Primary borderline epithelial neoplasm of ovary (ICD C56)	44 (100%)	24 (55%)	10 (23%)	5 (11%)	5 (11%)	0	
Primary invasive epithelial neoplasm of ovary (ICD C56)	244 (100%)	147 (60%)	21 (9%)	29 (12%)	44 (18%)	3 (1%)	
Primary invasive epithelial neoplasm of fallopian tube (ICD C57.0)	19 (100%)	13 (68%)	2 (11%)	0	4 (21%)	0	
Undesignated (unable to delineate if primary site ovary or fallopian tube or peritoneum)	20 (100%)	8 (40%)	3 (15%)	5 (25%)	4 (20%)	0	
Primary peritoneal cancer	16 (100%)	13 (81%)	3 (19%)	0	0	0	
USS (50 623 women, 548 825 women-years)							
Primary ovarian cancer	314 (100%)	161 (51%)	60 (19%)	46 (15%)	34 (11%)	13 (4%)	
Primary non-epithelial neoplasm of ovary (ICD C56)	12 (100%)	11 (92%)	0	1 (8%)	0	0	
Primary borderline epithelial neoplasm of ovary (ICD C56)	53 (100%)	48 (91%)	2 (4%)	1 (2%)	0	2 (4%)	
Primary invasive epithelial neoplasm of ovary (ICD C56)	220 (100%)	93 (42%)	48 (22%)	37 (17%)	31 (14%)	11 (5%)	
Primary invasive epithelial neoplasm of fallopian tube (ICD C57:0)	13 (100%)	4 (31%)	3 (23%)	3 (23%)	3 (23%)	0	
Undesignated (unable to delineate if primary site ovary or fallopian tube or peritoneum)	16 (100%)	5 (31%)	7 (44%)	4 (25%)	0	0	
Primary peritoneal cancer	10 (100%)	3 (30%)	3 (30%)	4 (40%)	0	0	
No screening (101299 women, 1097089 women-year	s)						
Primary ovarian cancer	630 (100%)		501 (80%)		129 (20%)		
Primary non-epithelial neoplasm of ovary (ICD C56)	8 (100%)		7 (88%)		1 (13%)		
Primary borderline epithelial neoplasm of ovary (ICD C56)	62 (100%)		50 (81%)		12 (19%)		
Primary invasive epithelial neoplasm of ovary (ICD C56)	493 (100%)		392 (80%)		101 (20%)		
Primary invasive epithelial neoplasm of fallopian tube (ICD C57.0)	28 (100%)		21 (75%)		7 (25%)		
Undesignated (unable to delineate if primary site ovary or fallopian tube or peritoneum)	38 (100%)		30 (79%)		8 (21%)		
Primary ovarian neoplasm (histology not available)	1 (100%)		1 (100%)		0		
Primary peritoneal cancer	15 (100%)		15 (100%)		0		

Data are n (%). MMS=multimodal screening. ICD=International Classification of Diseases. USS=ultrasound screening. *Includes in the MMS group, three screen-positive neoplasms of uncertain malignancy that have been recoded by outcomes review as non-epithelial ovarian cancer and one screen-positive borderline that has been recoded as invasive after the published incidence analysis; in the USS group, it includes five screen-positive neoplasms of uncertain malignancy that have been recoded by outcomes review as non-epithelial ovarian cancer stat were identified after the published prevalence analysis. †For the no screening group, this category applies during the screening phase. ‡Refers to more than 1 year after last potential screen in 2011 based on the anniversary of an individual's randomisation date.

Table 2: Ovarian and primary peritoneal cancers grouped by primary site and screening status

ovarian cancer, or exited the registry before recruitment. We did all statistical analyses using Stata (version 14), R (version 3.2.1; packages flexsurv and PwrGSD), and Stan (version 2.8.0).³⁴

This trial is registered with ClinicalTrials.gov, number NCT00058032.

Role of the funding source

The funders of the study had no role in study design, data collection, data analysis, data interpretation, or writing of the report. UM and AR extracted the dataset. MB, SJS, DJR, UM, and AR had full access to all the data in the study. IJJ, UM, SJS, and MP had final responsibility for the decision to submit for publication.

Results

We invited 1243 282 women to participate, recruiting 205090 between April 17, 2001, and Sept 29, 2005, randomising 202638 (16·3%) between June 1, 2001, and Oct 21, 2005: 101359 (50·0%) to no screening, 50640 (25·0%) to MMS, and 50639 (25·0%) to USS. By end of screening on Dec 31, 2011, women attended 345 570 MMS and 327775 USS annual screening 60 [<0·5%]; MMS 16 [<0·5%]; USS 16 [<0·5%]) from the primary analysis (figure 1). The final cohort eligible for analysis consisted of 202546 (>99·9%) women: 101299 (>99·9%) in the no screening group, 50624 (>99·9%) in the MMS group, and 50 623 (>99·9%) in the USS group. Baseline characteristics were balanced between study groups (table 1).

The last notification from the Health and Social Care Information Centre was received on March 25, 2015, and from Northern Ireland, for deaths, on April 9, 2015, and for cancer, on April 15, 2015. Complete follow-up until study completion (Dec 31, 2014) or death was possible in 200293 (98·9%) women (no screening 100149 [98·8%]; MMS 50084 [98·9%]; USS 50060 [98·9%]). Median followup time was 11·1 years (IQR 10·0–12·0) for all groups.

Of 3110 women investigated (appendix p 4), 1282 (41%) women were confirmed on outcomes review to have ovarian cancer (table 2). The overall sensitivity for detection of ovarian cancers, diagnosed within a year of a screening, was 84% (95% CI 79–88; 199 of 237) in the MMS group and 73% (66–79; 161 of 221) in the USS group. Of the primary peritoneal cancers, 81% (13 of 16) were screen detected with MMS and 30% (three of ten) were with USS. We noted evidence of a higher proportion of invasive epithelial ovarian and peritoneal cancer diagnosed with low-volume disease (stage I, II, and IIIa) in the MMS group (119 [40%] of 299; p<0.0001) than in the no screening group (149 [26%] of 574), but not in the USS group (62 [24%] of 259; p=0.57; appendix p 5).

At censorship, 649 (0.32%) women had died of ovarian cancer: 347 (0.34%) in the no screening group, 148 (0.29%) in the MMS group, and 154 (0.30%) in the USS group. The mortality reduction over years 0–14 with

	Number of women (n)	Deaths (n)	Mortality reduction 0–14 years (%)	p value	Mortality reduction 0–7 years (%)	Mortality reduction 7–14 years (%)
Ovarian cancer (primar	y analysis)					
Cox model						
MMS	50624	148	15% (-3 to 30)	0.10		
USS	50623	154	11% (-7 to 27)	0.21		
No screening	101299	347				
Royston-Parmar model						
MMS	50624	148	16% (-1 to 33)	0.11	8% (-20 to 31)	23% (1 to 46)
USS	50623	154	12% (-6 to 29)	0.18	2% (-27 to 26)	21% (-2 to 42)
No screening	101299	347				
Royston-Parmar model (excluding prevalent cases)						
MMS	50561	120	20% (-2 to 40)	0.021	8% (-27 to 43)	28% (-3 to 49)
No screening	101183	281				
Weighted log-rank (post-hoc)						
MMS	50624	148	22% (3 to 38)*	0.023		
USS	50623	154	20% (0 to 35)*	0.049		
No screening	101299	347				
Ovarian and primary pe	eritoneal can	cer (secon	ıdary analysis)			
Cox model						
MMS	50624	160	11% (-8 to 26)	0.23		
USS	50623	163	9% (-9 to 24)	0.31		
No screening	101299	358				
Royston-Parmar model						
MMS	50624	160	11% (-7 to 28)	0.15	4% (-25 to 27)	18% (-5 to 40)
USS	50623	163	10% (-8 to 27)	0.27	2% (-26 to 26)	17% (-8 to 38)
No screening	101299	358				
Royston-Parmar model (excluding prevalent cases)						
MMS	50561	131	16% (-6 to 35)	0.047	5% (-30 to 37)	24% (-7 to 45)
No screening	101191	298				
Weighted log-rank (post-hoc)						
MMS	50624	160	18% (-1 to 34)*	0.064		
USS	50623	163	17% (-3 to 33)*	0.097		
No screening	101299	358				

Data in parentheses are 95% CIs. MMS=multimodal screening. USS=ultrasound screening. *Mortality reduction from hazard ratio weighted by pooled cumulative ovarian cancer mortality.

Table 3: Summary of analyses of relative reduction of ovarian and primary peritoneal cancer deaths

the Cox model was 15% (95% CI –3 to 30; p=0.10) in the MMS group and 11% (–7 to 27; p=0.21) in the USS group (table 3). Figure 2 shows the Kaplan-Meier cumulative death rates; in the appendix (pp 12, 13), the ovarian curves are overlaid with RP fit. Figure 3 and the appendix (p 14) show that the no screening group hazard rate continues to rise throughout the study period, whereas the MMS hazard rate starts levelling off, becoming substantially lower than that of the no screening group at about 7 years, with the USS hazard rate levelling off at about 9 years, showing a potential delayed effect of screening.

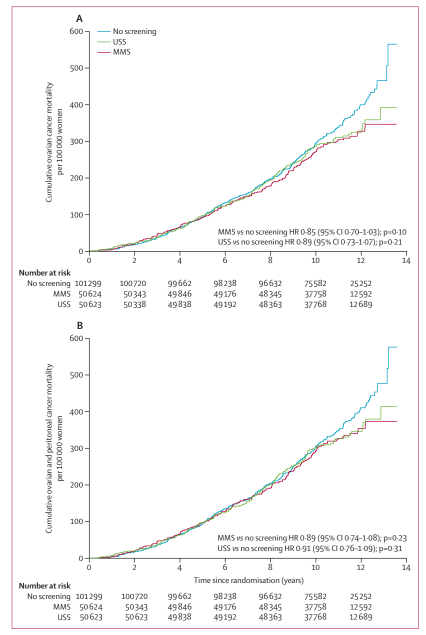


Figure 2: (A) Cumulative ovarian cancer and (B) ovarian and peritoneal cancer deaths

The Royston-Parmar model is shown in the appendix (p 12, 13). HR=hazard ratio. MMS=multimodal screening USS=ultrasound screening.

In the appendix (pp 15, 16), HRs plotted over time³² from the RP model, a non-parametric estimate of HRs, and averaged post-hoc HRs over 0–7 years and 7–14 years are shown. After year 7, the HRs decrease rapidly, showing non-proportional hazards (type II)³⁵ and a delayed mortality reduction for years 7–14 of 23% (95% CI 1–46) for MMS and 21% (–2 to 42) for USS. For years 0–7, the estimated mortality reduction was much smaller: 8% (–20 to 31) for MMS and 2% (–27 to 26) for USS (figure 2, table 3).

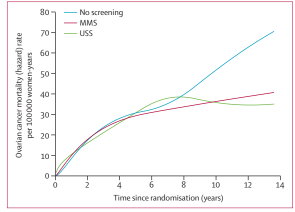


Figure 3: Rates of ovarian cancer The figure including confidence limits is in the appendix (p 14). MMS=multimodal screening. USS=ultrasound screening.

In the MMS group, we excluded an average of 63 (19%) prevalent ovarian cases from the 338 cases and we excluded an average of 116 (18%) prevalent cases from the 630 cases in the no screening group, showing the expected equal proportion of prevalent cases in the two groups. Hazards were not proportional (p=0.037), so we fitted separate RP models to the survival data for each group, showing excellent overlap with the non-parametric Kaplan-Meier cumulative mortality curves (figure 4). The hazards between the MMS and no screening groups were significantly different (p=0.021), showing that ovarian cancer mortality was significantly lower in the MMS group (20% [-2 to 40]) than in the no screening group (table 3). The mortality reduction was also higher for years 7–14 (28% [-3 to 49]) than for years 0–7 (8% [-27 to 43]).

Because the HRs were not constant over time and the Cox model has low power for detection of a late effect of this type,³⁵ we did a single post-hoc analysis using the WLR test inspired by the rationale in the PLCO Cancer Screening Trial report (appendix p 11).²⁴ The median delay from randomisation to death in the no screening group was over 8 years (randomisation to cancer diagnosis 6 years; diagnosis to death $2 \cdot 3$ years). This analysis suggested a significant reduction in ovarian cancer mortality in the MMS group compared with the no screening group, but not in the USS group (table 3).

In addition to the ovarian and fallopian tube deaths, we noted a further 11 deaths due to primary peritoneal cancer in the no screening group, 12 in the MMS group, and nine in the USS group (table 3). This resulted in smaller mortality reductions with the MMS versus no screening and USS versus no screening comparisons than with the ovarian cancer mortality reductions (figure 2, table 3). As with the primary outcome, the no screening group hazard rate continues to rise, whereas the MMS and USS hazard rates start levelling off (appendix p 17). The RP model also yielded higher mortality reductions for years 7–14 than for years 0–7, with the MMS reduction increasing if prevalent cases are excluded (figure 4). In the MMS group, of 427 448 eligible screening episodes, 345 570 were attended, giving a compliance of 80 \cdot 8%. In the USS group, of 420 047 eligible screening episodes, 327775 were attended, giving a 78 \cdot 0% compliance (appendix p 6). At censorship with a maximum follow-up of 13 \cdot 6 years, the preliminary number needed to screen to prevent one death from ovarian cancer was 641 (95% CI 375–1934; appendix p 18). Sensitivity analyses for source of ovarian cancer death, correlation within RCs, competing risks, and parametric estimation show only minor differences (appendix p 7).

We noted no evidence of a difference in deaths because of other causes between the MMS, USS, and no screening groups (appendix p 8). The age-standardised incidence and mortality rates for cancer and death from any cause increased with time from randomisation in the no screening group such that at the end of the trial, rates were similar to UK population rates (appendix p 23). Survival curves from date of diagnosis of ovarian cancers in the no screening group were similar to those of the agestandardised UK population, with a 5 year survival rate of 41% (95% CI 37–45) and a 10 year survival rate of 33% (28–38; appendix p 19).

The overall incidence of ovarian cancer was 57 (95% CI 53-62; 630 cancers; 1097089 women-years) per 100000 women-years in the no screening group, 62 (55-68; 338 cancers; 548553 women-years) per 100000 women-years in the MMS group, and 57 (51-64; 314 cancers; 548825 women-years) per 100000 womenyears in the USS group. We noted no evidence of a difference in incidence of ovarian cancer between the three groups (appendix pp 20-22). The incidence of invasive epithelial ovarian cancer and peritoneal cancer was also similar: 52 (48-61; 575 cancers; 1097089 womenyears) per 100000 women-years in the no screening group, 55 (48-57; 299 cancers; 548 553 women-years) per 100 000 women-years in the MMS group, and 47 (41-53; 259 cancers; 548825 women-years) per 100000 womenyears in the USS group.

We noted screening complications in 30 (<1%) women in the MMS group and 61 (<1%) in the USS group, giving a screening-related complication rate of 8.6 per 100 000 in the MMS group and 18.6 per 100000 in the USS group (appendix p 9). We noted benign adnexal pathology or normal adnexa in 488 (1.0%) women in the MMS group and 1634 (3.2%) women in the USS group who had screen-positive surgery (appendix p 9). This finding translates to 14 (345 572 annual screens) false-positive surgeries per 10000 screens in the MMS group and 50 (327775 annual screens) false-positive surgeries per 10000 screens in the USS group. For each ovarian and peritoneal cancer detected by screening, an additional two (<0.1%; 211 annual screens) women in the MMS group and ten (<0.1%; 164 annual screens) women in the USS group had false-positive surgery. Women in the MMS group had a complication rate of 3.1% (95% CI 1.7-5.0; 15 of 488) and those in the USS group had a rate

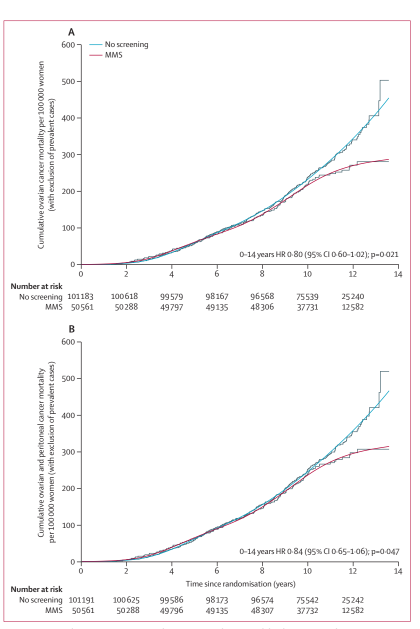


Figure 4: (A) Cumulative ovarian cancer and (B) ovarian and peritoneal deaths in MMS and no screening groups after exclusion of prevalent cases

HRs and mortality reductions for 0–7 years and 7–14 years calculated from the Royston-Parmar model. Cumulative mortality curves from the Royston-Parmar model are overlaid onto Kaplan-Meier curves. HR=hazard ratio. MMS=multimodal screening.

of $3 \cdot 5\%$ ($2 \cdot 7 - 4 \cdot 5$; 57 of 1634). Outside of the trial, during the same period, 792 ($0 \cdot 8\%$) women in the no screening group, 466 ($0 \cdot 9\%$) in the MMS group, and 441 ($0 \cdot 9\%$) in the USS group had both ovaries or the only remaining ovary removed for a range of indications and had benign pathology or normal adnexa. Data for complications associated with these surgeries are not available. The overall ratio of women who had surgery resulting in benign adnexal pathology or normal adnexa to women with ovarian and peritoneal cancer was $1 \cdot 2$ (792:645) in the no screening group, $2 \cdot 7$ (954:354) in the MMS group, and 6.4 (2075:324) in the USS group. Details of final follow-up questionnaire responses are provided in the appendix (p 25). Ovarian cancer screening outside of UKCTOCS was documented in 1660 (4.3% [95% CI 4.1-4.5]) of 38 238 women in the no screening group who completed the follow-up questionnaire in 2014.

Discussion

The relative mortality reduction was 15% in the MMS group and 11% in the USS group, but these reductions were not significant with the primary prespecified Cox analysis. In retrospect, it would have been preferable to specify a primary analysis that was weighted to reflect the predictable delay in mortality reduction in a screening trial of this type. Nevertheless, in view of the trend in mortality in the no screening and screening groups, significance using the Cox analysis could be achieved on further follow-up. However, the prespecified secondary subgroup analysis with exclusion of prevalent cases in the MMS group was significant, suggesting that the longterm effect of an MMS screening programme is about a 28% mortality reduction after 7 years of screening. This significant, yet delayed effect was supported by the sole post-hoc weighted analysis, which was also significant for the MMS group. These results are encouraging for various reasons. First, the mortality hazard rate in the no screening group seems to increase, whereas in the two screened groups, it levels off, resulting in decreasing HRs. This finding suggests that the difference in mortality between no screening and screening groups will increase with time and further follow-up. Second, the mortality difference was not constant across the trial. It was low for the first 7 years after randomisation, but the estimated mortality reductions increased during years 7-14. The mortality reduction in the MMS group for years 7-14 remained high in subgroup analyses with exclusion of prevalent cases for primary and secondary outcomes and in the secondary outcome analyses including peritoneal cancer deaths (albeit slightly lower). This late effect was predictable in view of the unavoidable time interval from randomisation to diagnosis and then death. For participants who died in the no screening group the median interval from randomisation to death was more than 8 years.

The late effect on mortality seen in this trial is often noted in screening trials in which survival is measured from date of randomisation. In the European Randomised Study of Screening for Prostate Cancer trial,³⁶ an effect of screening only emerged 7 years after randomisation. Similarly, investigators of the Norwegian Colorectal Cancer Prevention Trial³⁷ noted an effect 9 years from randomisation. The PLCO Cancer Screening Trial²⁴⁻²⁷ and National Lung Screening Trial²⁸ used the weighted log-rank test in anticipation of this delayed effect. Overall, the results suggested that an unequivocally significant difference in mortality might emerge after longer follow-up. The estimate of relative reduction in ovarian cancer mortality in UKCTOCS is in keeping with relative reductions noted in breast cancer randomised screening trials, which vary between 15% and 25%, and with those noted in meta-analyses of observational studies, which vary between 13% and 17%.³⁸ Using UK population data from 2007, Loberg and colleagues³⁸ reported that for 1000 women invited to biennial mammography screening for 20 years from 50 years of age, two to three women are prevented from dying of breast cancer. Findings from this trial suggest that for 641 women screened annually using the multimodal strategy for 14 years, one ovarian cancer death is prevented.

The only directly comparable trial with this trial is the ovarian component of the PLCO Cancer Screening Trial.²⁴ However, some differences exist between the two, including, in this trial, use of ROCA (as opposed to a CA125 cutoff) and detailed screening algorithms. We centrally managed implementation of the entire screening protocol with predefined pathways for all women detected to have abnormalities, fail-safe monitoring, and quality assurance. In the PLCO Cancer Screening Trial, no difference was noted in ovarian and peritoneal cancer deaths between the screening and control groups at a median follow-up of 12.4 years. In our secondary outcome analysis, we noted a mortality trend that was in keeping with our primary analysis and different to that noted in the PLCO Cancer Screening Trial. This finding follows on from the higher sensitivity and stage shift noted earlier in the MMS group of this trial.

Both venepuncture and transvaginal ultrasound were associated with minor complications and very low complication rates. Most postmenopausal women found transvaginal USS acceptable, with 3.5% reporting moderate or severe pain during the scan.³⁹ As previously reported from the UKCTOCS psychosocial study,15 screening did not increase general anxiety. The number of women having surgery for false-positive screen results was as predicted in the trial design and for MMS, was much fewer than the upper acceptable limit, whereas for USS, it was on the border of acceptability. The small number of operations associated with MMS has been noted in previous trials.^{9,40} During the course of this trial, the ratio of women who had surgery for which ovaries had benign pathology or were normal to those diagnosed with ovarian and peritoneal cancer was 2.3-times higher in the MMS group and 5.3-times higher in the USS group than in the no screening group. Complication rates of false-positive surgery of 3.1% in the MMS group and 3.5% in the USS group were similar to the major complication rate of 2.9% reported for benign surgery undertaken in NHS gynaecological oncology centres in a recent multicentre audit.41

Key strengths of this trial include scale, with more than 202 000 participants, more than 670 000 annual screening episodes, and more than 2.19 million women-years of follow-up; the multicentre setting within the UK NHS;

central management of screening protocols with use of a bespoke web-based trial management system; high compliance; low contamination in the no screening group; completeness of ascertainment of deaths through linkage to national registries; an outcomes review committee masked to group assignment; and treatment of all women within the NHS. The healthy volunteer effect, noted initially, waned over time, such that by the end of the trial, age-standardised cancer incidence and all-cause mortality rates in the no screening group were similar to the UK population, as were ovarian cancer survival rates (appendix p 23).

The main limitation of this trial was our failure to anticipate the late effect of screening in our statistical design. Had we done so, the weighted log-rank test could have been planned in line with many other large cancer screening trials, including the ovarian component of the PLCO Cancer Screening Trial.²⁴ As a result, we report, as planned, the Cox test as the primary analysis balanced by a significant preplanned analysis with exclusion of prevalent cases, with support from a sole post-hoc WLR test in the MMS group. Generalisability of the results will depend on central implementation of the screening protocols, with accompanying quality assurance processes. This implementation, although challenging, is achievable through the processes used in the NHS national screening programme. Further follow-up is needed to assess the extent of the mortality reduction before firm conclusions can be reached on the long-term efficacy and costeffectiveness of ovarian cancer screening.

Contributors

IJJ and UM are the principal investigators and MP, SJS, SC, AJM, and LF are co-investigators; all of these people conceived and designed the study. SJS and IJJ developed the risk of ovarian cancer algorithm. IJJ led trial design and secured funding. MP, SJS, MB, IJJ, and UM developed the statistical analysis plan. UM led implementation of the trial and development of the trial protocols, with oversight from IJI, MP, SJS, and SC and supported by the coordinating centre team (AD, GF, JF, AG-M, RG, JKK, MH, RH, WRL, AR, AS, and FW). UM and WRL designed and WRL wrote the bespoke trial management system. The trial centre leads (NNA, DC, SD, KG, JH, SL, AL, UM, TM, JM, DO, KR, IS, MWS, KW, RW. and HJ) oversaw recruitment and trial implementation at the regional trial centres. The laboratory and administrative teams at the coordinating centre supporting the trial and regional centre teams are listed in the appendix (pp 26-28; collated by JKK, AR, and DNC). UM, SA, DC, DNC, SKD, AD, SD, GF, JF, AG-M, KG, RG, JKK, MH, RH, JH, CK, SL, SJL, AL, TM, JM, DO, KR, AR, IS, MWS, AS, JT, FW, KW, and RW collected data. NS, EB, MW, and KR reviewed outcomes. SJL, MH, SA, RH, SKD, DNC, CK, JT, and AR contributed to the outcomes review process. SC, NNA, and MWS assisted the coordinating team with oversight of the ultrasound screening group. UM and AR prepared the dataset that was used for the analysis. MB, SJS, MP, and DJR did the statistical analysis. UM, AR, AG-M, and MB prepared the descriptive statistics. MB, DJR, SJS, AR, UM, and AG-M prepared the figures and tables. JKK, SJS, MW, MB, AG-M, and UM did the literature searches for the report. UM, JKK, IJJ, SJS, MP, AR, AG-M, MB, and AJM interpreted data and drafted the report. All authors critically reviewed the report and approved it before submission.

Declaration of interests

IJJ reports personal fees from and stock ownership in Abcodia as the non-executive director and consultant. He reports personal fees from Women's Health Specialists as the director. He has a patent for the Risk of Ovarian Cancer algorithm and an institutional licence to Abcodia with royalty agreement. He is a trustee (2012–14) and Emeritus Trustee (2015 to present) for The Eve Appeal. He has received grants from the Medical Research Council (MRC), Cancer Research UK, the National Institute for Health Research, and The Eve Appeal. UM has stock ownership in and research funding from Abcodia. She has received grants from the MRC, Cancer Research UK, the National Institute for Health Research, and The Eve Appeal. NNA is the founder of, owns stock in, and is a board member of MedaPhor, a spin-off company at Cardiff University. He has a patent for the ultrasound simulation training system MedaPhor. SA is funded by a research grant from Abcodia. AD reports personal fees from Abcodia. AL reports personal fees from Roche as a panel member and advisory board member and fees from Sanofi Pasteur Merck Sharp & Dohme (Gardasil) as an advisory board member. JM is involved in a private ovarian cancer screening programme after closure of this trial. MWS reports personal fees from Abcodia as a consultant. LF reports funding from the MRC for the UK Collaborative Trial of Ovarian Cancer Screening psychosocial study. She reports personal fees from GlaxoSmithKline, Amgen, AstraZeneca, Roche, Pfzier, Teva, Bristol-Myers Squibb, and Sanofi and grants from Boehringer Ingelheim and Roche. SJS reports personal fees from the LUNGevity Foundation and SISCAPA Assay Technologies as a member of their Scientific Advisory Boards. He reports personal fees from Abcodia as a consultant and AstraZeneca as a speaker honorarium. He has a patent for the Risk of Ovarian Cancer algorithm and an institutional license to Abcodia. All other authors declare no competing interests.

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References

- Cancer Research UK. Ovarian cancer survival statistics. One-, five- and ten-year survival for ovarian cancer. http://www. cancerresearchuk.org/health-professional/cancer-statistics/ statistics-by-cancer-type/ovarian-cancer/survival/#heading-Zero (accessed Nov 3, 2015).
- 2 Jacobs I, Stabile I, Bridges J, et al. Multimodal approach to screening for ovarian cancer. *Lancet* 1988; 1: 268–71.
- 3 Jacobs IJ, Skates S, Davies AP, et al. Risk of diagnosis of ovarian cancer after raised serum CA 125 concentration: a prospective cohort study. *BMJ* 1996; **313**: 1355–58.
- 4 Menon U, Talaat A, Jeyarajah AR, et al. Ultrasound assessment of ovarian cancer risk in postmenopausal women with CA125 elevation. Br J Cancer 1999; 80: 1644–47.
- 5 Jacobs IJ, Skates SJ, MacDonald N, et al. Screening for ovarian cancer: a pilot randomised controlled trial. *Lancet* 1999; 353: 1207–10.
- 6 Skates SJ, Pauler DK, Jacobs IJ. Screening based on the risk of cancer calculation from Bayesian hierarchical changepoint and mixture models of longitudinal markers. J Am Stat Assoc 2001; 96: 429–39.
 - Skates SJ, Menon U, MacDonald N, et al. Calculation of the risk of ovarian cancer from serial CA-125 values for preclinical detection in postmenopausal women. J Clin Oncol 2003; 21(suppl 10): 206s–10s.

- 8 Menon U, Talaat A, Rosenthal AN, et al. Performance of ultrasound as a second line test to serum CA125 in ovarian cancer screening. *BJOG* 2000; **107**: 165–69.
- 9 Menon U, Skates SJ, Lewis S, et al. Prospective study using the risk of ovarian cancer algorithm to screen for ovarian cancer. J Clin Oncol 2005; 23: 7919–26.
- 10 Cancer Research UK. Ovarian cancer statistics. Ovarian cancer mortality. http://www.cancerresearchuk.org/health-professional/ cancer-statistics/statistics-by-cancer-type/ovarian-cancer#heading-One (accessed Oct 6, 2015).
- 11 Cancer Research UK. Ovarian cancer survival statistics. Ovarian cancer survival by stage at diagnosis. http://www.cancerresearchuk. org/health-professional/cancer-statistics/statistics-by-cancer-type/ ovarian-cancer/survival#heading-Three (accessed Oct 6, 2015).
- 12 Menon U, Gentry-Maharaj A, Ryan A, et al. Recruitment to multicentre trials—lessons from UKCTOCS: descriptive study. BMJ 2008; 337: a2079.
- 13 Menon U, Gentry-Maharaj A, Hallett R, et al. Sensitivity and specificity of multimodal and ultrasound screening for ovarian cancer, and stage distribution of detected cancers: results of the prevalence screen of the UK Collaborative Trial of Ovarian Cancer Screening (UKCTOCS). *Lancet Oncol* 2009; **10**: 327–40.
- 14 Menon U, Ryan A, Kalsi J, et al. Risk algorithm using serial biomarker measurements doubles the number of screen-detected cancers compared with a single-threshold rule in the United Kingdom Collaborative Trial of Ovarian Cancer Screening. J Clin Oncol 2015; 33: 2062–71.
- 15 Barrett J, Jenkins V, Farewell V, et al. Psychological morbidity associated with ovarian cancer screening: results from more than 23,000 women in the randomised trial of ovarian cancer screening (UKCTOCS). *BJOG* 2014; **121**: 1071–79.
- 16 Sharma A, Burnell M, Gentry-Maharaj A, et al. Quality assurance and its impact on ovarian visualisation rates in the multicentre United Kingdom Collaborative Trial of Ovarian Cancer Screening (UKCTOCS). Ultrasound Obstet Gynecol 2015; published online June 19. DOI:10.1002/uog.14929.
- 17 Prat J, for the FIGO Committee on Gynecologic Oncology. Staging classification for cancer of the ovary, fallopian tube, and peritoneum. *Int J Gynaecol Obstet* 2014; **124**: 1–5.
- 18 Tavassoli FA, Devilee P, eds. World Health Organization Classification of Tumours. Pathology and genetics. Tumors of the breast and female genital organs. Lyon: World Health Organization, 2003.
- 19 Daya D, Cheung AN, Khunamornpong S, et al. Tumors of the peritoneum: epithelial tumors of Müllerian type. In: Kurman RJ, Carcangiu ML, Herrington CS, Young RH, eds. WHO classification of tumors of female reproductive organs, 4th edn. Lyon: International Agency for Research on Cancer, 2014: 92–93.
- 20 Self SG. An adaptive weighted log-rank test with application to cancer prevention and screening trials. *Biometrics* 1991; 47: 975–86.
- 21 Self SG, Etzioni R. A likelihood ratio test for cancer screening trials. Biometrics 1995; 51: 44–50.
- 22 Shen Y, Cai J. Maximum of the weighted Kaplan-Meier tests with application to cancer prevention and screening trials. *Biometrics* 2001; 57: 837–43.
- 23 Warwick J, Duffy SW. A review of cancer screening evaluation techniques, with some particular examples in breast cancer screening. J R Stat Soc Ser A 2005; 168: 657–77.

- 24 Buys SS, Partridge E, Black A, et al. Effect of screening on ovarian cancer mortality: the Prostate, Lung, Colorectal and Ovarian (PLCO) cancer screening randomized controlled trial. *JAMA* 2011; 305: 2295–303.
- 25 Andriole GL, Crawford ED, Grubb RL 3rd, et al. Mortality results from a randomized prostate-cancer screening trial. *N Engl J Med* 2009; **360**: 1310–19.
- 26 Oken MM, Hocking WG, Kvale PA, et al. Screening by chest radiograph and lung cancer mortality: the Prostate, Lung, Colorectal, and Ovarian (PLCO) randomized trial. *JAMA* 2011; 306: 1865–73.
- 27 Schoen RE, Pinsky PF, Weissfeld JL, et al. Colorectal-cancer incidence and mortality with screening flexible sigmoidoscopy. N Engl J Med 2012; 366: 2345–57.
- 28 National Lung Screening Trial Research Team, Aberle DR, Adams AM, et al. Reduced lung-cancer mortality with low-dose computed tomographic screening. N Engl J Med 2011; 365: 395–409.
- 29 Fleming TR, Harrington DP. Counting processes and survival analysis. Hoboken: John Wiley & Sons, 2011.
- 30 Izmirlian G. Estimation of the relative risk following group sequential procedure based upon the weighted log-rank statistic. *Stat Interface* 2014; **7**: 27–42.
- 31 Royston P, Parmar MK. Flexible parametric proportional-hazards and proportional-odds models for censored survival data, with application to prognostic modelling and estimation of treatment effects. *Stat Med* 2002; 21: 2175–97.
- 32 Liu ZA, Hanley JA, Strumpf EC. Projecting the yearly mortality reductions due to a cancer screening programme. J Med Screen 2013; 20: 157–64.
- 33 Fine JP, Gray RJ. A proportional hazards model for the subdistribution of a competing risk. J Am Stat Assoc 1999; 94: 496–509.
- 34 Carpenter B, Gelman A, Hoffman M, et al. Stan: a probabilistic programming language. J Stat Softw (in press).
- 35 Lagakos SW, Schoenfeld DA. Properties of proportional-hazards score tests under misspecified regression models. *Biometrics* 1984; 40: 1037–48.
- 36 Schroder FH, Hugosson J, Roobol MJ, et al. Screening and prostate-cancer mortality in a randomized European study. N Engl J Med 2009; 360: 1320–28.
- 37 Holme O, Loberg M, Kalager M, et al. Effect of flexible sigmoidoscopy screening on colorectal cancer incidence and mortality: a randomized clinical trial. JAMA 2014; 312: 606–15.
- 38 Loberg M, Lousdal ML, Bretthauer M, Kalager M. Benefits and harms of mammography screening. *Breast Cancer Res* 2015; 17: 63.
- 39 Gentry-Maharaj A, Sharma A, Burnell M, et al. Acceptance of transvaginal sonography by postmenopausal women participating in the United Kingdom Collaborative Trial of Ovarian Cancer Screening. Ultrasound Obstet Gynecol 2013; 41: 73–79.
- 40 Lu KH, Skates S, Hernandez MA, et al. A 2-stage ovarian cancer screening strategy using the risk of ovarian cancer algorithm (ROCA) identifies early-stage incident cancers and demonstrates high positive predictive value. *Cancer* 2013; **119**: 3454–61.
- 41 Iyer R, Gentry-Maharaj A, Nordin A, et al. Predictors of complications in gynaecological oncological surgery: a prospective multicentre study (UKGOSOC-UK gynaecological oncology surgical outcomes and complications). Br J Cancer 2015; 112: 475–84.

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Supplementary appendix

This appendix formed part of the original submission and has been peer reviewed. We post it as supplied by the authors.

Supplement to: Jacobs IJ, Menon U, Ryan A, et al. Ovarian cancer screening and mortality in the UK Collaborative Trial of Ovarian Cancer Screening (UKCTOCS): a randomised controlled trial. *Lancet* 2015; published online Dec 17. http://dx.doi. org/10.1016/S0140-6736(15)01224-6.

Appendix: Supplementary Material

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A. Supplementary Tables

ICD-10 code	Description
C56	Malignant neoplasm of ovary
C57·0	Malignant neoplasm of fallopian tube
C57·4	Uterine adnexa, unspecified
C57·7	Other specified female genital organs
C57·8	Malignant neoplasm of overlapping lesion of female genital organs
C57·9	Malignant neoplasm of female genital organ, unspecified
C48·0	Retroperitoneum
C48·1	Specified parts of peritoneum
C48·2	Malignant neoplasm of peritoneum, unspecified
C48·8	Overlapping lesions of retroperitneum and peritoneum
C76·2	Malignant neoplasm of abdomen
C76·3	Malignant neoplasm of pelvis
C80	Malignant neoplasm without specification of site
D07·3	Carcinoma in situ of other/unspecified female genital organ
D28·2	Benign neoplasm of fallopian tube
D28·9	Benign neoplasm of female genital organ, unspecified
D36·9	Benign neoplasm of unspecified site
D39·1	Neoplasm of uncertain or unknown behaviour of ovary
D39·9	Neoplasm of uncertain or unknown behaviour of female genital organ, unspecified

Abbreviations: ICD, International Classification of Diseases.

Web Table 1. ICD-10 Codes of Notes Reviewed by the Outcomes Committee

Source of notification	Women with source data available	Women with relevant ICD code/cancer*						
Cancer Registry	45 877 (202 625 flagged)	1474						
Death certificate	11 807 (202 625 flagged)	1314 [†]						
Hospital Episode Statistics [‡]	97 505 (158 077 before matching)	572						
National Cancer Intelligence Network (NCIN) [#]	32 846 (158 077 before matching)	871						
Follow-up Questionnaire 1	13 0581	171						
Follow-up Questionnaire 2	86 039	211						
Contacted by patient or relative		405						
Informed by physician / trial surgery		442						
Total number of women reviewed by Outcomes Committee	3110							
Data is n. *C56, C57·0, C57·4, C57·7, C57·8, C57·9, C48	Data is n. *C56, C57·0, C57·4, C57·7, C57·8, C57·9, C48·0, C48·1, C48·2, C48·8, C76·2, C76·3, C80, D07·3, D28·2, D28·9, D36·9,							
D39·1, and D39·9. [†] In absence of cancer registration. [‡] Only available for women resident in England.								

Web Table 2: Source of notification of cancer of women whose notes were submitted for Outcome Review

Characteristic	MMS	USS	No screening	Total
Morphology Ovarian cancers	338	314	630	1282
Invasive epithelial ovarian/tubal/undesignated cancer	283	249	559	1091
Type I invasive epithelial cancer	49	32	87	168
Low grade serous	12	7	17	36
Carcinoma, NOS	0	0	3	3
Mucinous	5	6	22	33
Endometrioid (low grade)	16	7	23	46
Clear cell	16	12	20	48
Brenner	0	0	1	1
Mixed	0	0	1	1
Type II invasive epithelial cancer	212	194	410	816
High grade serous	169	156	331	656
Carcinoma, NOS	105	20	34	73
High grade endometrioid	17	8	10	35
Carcinosarcoma	7	10	32	49
Adenocarcinoma	0	0	2	49 2
		0	2	
Neuroendocrine carcinoma	0		-	1
Type uncertain (grade unknown)	22	23	62	107
Serous	5	6	15	26
Carcinoma, NOS	16	17	46	79
Endometrioid	0	0	1	1
Small Cell Carcinoma	1	0	0	1
Primary borderline epithelial neoplasm of ovary	44	53	62	159
Serous	25	40	30	95
Mucinous	17	10	28	55
Endometrioid	2	1	1	4
Brenner	0	0	3	3
Mixed	0	2	0	2
Primary non-epithelial neoplasm of ovary	11	12	8	31
Granulosa cell	6	6	7	19
Carcinoid	1	3	0	4
Sarcoma	1	2	0	3
Sertoli-Leydig Cell	2	1	0	3
Squamous Cell Carcinoma	1	0	1	2
Primary ovarian neoplasm (histology not available)	0	0	1	1
Primary peritoneal cancer	16	10	15	41
Serous	13	8	12	33
Carcinoma, NOS	3	2	1	6
Carcinosarcoma	0	0	1	1
Clear cell	0	0	1	1
Stage		-		
	220	214	(20	1202
Ovarian cancers	338 283	314 249	630 559	1282 1091
Invasive epithelial ovarian/tubal/ undesignated cancer	-	00	0.1	
	76	39	91	206
la	29	13	35	77
Ib	4	2	1	7
Ic	43	24	55	122
II	31*	19	45	94
IIa	8	6	13	27
IIb	10	5	16	31
IIc	12	8	16	36
III	142	141	314	597
IIIa	11	4	12	27
IIIb	30	21	38	89
IIIc	101	116	264	481
IV	33	50	108	191
Unable to stage	1	0	1	2
Primary non-epithelial neoplasm of ovary	11	12	8	31
I	10	10	7	27
II	0	0	1	1
III	1	1	0	2
IV	0	1	0	1
Primary borderline epithelial neoplasm of ovary	44	53	62	159
I	41	50	49	140
II	0	0	4	4
III	3	3	9	15
Primary ovarian neoplasm (stage not available)	0	0	1	1
	16	10	15	41
			15 1	41 2
Primary peritoneal cancer		0		1.
Primary peritoneal cancer IIb	1	0		
Primary peritoneal cancer IIb III	1 14	9	10	33
Primary peritoneal cancer IIb	1			

Web Table 3: Morphology and stage of ovarian and primary peritoneal cancers

Annual Women due screen*		en due screen* Women ineligible [†] for Screen Women eligible for screen		gible for screen	Women who attended Compliance			Predicted co	Predicted compliance [‡]		Observed / predicted compliance			
screen	MMS	USS	MMS	USS	MMS	USS	MMS	USS	MMS	USS	MMS	USS	MMS	USS
1	50624	50623	10	67	50614	50556	49822	47955	98.4%	94.9%	100.0%	100.0%	0.98	0.95
2	50624	50623	265	1019	50359	49604	45893	44106	91.1%	88.9%	94.5%	91.0%	0.96	0.98
3	50624	50623	569	1458	50055	49165	43588	41951	87.1%	85.3%	87.5%	85.4%	1.00	$1 \cdot 00$
4	50624	50623	898	1862	49726	48761	41669	40025	83.8%	82.1%	83.6%	81.9%	1.00	$1 \cdot 00$
5	50624	50623	1275	2239	49349	48384	39925	38286	80.9%	79.1%	80.4%	78.8%	1.01	1.00
6	50624	50623	1660	2624	48964	47999	38283	36345	78.2%	75.7%	77.6%	76.0%	1.01	1.00
7	50622	50615	2064	3021	48558	47594	35170	32969	72.4%	69.3%	75.0%	72.7%	0.97	0.95
8	40430	40406	2043	2842	38387	37564	26091	23950	68·0%	63.8%	69.5%	66.5%	0.98	0.96
9	28235	28227	1666	2240	26569	25987	16878	15186	63.5%	58.4%	65.2%	61.2%	0.97	0.95
10	13864	13849	954	1278	12910	12571	7328	6334	56.8%	50.4%	61.0%	56.1%	0.93	0.90
11	2132	2115	175	253	1957	1862	923	668	47.2%	35.9%	54.5%	48.3%	0.87	0.74
Total	439027	438950	11579	18903	427448	420047	345570	327775	80.8%	78·0%	82.0%	80.0%	0.99	0.98
Data are n	umbers or p	ercentages. *	Based on a	nniversary of	Randomisa	tion date. [†] Dea	d/ovaries re	moved or ov	arian cancer	diagnosed j	prior to screen	. [‡] Four percent	attrition yea	r on year.

Web Table 4: Compliance with annual screening

Description	Group No. of Women		Deaths	Unweighted	Unweighted		
		women		Mortality reduction (95% CI)	p-value	Mortality reduction (95% CI)	p-value
Outcome death due to ovarian cancer							
1) Events restricted to those with either death or cancer registration	MMS USS No screening	50624 50623 101299	148 154 347	15·7 (-2·3, 30·6) 11·1 (-7·5, 26·5) 0	0·081 0·222	23·8 (4·4, 39·4) 11·1 (0·0, 35·8) 0	0·020 0·057
2) Accounting for potential within- centre correlation: by allowing different baseline hazard for each RC	MMS USS No screening	50624 50623 101299	148 154 347	14·7 (-3·4, 29·6) 11·3 (-7·2, 26·7) 0	0·102 0·210	23·2 (3·7, 38·7) 20·3 (0·4, 36·2) 0	0·022 0·046
 Accounting for potential within- centre correlation by use of cluster- robust standard errors 	MMS USS No screening	50624 50623 101299	148 154 347	14·7 (-3·9, 30·0) 11·3 (-13·7, 30·8) 0	0·115 0·344	23·2 (2·7, 39·4) 20·3 (7·3, 40·8) 0	0·029 0·136
4) Competing risks regression model treating 'other deaths' as competing risk rather than censored	MMS USS No screening	50624 50623 101299	148 154 347	14·7 (-3·4, 29·6) 11·3 (-7·3, 26·6) 0	0·105 0·217	23·2 (3·7, 38·7) 20·2 (0·3, 36·1) 0	0·022 0·047
5) Parametric estimation: proportional hazards Weibull model [†]	MMS USS No screening	50624 50623 101299	148 154 347	14·7 (-3·4, 29·6) 11·3 (-7·2, 26·6) 0	0·102 0·212	23·2 (3·7, 38·7) 20·3 (0·4, 36·2) 0	0·022 0·046

Data is n and % (95% CI). *Mortality reduction and CI's from hazard ratio weighted by pooled cumulative ovarian cancer mortality. *For the Weibull model shape parameter p = 1.62 (95% CI: 1.51, 1.75) and baseline (log-hazard) rate = -9.56 (95% CI: -9.87, -9.27) [unweighted] and p=3.02 (95% CI: 2.80, 3.23) with baseline (log-hazard) rate = -13.62 (95% CI: -13.17, -14.07) [weighted]

Web Table 5: Sensitivity analysis for primary analysis

Original underlying cause of death listed on death certificate excluding		Group				
ovarian/primary peritoneal (ICD-10 code)*	No screening	MMS	USS	RR [†]	p value	
Malignant neoplasm of uncertain origin (C80)	243 (22.1)	126 (23)	111 (20.2)	0.98	0.78	
Other cancers - not ovarian/primary peritoneal or C80 (C00-C99 [‡])	3144 (286.6)	1589 (289.7)	1554 (283.2)	1.00	0.98	
Diseases of the circulatory system (I00-I99)	1471 (134-1)	728 (132.7)	703 (128-1)	0.97	0.45	
Diseases of the digestive system (K00-K99)	286 (26.1)	134 (24.4)	151 (27.5)	1.00	0.96	
Diseases of the nervous system (G00-G99)	249 (22.7)	127 (23.2)	110 (20)	0.95	0.58	
Diseases of the respiratory system (J00-J99)	578 (52.7)	310 (56.5)	286 (52.1)	1.03	0.60	
Mental and behavioural disorders (F00-F99)	160 (14.6)	81 (14.8)	65 (11.8)	0.91	0.42	
Other	493 (44.9)	263 (47.9)	269 (49)	1.08	0.22	
Missing	34 (3.1)	18 (3.3)	13 (2.4)	0.91	0.71	
Total deaths	6658	3376	3262	0.99	0.65	
Total women years	1097089	548533	548825			
Data are n (death per 100,000 women years) *Cause of death was cat	egorised accordi	ing to the codes in	the International	Classificatio	on of	
Diseases, 10th Revision (ICD-10). [†] RR mortality rate ratio for no scr	eening group vs	MMS & USS con	nbined. [‡] Excluding	g C56, C57∙	0, C57·7,	
C57·9, C48·1, C48·2 and C80.						

Web Table 6: Original underlying cause of death as per Death Certificate excluding ovarian/primary peritoneal cancer by group

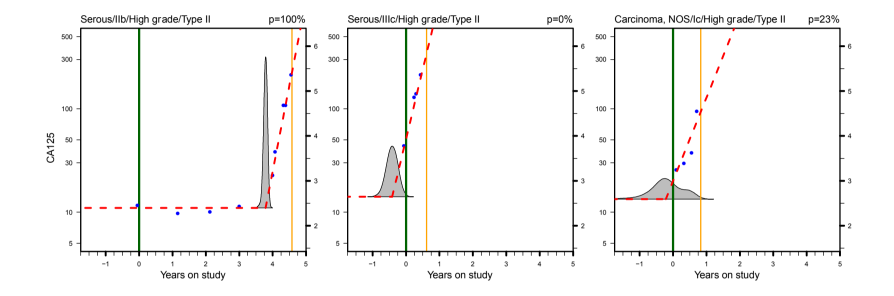
Complications related to screening			
MMS		USS	
Complication type	No. of women	Complication type	No. of women
Bruising	13	Pain	20
Pain	8	Cystitis/infection	11
Haematoma	3	Discomfort	5
Fainting	1	Bruising	2
Cystitis/infection	1	Fainting	1
Other	4	Other	22
Total	30	Total	61
Rate	8.6/100 000	Rate	18.6/100 000

Complications related to screen-positive surgery

MMS		USS				
Complication type	No. of women	Complication type	No. of women			
Anaesthetic	1	Injury to hollow viscus (4 GI, 3 bladder, 4 ureter)	11			
Injury to hollow viscus (2 GI, 1 bladder)	3	Haemorrhage	11			
Haemorrhage	2	Anaesthetic/Myocardial Infarction	3			
Deep Vein Thrombosis	1	Hernia	6			
Bowel obstruction	4	Deep Vein Thrombosis/Pulmonary Embolism	3			
Wound breakdown - total dehiscence	1	Wound breakdown	6			
Significant ileus	1	Bowel obstruction	4			
Uterine perforation	1	Wound/supravaginal haematoma	4			
Infection	1	Infection	6			
		Pain - ward readmission/further operation	3			
Total	15	Total	57			
Rate	3.1% (15/488)		3.5% (57/1634)			
CL Costro Interting In warman who had more than and complication, the most aview was reported						

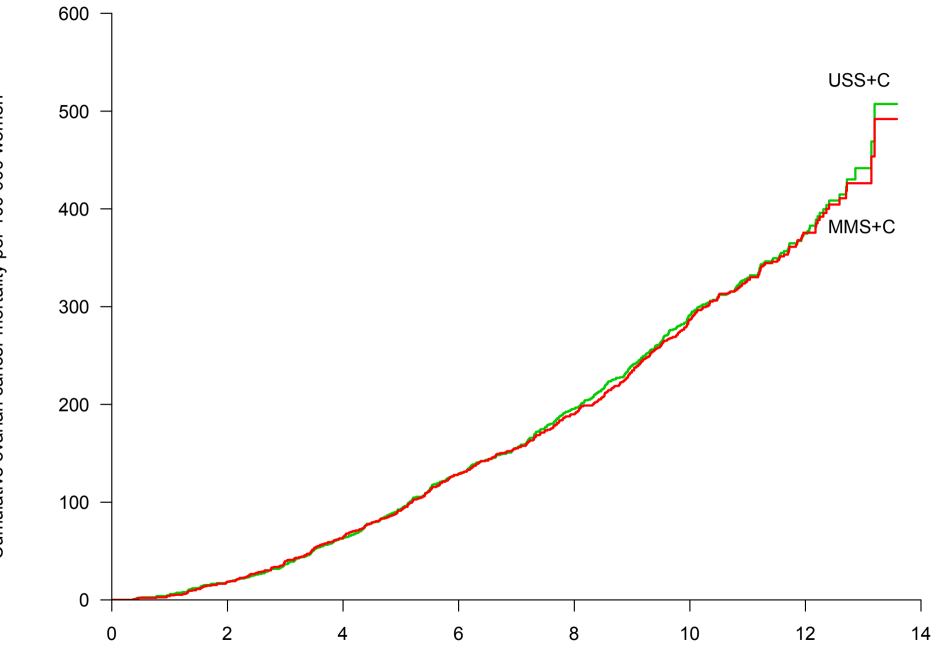
GI – Gastro Intestinal. In women who had more than one complication, the most serious was reported.

Web Table 7: Complications related to screening and screen-positive surgery in women with benign or normal adnexa



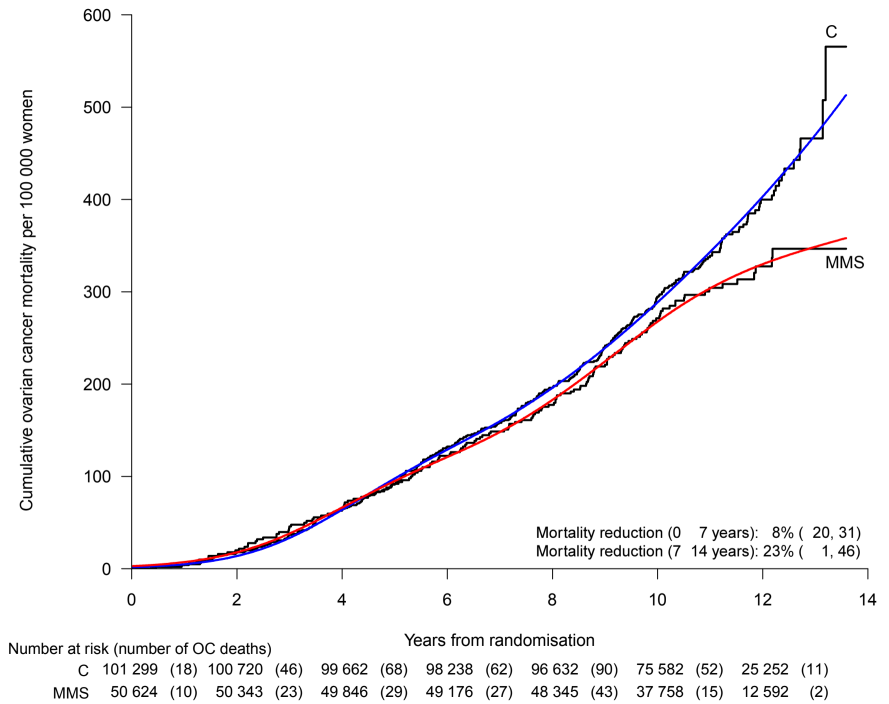
Web Figure 1: Change point distribution in three cases with uniform AUC.

Change point distribution in three cases with green vertical line start of screening, gold vertical line is diagnosis, blue dots are CA125 test results (left vertical scale) and log(CA125) right vertical scale, red dashed line is best change-point model, and distribution of change-point is in grey where: (a) the change-point is after screening starts – case included in 100% of multiple imputation analyses, (b) the change-point is before screening starts and case is excluded from 100% of multiple imputation analyses, and (c) the change-point relative to start of screening is uncertain with 23% of multiple imputations including this case, since 23% of the change-point distribution is after the start of screening – right of green line.



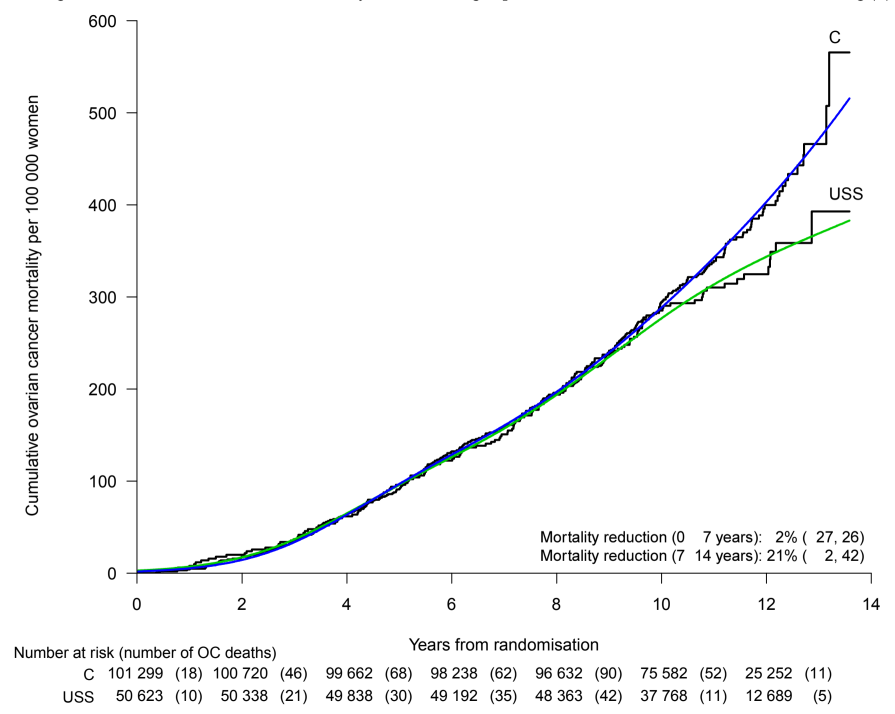
Years from randomisation

Cumulative ovarian cancer mortality per 100 000 women

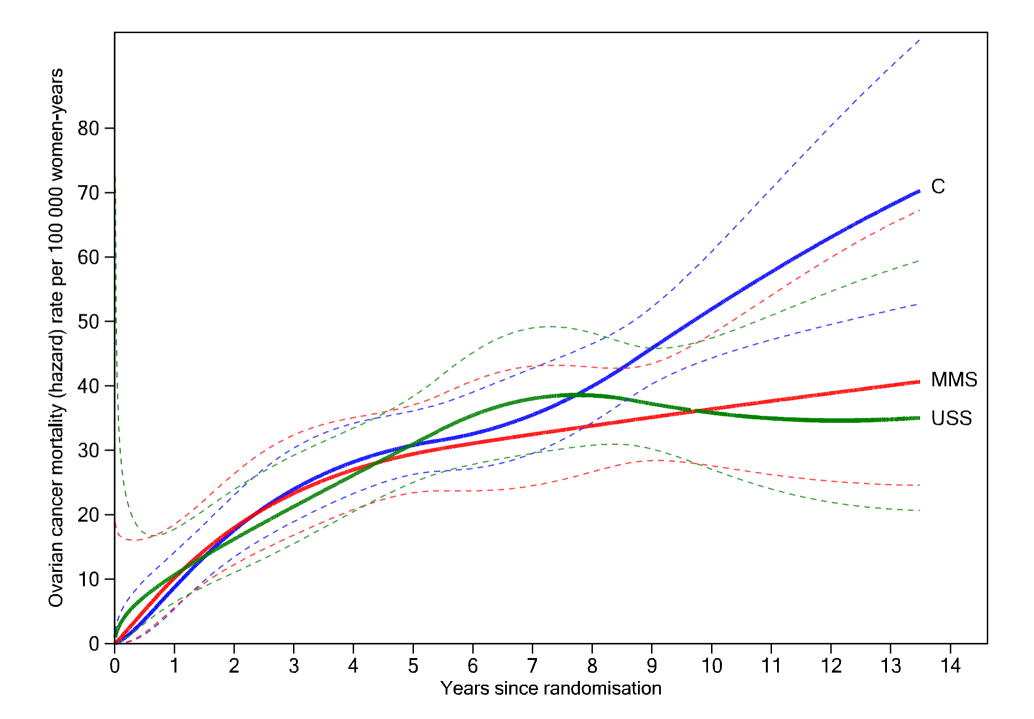


Web Figure 3a: Cumulative ovarian cancer deaths by randomisation group with RP models overlaid - MMS versus no screening (C).

12

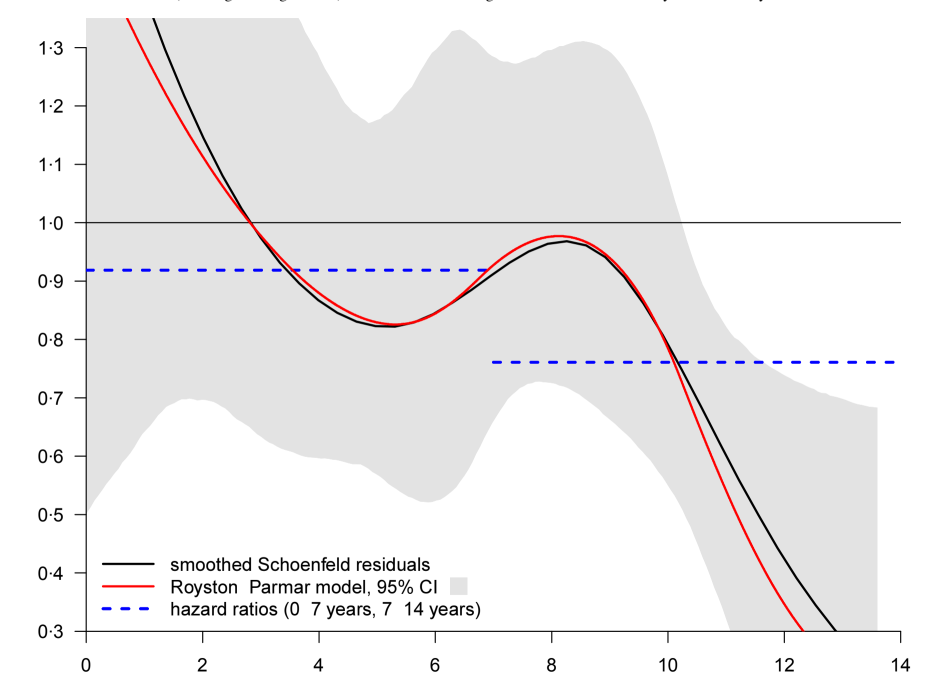






Web Figure 4: Ovarian cancer death rates with confidence limits by randomisation group.

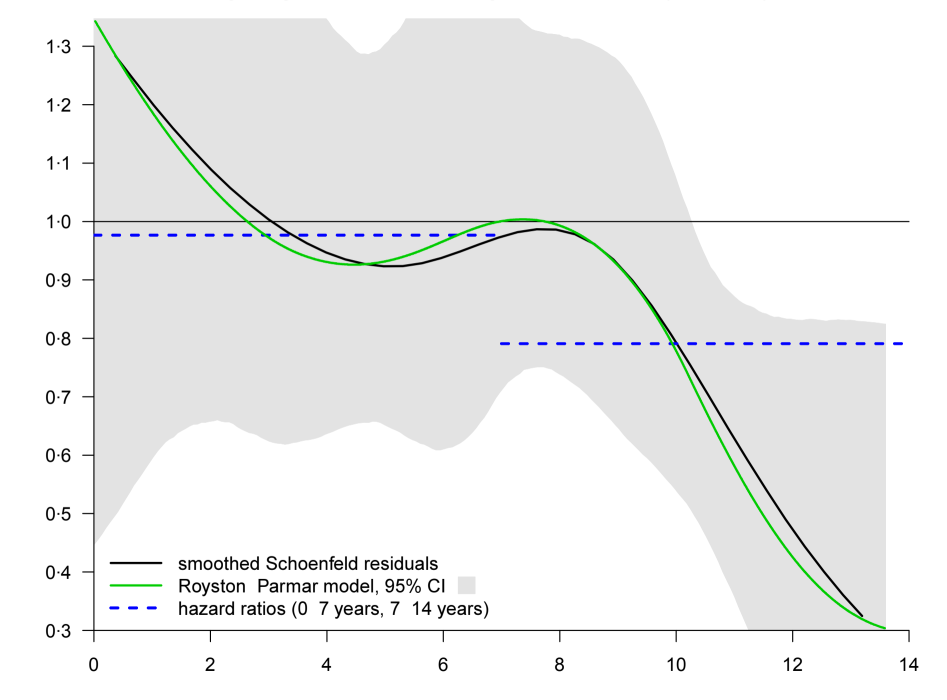
Web Figure 5a: Non-parametric estimate of hazard ratio from smoothed scaled Schoenfeld residuals and parametric estimate from Royston-Parmar model with 95% confidence bands (showing close agreement) and estimates of average hazard ratio between 0-7 years and 7-14 years: MMS versus C



Hazard ratio of MMS to C

Years from randomisation

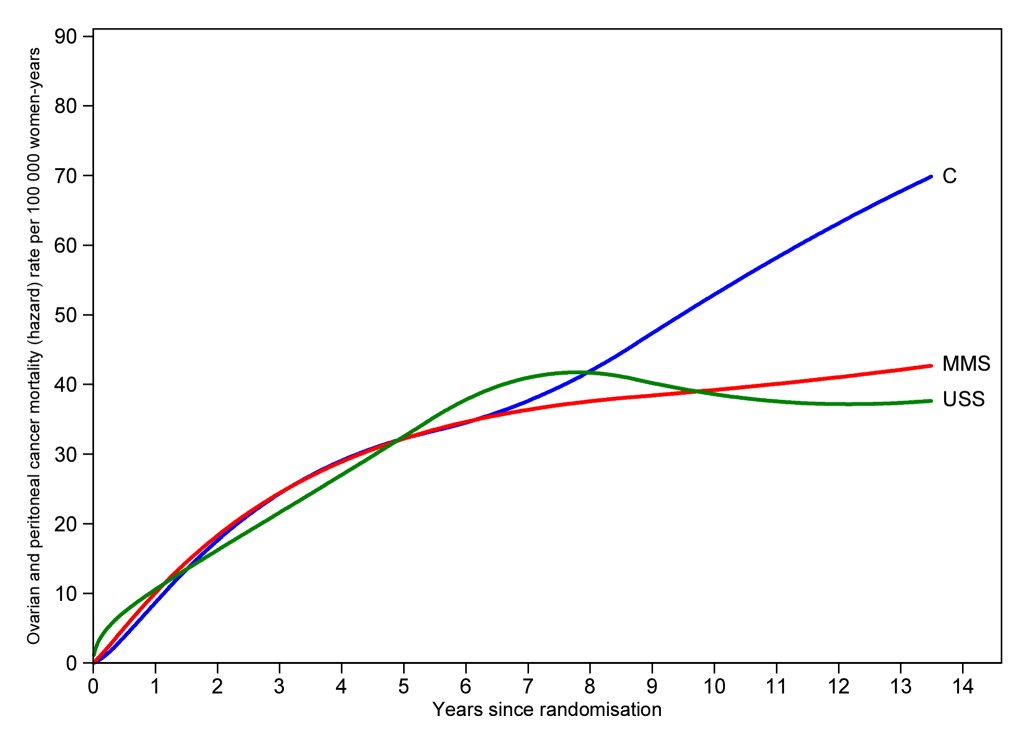
Web Figure 5b: Non-parametric estimate of hazard ratio from smoothed scaled Schoenfeld residuals and parametric estimate from Royston-Parmar model with 95% confidence bands (showing close agreement) and estimates of average hazard ratio between 0-7 years and 7-14 years: USS versus C



Hazard ratio of USS to C

Years from randomisation

Web Figure 6: Rates of ovarian and peritoneal cancer



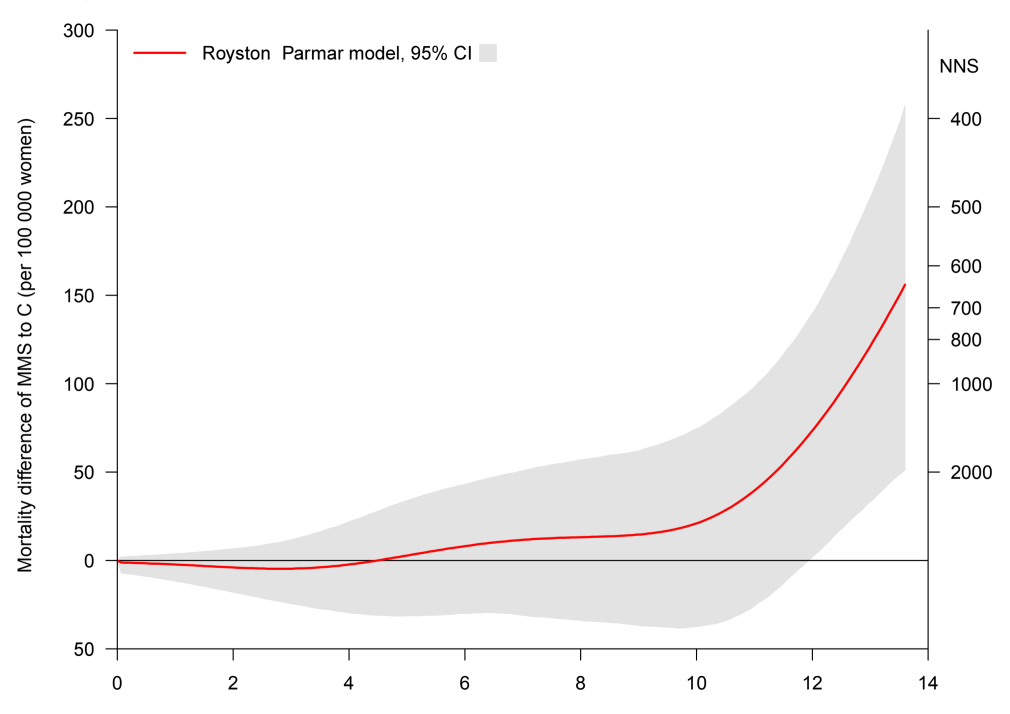
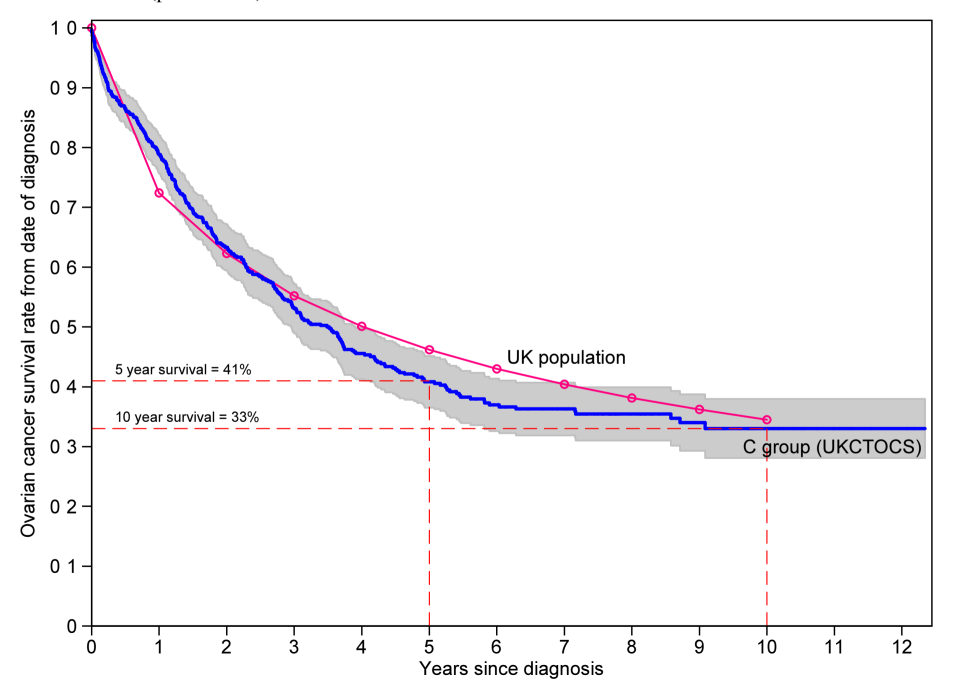
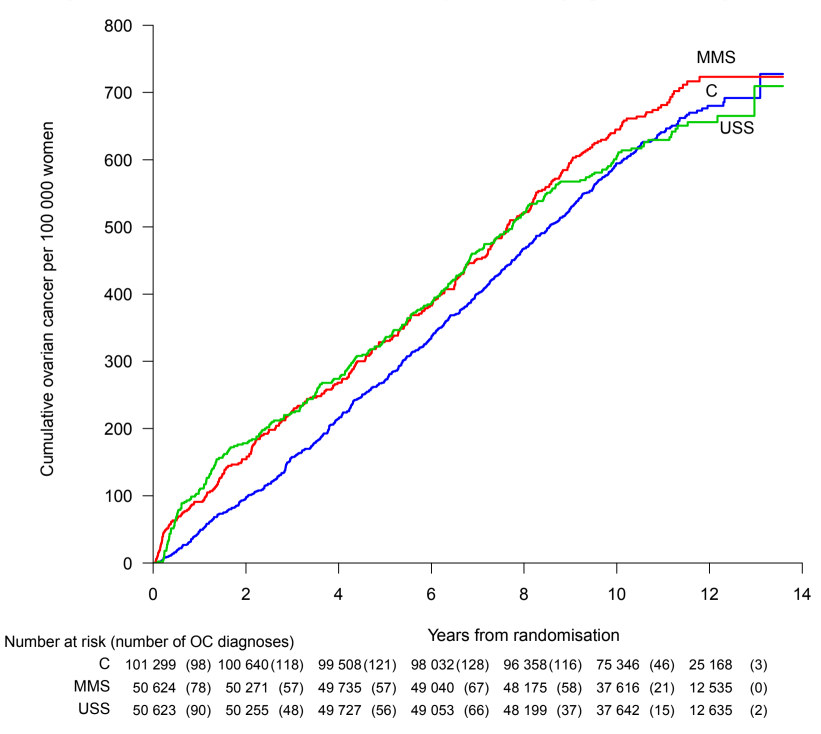


Figure 5: Mortality difference and numbers needed to screen (NNS) to prevent one death from ovarian cancer.

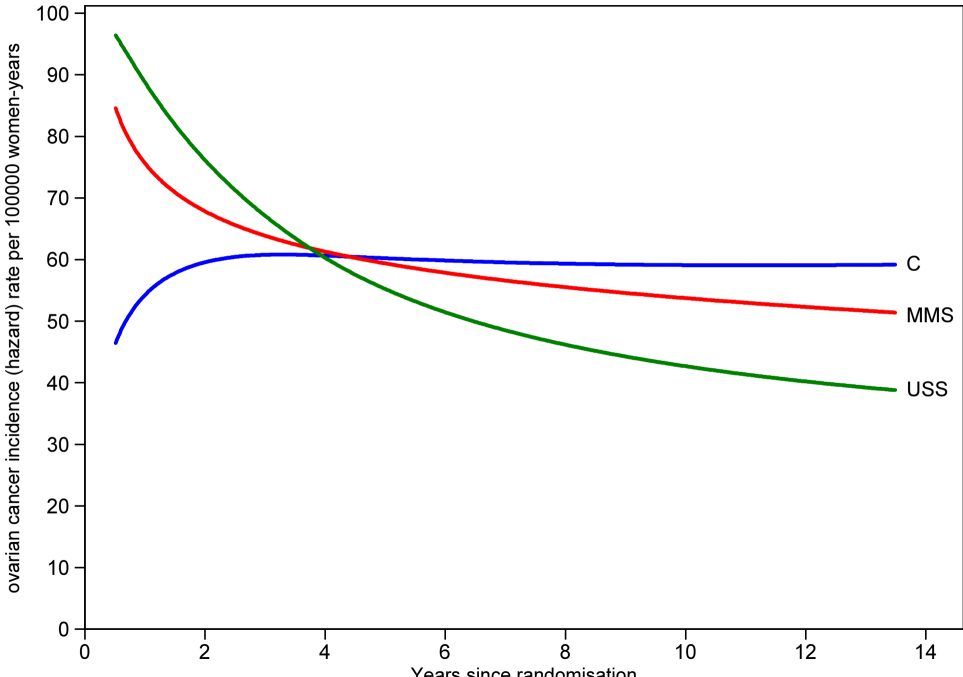
Years from randomisation

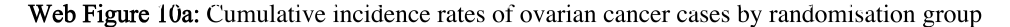
Web Figure 6: Ovarian cancer survival in women with ovarian cancer in the Control group. Overlaid are UKCTOCS 5 and 10 year survival rates (red dash lines) and age-standardized UK population yearly survival rates (pink circles).





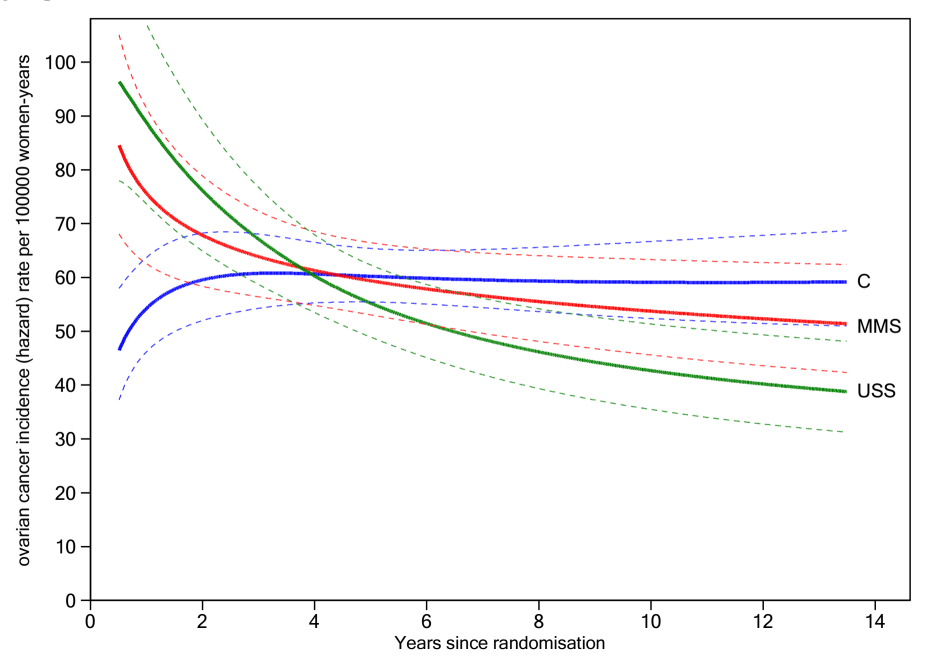
Web Figure 9: Cumulative incidence of ovarian cancer cases by randomization group. (C = no screening)



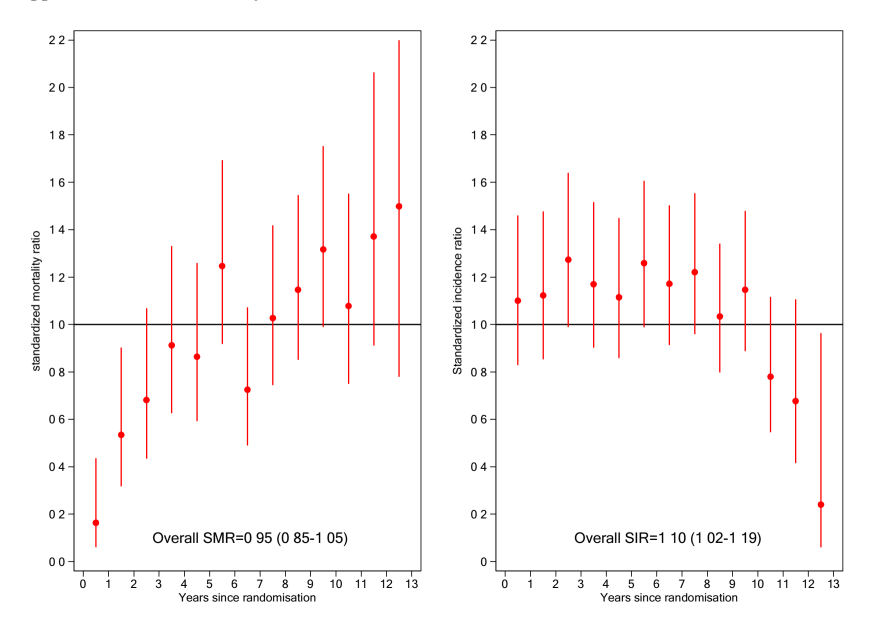


Years since randomisation

Web Figure 10b: Cumulative incidence rates of ovarian cancer cases by randomisation group with 95% confidence bands using Royston-Parmar models with 1 cubic spline knot for each group



Web Figure 11: Standardized mortality ratio estimates (left panel) and standardized incidence ratio estimates (right panel) with 95% confidence limits over time for the Control group. Observed rates have been compared to age-adjusted expected rates based on UK population data. Note, for presentational purposes, the upper-bound for the 13th year SMR estimate has been capped at 2.2, but is actually 2.88.



C. Estimation of CA125 Change-point

To identify the likelihood of a case being prevalent, we used the serial CA125 pattern to estimate whether each cancer's change-point occurred prior to randomisation (T_0). The intersection of the baseline CA125 line and the rising CA125 line estimated the change-point (Supplementary Figure 1a).¹ Cases where the change-point was before T_0 were excluded (Supplementary Figure 1b). This approach provides an objective assessment of which cases are prevalent based on a case's CA125 profile instead of assuming an arbitrary interval of time from T₀ to diagnosis to define prevalent cases. To estimate the change-point for ovarian cases in the no screening group, banked baseline serum samples were available in 517 of the 630 women. These were retrieved and CA125 measured in the CC laboratory using the same assay/analyser as during the trial. 105 baseline MMS samples were also re-assayed for CA125 revealing no significant difference due to long term storage between the new result and the measurement at randomisation. The results combined with CA125 at diagnosis in no screening group cases and the distribution of rate of rise from MMS cases were used to estimate change-points for no screening group cases using the same method as for the MMS cases.¹ For the no screening group cases and some MMS cases, there was uncertainty as to whether the change-point occurred after T_0 (Supplementary Figure 1c). Multiple imputation² of the change-points with 100 imputations accounted for this uncertainty. RP methods³ modelled the cumulative hazards for each arm as a smooth spline function of time with the number of equally spaced knots chosen to minimize the Akaike information criterion (AIC).³ With an RP model we assessed proportionality of hazards and effect of screening with Wald tests³ combined over multiple imputations².

References

- Skates SJ, Pauler DK, Jacobs IJ. Screening based on the risk of cancer calculation from Bayesian hierarchical changepoint and mixture models of longitudinal markers. *Journal of the American Statistical Association* 2001; 96(454): 429-39.
- 2. Rubin DB. Multiple imputation for nonresponse in surveys. New York: John Wiley & Sons, Inc; 1987.
- Royston P, Parmar MK. Flexible parametric proportional-hazards and proportional-odds models for censored survival data, with application to prognostic modelling and estimation of treatment effects. *Statistics in medicine* 2002; 21(15): 2175-97.

D. Outcomes review

Ascertainment of outcomes involved interrogation of all available data sources to identify women diagnosed postrandomisation with any of 19 ICD-10 codes14 (Supplementary Table1). Copies of medical notes were retrieved for all except women who had an ICD10-C80 (malignant neoplasm of uncertain origin) and also had another specific non ovarian/peritoneal cancer registration. The table below details the proportion of women for whom we were able to provide information to the Outcomes review committee in the form of copies of medical notes in addition to cancer and/or death registration. Excluding women with ICD10-C80 (malignant neoplasm of uncertain origin) who also had another specific non ovarian/peritoneal cancer registration, we were able to provide additional information in 99% (1757/1767) of women with regards to cancer diagnosis and 97% (876/900) with regard to death diagnosis. There was no significant differences in the proportion of women with missing data between the randomisation groups.

Availability of information in addition to cancer and/or death registration

Review for cancer diagnosis

	Cases submitted to outcomes review				
	No screening	MMS	USS	Total	
Total	812	495	460	1767	
Additional data available	806	493	458	1757	
No additional data	6 (0.74%)	2 (0.40%)	2 (0.43%)	10 (0.57%)	
Comparison with no screening group (p value)		0.718	0.718		

Review for Death diagnosis

Details -	Cases submitted to outcomes review			
Details	No screening	MMS	USS	Total
Total	450	234	216	900
Additional data available	439	229	208	876
No additional data	11 (2.44%)	5 (2.14%)	8 (3.70%)	24 (2.67%)
Comparison with no screening group (p value)		0.801	0.61	

E. Details of final follow up questionnaire

In 2014 the final follow-up questionnaire was sent to 169 762 (88 743 no screening group, 41 556 MMS, 39 463 USS) women. Questionnaires were not sent to 32 784 (12 556 no screening group, 9 068 MMS, 11 160 USS) women as they had died (12 035), been diagnosed with ovarian cancer (156) or requested not to be contacted (20 593). The response rate was 43 · 1% (38 238/ 88 743) in the no screening group; 55 · 3% (22 975/41 556) in the MMS group and 56 · 5% (22 297/39 463) in the USS group.

F. UKCTOCS committees and teams

DMEC Committee

Prof P Boyle (Chair), Prof APM Heintz, S Kjaer, EL Trimble.

Trial Steering Committee

Prof D Luesley (Chair; independent member), L Bayne (independent member), Prof J Cuzick (independent member), Prof L Fallowfield, Prof I Jacobs, Prof U Menon, Prof M Parmar, Prof J Patnick (independent member).

Trial Management Committee

Prof I Jacobs (Chair), M Burnell, S Campbell, S Davies, Anne Dawnay, A Gentry-Maharaj, J Kalsi, Prof Lesley Fallowfield, Prof A McGuire, Prof U Menon, T Mould, Prof M Parmar, A Ryan, S Skates, R Woolas.

Ultrasound Sub-Committee

Prof U Menon (Chair), Prof N Amso, C Brunell, Prof S Campbell, G Fletcher, K Ford, A Gentry-Maharaj, J Kalsi, R Rangar, A Ryan, M Seif, G Turner.

Outcome Review Committee

N Singh (chair), E Benjamin, K Reynolds, , Prof M Widschwendter.

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Regional Trial Centre Teams

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Barts

D Oram (Lead), Usha Menon (Lead) J Ademi, C Amarasinghe, CM Baque-Juxton, S Bhola, J Bramble, J Chapman, J Charalambous, A Clough, L Cole, L Crosby, J Cunningham, E d'Tisi, E Ferrier, E Forde, P Goulding, B Heyer, J Jonsson, A Knowles, E Liu-Koo, AM Mackinson, V Medic, U Menon, A Relf, K Reynolds, B Rufford, E Ryan, S Sheik, C Stubbs, L Walters, D Warrington, J Webb.

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Hailward, J Hazelton, M Herod, S Inwood, S Jones, V Jones, L Korb, H Lee, L Limbert, K Lord, J Maloney, M Maraj, J McCarthy, L McGlynn, D Ndlela, J Newman, A Nicolson, K Pearson, S Pennington, D Petter, P Stewart, A Tannock, B Thomson, A Webster, J Webster, S West, H Wright, G Zabroski.

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Manchester

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Derby

H Jenkins[§] (Lead), I Scott (Lead)A Bali, J Barke, C Benson, C Bower-Smith, H Bullock, J Caborn, S Crockett, A Ferguson, G Forbes, J Gomes, R Harrison, C Hollins, M Jones, A North, R Rock, M Scott, H Stanton, S Thompson, M Tudge, G Turner, J Weston, C Williams.

Royal Free

T Mould (Lead), I Aitken, S Amin, I Beal, S Bhola, S Blackmoore, K Borroughs, H Brown, S Burke, D Colia, L Crosby, G Desai, C Efueye, H Evans, E Ferrier, K Fitzgerald, G Fletcher, C Fox, G Gaston, K Harvey, B Heyer, T Hitchen, K Isherwood, E Izard, E Koo, M Lagos, K Lakhani, AM Mackinson, L McKenzie, V Medic, S Mohamed, S More, K Muir, W Myburg, N Nayak, O Ojo, N Old, A Oldham, S Porcherot, E Rawstron, M Sharma, T Stevens, C Sundstrom, E Sweeney, J Terwin, D Townshend, F Turner, S Wellington, D White, J Wickes, L Young, E Zard.

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R Woolas (Lead), S Aldcock, M Anderson, E Barclay, ? Bell, R Bonner, E Bowes, D Brinkmann, J Burns, K Chorley, C Dhar, K Fairley, F Gardner, B Gibbs, Y Griffiths, R Harrison, L Hayward, C Ihezue, C Isaac, D James, F Jones, D Mason, E Merritt, R Morris, M Oakey, J Skinner, S Tilbury, J Turpitt, A Webb, C West, J Woolas.

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Belfast

S Dobbs (Lead), M Alkalbani, M Carey, J Caruth, M Clarke, J Forbes, M Gallagher, B Gibbs, M Hendron, R McClelland, M McComskey, D Morgan, M Murray, A O'Donnell, C Poh, J Price, H Sheldon, M Yoong, A Zawislak.

Cardiff

N Amso (Lead), S Basu, D Bell, H Clarke, A Evans, J Evans, T Griffiths, P Henderson, R Howells, R Jones, G Jose, G Looker, C Morgan, G Rieck, A Rogers, A Sharma, A Sims, S Underwood, D Williams, G Williams.

North Wales

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Middlesbrough

D Cruickshank (Lead), A Bullen, V Chadwick, K Chapman, J Francis, R Goldie, C Ikwan-McCabe, K Jan, D Khan, L Lewis, J Nevin, L Prentis, J Proll, R Shanbhag, G Tarr