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Intraoperative frozen section analysis for the diagnosis of early stage ovarian cancer in suspicious pelvic masses

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ABSTRACT

Background

Women with suspected early-stage ovarian cancer need surgical staging which involves taking samples from areas within the abdominal cavity and retroperitoneal lymph nodes in order to inform further treatment. One potential strategy is to surgically stage all women with suspicious ovarian masses, without any histological information during surgery. This avoids incomplete staging, but puts more women at risk of potential surgical over-treatment.

A second strategy is to perform a two-stage procedure to remove the pelvic mass and subject it to paraffin sectioning, which involves formal tissue fixing with formalin and paraffin embedding, prior to ultrathin sectioning and multiple site sampling of the tumour. Surgeons may then base further surgical staging on this histology, reducing the rate of over-treatment, but conferring additional surgical and anaesthetic morbidity.

A third strategy is to perform a rapid histological analysis on the ovarian mass during surgery, known as 'frozen section'. Tissues are snap frozen to allow fine tissue sections to be cut and basic histochemical staining to be performed. Surgeons can perform or avoid the full surgical staging procedure depending on the results. However, this is a relatively crude test compared to paraffin sections, which take many hours to perform. With frozen section there is therefore a risk of misdiagnosing malignancy and understaging women subsequently found to have a presumed early-stage malignancy (false negative), or overstaging women without a malignancy (false positive). Therefore it is important to evaluate the accuracy and usefulness of adding frozen section to the clinical decision-making process.

Objectives

To assess the diagnostic test accuracy of frozen section (index test) to diagnose histopathological ovarian cancer in women with suspicious pelvic masses as verified by paraffin section (reference standard).

Search methods

We searched MEDLINE (January 1946 to January 2015), EMBASE (January 1980 to January 2015) and relevant Cochrane registers.

Selection criteria

Studies that used frozen section for intraoperative diagnosis of ovarian masses suspicious of malignancy, provided there was sufficient data to construct 2 x 2 tables. We excluded articles without an available English translation.

Data collection and analysis

Authors independently assessed the methodological quality of included studies using the Quality Assessment of Diagnostic Accuracy Studies tool (QUADAS-2) domains: patient selection, index test, reference standard, flow and timing. Data extraction converted 3 x 3 tables of per patient results presented in articles into 2 x 2 tables, for two index test thresholds.

Main results

All studies were retrospective, and the majority reported consecutive sampling of cases. Sensitivity and specificity results were available from 38 studies involving 11,181 participants (3200 with invasive cancer, 1055 with borderline tumours and 6926 with benign tumours, determined by paraffin section as the reference standard). The median prevalence of malignancy was 29% (interquartile range (IQR) 23% to 36%, range 11% to 63%). We assessed test performance using two thresholds for the frozen section test. Firstly, we used a test threshold for frozen sections, defining positive test results as invasive cancer and negative test results as borderline and benign tumours. The average sensitivity was 90.0% (95% confidence interval (CI) 87.6% to 92.0%; with most studies typically reporting range of 71% to 100%), and average specificity was 99.5% (95% CI 99.2% to 99.7%; range 96% to 100%).

Similarly, we analysed sensitivity and specificity using a second threshold for frozen section, where both invasive cancer and borderline tumours were considered test positive and benign cases were classified as negative. Average sensitivity was 96.5% (95% CI 95.5% to 97.3%; typical range 83% to 100%), and average specificity was 89.5% (95% CI 86.6% to 91.9%; typical range 58% to 99%).

Results were available from the same 38 studies, including the subset of 3953 participants with a frozen section result of either borderline or invasive cancer, based on final diagnosis of malignancy. Studies with small numbers of disease-negative cases (borderline cases) had more variation in estimates of specificity. Average sensitivity was 94.0% (95% CI 92.0% to 95.5%; range 73% to 100%), and average specificity was 95.8% (95% CI 92.4% to 97.8%; typical range 81% to 100%).

Our additional analyses showed that, if the frozen section showed a benign or invasive cancer, the final diagnosis would remain the same in, on average, 94% and 99% of cases, respectively.

In cases where the frozen section diagnosis was a borderline tumour, on average 21% of the final diagnoses would turn out to be invasive cancer.

In three studies, the same pathologist interpreted the index and reference standard tests, potentially causing bias. No studies reported blinding pathologists to index test results when reporting paraffin sections.

In heterogeneity analyses, there were no statistically significant differences between studies with pathologists of different levels of expertise.

Authors' conclusions

In a hypothetical population of 1000 patients (290 with cancer and 80 with a borderline tumour), if a frozen section positive test result for invasive cancer alone was used to diagnose cancer, on average 261 women would have a correct diagnosis of a cancer, and 706 women would be correctly diagnosed without a cancer. However, 4 women would be incorrectly diagnosed with a cancer (false positive), and 29 with a cancer would be missed (false negative).

If a frozen section result of either an invasive cancer or a borderline tumour was used as a positive test to diagnose cancer, on average 280 women would be correctly diagnosed with a cancer and 635 would be correctly diagnosed without. However, 75 women would be incorrectly diagnosed with a cancer and 10 women with a cancer would be missed.

The largest discordance is within the reporting of frozen section borderline tumours. Investigation into factors leading to discordance within centres and standardisation of criteria for reporting borderline tumours may help improve accuracy. Some centres may choose to perform surgical staging in women with frozen section diagnosis of a borderline ovarian tumour to reduce the number of false positives. In their interpretation of this review, readers should evaluate results from studies most typical of their population of patients.

PLAIN LANGUAGE SUMMARY

Is a 'quick diagnosis' test on an ovarian mass during surgery accurate?

The issue

When women go to their doctor with a mass that could be ovarian cancer, they are normally referred for surgery, since the mass may need to be removed and examined microscopically in a laboratory in a procedure known as paraffin section histopathology. A third of women with ovarian cancer present with a cyst or mass without any visible evidence of spread elsewhere. However, in these apparently early-stage cancers (confined to the ovary) surgical staging is required to decide if chemotherapy is required. This staging consists of sampling tissues within the abdomen, including lymph nodes.

Different staging strategies exist. One is to perform surgical staging for all women who might have a cancer, to get information about spread. This may result in complications due to additional surgical procedures that may turn out to be unnecessary in approximately two thirds of women.

A second strategy is to perform an operation to remove just the suspicious mass and await the paraffin section diagnosis. This may result in needing a further operation in one third of women if cancer is confirmed, putting them at increased risks from another operation.

A third strategy is to send the mass to the laboratory during the operation for a quick diagnosis, known as 'frozen section'. This helps the surgeon decide if further surgical treatment is required during a single operation.

Why is this review important?

Frozen section is not as accurate as the traditional slower paraffin section examination, and it entails a risk of incorrect diagnosis, meaning that some women may not have all the samples taken at the initial surgery and may need to undergo a second operation; and others may undergo unnecessary surgical sampling.

How was this review conducted?

We searched all available studies reporting use of frozen section in women with suspicious ovarian masses. We excluded studies without an English translation and studies without enough information to allow us to analyse the data.

What are the findings?

We included 38 studies (11,181 women), reporting three types of diagnoses from the frozen section test.

1. Cancer, which occurred in an average of 29% of women.
2. Borderline tumour, which occurred in 8% of women.
3. Benign mass.

In a hypothetical group of 1000 patients where 290 have cancer and 80 have a borderline tumour, 261 women would receive a correct diagnosis of a cancer and 706 women would be correctly diagnosed without a cancer based on a frozen section result. However, 4 women would be incorrectly diagnosed as having a cancer where none existed (false positive), and 29 women with cancer would be missed and potentially need further treatment (false negative).

If surgeons used a frozen section result of either a cancer or a borderline tumour to diagnose cancer, 280 women would be correctly diagnosed with a cancer and 635 women would be correctly diagnosed without a cancer. However, 75 women would be incorrectly diagnosed as having a cancer, and 10 women with cancer would be missed on the initial test and found to have a cancer after surgery.

If the frozen section result reported the mass as benign or malignant, the final diagnosis would remain the same in, on average, 94% and 99% of the cases, respectively.

In cases where the frozen section diagnosis was a borderline tumour, there is a chance that the final diagnosis would turn out to be a cancer in, on average, 21% of women.

What does this mean?

Where the frozen section diagnosis is a borderline tumour, the diagnosis is less accurate than for benign or malignant tumours. Surgeons may choose to perform additional surgery in this group of women at the time of their initial surgery in order to reduce the need for a second operation if the final diagnosis turns out to be a cancer, as it would on average in one out of five of these women.

BACKGROUND

Target condition being diagnosed

In 2012, 238,719 women worldwide were diagnosed with epithelial ovarian cancer (EOC), and 151,905 died from the disease, corresponding to an annual incidence of 6.1 cases per 100,000 women, an annual mortality rate of 4.3 deaths per 100,000 and a cumulative lifetime risk of 0.5% (GLOBOCAN 2012). Ovarian cancer is the sixth most common cancer and the seventh most common cause of cancer death in women. A woman's risk of developing cancer of the ovary by age 75 years varies between countries, ranging from 0.5% to 1.6%. This corresponds to an age-standardised rate of ovarian cancer from 5 to 14 cases per year in 100,000 women under 75. In Europe, 30% to 44% of women with ovarian cancer are alive five years after diagnosis (EUROCORE 2003). The poor survival associated with ovarian cancer is largely because most women are diagnosed when the cancer is already at an advanced stage (Jemal 2008), with only 30% being early-stage, that is, confined to the ovary (NCIN 2015).

Early-stage ovarian cancer, or stage I and II according to the International Federation of Gynecology and Obstetrics (FIGO 2015), has a combined incidence of less than 35%, with five-year survival rates of 92% and 55% for stage I and stage II, respectively (Cancer Research UK 2012) (Appendix 1).

Currently, women presenting with a pelvic mass suspected of being ovarian cancer are triaged according to the risk of malignancy index (RMI) (Bailey 2006; Jacobs 1990). RMI is a product ($RMI = U \times M \times CA125$) of suspicious ultrasound features of the mass (multilocular cysts, solid areas, metastases, ascites and bilateral lesions), menopausal status (postmenopausal = 3) and serum CA125 levels (IU/ml). There is some uncertainty as to the optimal threshold for the RMI; however, women with high RMIs (e.g. > 200) are usually scheduled for staging laparotomy at cancer centres.

Unlike advanced disease, early disease may not be obvious at surgery. It is up to gynaecological cancer centres to decide how they manage these masses. Some may choose to await final histology before planning surgical staging; others may opt to surgically stage those with high RMI; and others may employ frozen section analysis to provide an intraoperative diagnosis.

The value of surgical staging is to detect micrometastases, present in approximately 25% cases of invasive epithelial ovarian cancer, which are not macroscopically evident and would warrant adjuvant chemotherapy (Helewa 1986).

Index test(s)

Intraoperative frozen section histopathological analysis of a suspicious pelvic mass may facilitate the appropriate selection of women requiring surgical staging. Frozen sections are not routinely performed in all gynaecological cancer units in the United Kingdom;

as a result, optimal surgical staging may be omitted at primary laparotomy, particularly in early-stage disease. In the ICON 1 trial, adjuvant platinum-based chemotherapy offered improved overall survival in clinically stage I disease (ICON 1). This study included 93% of cases with clinically early-stage disease who underwent hysterectomy, bilateral salpingo-oophorectomy and omentectomy as a minimum surgical procedure. Optimal staging was therefore not performed.

The importance of optimal surgical staging was further highlighted in the ACTION study (Trimbos 2003), which showed that in a subgroup analysis on the effect of surgical staging, the benefit of adjuvant chemotherapy appeared to be limited to patients who underwent suboptimal staging and so had a higher risk of undetected residual disease. In a subgroup analysis of patients with optimal surgical staging, adjuvant chemotherapy was not associated with overall or recurrence-free survival. Optimal staging was shown to be an independent prognostic factor for progression-free and overall survival. Optimal staging included omentectomy, washings, peritoneal biopsies, and pelvic and paraaortic lymph node sampling. Women with early stage epithelial ovarian cancer who undergo optimal surgical staging survive longer than those suboptimally staged (Trimbos 2010). However, this benefit of surgical staging did not reach significance in a recent Cochrane review update of these studies with 10-year outcome data (Lawrie 2015). If the frozen section shows a suspicious pelvic mass to be benign, a full staging procedure is not necessary, and fertility-sparing surgery could be offered if appropriate. Unnecessary surgical staging can lead to lymphoedema, lymphocyst formation, and visceral and neurovascular injury. Lymphoedema and lymphocyst formation are often chronic conditions that can negatively affect quality of life.

If the frozen section shows a borderline or malignant ovarian tumour, surgeons often perform optimal staging. Therefore, the potential benefits of performing intraoperative frozen section include: reducing surgical morbidity associated with unnecessary optimal staging; reducing the need for a second surgical procedure to complete surgical staging where it has been suboptimal; and reducing operating costs.

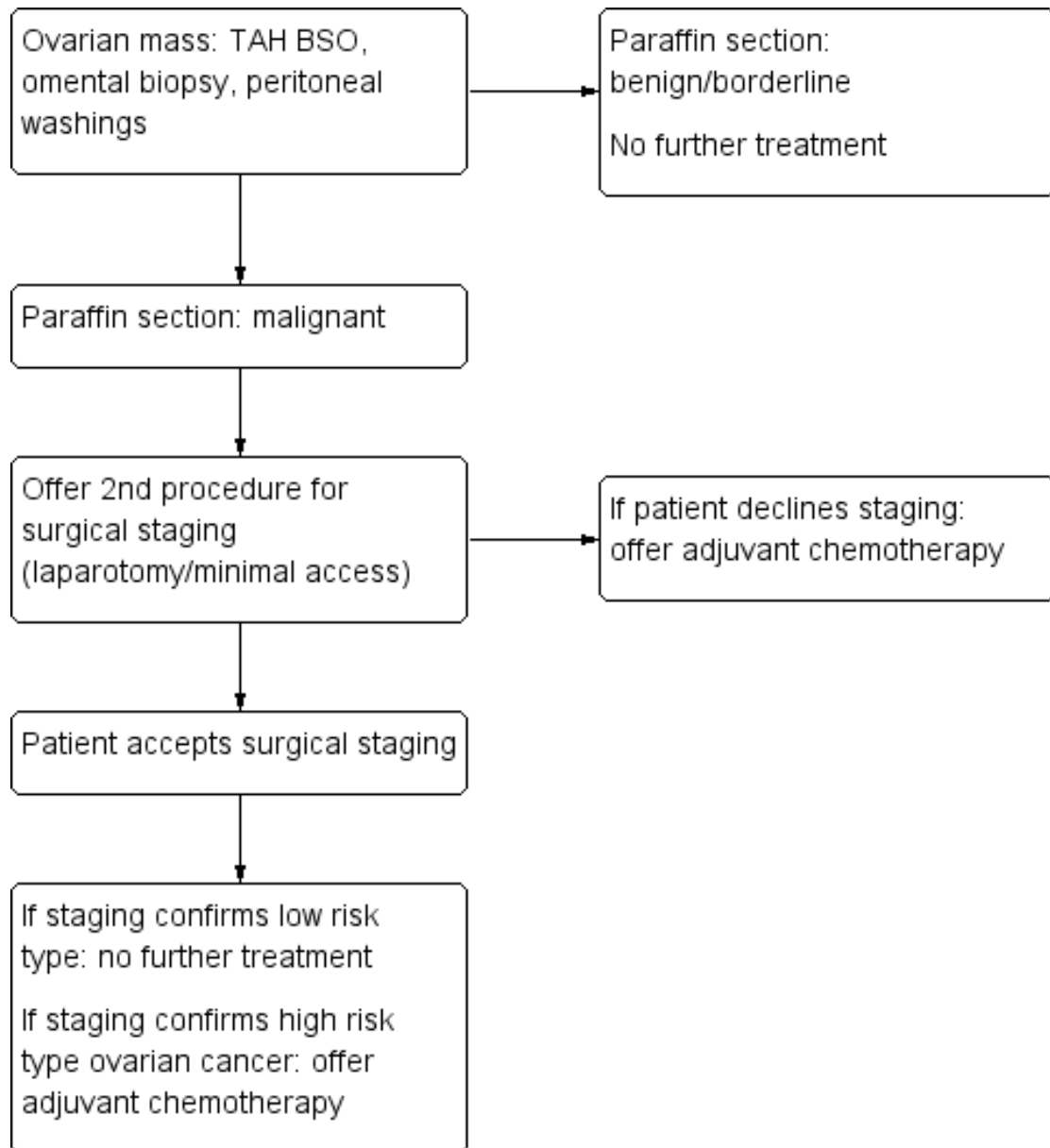
Several studies and reviews have reported high sensitivity, specificity and overall accuracy when comparing intraoperative frozen section with paraffin section examination (Cross 2012; Gol 2003; Medeiros 2005; Naik 2006).

Clinical pathway

Most women with suspected early ovarian cancer undergo surgical staging to identify metastases. This optimally includes inspection and palpation of peritoneal cavity and organs, biopsy of peritoneum and suspicious nodules, peritoneal washings, total abdominal hysterectomy (TAH), bilateral salpingo-oophorectomy (BSO), omentectomy, and retroperitoneal lymph node assessment and sampling. An appendicectomy can be performed if

the tumour is mucinous. Failure to complete the above staging in cases of malignancy is called suboptimal staging. Diagnosis is confirmed by paraffin section examination of surgical specimens, which is usually reported a few days after the surgery. This is the 'gold standard' of histopathological reporting (Figure 1).

Figure 1. Flow diagram showing clinical pathway if no frozen section available and staging offered based on paraffin section



When centres use paraffin sections to guide management rather than frozen sections, a second surgical procedure may be required in order to complete staging in women with confirmed ovarian cancer. This may increase anxiety in addition to increasing the risk of surgical and anaesthetic morbidity.

Where frozen section is used in the clinical pathway, it is used on the same tissue samples as will be used for paraffin section, but allows decisions about the need for further surgical staging to be made within the same operation. Where the frozen section result is benign, patients need only be offered further surgical staging if the subsequent paraffin section result is malignant (

Figure 2). Where the frozen section result is malignant, patients can be offered immediate surgical staging without the need for a second surgery (Figure 2). Where the frozen section result is borderline, there are two options (Figure 3). In option 1, the clinical team and patient agree in advance to await the paraffin section result, with further surgical staging or adjuvant chemotherapy offered if subsequent paraffin result is malignant. In option 2, they agree in advance to proceed to immediate surgical staging. Adjuvant chemotherapy decisions are made on the basis of paraffin section test results (Figure 1, Figure 2, Figure 3).

Figure 2. Flow diagram showing clinical pathway for frozen section benign or malignant and surgical staging offered. Example average numbers are shown for a hypothetical population of 1000 women, with prevalences of malignancy 29%, borderline 8%, benign 63%. Prevalences are based on averages across all included studies.

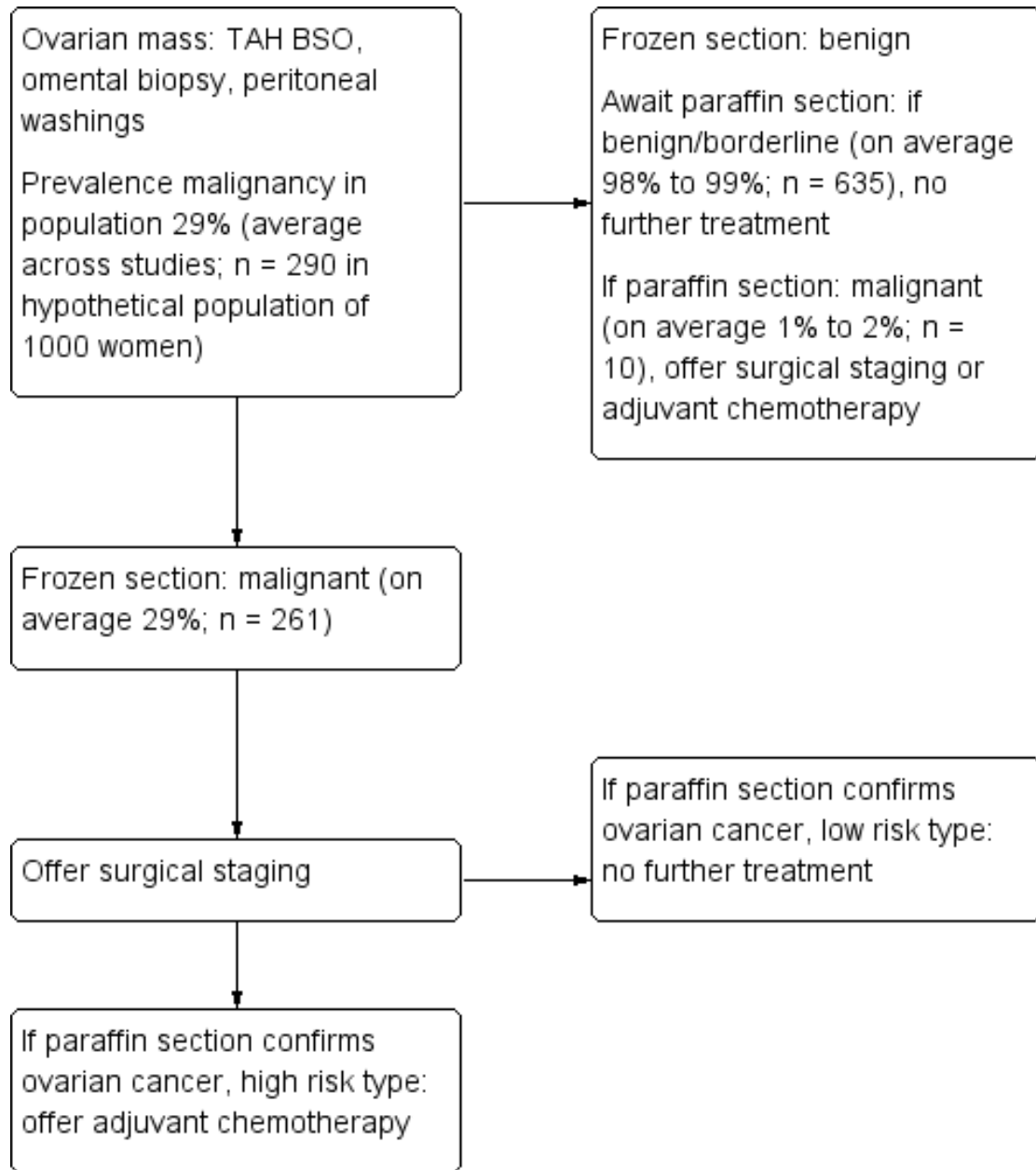
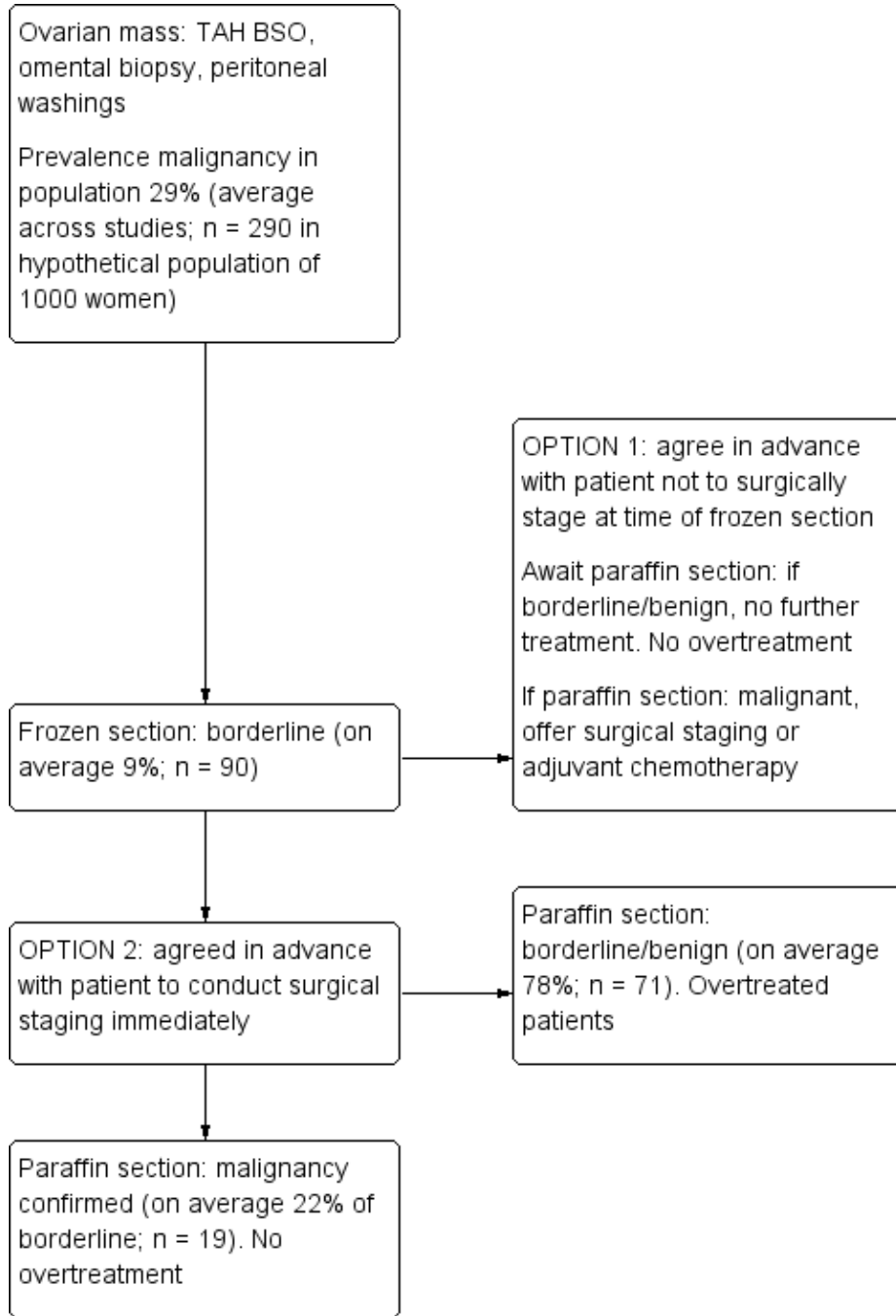


Figure 3. Flow diagram showing clinical pathway options (stage or not to stage) for borderline frozen section diagnosis. Example average numbers are shown for a hypothetical population of 1000 women, with prevalences of malignancy 29%, borderline 8%, benign 63%. Prevalences are based on averages across all included studies.



In women with macroscopically evident stage III disease, there is no need to use frozen section to confirm malignancy. However, as discussed earlier, there is a proportion of women with apparently stage I disease who have microscopic involvement of paraaortic lymph nodes or omentum and are upstaged after surgical staging. These women are offered dual-agent chemotherapy with a taxane. In women with low risk disease, that is, stage IA grade 1 disease; or in those with comprehensively staged stage IB grade 1-2 disease, experts thought until recently that there was no survival advantage associated with adjuvant chemotherapy (Winter-Roach 2012). However, an update of these data suggest that there may be a longer term advantage of chemotherapy, even in these women (Lawrie 2015). For women who have had suboptimal staging, clinicians should discuss a second surgical staging procedure or adjuvant chemotherapy (NICE 2011). Therefore, the clinical consequence of suboptimal surgical staging is that women who appear to have low-risk stage I ovarian cancer may require adjuvant chemotherapy.

In the cases of high risk disease, that is, stage IB grade 3; stage IC and higher; and clear cell cancers, six cycles of adjuvant platinum-based chemotherapy are recommended (NICE 2011). Due to the good response rate to chemotherapy, early-stage serous ovarian cancers are often treated with six cycles of adjuvant dual-agent chemotherapy, including a taxane. With regard to clear cell cancers, as the response to chemotherapy is often poor, there may be a therapeutic benefit to performing lymphadenectomy in case of micrometastases. It should be noted, however, that NICE guidance recommends assessment of retroperitoneal lymph nodes but not systematic or block dissection of retroperitoneal lymph nodes in women with clinically apparent stage I disease. Maggioni 2006 demonstrated that patients undergoing systematic pelvic and paraaortic lymphadenectomy compared to sampling were found to have more micrometastases (22% versus 9%), leading to up-staging of apparent stage I disease to stage IIIC. However, there was greater morbidity, operating time and hospital cost with no demonstrable overall survival advantage.

Prior test(s)

Serum CA125 and abdominal ultrasound are performed as part of the RMI assessment. Women presenting with a pelvic mass and a high RMI score will usually undergo a computed tomography (CT) scan, magnetic resonance imaging (MRI) of the pelvis and abdomen, or both to establish the extent of disease. Interpretation of the histology slides at frozen section is made independently of these prior tests, and so these bear no relevance to the diagnostic accuracy of the frozen section test.

Role of index test(s)

Intraoperative frozen section analysis may allow appropriate selection of women with suspicious pelvic masses who would benefit from optimal surgical staging.

Alternative test(s)

The 'gold standard' for the diagnosis of ovarian cancer is histopathological examination of surgical specimens by paraffin section after laparotomy. In women not undergoing frozen section, surgeons may choose to await paraffin section histology prior to performing staging; to stage all women deemed to have a high risk of malignancy; or to stage according to clinical suspicion, for example by performing biopsy on peritoneal adhesions or sampling enlarged lymph nodes.

Rationale

The importance of optimal surgical staging in ovarian cancer is now well established. Frozen section analysis at diagnostic laparotomy may allow the surgeon to accurately identify those women with early stage ovarian cancer (who may otherwise not have been identified during the initial procedure) who will benefit from optimal surgical staging. This may avoid the need for a subsequent restaging procedure or adjuvant chemotherapy (Trimbos 2003). The role of intraoperative frozen section analysis in the diagnosis and management of early stage disease is particularly topical at present, with many recent studies reporting high sensitivities and specificities for this diagnostic test. We decided to review the evidence for and against frozen section as an accurate test to diagnose early ovarian cancer.

OBJECTIVES

Primary objectives

To assess the diagnostic test accuracy of frozen section (index test) in the histopathological diagnosis of ovarian cancer in women with suspicious pelvic masses as verified by paraffin section (reference standard).

Within our review we aimed to establish the diagnostic accuracy of frozen section in comparison to a reference standard diagnosis of cancer from paraffin section, using measures of sensitivity and specificity. There were two primary objectives.

1. To determine the accuracy of frozen section to identify cancer cases, using a test threshold for frozen section that defines cancer as a positive test result and considers both borderline and benign results as test negative (Table 1). The rationale is that clinical and surgical management is different where a case of malignancy is identified.

2. To assess the accuracy of frozen section to identify cancer, using a test threshold for frozen section that defines both cancer and borderline cases as positive test results and considers benign results as test negative (Table 2). The rationale is that the literature reports a high rate of cases where the frozen section result was borderline, but the final result from paraffin section was malignant. There are

potentially serious repercussions from managing patients with a cancer outside a cancer pathway when 'under staging' occurs, that is, if patients with malignancy do not receive surgical staging, including lymphadenectomy. This is particularly relevant in women found to have borderline ovarian masses at frozen section, as many will receive a final paraffin section diagnosis of malignancy. Which threshold is considered most useful in practice depends on the clinicians' judgement.

Secondary objectives

1. To establish if intraoperative frozen section analysis allows the surgeon to accurately identify the cases of early stage ovarian cancer that may benefit from optimal surgical staging.
2. To assess the accuracy of final diagnosis of malignancy, in a subgroup of women with a frozen section result of either borderline or cancer. This corresponds to one strategy for referral for cancer treatment.

METHODS

Criteria for considering studies for this review

Types of studies

We included studies published in any language and, where possible, had non-English articles translated. We excluded studies that involved ten or fewer patients.

Studies were eligible if:

1. Both frozen section analysis and paraffin section analysis were performed in the same patient;
 2. The absolute numbers of observations of true positives, false positives, false negatives and true negatives were available or derivable from the data reported in the primary studies.
- We included both prospective and retrospective studies. However, we excluded retrospective studies that collected data for a specific histological type only, such as borderline tumours, due to the risk of reporting bias.

We excluded studies in which frozen section analysis was performed for conditions other than ovarian malignancy as well as studies for which no English translation was available.

Participants

Women presenting to a secondary or tertiary care setting with a pelvic mass suspicious of ovarian cancer, in whom physicians employed frozen section analysis prior to paraffin section analysis.

Index tests

Intraoperative frozen section histopathological analysis. Test results were classified as malignant, borderline or benign. We present results using two different thresholds for the index test; malignant vs borderline/benign, and malignant/borderline vs benign. The reference standard remains diagnosis of malignancy in all analyses in the review.

The diagnostically important distinction to make is between malignant/borderline and benign frozen section, because although only women with malignant disease require surgical staging, studies have found the risks of borderline frozen section returning as malignant to be high, and inadequately staging these women at primary laparotomy may be deemed unacceptable (Cross 2012; Puls 1997). However, many would argue that performing unnecessary staging on women with borderline disease confers unnecessary morbidity.

Target conditions

Ovarian malignancy, not obvious at a surgically or radiologically advanced stage.

Reference standards

Paraffin section histopathological analysis. Test results are classified as malignant, borderline or benign. We present results using the threshold for women classified as having ovarian cancer as 'malignant' versus women not having cancer as 'borderline or benign'.

Search methods for identification of studies

Electronic searches

We identified eligible studies by searching the following electronic databases.

- The Cochrane Gynaecological Cancer Group Specialised Register January 2015.
- Cochrane Central Register of Controlled Trials (CENTRAL), (2015, Issue 1).
- MEDLINE - Ovid (January 1946 to January 2015).
- EMBASE - Ovid (January 1980 to January 2015).
- Database of Abstracts of Reviews of Effects (DARE) (Issue 4, 2014).
- Health Technology Assessments (HTA) Database (Issue 4, 2014).

The MEDLINE search strategy included both subject headings (MeSH terms) and text words for the target condition (ovarian malignancy) and the histological technique under investigation ('Frozen Section analysis'). We did not apply language restrictions. We adapted the MEDLINE search to search CENTRAL, EMBASE, DARE and HTA databases. In particular, we adapted the

MEDLINE MeSH terms into the corresponding terms available in the EMTREE vocabulary. We present full details of the MEDLINE and EMBASE search strategies, together with a brief summary of the MEDLINE search strategy, in [Appendix 2](#). We imported all citations identified by the MEDLINE and EMBASE search strategies into an electronic database. We identified all potentially eligible articles on PubMed and used the 'related articles' feature to carry out a further search for newly published papers.

Searching other resources

Unpublished and grey literature

We searched for ongoing trials in the following trial registers and contacted experts in the field to identify any further ongoing trials.

- *meta*Register of Controlled Trials (*mRCT*) (<http://www.controlled-trials.com/mrct/>).
- Physicians Data Query (PDQ) (<http://www.cancer.gov/cancertopics/pdq>).
- ClinicalTrials.gov (<http://clinicaltrials.gov/>).
- National Cancer Institute (<http://www.cancer.gov/clinicaltrials/search>).

Handsearching

We handsearched the citation lists of included studies, key textbooks and existing systematic reviews and checked their references. When we retrieved relevant studies (even if we finally excluded them), we also searched their references in order to minimise the potential for missing relevant studies. We handsearched conference reports in the following sources.

- *Gynecologic Oncology* (Annual Meeting of the American Society of Gynecologic Oncologists).
- *International Journal of Gynecologic Cancer* (Annual Meeting of the International Gynecologic Cancer Society).
- British Gynaecological Cancer Society.
- European Society of Gynaecological Oncology.
- Society of Gynaecological Oncologists.

Data collection and analysis

Selection of studies

We downloaded all titles and abstracts retrieved by electronic searching to Endnote and removed duplicates. Two authors (NR and AP) independently examined the remaining reference titles and abstracts to retrieve the full text of all potentially relevant reports. Three authors (NR, AB and CF) independently reviewed all relevant reports according to the pre-defined inclusion criteria to determine eligibility. We resolved any disagreements through arbitration by another author (RS), and we documented reasons for exclusions.

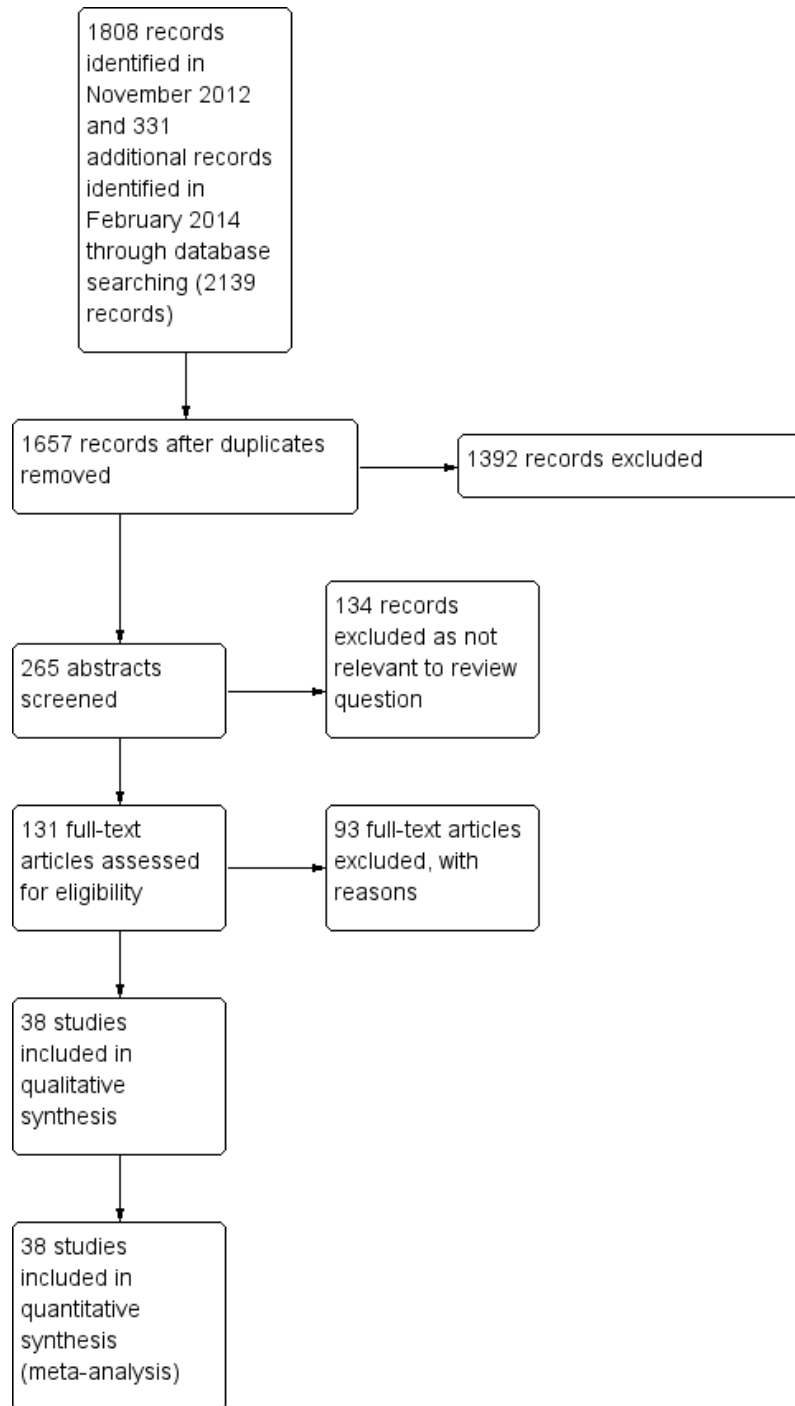
Data extraction and management

One author (TL) designed and trialled a data extraction form specifically to collect details from selected studies. Two authors (NR and AB) recorded the relevant information for each individual study, without concealing the study authorship or publication details. This information included: lead author, year of publication, accrual dates, country and setting, study design, method of recruitment, setting, number and characteristics of participants, any additional preoperative investigations performed, the reference standard used, any comparator tests used, follow-up, and information related to the pathologists interpreting the specimens (background specialty, level of expertise). Two authors (NR and AB) extracted data from the selected reports, and two authors (RS and SM) checked the data extractions.

Assessment of methodological quality

Two review authors (NR and AB) independently assessed the methodological quality of each included study using the revised Quality Assessment of Diagnostic Accuracy Studies tool (QUADAS-2) ([Whiting 2011](#)). We consulted a third author (RS) in case of discrepancy between authors. The QUADAS-2 tool is structured into a series of questions in four domains that should be answered 'yes', 'no' or 'unclear', and it aims to evaluate the spectrum of bias. We resolved any disagreements by discussion. [Figure 4](#) is a graphic summary of the methodological quality of included studies.

Figure 4. Study flow diagram.



We assessed the core QUADAS items in the following domains: patient selection, index test, reference standard, flow and timing.

Patient selection

Could the selection of patients have introduced bias?

1. Was a consecutive or random sample of patients enrolled?
 - i) Yes; a study ideally should enrol a consecutive or random sample of eligible patients with suspected disease to prevent the potential for bias.
 - ii) No, a non-consecutive sample of patients was used.
 - iii) Unclear.
2. Was a case-control design avoided?
 - i) Yes; studies enrolling participants with known disease and a control group without the condition may exaggerate diagnostic accuracy.
 - ii) No.
 - iii) Unclear.
3. Did the study avoid inappropriate exclusions?
 - i) Yes; studies that make inappropriate exclusions (for example, not including 'difficult-to-diagnose' patients) may result in overestimation of diagnostic accuracy.
 - ii) No.
 - iii) Unclear.
4. Were the patients selected representative of the patient population the index test would apply to?
 - i) Yes; patients with a high risk of malignancy index (RMI > 200) are usually the subjects who would benefit from this index test.
 - ii) No; use of the index test in patients at low risk of malignancy or in those with incidental finding at laparotomy for other condition may bias the results.
 - iii) Unclear.

Index test

Could the conduct or interpretation of the index test have introduced bias?

1. Were the index test results interpreted without knowledge of the results of the reference standard?
 - i) Yes; knowledge of the reference standard may influence interpretation of index test results. The potential for bias is related to the subjectivity of interpreting index test and the order of testing. If the index test is always conducted and interpreted before the reference standard, this item can be rated 'yes'.
 - ii) No; if there was a previous histological diagnosis of malignancy made during investigation of the same cyst, this item can be rated 'no'.
 - iii) Unclear.
2. Were the index test results interpreted by a pathologist specialising in gynaecological oncology?

- i) Yes; specialist centres employing dedicated gynaecological oncology pathologists may perform better in interpreting frozen section slides and thereby improve the sensitivity and specificity of the test.

- ii) No.
- iii) Unclear.

Reference standard

Could the reference standard, its conduct or its interpretation have introduced bias?

1. Is the reference standard likely to correctly classify the target condition?
 - i) Yes; estimates of test accuracy are based on the assumptions that the reference standard is 100% sensitive and that specific disagreements between the reference standard and index test result from incorrect classification by the index test.
 - ii) No.
 - iii) Unclear.
2. Were the reference standard results interpreted without knowledge of the results of the index test?
 - i) Yes; knowledge of the index test results may influence interpretation of the reference standard results. Potential for bias is related to the potential influence of previous knowledge on the interpretation of the reference standard.
 - ii) No.
 - iii) Unclear.

Flow and timing

Could the patient flow have introduced bias?

1. Did all patients receive a reference standard, and if so did they receive the same reference standard?
 - i) Yes; verification bias occurs when only a proportion of the study group receives confirmation of the diagnosis by the reference standard, or if some patients receive a different reference standard. If the results of the index test influence the decision on whether to perform the reference standard or which reference standard is used, estimated diagnostic accuracy may be biased. Accepted best practice is to verify all frozen section diagnoses with paraffin section histology.
 - ii) No.
 - iii) Unclear.
2. Were all patients included in the analysis?
 - i) Yes; all participants recruited to the study should be included in the analysis. A potential for bias exists if the number of patients enrolled differs from the number of patients included in the 2 x 2 table of results, because patients lost to follow-up differ systematically from those who remain.

- ii) No.
- iii) Unclear.

Statistical analysis and data synthesis

We entered data into Cochrane's statistical software, [Review Manager 2014](#), to calculate sensitivity and specificity for each study (we also present 95% confidence intervals of these point estimates in a forest plot). We present individual study results graphically by plotting estimates of sensitivities and specificities in receiver operating characteristic (ROC) space. All studies reported 3 x 3 tables per patient enabling extraction of 2 x 2 tables from all studies for three analyses of accuracy:

1. Reference test (paraffin test): positive result malignancy, negative result borderline or benign. Index test (frozen section): positive result malignancy, negative result borderline or benign ([Table 1](#)).
2. Reference test: positive result malignancy, negative result borderline or benign. Index test: positive result malignancy or borderline, negative result benign ([Table 2](#)).
3. Subgroup analysis of malignant and borderline by index test. Reference test: positive result malignancy, negative result borderline or benign. Index test: positive result malignancy, negative result borderline or benign.

We used `xtmelogit` commands in the Stata 13.1 statistical package ([Stata 2013](#)) to meta-analyse pairs of sensitivity and specificity using a bivariate random-effects approach ([Reitsma 2005](#)). The bivariate approach was suitable for test results from 2 x 2 tables based on categorical test thresholds. This approach enabled us to calculate summary estimates of sensitivity and specificity, while correctly dealing with any correlation that might exist between sensitivity and specificity as well as the following sources of variation.

1. Imprecision in measurement of sensitivity and specificity within each study.
2. Variation beyond chance in sensitivity and specificity between studies.

We incorporated covariates in the bivariate model in order to examine the effect of potential sources of heterogeneity on sensitivity and specificity. We used the results of the bivariate model to calculate likelihood ratio tests in order to assess the statistical significance of covariates.

Investigations of heterogeneity

We constructed a ROC plot of sensitivity versus 1 – specificity and explored the heterogeneity of the sensitivity and specificity estimates by examining both the ROC plot and forest plot. In the protocol, we proposed to run a separate heterogeneity analysis for the following situations if there were sufficient studies reporting differences in these study characteristics.

- Preoperative investigation including a combination of imaging and tumour markers (CA 125 +/- HE4).

- Preoperative imaging including CT or MRI scans.
- RMI score > 200.
- High risk study population, for example in a tertiary referral centre.
- Size of ovarian cyst.
- Ovarian cyst histological type, for example mucinous or serous.
- Expertise of pathologist reporting.

However, studies reported only two of these characteristics: expertise of reporting pathologist and whether there was a high risk study population, for example, in a tertiary referral centre. All studies took place in university hospitals or tertiary referral centres, so we could not examine for heterogeneity of study setting. We were able to conduct a heterogeneity analysis for expertise of pathologist reporting, for primary objective #1 and secondary objective #2, although the model did not converge in a heterogeneity analysis of primary objective #2. For primary outcome #2, the model did not converge, as there were only four studies in the less experienced group, one of which was [Toneva 2012](#), where specificity was low (mostly likely due to small study size bias). Data extraction grouped pathologist expertise into four categories as described in the studies (specialist gynaecological pathologist, consultant pathologist, general pathologist or reader expertise not recorded). For heterogeneity analyses, we divided readers into more experienced (specialist gynaecological pathologist or consultant pathologist) and less experienced/unknown expertise (general pathologist or reader expertise not recorded). We conducted covariate analysis specifying reader expertise as a covariate in STATA as recommended in the Methods of the *Cochrane Handbook for Diagnostic Test Accuracy Reviews* ([Macaskill 2010](#)).

Sensitivity analyses

We had planned sensitivity analyses for studies without verification bias and those without missing data.

Assessment of reporting bias

We documented data regarding loss to follow-up and any loss of data from pre-specified outcomes. As recommended in , we did not conduct analyses to test for reporting bias ([Macaskill 2010](#)).

RESULTS

Results of the search

Results of the combined CENTRAL, MEDLINE, EMBASE, DARE and HTA searches until January 2015 yielded 1657 records. Four review authors (NR, AB, AP and CF) independently screened and reviewed the titles and abstracts. Of these, 131 were selected

for classification. Two authors (NR and AB) read the full-text articles and assessed eligibility for the review. We discussed any dispute with a third author (RS) (Figure 4). We excluded 93 studies for the reasons summarised below. Some were excluded for more than one reason.

- They were reviews, editorials, commentaries, case reports, surveys, letters to the editor or conference abstracts (26).
- They were meta-analyses (2).
- An English translation was not available (6).
- We were unable to construct 2 x 2 tables from the results (13).
- They were not studies using frozen section intraoperative diagnosis (33).
- They reported only certain histologies (epithelial, serous or mucinous) (11).
- They reported only borderline diagnoses (8); these studies were not representative of the preselected population and did not meet the inclusion criteria.
- They did not represent the population studied by this review (20); these studies included predominantly benign populations and populations in which evidence of extra-ovarian spread was present at time of frozen section.

For further details see [Characteristics of excluded studies](#).

We included 38 studies in 11,181 women. All studies evaluated

the index test of frozen section in comparison to the reference standard of paraffin section.

Methodological quality of included studies

Of the 38 included studies, we considered 1 study to be at high risk of bias (Wang 1998), and we had concerns regarding the applicability in 2 studies (Ilvan 2005; Wang 1998). Wang 1998 reported outcomes of 792 consecutive gynaecological frozen sections, which included 299 samples from ovarian tissue, 360 samples from lymph nodes, 56 from uterine tissue and 77 samples from other sites. The same pathologist reported the paraffin and frozen sections. Ilvan 2005 reported making 7.5% of their frozen section diagnoses on gross/macrosopic inspection alone. In fact, grossly benign specimens were submitted in 46 cases. In some cases, two experienced pathologists in gynaecological pathology employed touch imprint methodology for diagnosis as well.

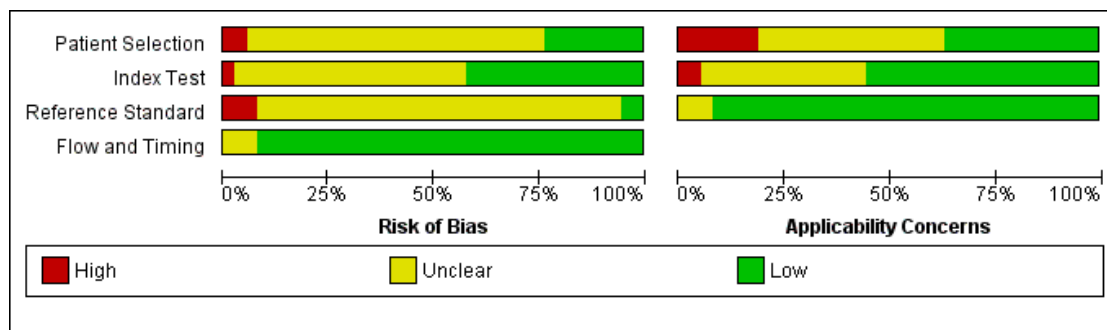
Overall, we found that the quality of the included studies was acceptable with a low or unclear risk of bias (Figure 5; Figure 6). However, we note that in many studies our assessment of risk of bias was unclear; for example, it is not clear if pathologists interpreted the reference test (paraffin) without knowledge of the index test (frozen section), but this is unlikely to have introduced bias in the diagnosis of ovarian cancer.

Figure 5. Risk of bias and applicability concerns summary: review authors' judgements about each domain for each included study

| | Risk of Bias | | | | Applicability Concerns | | |
|------------------------|-------------------|------------|--------------------|-----------------|------------------------|------------|--------------------|
| | Patient Selection | Index Test | Reference Standard | Flow and Timing | Patient Selection | Index Test | Reference Standard |
| Agikalın 2014 | ? | ? | ? | ● | ? | ? | ? |
| Bazot 2006 | ? | ● | ? | ● | ? | ? | ● |
| Bige 2011 | ? | ● | ? | ● | ● | ● | ● |
| Boriboonhirunsarn 2004 | ? | ● | ? | ● | ? | ● | ● |
| Canis 2004 | ● | ? | ? | ● | ● | ● | ● |
| Cross 2012 | ● | ● | ? | ● | ● | ● | ● |
| Cuello 1999 | ? | ? | ? | ● | ● | ● | ● |
| Fanfani 2007 | ● | ● | ? | ● | ● | ● | ● |
| García 1997 | ? | ? | ? | ● | ? | ? | ● |
| Gorisek 2009 | ? | ? | ? | ? | ? | ? | ● |
| Hamed 1993 | ● | ? | ? | ● | ● | ? | ? |
| Ilker 2011 | ● | ● | ? | ● | ● | ● | ● |
| Ivan 2005 | ? | ● | ? | ● | ● | ● | ● |
| Kokka 2009 | ● | ? | ? | ● | ● | ● | ● |
| Lim 1997 | ? | ● | ? | ● | ● | ● | ● |
| Maheshwari 2006 | ? | ? | ? | ● | ? | ? | ● |
| Malipattil 2013 | ? | ● | ? | ● | ? | ● | ● |
| Naik 2006 | ● | ? | ? | ● | ● | ● | ● |
| Pavakis 2009 | ? | ? | ? | ● | ● | ? | ? |
| Pinto 2001 | ? | ? | ? | ● | ? | ? | ● |
| Puls 1997 | ● | ? | ● | ● | ● | ● | ● |
| Rakhshan 2009 | ? | ● | ? | ● | ? | ● | ● |
| Rose 1994 | ? | ? | ? | ● | ? | ? | ● |
| Stewart 2006 | ● | ● | ● | ● | ● | ● | ● |
| Subbian 2013 | ? | ● | ? | ? | ● | ● | ● |
| Sukumaran 2014 | ? | ? | ? | ● | ● | ● | ● |
| Suprasert 2008 | ? | ? | ? | ● | ? | ? | ● |
| Tangjitgamol 2004 | ? | ? | ? | ● | ? | ? | ● |
| Taskiran 2008 | ● | ? | ? | ● | ● | ? | ● |
| Toneva 2012 | ? | ● | ? | ● | ● | ● | ● |
| Torres 1998 | ? | ? | ? | ● | ? | ? | ● |
| Twaalfhoven 1991 | ? | ? | ? | ? | ? | ? | ● |
| Wakahara 2001 | ? | ● | ? | ● | ? | ● | ● |
| Wang 1998 | ? | ● | ● | ● | ● | ● | ● |
| Wasinghon 2008 | ● | ? | ? | ● | ● | ● | ● |
| Wootpoom 2006 | ? | ? | ? | ● | ? | ? | ● |
| Yarandi 2008 | ? | ● | ? | ● | ● | ● | ● |
| Yeo 1998 | ? | ● | ● | ● | ? | ● | ● |

● High ? Unclear ● Low

Figure 6. Risk of bias and applicability concerns graph: review authors' judgements about each domain presented as percentages across included studies



Types of studies

We included 38 retrospective studies. There were no case-control studies. The sampling methods were consecutive in most studies and unreported in the rest. All took place in university hospitals or tertiary care settings (see [Characteristics of included studies](#)). We excluded eight studies because we could not extract 2 x 2 tables; these studies only included cases with borderline results by frozen section.

The interest in borderline ovarian tumours at frozen section diagnosis arises from the fact that this diagnostic group is most likely to see a change in diagnosis at paraffin section. This fact has been attributed to various factors, namely ovarian histology, size of mass and expertise of pathologist. In this review, two studies discussed the ability of frozen section to predict malignancy depending on histology ([Cross 2012](#); [Puls 1997](#)). [Puls 1997](#) included only serous or mucinous ovarian masses, analysing the effect of weight on interpretation of frozen section and reporting the greatest discordance between frozen and paraffin section in frozen section-reported borderline mucinous masses weighing over 1360g, with 50% (four out of eight) being upgraded to malignant at paraffin section. [Cross 2012](#) reported the majority of sampling errors in serous and mucinous tumours, which accounted for 52.3% of their 1439 ovarian masses submitted for frozen section. The false negative rate for serous tumours was 0.7% and for mucinous tumours 3.8%. Furthermore, 47.2% of all borderline tumours were reclassified as malignant on paraffin section, and these were evenly distributed amongst the serous and mucinous categories.

Patient selection

We considered participants in the included studies to be representative of patients receiving the index test in clinical practice. The majority of studies reported women with pelvic masses, although none provided information regarding tumour markers, such as CA125, or preoperative imaging.

Index test methods

All patients in the included studies received the index test, namely frozen section, and a number of studies provided details of frozen sectioning. Typically, this involved taking between 1 and 7 sections from the ovarian mass, cut into 5 μ m thick frozen sections. Pathologists of varying expertise performed analyses of the frozen section: specialist gynaecological pathologist (6), consultant pathologist (8) or general pathologist (4); studies did not record expertise in 20 cases.

Several studies reported 'deferred' or unclear diagnoses at frozen section, where the pathologist was unable to make a diagnosis on the submitted material. We excluded these results from 3 x 3 tables. Pathologists may defer diagnosis to paraffin section for 3 reasons: not enough tissue is submitted for analysis; the pathologist is unable to make a diagnosis; or there are technical issues.

Reference standard methods

All patients received the reference standard, namely paraffin section. In three studies ([Puls 1997](#); [Wang 1998](#); [Yeo 1998](#)), the same pathologist interpreted the paraffin section and the frozen section. The other included studies did not mention whether they employed the same pathologist to interpret both index and reference

standard tests. No studies reported blinding of pathologists to index test results when reporting paraffin sections. There were no biases with flow or timing, as paraffin section was always performed after frozen section, with both tests conducted on samples taken at the same time.

Flow and timing

All patients who received frozen section then received paraffin section. There was no bias in flow or timing amongst included studies. The only potential source of bias was interpretation of both reference and index tests by the same reporting pathologist, as indicated in the studies of [Puls 1997](#), [Wang 1998](#) and [Yeo 1998](#). In clinical practice, it is likely that most surgeons, at the time of submitting the surgical specimen for paraffin section, will indicate to the pathologist that a frozen section has already been performed and detail the results of the frozen section. As blinding from this has not been reported in any of the included studies, it is entirely reasonable to believe that pathologists were aware of the frozen section results when interpreting the paraffin section. Paraffin section analysis was performed on the same submitted mass as the frozen section, and therefore time interval to paraffin section was not an issue, as there was no risk of disease progression between tests.

Investigations of heterogeneity

Unfortunately, only one study gave adequate information about histology of all frozen sections performed ([Cross 2012](#)). [Puls 1997](#) gave enough information for 2 x 2 tables to be constructed for serous, mucinous and endometrioid tumours only. We were therefore unable to perform a heterogeneity analysis according to histopathological type.

None of the included studies provided sufficient information regarding preoperative investigations or imaging, RMI value, or size of mass to conduct heterogeneity analyses.

We investigated variability between studies to establish whether levels of expertise of pathologists reading the frozen section results could explain heterogeneity between studies. We found that there was no statistically significant difference between studies with different levels of expertise of pathologists in primary outcome #1 and secondary outcome #2. For primary outcome #2, the model did not converge, as there were only four studies in the less experienced group, one of which was [Toneva 2012](#), where specificity was low (mostly likely due to small study size bias). The lack of heterogeneity due to expertise of pathologists may be due to the fact that the included studies originated from university hospitals or tertiary centres.

Sensitivity analyses

We did not conduct sensitivity analyses, as all studies excluded verification bias. A sensitivity analysis based on missing data will be included in a review update, but we note there was only a small amount of missing data.

Findings

Thirty-eight studies were suitable for addressing the review objectives, as we were able to extract 3 x 3 tables from all studies based on thresholds of cancer, borderline and benign for both frozen and paraffin section results. There were a total of 169 deferred diagnoses excluded from 11,350 cases (1.5% of all cases), leaving 11,181 cases for analysis. Unfortunately, only one study commented on surgical staging with regard to frozen section, and therefore we could not perform an analysis to address secondary objective #1 ([Naik 2006](#)). The results addressing the two primary objectives and secondary objective #2 are detailed below. We summarise these results in the [Summary of findings](#), giving different examples of pre-test prevalences of malignancy to allow clinicians to infer the relevance of the data according to their population.

Primary objective #1: accuracy of frozen section cancer results to identify women with cancer

Sensitivity and specificity results were available from 38 studies involving 11,181 participants (3200 identified with cancer, 1055 as borderline and 6926 as benign by paraffin section reference standard). We used a test threshold for frozen sections to define cancer as a positive test result and borderline and benign results as negative test results. The prevalence of cancer ranged from 11% to 63%. The average sensitivity was 90.0% (95% confidence interval (CI) 87.6% to 92.0%; typical range 71% to 100%, with one small study, [García 1997](#), reporting it as 64%), and the average specificity was 99.5% (95% CI 99.2% to 99.7%: range 96% to 100%).

[Figure 7](#) is a forest plot of sensitivity and specificity with 95% confidence intervals for all studies, ordered by the percentage of cancer cases, that is, disease positive (DP) in each study to give insight into the representativeness of the study. [Figure 8](#) shows the results from all studies in a ROC plot. Both figures show that the data are homogeneous enough to combine by meta-analysis and give summary estimates. The average sensitivity was 90.0% (95% CI 87.6% to 92.0%; range 64% to 100%), and the average specificity was 99.5% (95% CI 99.2% to 99.7%: range 96% to 100%). Results for specificity were more homogenous than for sensitivity, where we have ordered studies in the forest plot by the number of cancer cases. Studies with small numbers of cancer cases have wider confidence intervals for sensitivity, and study estimates are likely to be less reliable. This is particularly pertinent for [García 1997](#), with 11 cancer cases.

Figure 7. Forest plot: frozen section threshold malignant vs borderline or benign

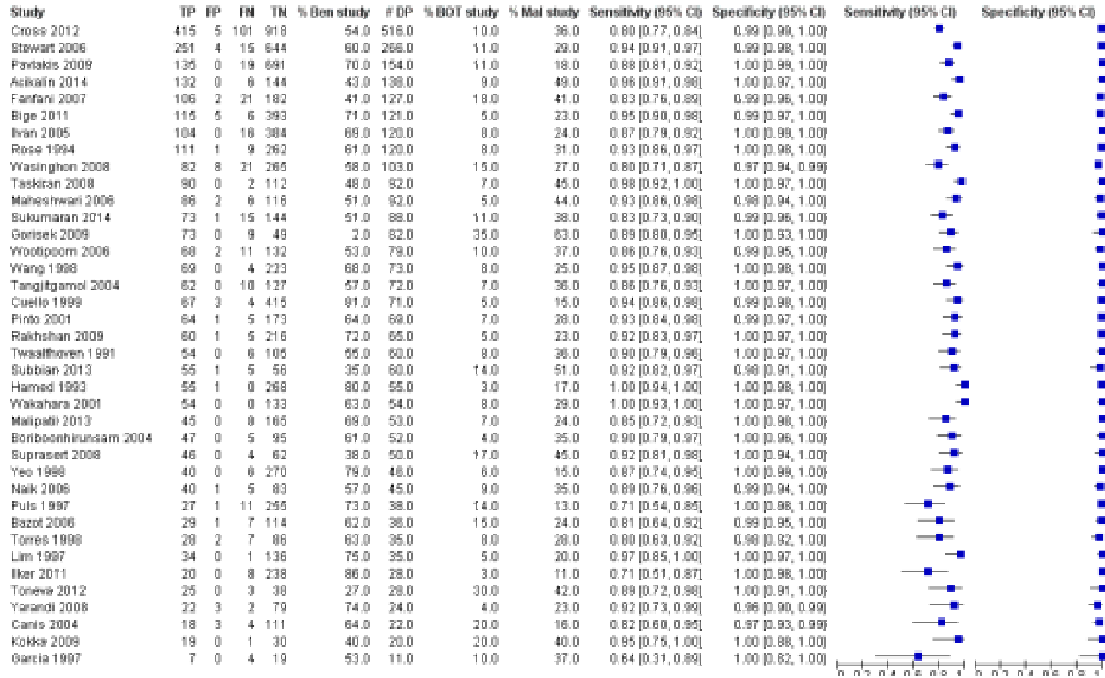
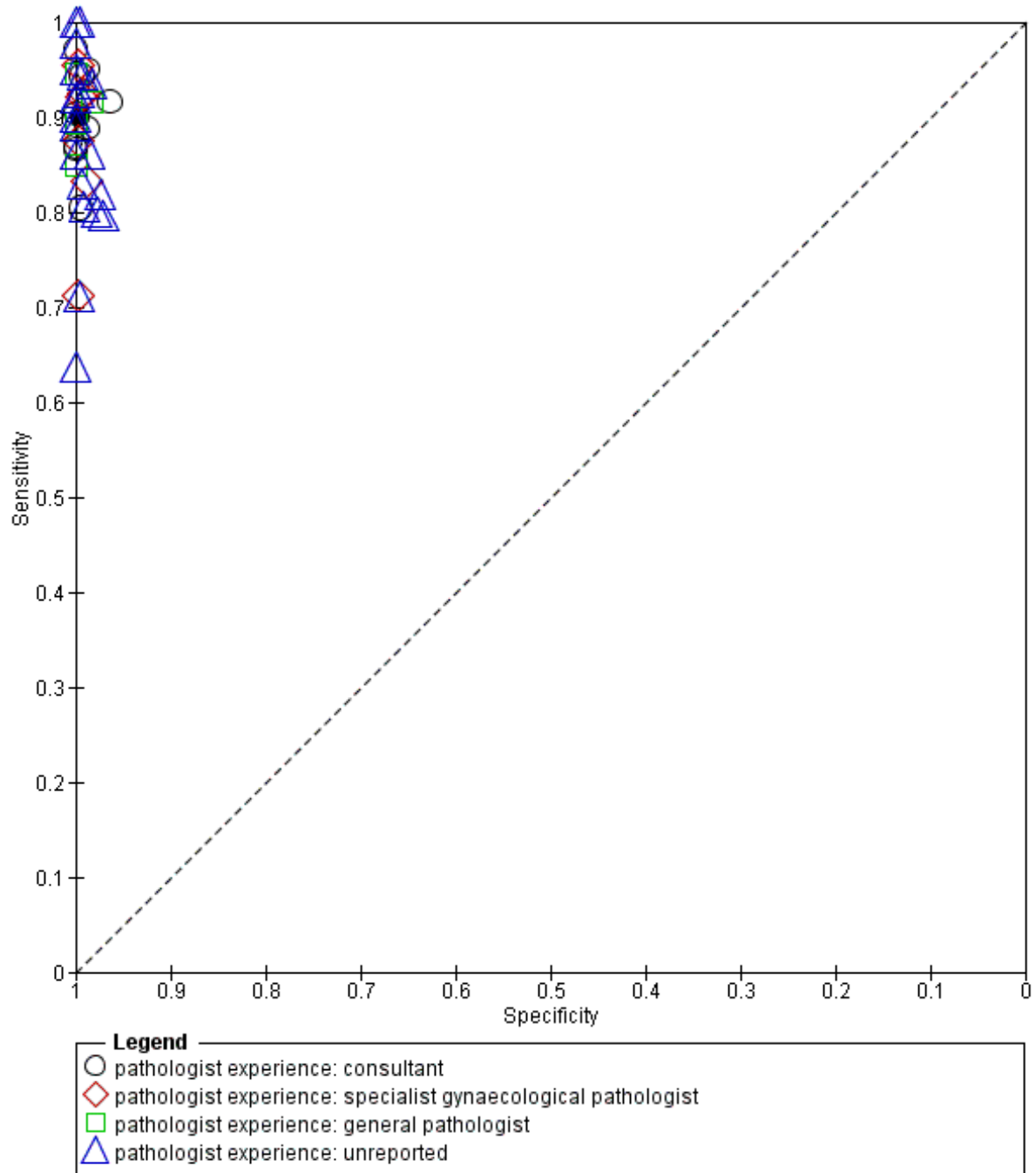


Figure 8. Summary ROC plot of I frozen section: threshold malignancy vs borderline or benign



We completed a pre-specified analysis of heterogeneity based on pathologist reader expertise, defining four reader groups that we then grouped into two categories to enable analysis to have a sufficient number of studies in each group: more specialised (consultant, and specialist gynaecological pathologist) and other (general pathologist or expertise not reported). Figure 8 shows the expertise of pathologists for each study using different symbols. Statistical analysis did not show a statistically significant difference in sensitivity and specificity based on reader expertise.

Primary objective #2: accuracy of frozen section cancer or borderline results to identify women with cancer

Sensitivity and specificity results were available from the same 38 studies using the test threshold for frozen section where we considered both cancer and borderline cases to be positive and benign cases to be negative. The average sensitivity was 96.5% (95% CI 95.5% to 97.3%; typical range 83% to 100%, with one very small

study, García 1997, reporting a sensitivity of 0%) and the average specificity was 89.5% (95% CI 86.6% to 91.9%: typical range 58% to 99%, with one study, Gorisek 2009, reporting a specificity of 29%). Results were reasonably homogeneous except for differences likely to be due to small sample sizes.

Figure 9 is a forest plot of sensitivity and specificity with 95% confidence intervals for all studies, ordered by the number of disease negative cases (DN = benign) with the studies reporting the largest numbers of benign cases at the top. In addition, we show the percentage of cancer, borderline and benign in each study to give insight into the representativeness of the study. The percentage of borderline cases is likely to influence the specificity results, as many of these cases are not found to be malignant by the reference test of paraffin section. This is well demonstrated by Gorisek 2009, with only 2% of benign cases, where specificity is reduced due to the high proportion of borderline cases in the study population.

Figure 9. Forest plot: frozen section threshold malignant or borderline vs benign

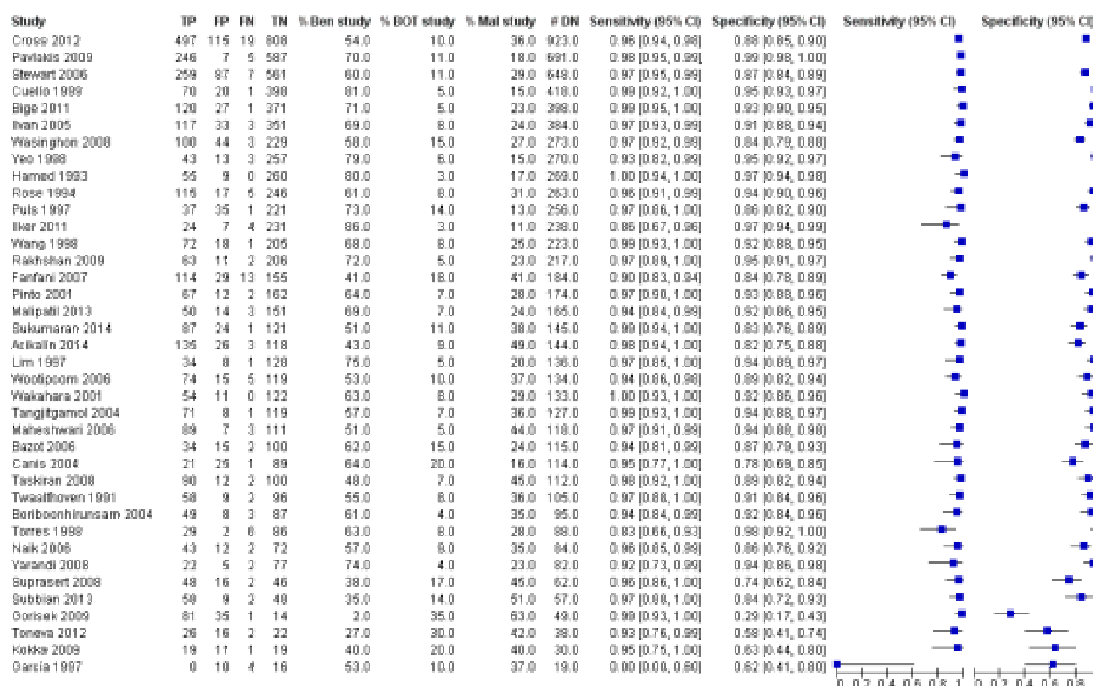
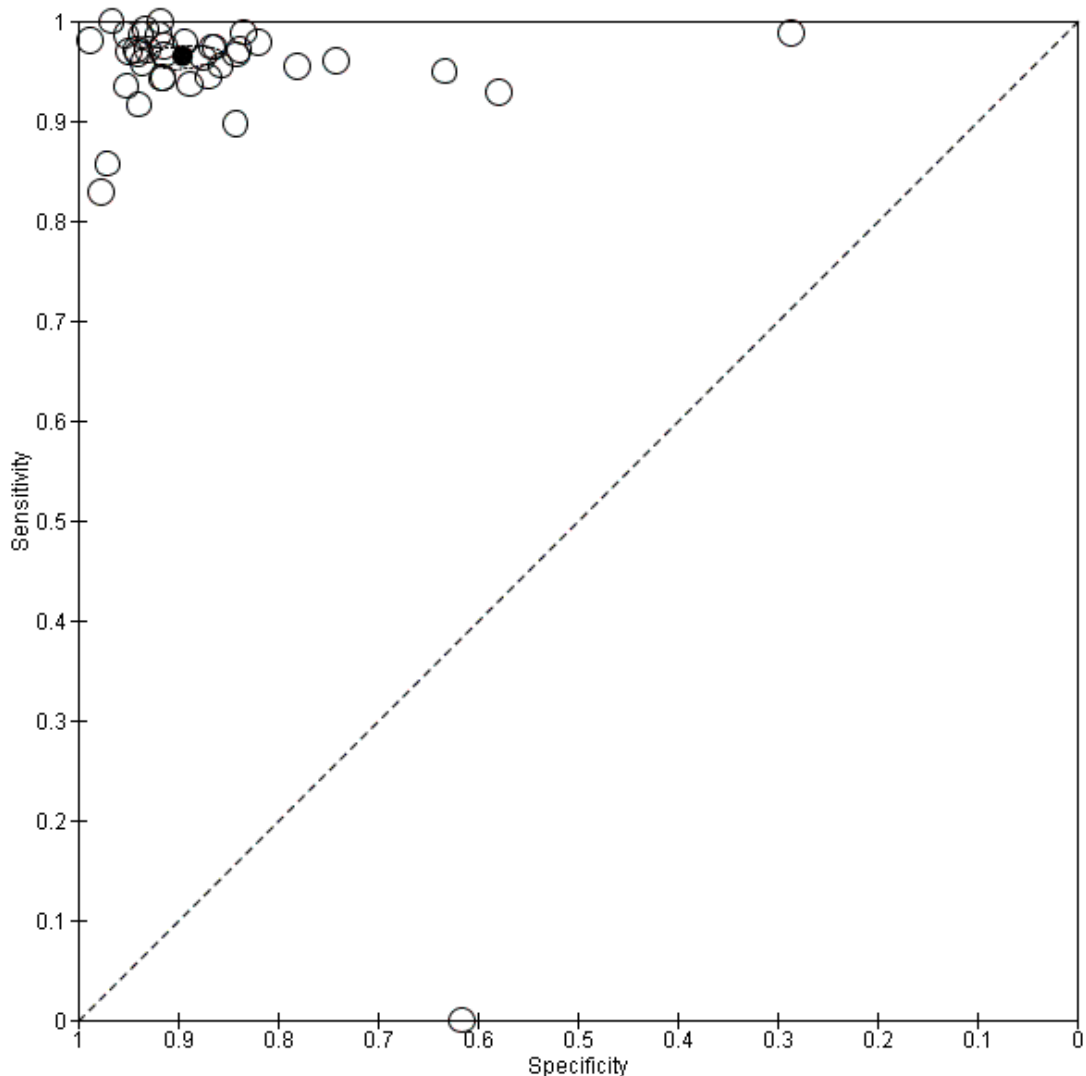


Figure 10 shows the results from all studies in a ROC plot.

Figure 10. Summary ROC plot of 2 frozen section: threshold malignancy or borderline vs benign



Both figures show that sensitivity is reasonably homogeneous, but as expected, specificity is more variable in studies with a relatively high percentage of borderline cases and a low number of disease negative (i.e. benign) cases (García 1997; Gorisek 2009; Kokka 2009; Toneva 2012). We have used bivariate meta-analysis to obtain estimates for both average sensitivity and average specificity, as there are a reasonable number of studies. The average sensitivity was 96.5% (95% CI 95.5% to 97.3%; typical range 83% to 100%, with one very small study, García 1997, reporting a sensitivity of 0%), and the average specificity was 89.5% (95% CI 86.6% to 91.9%: typical range 58 to 99, with one study, Gorisek 2009, reporting a specificity of 29%).

We attempted a pre-specified analysis of heterogeneity based on reader expertise, but models did not converge.

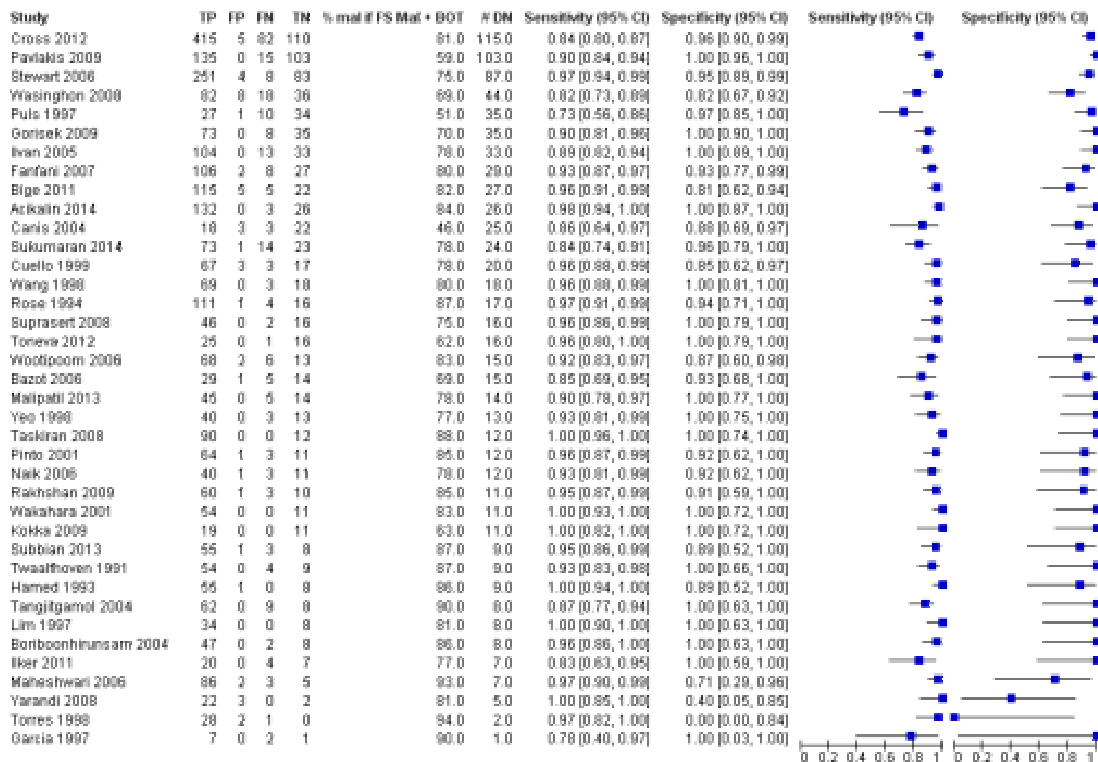
Secondary objective #2: accuracy of final diagnosis of malignancy in women with a frozen section result of either borderline or cancer

Sensitivity and specificity results were available from the same 38 studies, including the subset of 3953 participants with a frozen section result of either borderline or invasive cancer, based on the accuracy of referral for cancer management, that is, surgical staging

in invasive cancer.

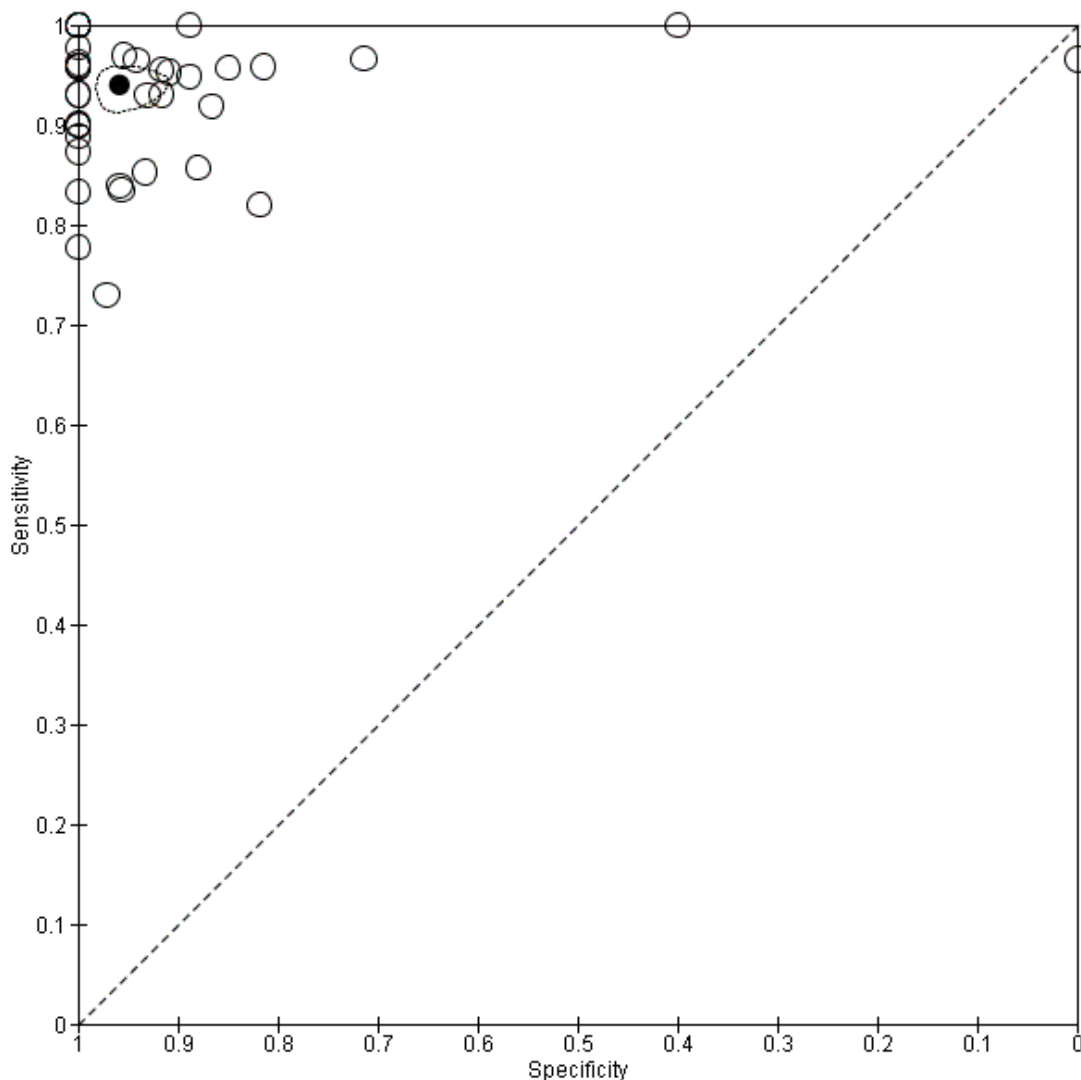
Figure 11 is a forest plot of sensitivity and specificity with 95% confidence intervals for all studies, ordered by the number of borderline cases in each study, with the studies reporting the highest number of borderline cases shown at the top.

Figure 11. Forest plot: frozen section malignant or borderline only. Threshold malignant



Studies with small numbers of disease negative cases (borderline cases) at the bottom of the plot have more variation in estimates of specificity, most likely due to small numbers in a study, likely overriding any other potential sources of bias in these studies. We include the percentage of malignant cancer in these patients with frozen section results of either cancer or borderline to aid understanding of how studies may compare to other centres. The percentage of borderline cases is likely to influence the specificity results, as many of these cases are not malignant according to the reference test of paraffin section. Figure 12 shows the results from all studies in a ROC plot.

Figure 12. Figure 8 (Analysis 3)

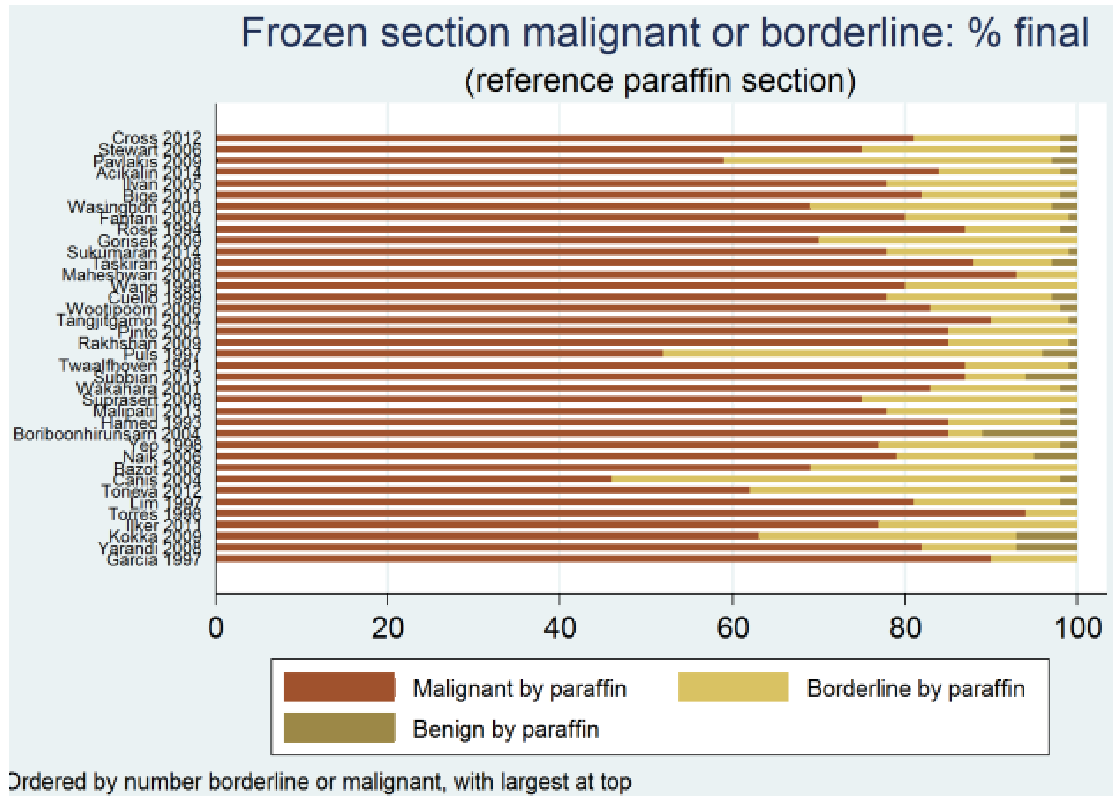


Both figures show that sensitivity and specificity are more heterogeneous than other analyses. Results from studies with low numbers of borderline cases are particularly heterogeneous due to small sample sizes of two, five and seven borderline cases (in Torres 1998, Maheshwari 2006 and Yarandi 2008, respectively). We used bivariate meta-analysis to obtain estimates both for average sensitivity and average specificity, as there were a reasonable number of studies. The average sensitivity was 94.0% (95% CI 92.0% to 95.5%; range 73% to 100%), and the average specificity was 95.8% (95% CI 92.4% to 97.8%; typical range 81% to

100%, with three outlier studies, Torres 1998, Yarandi 2008 and Maheshwari 2006, showing specificities of 0% , 40% and 71%, respectively).

Figure 13 presents the reference standard result for all studies for frozen section diagnoses of either cancer or borderline. Across all studies, an average of 81% of results were malignant by the reference standard (median 81%, interquartile range (IQR) 78% to 84%), 17% were borderline (IQR 14% to 21%) and 2% were benign (IQR 2% to 5%).

Figure 13. Frozen section result malignant or borderline: final diagnosis by reference standard



Additional analyses

Figure 14, Figure 15 and Figure 16 provide a breakdown of frozen section results by postsurgical reference standard for benign, cancer and borderline results, respectively, to provide additional insight on the correspondence between test results. This provides further information to help understand how frozen section results were updated following postsurgical paraffin section confirmation in our included studies.

Figure 14. Frozen section result benign: final diagnosis by reference standard

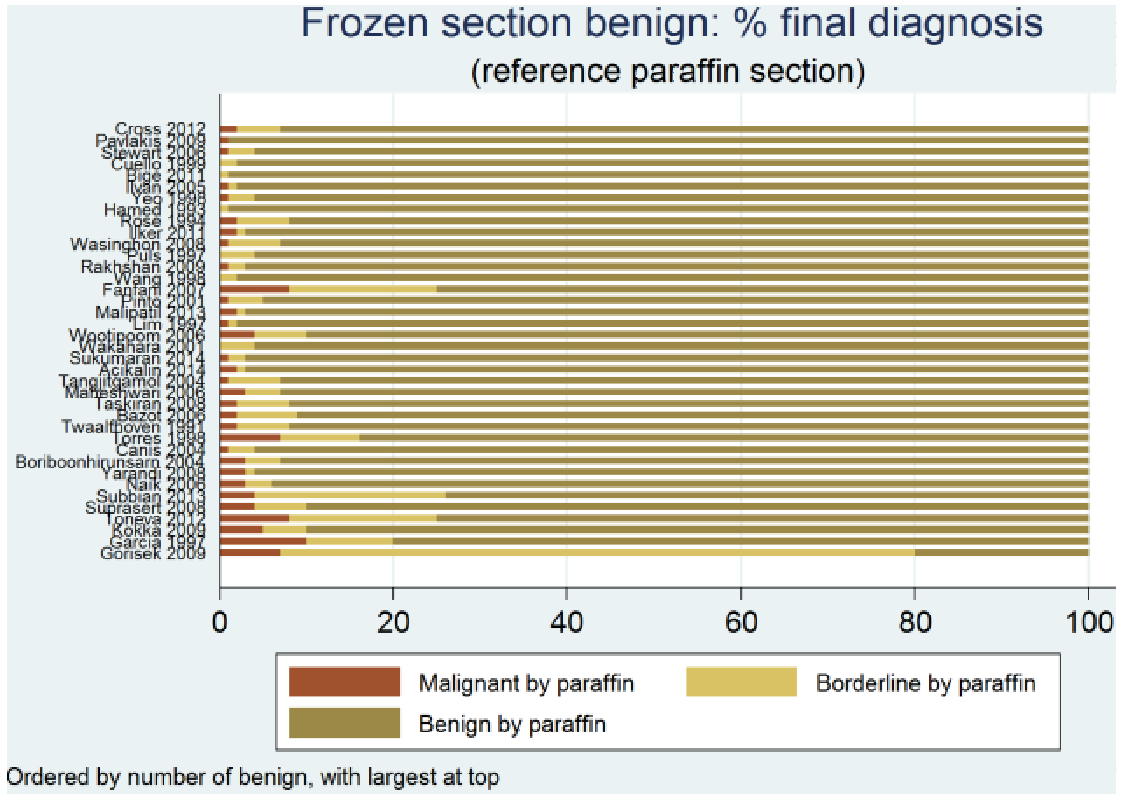


Figure 15. Frozen section result malignant: final diagnosis by reference standard

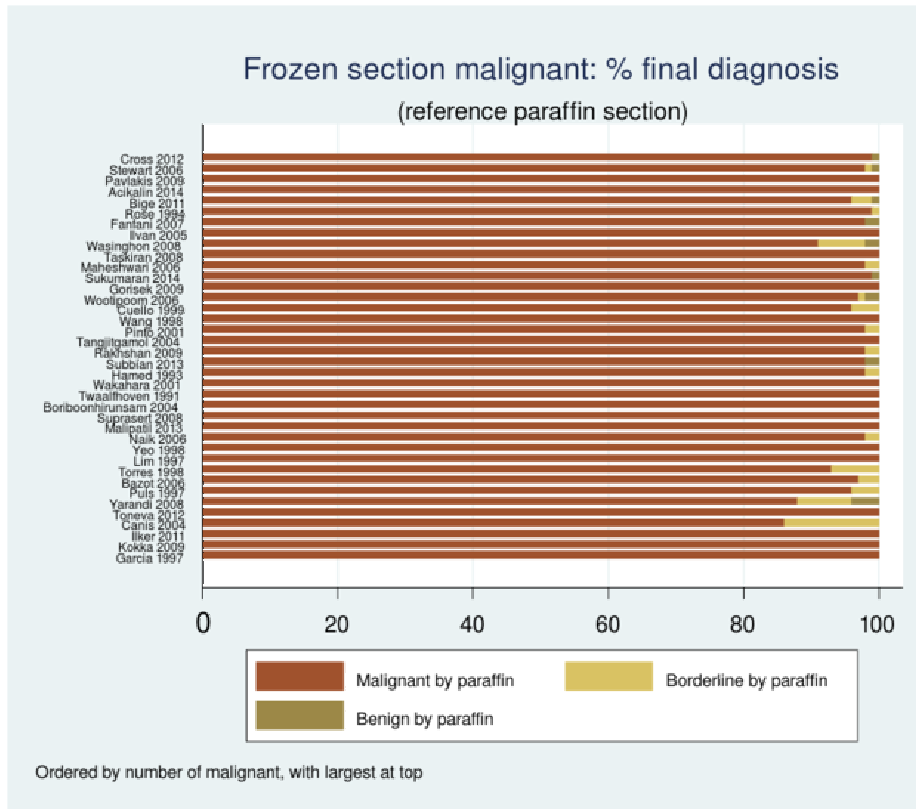
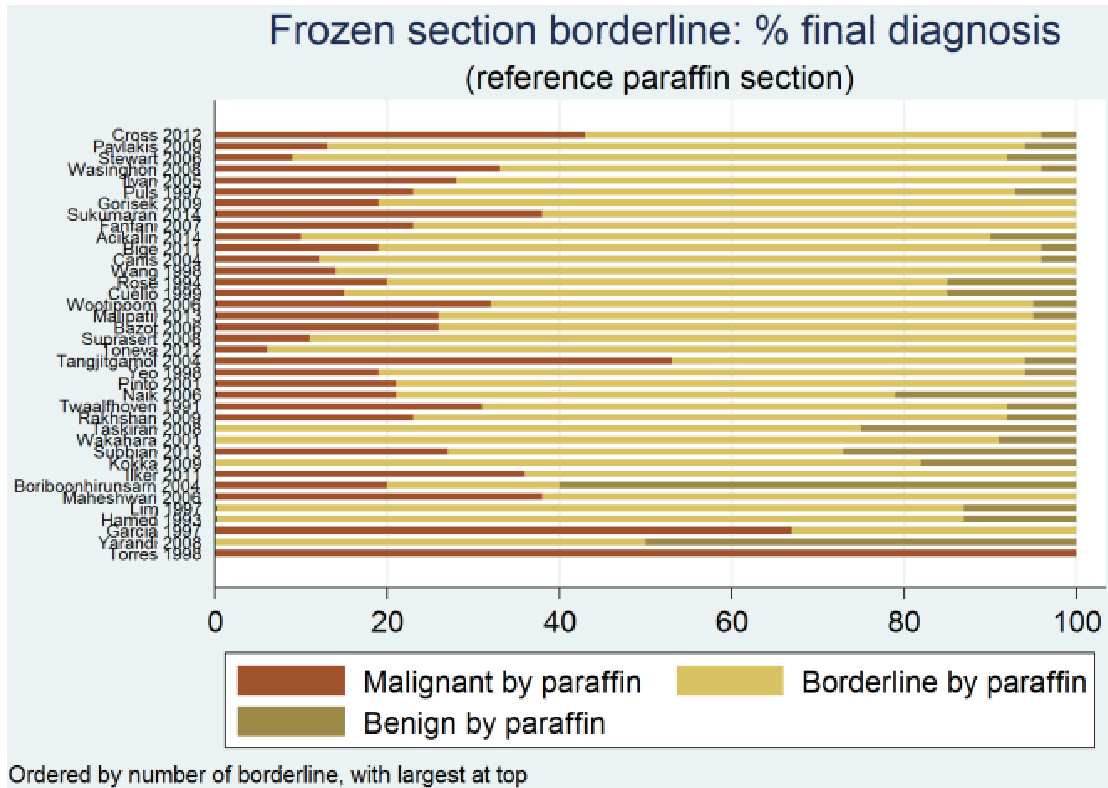


Figure 16. Frozen section result borderline: final diagnosis by reference standard



On average, 94% (IQR 92% to 96%) of benign diagnoses from frozen section were found to be benign on paraffin section.

On average, 99% (IQR 98% to 100%) of cancer results from frozen section were found to be cancerous on paraffin section.

However, for borderline results from frozen section, on average only 73% (IQR 63% to 78%) remained borderline, but 21% (IQR 14% to 26%) were upgraded to cancer, and 6% (IQR 2 to 8%) were downgraded to benign.

Summary of findings

Review question: to establish the accuracy and other diagnostic parameters (sensitivity and specificity) of intraoperative frozen section analysis in the histopathological diagnosis of ovarian cancer, in comparison to paraffin section reporting

Patients/population: women presenting to a secondary or tertiary care setting with a pelvic mass suspicious of ovarian cancer, in whom frozen section analysis was employed prior to paraffin section analysis

Role: intraoperative diagnosis of ovarian mass to guide surgery

Index tests: intraoperative frozen section histopathological analysis

Reference standards: paraffin section histopathological analysis

Studies: retrospective cohort; no prospective cohort studies identified

Clinical setting: any clinical setting. In this review, all included studies took place in university hospitals or tertiary centres

| Frozen section test | Effect (96% CI) | No of participants, DP (studies) | Test result | Number of results per 1000 participants tested ^d (95% CI) | | |
|---|--|----------------------------------|-----------------|--|------------------|------------------|
| | | | | Prevalence of malignancy in all patients (where frozen section is used) | | |
| | | | | Prevalence 23% | Prevalence 29% | Prevalence 36% |
| Primary objective #1: Frozen section: malignant versus borderline/benign All patients (8% borderline) | Sensitivity 90.0 (87.6 to 92.0) Specificity 99.5 (99.2 to 99.7) | 11,181, 3200 (38) | True positives | 207 (201 to 212) | 261 (254 to 267) | 324 (315 to 331) |
| | | | False negatives | 23 (18 to 29) | 29 (23 to 36) | 36 (29 to 45) |
| | | | False positives | 4 (2 to 6) | 4 (2 to 6) | 3 (2 to 5) |
| | | | True negatives | 766 (764 to 768) | 706 (704 to 708) | 637 (635 to 638) |
| Primary objective #2: Frozen section: malignant/borderline versus benign All patients (8% borderline) | Sensitivity 96.5 (95.5 to 97.3) Specificity 89.5 (86.6 to 91.9) | 11,181, 3200 (38) | True positives | 222 (220 to 224) | 280 (277 to 282) | 347 (344 to 350) |
| | | | False negatives | 8 (6 to 10) | 10 (8 to 13) | 13 (10 to 16) |
| | | | False positives | 81 (62 to 103) | 75 (58 to 95) | 67 (52 to 86) |
| | | | True negatives | 689 (667 to 708) | 635 (615 to 652) | 573 (554 to 588) |
| | | | | Prevalence of malignancy in subgroup of patients with frozen section malignant/borderline ^b | | |
| | | | | Prevalence 75% | Prevalence 80% | Prevalence 86% |

| | | | | | | |
|--|---------------------------------|-----------------|-----------------|------------------|-------------------|------------------|
| Secondary objective #2: Frozen section: malignant versus borderline/benign in subgroup of patients with frozen section result malignant/borderline | Sensitivity 94.0 (92.0 to 95.5) | 3953, 3084 (38) | True positives | 705 (690 to 716) | 752 (736 to 764) | 808 (791 to 821) |
| | Specificity 95.8 (92.4 to 97.8) | | False negatives | 45 (34 to 60) | 48 (36 to 64) | 52 (39 to 69) |
| | | | False positives | 11 (6 to 19) | 8 (4 to 15) | 6 (3 to 11) |
| | | | True negatives | 240 (231 to 245) | 192 (185 to 196) | 134 (129 to 137) |

Attributes of tests contributing to benefits and risks

| | |
|----------------|--|
| Frozen section | <p>Intraoperative frozen section enables surgeons to perform surgical staging appropriately in patients with frozen section diagnosis of ovarian malignancy, thereby reducing the need for a second surgical procedure. If surgical staging confirms there is no extra ovarian disease, chemotherapy may not be required</p> <p>Other benefits of frozen section include diagnosis of tumour origin and diagnosis of metastatic disease</p> <p>Risks of frozen section relate to overstaging/overtreatment for malignancy in false positives</p> |
|----------------|--|

Overall quality of evidence/risk of bias:

Patient selection: included studies are retrospective. The majority report consecutive cases, but in several the sampling is unclear

Index test: deferred and unclear frozen section results reported and excluded from 3 x 3 tables

Reference standard: Pathologists must have been aware of frozen section results at time of performing paraffin section. Four studies reported the same pathologist interpreting both tests

Flow and timing: paraffin section takes place after frozen section so no bias in timing

Precision: Average estimates of both sensitivity and specificity have good precision

CI: confidence interval; **DP:** disease positive.

^aPrevalence of malignancy from included studies. median, lower and upper interquartile range values of 23%, 29% and 36% respectively.

^bPrevalence of malignancy from included studies from subgroup of cases where frozen section of malignant or borderline may be used to refer to cancer surgery: median, lower and upper interquartile range values of 75%, 80% and 86% respectively.

DISCUSSION

Summary of main results

We report the largest review of frozen section accuracy in ovarian masses to date, with a median prevalence of 29% cancers (IQR 23% to 36%) across the included studies. In the [Summary of findings](#) we have provided examples of prevalence of malignancy to help clinicians to interpret presented results and inform their practice. Accuracy results were relatively consistent between studies, except for studies with small numbers of cases. All studies were retrospective, with the majority reporting consecutive cases. We excluded deferred and unclear frozen section results from analysis. We expect that reference standard interpretation was not blinded to frozen section results.

In a hypothetical study of 1000 patients, of whom 290 had cancer and 80 were borderline, on average 261 women (95% CI 254 to 267) would receive a correct diagnosis of cancer based on a frozen section result of cancer, and 706 women (95% CI 704 to 708) would be correctly diagnosed without cancer. However, on average 4 women (95% CI 2 to 6) would be incorrectly diagnosed as having a cancer (false positive), and 29 women (95% CI 23 to 36) with cancer would be missed at the time of surgery (false negative).

Likewise, in a hypothetical population of 1000 women, of whom 290 had cancer and 80 were borderline, based on a frozen section result of either cancer or borderline to diagnose cancer, on average 280 women (95% CI 277 to 282) would be correctly diagnosed with a cancer and correctly receive surgical staging. Six hundred and thirty-five women (95% CI 615 to 652) would be correctly diagnosed without cancer. However, on average 75 women (95% CI 58 to 95) would be incorrectly diagnosed as having a cancer on frozen section and would be over treated with surgical staging. Ten women (95% CI 8 to 13) with cancer would be missed at the time of surgery and might require a second surgical procedure for staging.

Our additional analyses showed that if the frozen section was benign or cancerous, then the final diagnosis would remain the same in, on average, 94% and 99% of cases, respectively ([Figure 14](#); [Figure 15](#)).

In cases where the frozen section diagnosis was borderline, there is a chance that the final diagnosis would turn out to be cancer in, on average, 21% of women ([Figure 16](#)).

On investigating the factors that could lead to variability between studies, we found that there was no difference in diagnostic accuracy between levels of expertise of pathologists. In cases where there was a discordance between frozen section and paraffin section, most studies tabulated reasons for discordance that fell into one of two categories: tissue sampling error (where the sampled portions of the mass failed to give the paraffin section diagnosis); or interpretation error (where the pathologist incorrectly reported

the frozen section samples). Tissue sampling error was more commonly reported in borderline frozen section diagnoses.

Strengths and weaknesses of the review

Strengths

This review presents a meticulous analysis of existing literature and interprets the data with presentation of clinical relevance. By applying previously defined, clear criteria for eligibility, we aimed to minimise heterogeneity in included studies. We excluded studies that did not represent the population in which frozen section might be used to assess suspicious ovarian masses. We assessed methodological quality and risk of bias. Although several studies had small sample sizes, the number of studies included in the review (N = 38) and the number of patients (N = 11,181) increased the power of the meta-analyses.

Analysis of the data by varying the test positive response ('malignant'; and 'malignant and borderline') facilitates interpretation of the test data to guide surgical management. Specifically, the pooled analysis of borderline cases compared to test positive malignant cases provides valuable information to aid not only intraoperative decision-making but also perioperative counselling of patients about likely outcomes.

Weaknesses

There are three weaknesses regarding pathology reporting within this review. Firstly, although this review addresses the effect of pathologist expertise on frozen section interpretation, all included studies were conducted in university hospitals or tertiary centres, which may introduce a reporting bias within this review. Secondly, given that no studies reported pathologist blinding, we have to assume that all included studies in this review were unblinded. The extent of bias is somewhat limited given the flow and timing of the tests, in that the index test always precedes the reference standard. The implications of unblinded testing might potentially mean that a pathologist reporting the paraffin section would be more likely to agree with the frozen section, especially if it is the same pathologist reporting both. However, in clinical practice the Pathology department usually receives the result of the frozen section as part of required clinical information in submitting tissue for histological processing. Thirdly, the criteria used for diagnosis of borderline ovarian tumours varies internationally. The included studies did not report the criteria used.

Very few studies reported the use of preoperative imaging or tumour markers. It was therefore difficult to make inferences about these variables.

Applicability of findings to the review question

Frozen section is a useful tool in aiding intraoperative management of suspicious ovarian masses. This review finds that if the frozen section is benign or cancerous, the paraffin section will be concordant in 94% and 99% of cases, respectively. In these groups there is a high likelihood patients will receive the appropriate surgery based on frozen section results, as indicated by Naik 2006, thereby avoiding unnecessary staging in those with benign histology according to paraffin section and without compromising those with true stage I ovarian cancer. Lawrie 2015 demonstrated in a subgroup analysis of three trials that, at a median follow-up of 5 years, there was no apparent additional benefit to overall survival from adjuvant chemotherapy in the group that was optimally staged (Bolis 1995; ICON 1; Trimbos 2003). However, they had concerns about selective reporting of the 10-year survival data and performed an exploratory analysis of 'deaths from ovarian cancer' at 10 years. This analysis suggested that "the difference between subgroups (optimally versus suboptimally staged) in deaths from ovarian cancer was not statistically significant (test for subgroup differences: Chi^2 test = 2.75, degrees of freedom (df) = 1, $P = 0.10$; I^2 statistic = 63.6%)".

The prevalence of cancer in a referred population is particularly relevant for borderline ovarian tumours given the degree of discordance with paraffin section diagnosis. In this review, with an average prevalence of cancer of 29%, the chance of a patient with a borderline tumour being appropriately treated with surgical staging is 21%. This would in turn mean that, should all women with borderline frozen section undergo a surgical staging procedure including hysterectomy, pelvic +/- paraaortic lymphadenectomy and omentectomy, 79% of them would be over treated. This confers unnecessary risk of morbidity, which includes lymphoedema, lymphocyst formation, visceral and neurovascular injury. The benefit, however, is a reduction in morbidity associated with a second surgical procedure should low-risk ovarian cancer be diagnosed on paraffin section.

In their interpretation of this review, readers should evaluate the presented results by comparing the prevalence of test positive (cancer) in their population to examples provided in the Summary of findings. The clinicopathological considerations to be taken into account when using frozen section include the following: women with high-risk disease will require adjuvant platinum-based chemotherapy; optimal staging in true stage I disease confers prognostic benefit; staging will detect stage III disease in a quarter of women who will require dual agent chemotherapy; and women need to be well counselled regarding the risks of over treatment and under treatment if physicians rely on frozen section results.

In addition, although outside the scope of this review, the clinical benefits of frozen section analysis include the ability to diagnose metastatic disease and, in some cases, site of origin. This may lead to better exploration of other organs at laparotomy for site of primary tumour and avoid unnecessary surgical staging in non-ovarian malignancy.

AUTHORS' CONCLUSIONS

Implications for practice

Frozen section testing of ovarian masses can be used intraoperatively in gynaecological cancer centres for investigation of women with ovarian masses suspected to be early-stage malignancy. In practice, use of frozen section depends on a number of factors.

- The clinical suspicion of cancer. This is usually reflected by the prevalence of cancer within a referred population, that is, a tertiary centre will report higher rates of cancer than a secondary centre. Women undergoing the index test can be counselled about the advantages and disadvantages of undergoing surgical staging if the frozen section result is borderline.

- The expertise of the gynaecologist to perform a surgical staging procedure should the frozen section result prove to be cancer. The value of the index test depends on the ability of the surgeon to appropriately manage the case.

- The ability of the pathologist to interpret frozen sections and for histopathology departments to provide the frozen section service.

Implications for research

The largest discordance is within the reporting of frozen section borderline tumours. The authors would encourage future publications to include all reported frozen section results and their histological subtypes so that further subgroup analyses on the borderline population can be performed to minimise reporting bias and heterogeneity analyses can be performed on histological subtypes. Investigation into factors leading to discordance within centres and standardisation of criteria for reporting borderline tumours may help further improve accuracy.

Further research is also warranted, perhaps in the form of a randomised clinical trial, to evaluate the oncological and surgical outcomes from surgical staging in cases of apparent stage I ovarian cancer. This would help establish whether there is a place for frozen section analysis in gynaecological cancer centres and further inform clinical practice by addressing not only survival but also morbidity associated with under- and over-staging in the borderline population.

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* Indicates the major publication for the study

CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Açikalin 2014

| Study characteristics | | | |
|--|---|--------------|------------------------|
| Patient sampling | Design: Retrospective Setting: Turkey Accrual dates: July 2006 - January 2013 No participants: 282 No assessed: 282 Inclusion criteria: Re-analysis of charts of 282 women with an ovarian neoplasm (42.8% of all gynaecologic FSs) with intraoperative FS reports. Paraffin section diagnoses with non-tumoural ovarian lesions (massive ovarian edema, hemorrhagic necrosis, benign cysts, infections) were excluded. Included previous histological diagnosis: unclear | | |
| Patient characteristics and setting | Excluded non-tumoral ovarian masses | | |
| Index tests | "All fresh gross specimen were examined by a resident and a pathologist or particularly gynecopathologist, in terms of localization, size, colour, content, heterogeneity, infiltration pattern of the tumour and condition of the ovarian capsule. One to four sections depending on the size and heterogeneity of the tumour were sampled in a cryostat and sections were stained by hematoxylin-eosin. Slides were evaluated and reported to the surgeon by the pathologist. Final PS diagnosis reported by an experienced gynecopathologist was accepted as accurate diagnose." | | |
| Target condition and reference standard(s) | Malignancy; paraffin section | | |
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | Re-analysis of charts of 282 women with an ovarian neoplasm (42.8% of all gynaecologic FSs) with intraoperative FS reports diagnosed between July 2006 and January 2013. Paraffin section diagnoses with non-tumoural ovarian lesions (massive ovarian edema, hemorrhagic necrosis, benign cysts, infections) were excluded | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Unclear | | |

Açikalin 2014 (Continued)

| | | | |
|--|---------|--|--|
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| Unclear | | | |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Unclear | | |
| Unclear | | | |
| DOMAIN 3: Reference Standard | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | |
| Unclear | | | |
| DOMAIN 4: Flow and Timing | | | |
| Did all patients receive a reference standard? | Yes | | |
| Did all patients receive the same reference standard? | Yes | | |
| Were un-interpretable/intermediate test results reported? | Yes | | |

Açikalin 2014 (Continued)

| | | | |
|--|-----|--|--|
| Were withdrawals from the study explained? | Yes | | |
| | | | |

Bazot 2006

Study characteristics

| | |
|--|---|
| Patient sampling | Design: Retrospective Setting: France Accrual dates: Jan 1999 - Dec 2003 No participants: 136 No assessed: 151* Inclusion criteria: Complex / suspicious adnexal masses referred for pre-op MRI Included previous histological diagnosis: unclear |
| Patient characteristics and setting | Complex / suspicious adnexal masses referred for pre-op MRI. France |
| Index tests | FS prepared from vegetations in cyst walls or solid areas selected by macroscopic exam |
| Target condition and reference standard(s) | Malignant or borderline on paraffin section. |
| Flow and timing | FS before PS. |
| Comparative | |
| Notes | Compares accuracy of pre-op MRI and FS. * 136 women, 32 with bilateral masses (15 bilateral and 17 unilateral FS) - total 151 FS analyses included |

Methodological quality

| Item | Authors' judgement | Risk of bias | Applicability concerns |
|--|--------------------|--------------|------------------------|
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Yes | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |

Bazot 2006 (Continued)

| | | | | |
|--|---------|--|--|--|
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | | |
| Unclear | | | | |
| DOMAIN 2: Index Test All tests | | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | | |
| Were the index tests Interpreted by consultant or specialist gynec onc pathologist? | Unclear | | | |
| Unclear | | | | |
| DOMAIN 3: Reference Standard | | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | | |
| Low | | | | |
| DOMAIN 4: Flow and Timing | | | | |
| Did all patients receive a reference standard? | Yes | | | |
| Did all patients receive the same reference standard? | Yes | | | |
| Were un-interpretable/intermediate test results reported? | No | | | |
| Were withdrawals from the study explained? | No | | | |

| Study characteristics | | | |
|--|---|--------------|------------------------|
| Patient sampling | Design: Retrospective Setting: Turkey Accrual dates: Jan 2002 - Dec 2010 No participants: 578 No assessed: 519 Inclusion criteria: Indications for FS - radiologically or macroscopically benign appearing ovarian masses with high CA125, history of malignancy other than ovary and fertility preserving surgery for young cases. 59 exclusions - 14 definitive diagnosis not obtained by FS, 23 no ovarian tissue identified in masses, 22 metastases to ovaries Included previous histological diagnosis: unclear | | |
| Patient characteristics and setting | University hospital in Turkey | | |
| Index tests | 2-5 sections esp from solid areas, examined by at least 2 consultant pathologists (gynae 83.2%, non-gynae 16.8%) | | |
| Target condition and reference standard(s) | Malignant or borderline disease; paraffin section | | |
| Flow and timing | PS performed after FS. 59 patients excluded: in 14, definitive diagnosis could not be obtained at FS; in 23, no ovarian tissue identified in masses; in 22, metastases to ovary | | |
| Comparative | | | |
| Notes | FS reported by at least 2 consultant pathologists (gynae pathologists 83.2%, non-gynae 16.8%). Unclear if same pathologists reported PS or if FS results masked. Possible interpretation bias | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Unclear | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |

Bige 2011 (Continued)

| | | |
|--|---------|-------------|
| | | High |
| DOMAIN 2: Index Test All tests | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Yes | |
| | | Low |
| DOMAIN 3: Reference Standard | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | |
| | | Low |
| DOMAIN 4: Flow and Timing | | |
| Did all patients receive a reference standard? | Yes | |
| Did all patients receive the same reference standard? | Yes | |
| Were un-interpretable/intermediate test results reported? | Yes | |
| Were withdrawals from the study explained? | Yes | |
| | | |

Boriboonhirunsarn 2004

| Study characteristics | | | |
|--|---|---------------------|-------------------------------|
| Patient sampling | Design: Prospective diagnostic test accuracy Setting: Thailand Accrual dates: July 2001 to March 2002 No participants: 147 No assessed: 147 Inclusion criteria: Included women with ovarian tumours for surgery. Excluded if give prior treatment for cancer (radiotherapy or chemotherapy). Included previous histological diagnosis: NR | | |
| Patient characteristics and setting | Included women with ovarian tumours for surgery. Excluded if give prior treatment for cancer (radiotherapy or chemotherapy).Thailand | | |
| Index tests | Number of slides and area to be sectioned were determined by one experienced pathologist, who also examined and interpreted all slides, 'Slides were interpreted without knowledge of the results of those prepared by the other technique' | | |
| Target condition and reference standard(s) | Malignant or borderline on PS. | | |
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | Surgical extent not reported. | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Yes | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Yes | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| | | | Unclear |
| DOMAIN 2: Index Test All tests | | | |

Boriboonthirunsarn 2004 (Continued)

| | | | | |
|--|---------|--|--|------------|
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Yes | | | |
| | | | | Low |
| DOMAIN 3: Reference Standard | | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | | |
| | | | | Low |
| DOMAIN 4: Flow and Timing | | | | |
| Did all patients receive a reference standard? | Yes | | | |
| Did all patients receive the same reference standard? | Yes | | | |
| Were un-interpretable/intermediate test results reported? | Yes | | | |
| Were withdrawals from the study explained? | Unclear | | | |
| | | | | |

Canis 2004

| | |
|------------------------------|---|
| Study characteristics | |
| Patient sampling | Design: Retrospective Setting: Canada Accrual dates: 5 years No participants: 141 No assessed: 141 Inclusion criteria: All women undergoing laparoscopy in order to treat ovarian or paraovarian tu- |

Canis 2004 (Continued)

| | |
|--|--|
| | mours. Excluded obvious malignancy and benign masses (uterine, peritoneal cysts, hydrosalpinges) . Included previous histological diagnosis: unclear |
| Patient characteristics and setting | Macroscopically suspicious ovarian tumours. Large tertiary care centre |
| Index tests | Details not reported |
| Target condition and reference standard(s) | Malignant or borderline disease |
| Flow and timing | FS before PS. |
| Comparative | |
| Notes | Laparoscopic management with staging procedure to FS (borderline or malignant) by either laparoscopy or laparotomy |

Methodological quality

| Item | Authors' judgement | Risk of bias | Applicability concerns |
|--|--------------------|--------------|------------------------|
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Yes | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Yes | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Yes | | |
| | | | Low |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Unclear | | |

| | | |
|--|---------|------------|
| | | Low |
| DOMAIN 3: Reference Standard | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | |
| | | Low |
| DOMAIN 4: Flow and Timing | | |
| Did all patients receive a reference standard? | Yes | |
| Did all patients receive the same reference standard? | Yes | |
| Were un-interpretable/intermediate test results reported? | Yes | |
| Were withdrawals from the study explained? | No | |

Cross 2012

| | |
|-------------------------------------|---|
| Study characteristics | |
| Patient sampling | Design: Prospective diagnostic test accuracy Setting: UK Accrual dates: Jan 2000 to Dec 2010 No participants: 1445 No assessed: 1439 Inclusion criteria: Women with possible ovarian malignancy were included. Women with obvious disseminated malignancy (FIGO stage III/IV) were excluded. Included previous histological diagnosis: yes, 6 cases |
| Patient characteristics and setting | Women with possible ovarian malignancy were included. Women with obvious disseminated malignancy (FIGO stage III/IV) were excluded. Tertiary centre, UK |

| | | | |
|--|--|---------------------|-------------------------------|
| Index tests | Two pieces of tissue taken for FS staining and reporting by consultant pathologist. Report was then phoned through to surgeon who used the info to determine the extent of surgery | | |
| Target condition and reference standard(s) | Malignant or borderline on PS. | | |
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | Women with borderline and malignant diagnosis underwent surgical staging | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Yes | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Yes | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Yes | | |
| | | | Low |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Yes | | |
| | | | Low |
| DOMAIN 3: Reference Standard | | | |

Cross 2012 (Continued)

| | | | |
|--|---------|--|--|
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | |
| Low | | | |
| DOMAIN 4: Flow and Timing | | | |
| Did all patients receive a reference standard? | Yes | | |
| Did all patients receive the same reference standard? | Yes | | |
| Were un-interpretable/intermediate test results reported? | Yes | | |
| Were withdrawals from the study explained? | Yes | | |

Cuello 1999

Study characteristics

| | |
|--|--|
| Patient sampling | Design: Retrospective Setting: Chile Accrual dates: Jan 1988-Oct 1998 No participants: 842 No assessed: 489 Inclusion criteria: Ovarian masses. 2x2 data available only for epithelial ovarian masses. Included previous histological diagnosis: unclear |
| Patient characteristics and setting | Epithelial ovarian masses. Chile. |
| Index tests | Sections were taken every 3-4cm, measuring 2-3mm. Frozen sections measuring 5microns were taken |
| Target condition and reference standard(s) | Malignant or borderline on PS. 2 x 2 data only available for epithelial ovarian masses |

Cuello 1999 (Continued)

| | | | |
|--|---------------------------|---------------------|-------------------------------|
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | - | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Unclear | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| | | | Low |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Unclear | | |
| | | | Low |
| DOMAIN 3: Reference Standard | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | |

| DOMAIN 1: Patient Selection | | | |
|--|---------|--|------------|
| Was a consecutive or random sample of patients enrolled? | Yes | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| | | | Low |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Yes | | |
| | | | Low |
| DOMAIN 3: Reference Standard | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | |
| | | | Low |
| DOMAIN 4: Flow and Timing | | | |
| Did all patients receive a reference standard? | Yes | | |
| Did all patients receive the same reference standard? | Yes | | |

Fanfani 2007 (Continued)

| | | | |
|---|-----|--|--|
| Were un-interpretable/intermediate test results reported? | Yes | | |
| Were withdrawals from the study explained? | Yes | | |
| | | | |

García 1997

| Study characteristics | | | |
|--|--|---------------------|-------------------------------|
| Patient sampling | Design: Retrospective Setting: Valencia Accrual dates: Jan 1994 to Oct 1995 No participants: 30 No assessed: 30 Inclusion criteria: Women with adnexal mass undergoing FS. None other given. Included previous histological diagnosis: unclear | | |
| Patient characteristics and setting | Uncertain setting in Valencia | | |
| Index tests | FS between 6-7 sections per specimen. | | |
| Target condition and reference standard(s) | Malignant or borderline on PS. | | |
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | - | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Unclear | | |
| Was a case-control design avoided? | Unclear | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |

García 1997 (Continued)

| | | | | |
|--|---------|--|--|--|
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | | |
| Unclear | | | | |
| DOMAIN 2: Index Test All tests | | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Unclear | | | |
| Unclear | | | | |
| DOMAIN 3: Reference Standard | | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | | |
| Low | | | | |
| DOMAIN 4: Flow and Timing | | | | |
| Did all patients receive a reference standard? | Yes | | | |
| Did all patients receive the same reference standard? | Yes | | | |
| Were un-interpretable/intermediate test results reported? | No | | | |
| Were withdrawals from the study explained? | No | | | |

Gorisek 2009

| Study characteristics | |
|--|---|
| Patient sampling | Design: Retrospective Setting: Slovenia Accrual dates: 1 January 1993 - 31 December 2001 No participants: 131 No assessed: 131 Inclusion criteria: Women treated for benign, borderline and malignant ovarian tumours Included previous histological diagnosis: unclear |
| Patient characteristics and setting | Women treated for benign, borderline and malignant ovarian tumour. Slovenia |
| Index tests | FS "After tumour removal, the fresh surgical specimen was immediately taken to the Department of Pathologic Morphology at the Maribor Teaching Hospital (now the University Clinical Centre Maribor). A pathologist prepared specimens from representative regions, froze them in a cryostat and cut slices with a microtome. The slices were mounted on a glass slide, stained with haematoxylin and eosin, and were then ready for microscopic evaluation." |
| Target condition and reference standard(s) | Malignant or borderline on PS. |
| Flow and timing | FS before PS. |
| Comparative | |
| Notes | - |

Methodological quality

| Item | Authors' judgement | Risk of bias | Applicability concerns |
|--|---------------------------|---------------------|-------------------------------|
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Unclear | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| | | | Unclear |

Gorisek 2009 (Continued)

| DOMAIN 2: Index Test All tests | | | |
|--|---------|--|----------------|
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Unclear | | |
| | | | Unclear |
| DOMAIN 3: Reference Standard | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | |
| | | | Low |
| DOMAIN 4: Flow and Timing | | | |
| Did all patients receive a reference standard? | Yes | | |
| Did all patients receive the same reference standard? | Yes | | |
| Were un-interpretable/intermediate test results reported? | No | | |
| Were withdrawals from the study explained? | Unclear | | |
| | | | |

Hamed 1993

| Study characteristics | |
|------------------------------|---|
| Patient sampling | Design: Retrospective Setting: Chile Accrual dates: Jan 1987-Oct 1992 No participants: 324 |

Hamed 1993 (Continued)

| | | | |
|--|--|---------------------|-------------------------------|
| | No assessed: 324 Inclusion criteria: Women with peristent pelvic masses aged 9-81years Included previous histological diagnosis: unclear | | |
| Patient characteristics and setting | Women with peristent pelvic masses aged 9-81years. Chile. | | |
| Index tests | FS. 5micron sections of tissue 3-4cmx2-3mm from mass | | |
| Target condition and reference standard(s) | Malignant or borderline on PS. | | |
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | Large age group, not certain representtaive of suspicious masses | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Unclear | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | No | | |
| | | | High |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Unclear | | |

Hamed 1993 (Continued)

| | | | | |
|--|---------|--|--|---------|
| | | | | Unclear |
| DOMAIN 3: Reference Standard | | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | | |
| | | | | Unclear |
| DOMAIN 4: Flow and Timing | | | | |
| Did all patients receive a reference standard? | Yes | | | |
| Did all patients receive the same reference standard? | Yes | | | |
| Were un-interpretable/intermediate test results reported? | Yes | | | |
| Were withdrawals from the study explained? | Yes | | | |
| | | | | |

Ilker 2011

| | |
|-------------------------------------|--|
| Study characteristics | |
| Patient sampling | Design: Retrospective Setting: Turkey Accrual dates: Jan 2002 - Dec 2008 No participants: 278 No assessed: 266 Inclusion criteria: Patients undergoing surgery for ovarian mass where FS performed Included previous histological diagnosis: unclear |
| Patient characteristics and setting | Ovarian mass. Turkey. |
| Index tests | FS 2-5 (5µm) slides from suspicious areas, reported by "expert" pathologist |

| | | | |
|--|--------------------------------|---------------------|-------------------------------|
| Target condition and reference standard(s) | Malignant or borderline on PS. | | |
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | 12 deferred cases. | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Unclear | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | No | | |
| | | | High |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Yes | | |
| | | | Low |
| DOMAIN 3: Reference Standard | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | |

Ilker 2011 (Continued)

| | | | |
|--|---------|--|--|
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | |
| Low | | | |
| DOMAIN 4: Flow and Timing | | | |
| Did all patients receive a reference standard? | Yes | | |
| Did all patients receive the same reference standard? | Yes | | |
| Were un-interpretable/intermediate test results reported? | Yes | | |
| Were withdrawals from the study explained? | Yes | | |

Ilvan 2005

| | |
|--|--|
| Study characteristics | |
| Patient sampling | Design: Retrospective Setting: Turkey Accrual dates: Jan 1995 to Dec 2003 No participants: 617 No assessed: 617 Inclusion criteria: Ovarian masses sent for FS. No exclusions given. Included grossly benign tumours (22 endometriotic, 3 follicles, 12 mature teratomas, 9 benign serous cystadenomas). Included previous histological diagnosis: unclear |
| Patient characteristics and setting | Pelvic masses in women referred to a tertiary centre. Grossly benign specimens submitted in 46 cases |
| Index tests | FS. Gross examination, touch imprints, sections (between 1 and 4) of ovary. 2 pathologists in gynaecology |
| Target condition and reference standard(s) | Malignant and borderline disease. PS. 2 pathologists experienced in gynaecological pathology interpreted FS. Pathologists also employed touch imprint technique |
| Flow and timing | Some FS diagnosis (7.5%) made on gross inspection only FS before PS |
| Comparative | |

| | | | |
|--|--|---------------------|-------------------------------|
| Notes | 46 grossly benign masses including 22 endometriotic cyst, 3 follicles, 12 mature teratomas, 9 benign serous cystadenomas | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Unclear | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| | | | High |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Yes | | |
| | | | High |
| DOMAIN 3: Reference Standard | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | |
| | | | Low |

Ilvan 2005 (Continued)

| DOMAIN 4: Flow and Timing | | | |
|---|---------|--|--|
| Did all patients receive a reference standard? | Yes | | |
| Did all patients receive the same reference standard? | Yes | | |
| Were un-interpretable/intermediate test results reported? | No | | |
| Were withdrawals from the study explained? | Unclear | | |
| | | | |

Kokka 2009

| Study characteristics | | | |
|--|---|--------------|------------------------|
| Patient sampling | Design: Retrospective Setting: UK Accrual dates: Oct 2006 - May 2008 No participants: 61 No assessed: 50 Inclusion criteria:- Included previous histological diagnosis: unclear | | |
| Patient characteristics and setting | Unclear inclusion criteria. | | |
| Index tests | FS. No details given. | | |
| Target condition and reference standard(s) | Malignant or borderline on PS. | | |
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | FS was considered in 71 patients; ten cases were excluded because of valid reasons; in 11 of 31 benign tumours FS was not requested by the surgeon | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |

| DOMAIN 1: Patient Selection | | | |
|--|---------|--|------------|
| Was a consecutive or random sample of patients enrolled? | Yes | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| | | | Low |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Unclear | | |
| | | | Low |
| DOMAIN 3: Reference Standard | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | |
| | | | Low |
| DOMAIN 4: Flow and Timing | | | |
| Did all patients receive a reference standard? | Yes | | |
| Did all patients receive the same reference standard? | Yes | | |

Kokka 2009 (Continued)

| | | | |
|---|-----|--|--|
| Were un-interpretable/intermediate test results reported? | Yes | | |
| Were withdrawals from the study explained? | Yes | | |
| | | | |

Lim 1997

| Study characteristics | | | |
|--|--|--------------|------------------------|
| Patient sampling | Design: Retrospective Setting: Singapore Accrual dates: Jan 1988 to Dec 1994 No participants: 173 No assessed: 171 Inclusion criteria: Women with ovarian tumours and laparotomy and FS. No other inclusion details. Included previous histological diagnosis: unclear | | |
| Patient characteristics and setting | University hospital, Singapore. Pelvic masses. Uncertain if previous diagnosis cancer. Majority of cases benign. Authors describe liberal use of frozen section in their hospital, even if mass thought to be benign | | |
| Index tests | FS. If discordant, pathologist reassessed if sampling or interpretational error | | |
| Target condition and reference standard(s) | Malignant or borderline on PS | | |
| Flow and timing | FS before PS. Ultrasound features correlated to final outcome | | |
| Comparative | | | |
| Notes | 2 cases FS diagnosis deferred and not included in analysis. Both were interpretational errors. When accuracy of FS was reassessed for cases where FS was clinically indicated, accuracy was 95.5% (105 out of 110) for benign, borderline or malignant | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Unclear | | |

Lim 1997 (Continued)

| | | | |
|--|---------|--|-------------|
| Was a case-control design avoided? | Unclear | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | No | | |
| | | | High |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Yes | | |
| | | | Low |
| DOMAIN 3: Reference Standard | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | |
| | | | Low |
| DOMAIN 4: Flow and Timing | | | |
| Did all patients receive a reference standard? | Yes | | |
| Did all patients receive the same reference standard? | Yes | | |
| Were un-interpretable/intermediate test results reported? | Yes | | |

Lim 1997 (Continued)

| | | | |
|--|-----|--|--|
| Were withdrawals from the study explained? | Yes | | |
| | | | |

Maheshwari 2006

| Study characteristics | | | |
|--|---|--------------|------------------------|
| Patient sampling | Design: Retrospective Setting: India Accrual dates: 1997-2001 No participants: 241 No assessed: 210 Inclusion criteria: Excluded non-ovarian FS and deferred FS. Included previous histological diagnosis: unclear, included cases with previous cancer at another site | | |
| Patient characteristics and setting | Unclear inclusion criteria. India. | | |
| Index tests | FS. 1-4 sections at 7-8micrometer intervals. | | |
| Target condition and reference standard(s) | Malignant or borderline on PS. | | |
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | Included 'clinically benign tumours with raised CA125' | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Unclear | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |

| | | | | |
|--|---------|--|--|--|
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | | |
| Unclear | | | | |
| DOMAIN 2: Index Test All tests | | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | | |
| Were the index tests Interpreted by consultant or specialist gyn-onc pathologist? | Unclear | | | |
| Unclear | | | | |
| DOMAIN 3: Reference Standard | | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | | |
| Low | | | | |
| DOMAIN 4: Flow and Timing | | | | |
| Did all patients receive a reference standard? | Yes | | | |
| Did all patients receive the same reference standard? | Yes | | | |
| Were un-interpretable/intermediate test results reported? | Yes | | | |
| Were withdrawals from the study explained? | Yes | | | |

Malipatil 2013

| Study characteristics | | | |
|--|--|--------------|------------------------|
| Patient sampling | Design: Retrospective Setting: India Accrual dates: 1999 - 2008 No participants: 223 No assessed: 218 Inclusion criteria: 5 exclusions - FS diagnosis deferred due to extensive necrosis / haemorrhage Included previous histological diagnosis: unclear | | |
| Patient characteristics and setting | Unclear inclusion criteria. India. | | |
| Index tests | FS. At least 2 general surgical pathologists reporting FS. Mean number of FS 2 (1-5) and PS 7 (1-33) | | |
| Target condition and reference standard(s) | Malignant or borderline on PS. | | |
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | 377 referrals for diagnosis; intraoperative diagnosis sought in 233 (apparently 223?) cases; diagnosis deferred in five cases due to extensive areas of haemorrhage and necrosis and was excluded from further analysis; 218 cases analysed | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Unclear | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| | | | Unclear |
| DOMAIN 2: Index Test All tests | | | |

Malipatil 2013 (Continued)

| | | | | |
|--|---------|--|--|------------|
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Yes | | | |
| | | | | Low |
| DOMAIN 3: Reference Standard | | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | | |
| | | | | Low |
| DOMAIN 4: Flow and Timing | | | | |
| Did all patients receive a reference standard? | Yes | | | |
| Did all patients receive the same reference standard? | Yes | | | |
| Were un-interpretable/intermediate test results reported? | Yes | | | |
| Were withdrawals from the study explained? | Yes | | | |
| | | | | |

Naik 2006

| | |
|------------------------------|---|
| Study characteristics | |
| Patient sampling | Design: Retrospective Setting: UK Accrual dates: July 2002 to June 2003 No participants: 130 No assessed: 130 inc. 1 deferred Inclusion criteria: Suspicious pelvic masses |

Naik 2006 (Continued)

| | | | |
|--|--|---------------------|-------------------------------|
| | Included previous histological diagnosis: yes | | |
| Patient characteristics and setting | Suspicious pelvic masses. Tertiary centre, UK. | | |
| Index tests | FS. No details given. | | |
| Target condition and reference standard(s) | Malignant or borderline on PS. | | |
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | 1 deferred was benign. | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Yes | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Yes | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Yes | | |
| | | | Low |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Unclear | | |
| | | | Low |

| DOMAIN 3: Reference Standard | | | |
|--|-----|--|------------|
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Yes | | |
| | | | Low |
| DOMAIN 4: Flow and Timing | | | |
| Did all patients receive a reference standard? | Yes | | |
| Did all patients receive the same reference standard? | Yes | | |
| Were un-interpretable/intermediate test results reported? | Yes | | |
| Were withdrawals from the study explained? | Yes | | |
| | | | |

Pavlakis 2009

| Study characteristics | |
|--|---|
| Patient sampling | Design: Retrospective Setting: Greece Accrual dates: Jan 2000 to Oct 2006 No participants: 932 No assessed: 932 Inclusion criteria: Ovarian or related masses submitted for FS. No other details given. Included previous histological diagnosis: unclear |
| Patient characteristics and setting | FS received from various centres. Uncertain if suspicious of malignancy when submitted. Ovarian and related masses sent. 594 of 932 specimens benign on PS |
| Index tests | FS. 1 to3 sections per ovarian or related mass. Examined by gynaecological pathologist |
| Target condition and reference standard(s) | Malignant or borderline on PS. |

| | | | |
|--|---|---------------------|-------------------------------|
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | Uncertain if only suspicious ovarian masses included. Risk of selection bias. Also, only epithelial tumours reported in 3 x 3 table | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Unclear | | |
| Was a case-control design avoided? | Unclear | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| | | | High |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Yes | | |
| | | | Unclear |
| DOMAIN 3: Reference Standard | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge | Unclear | | |

Pavlakis 2009 (Continued)

| | | | |
|---|-----|--|---------|
| of the results of the index tests? | | | |
| | | | Unclear |
| DOMAIN 4: Flow and Timing | | | |
| Did all patients receive a reference standard? | Yes | | |
| Did all patients receive the same reference standard? | Yes | | |
| Were un-interpretable/intermediate test results reported? | No | | |
| Were withdrawals from the study explained? | No | | |
| | | | |

Pinto 2001

| | |
|--|--|
| Study characteristics | |
| Patient sampling | Design: Retrospective Setting: Brazil Accrual dates: jan 1994 to April 1999 No participants: 243 No assessed: 243 Inclusion criteria: Ovarian tumours. No other details given. Included previous histological diagnosis: unclear |
| Patient characteristics and setting | Pathology laboratory in Brazil. Uncertain if tertiary referral centre for gynaecological malignancies |
| Index tests | FS. 1 to 3 sections per specimen reported by general pathologist. All slides reviewed by specialist gynaecological pathologist |
| Target condition and reference standard(s) | Malignant or borderline on PS. Histological type noted. |
| Flow and timing | FS before PS. |
| Comparative | |
| Notes | All slides (FS and PS) checked by a specialist gynaecological pathologist |
| Methodological quality | |

| Item | Authors' judgement | Risk of bias | Applicability concerns |
|--|--------------------|--------------|------------------------|
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Unclear | | |
| Was a case-control design avoided? | Unclear | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| | | | Unclear |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Yes | | |
| | | | Unclear |
| DOMAIN 3: Reference Standard | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | |
| | | | Low |
| DOMAIN 4: Flow and Timing | | | |
| Did all patients receive a reference standard? | Yes | | |

Pinto 2001 (Continued)

| | | | |
|---|-----|--|--|
| Did all patients receive the same reference standard? | Yes | | |
| Were un-interpretable/intermediate test results reported? | No | | |
| Were withdrawals from the study explained? | No | | |
| | | | |

Puls 1997

| Study characteristics | | | |
|--|--|--------------|------------------------|
| Patient sampling | Design: Retrospective Setting: South Carolina Accrual dates: 12 years No participants: 294 No assessed: 294 Inclusion criteria: Included all women with both FS and PS analysis of ovarian tumour (serous or mucinous). Included previous histological diagnosis: nr | | |
| Patient characteristics and setting | Ovarian tumour. Preoperative diagnosis unclear. South Carolina | | |
| Index tests | FS. One section per centimetre of wall. In most cases, same pathologist reported FS and PS. Pool of 8 pathologists reporting, expertise not given | | |
| Target condition and reference standard(s) | Malignant and borderline serous and mucinous ovarian tumours on PS | | |
| Flow and timing | Only PS which had conclusive FS were included FS before PS. | | |
| Comparative | | | |
| Notes | Of 632 operations, 294 were selected for having both FS and PS. Inconclusive FS is likely to have been excluded | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |

Puls 1997 (Continued)

| | | | | |
|--|---------|--|--|------------|
| Was a consecutive or random sample of patients enrolled? | Yes | | | |
| Was a case-control design avoided? | Yes | | | |
| Did the study avoid inappropriate exclusions? | Unclear | | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | | |
| | | | | Low |
| DOMAIN 2: Index Test All tests | | | | |
| Were the index tests interpreted without knowledge of the reference standard? | No | | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Unclear | | | |
| | | | | Low |
| DOMAIN 3: Reference Standard | | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | No | | | |
| | | | | Low |
| DOMAIN 4: Flow and Timing | | | | |
| Did all patients receive a reference standard? | Yes | | | |
| Did all patients receive the same reference standard? | Yes | | | |

Puls 1997 (Continued)

| | | | |
|---|-----|--|--|
| Were un-interpretable/intermediate test results reported? | No | | |
| Were withdrawals from the study explained? | Yes | | |
| | | | |

Rakhshan 2009

| Study characteristics | | | |
|--|--|--------------|------------------------|
| Patient sampling | Design: Retrospective Setting: Iran Accrual dates: March 1994 - May 2008 No participants: 282 No assessed: 282 Inclusion criteria: Ovarian masses submitted for frozen section Included previous histological diagnosis: unclear | | |
| Patient characteristics and setting | Ovarian masses. Iran. | | |
| Index tests | FS. 1-5 (5µm) sections interpreted by 1 of 5 attending general pathologists. All FS specimens reviewed by specialist gynae pathologists for study | | |
| Target condition and reference standard(s) | Malignant or borderline on PS. | | |
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | No deferred cases. Large proportion of cases benign. | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Unclear | | |
| Was a case-control design avoided? | Yes | | |

| | | | |
|--|---------|--|--|
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| Unclear | | | |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Yes | | |
| Low | | | |
| DOMAIN 3: Reference Standard | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | |
| Low | | | |
| DOMAIN 4: Flow and Timing | | | |
| Did all patients receive a reference standard? | Yes | | |
| Did all patients receive the same reference standard? | Yes | | |
| Were un-interpretable/intermediate test results reported? | Yes | | |
| Were withdrawals from the study explained? | Yes | | |

Rose 1994

| Study characteristics | | | |
|--|---|---------------------|-------------------------------|
| Patient sampling | Design: Retrospective Setting: USA Accrual dates: June 1983-1993 No participants: 383 No assessed: 383 Inclusion criteria: None given Included previous histological diagnosis: unclear | | |
| Patient characteristics and setting | Unclear inclusion criteria. USA. | | |
| Index tests | FS. 0-4 sections at 2-3mm intervals | | |
| Target condition and reference standard(s) | Malignant or borderline on PS. | | |
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | - | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Unclear | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| | | | Unclear |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the refer- | Yes | | |

Rose 1994 (Continued)

| | | | |
|--|---------|--|----------------|
| ence standard? | | | |
| Were the index tests Interpreted by consultant or specialist gynec onc pathologist? | Unclear | | |
| | | | Unclear |
| DOMAIN 3: Reference Standard | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | |
| | | | Low |
| DOMAIN 4: Flow and Timing | | | |
| Did all patients receive a reference standard? | Yes | | |
| Did all patients receive the same reference standard? | Yes | | |
| Were un-interpretable/intermediate test results reported? | No | | |
| Were withdrawals from the study explained? | No | | |

Stewart 2006

Study characteristics

| | |
|------------------|---|
| Patient sampling | Design: retrospective Setting: tertiary hospital, Australia Accrual dates: Jan 1999 to Dec 2003 No participants: 914 No assessed: 914 Inclusion criteria: FS of omentum and lymph node included. Included previous histological diagnosis: NR |
|------------------|---|

Stewart 2006 (Continued)

| | | | |
|--|--|---------------------|-------------------------------|
| Patient characteristics and setting | Tertiary centre pelvic masses. FS of omentum and lymph node included | | |
| Index tests | FS of omentum and lymph node included. Pathologists in general surgery and gyn oncology employed | | |
| Target condition and reference standard(s) | Malignant or borderline on PS. | | |
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | Good study with useful 2 x 2 table | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Yes | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Yes | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| | | | Low |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests interpreted by consultant or specialist gynec pathologist? | Yes | | |
| | | | Low |

Stewart 2006 (Continued)

| DOMAIN 3: Reference Standard | | | |
|--|---------|--|------------|
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | |
| | | | Low |
| DOMAIN 4: Flow and Timing | | | |
| Did all patients receive a reference standard? | Yes | | |
| Did all patients receive the same reference standard? | Yes | | |
| Were un-interpretable/intermediate test results reported? | No | | |
| Were withdrawals from the study explained? | Yes | | |
| | | | |

Subbian 2013

| Study characteristics | |
|-------------------------------------|--|
| Patient sampling | Design: Retrospective Setting: India Accrual dates: March 2004 - January 2006 No participants: 135 No assessed: 117 Inclusion criteria: Retrospective analysis of reports of frozen section and paraffin block diagnoses of patients undergoing surgery as primary line of therapy for suspected ovarian neoplasms Included previous histological diagnosis: unclear |
| Patient characteristics and setting | Suspected ovarian neoplasms, India. |
| Index tests | FS. "All the frozen section diagnoses were made by a team of expert onco- pathologists at the institute. Before sectioning, gross examination of the tumor was carried out and frozen section samples were taken from solid or suspicious areas. The number of bits sampled varied from one to |

Subbian 2013 (Continued)

| | | | |
|--|--|---------------------|-------------------------------|
| | three (average of two). The frozen section and the permanent section reports of each patient were compared. The frozen section results were divided into the following groups: Deferred, benign, borderline and malignant. Reports mentioned as 'suggestive of', 'suspicious of' or 'compatible with' were included in the diagnoses mentioned." | | |
| Target condition and reference standard(s) | Malignant or borderline on PS. | | |
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | Retrospective selection based on having had FS and PS . Deferred cases: 8/135 (5.9%). Ten patients diagnosed with non- neoplastic conditions were also excluded | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Unclear | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| | | | Low |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Yes | | |
| | | | Low |
| DOMAIN 3: Reference Standard | | | |

Subbian 2013 (Continued)

| | | | |
|--|---------|--|--|
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | |
| Low | | | |
| DOMAIN 4: Flow and Timing | | | |
| Did all patients receive a reference standard? | Yes | | |
| Did all patients receive the same reference standard? | Yes | | |
| Were un-interpretable/intermediate test results reported? | No | | |
| Were withdrawals from the study explained? | Yes | | |

Sukumaran 2014

Study characteristics

| | |
|--|---|
| Patient sampling | Design: Retrospective Setting: India Accrual dates: 2009-2012 No participants: 237 No assessed: 233 (4 deferred) Inclusion criteria: Excluded torsion Included previous histological diagnosis: unclear |
| Patient characteristics and setting | Torted pelvoic masses excluded. India. |
| Index tests | FS. 2-5 sections each 4-5microns |
| Target condition and reference standard(s) | Malignant or borderline on PS. |
| Flow and timing | FS before PS. |

| | | | |
|--|---------------------------|---------------------|-------------------------------|
| Comparative | | | |
| Notes | 4 deferred on FS. | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Unclear | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| | | | Low |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Unclear | | |
| | | | Low |
| DOMAIN 3: Reference Standard | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | |
| | | | Low |

| DOMAIN 4: Flow and Timing | |
|---|-----|
| Did all patients receive a reference standard? | Yes |
| Did all patients receive the same reference standard? | Yes |
| Were un-interpretable/intermediate test results reported? | Yes |
| Were withdrawals from the study explained? | Yes |
| | |

Suprasert 2008

| Study characteristics | |
|--|---|
| Patient sampling | Design: Retrospective review Setting: Thailand Accrual dates: Jan 2001 to Dec 2005 No participants: 127 No assessed: 112 Inclusion criteria: Women with pelvic masses. Excluded infarcted masses (4) or deferred (18) frozen section analysis. Included previous histological diagnosis: no |
| Patient characteristics and setting | Pelvic masses. Thailand. |
| Index tests | FS. Number of frozen sections determined by attending Pathologist. Deferred FS when suspicion of borderline or malignant considered not definitely diagnostic |
| Target condition and reference standard(s) | Malignant or borderline on PS. |
| Flow and timing | Specimens submitted to FS at surgeon's discretion - no clear protocol. FS before PS |
| Comparative | |
| Notes | Retrospective review. 15 excluded due to infarction and deferred FS No stage given. Variable sections were performed by Pathologist |
| Methodological quality | |

| Item | Authors' judgement | Risk of bias | Applicability concerns |
|--|--------------------|--------------|------------------------|
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Yes | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | No | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| | | | Unclear |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Unclear | | |
| | | | Unclear |
| DOMAIN 3: Reference Standard | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | |
| | | | Low |
| DOMAIN 4: Flow and Timing | | | |
| Did all patients receive a reference standard? | Yes | | |

Suprasert 2008 (Continued)

| | | | |
|---|-----|--|--|
| Did all patients receive the same reference standard? | Yes | | |
| Were un-interpretable/intermediate test results reported? | Yes | | |
| Were withdrawals from the study explained? | Yes | | |
| | | | |

Tangjitgamol 2004

| Study characteristics | | | |
|--|---|--------------|------------------------|
| Patient sampling | Design: Retrospective Setting: Thailand Accrual dates: Jan 1992 to Jan 2002 No participants: 212 No assessed: 212 inc. 13 deferred Inclusion criteria: Intact ovarian masses submitted. Included previous histological diagnosis: unclear | | |
| Patient characteristics and setting | Intact ovarioan masses. Thailand. | | |
| Index tests | FS. No details given. | | |
| Target condition and reference standard(s) | Malignant or borderline on PS. | | |
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | Included 13 deferred (7 benign, 76 borderline, 121 malignant). High percentage malignant PS. Risk selection bias | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Unclear | | |

Tangjitgamol 2004 (Continued)

| | | | |
|--|---------|--|----------------|
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| | | | Unclear |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Unclear | | |
| | | | Unclear |
| DOMAIN 3: Reference Standard | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | |
| | | | Low |
| DOMAIN 4: Flow and Timing | | | |
| Did all patients receive a reference standard? | Yes | | |
| Did all patients receive the same reference standard? | Yes | | |
| Were un-interpretable/intermediate test results reported? | Yes | | |

Tangjitgamol 2004 (Continued)

| | | | |
|--|-----|--|--|
| Were withdrawals from the study explained? | Yes | | |
| | | | |

Taskiran 2008

Study characteristics

| | |
|--|--|
| Patient sampling | Design: Retrospective Setting: Turkey Accrual dates: 1997 - 2006 No participants: 207 No assessed: 207 inc 3 deferred Inclusion criteria: Consecutive exploratory laparotomies for pelvic mass Included previous histological diagnosis: unclear |
| Patient characteristics and setting | Pelvic masses. Turkey. |
| Index tests | FS (5 μ m) from most suspected areas of mass, solid / papillary areas of tumour wall. No info on number of slides or reporting pathologists |
| Target condition and reference standard(s) | Malignant or borderline on PS. |
| Flow and timing | FS before PS. |
| Comparative | |
| Notes | 3 deferred cases. |

Methodological quality

| Item | Authors' judgement | Risk of bias | Applicability concerns |
|--|--------------------|--------------|------------------------|
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Yes | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |

Taskiran 2008 (Continued)

| | | | | |
|--|---------|--|--|----------------|
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | | |
| | | | | Low |
| DOMAIN 2: Index Test All tests | | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | | |
| Were the index tests Interpreted by consultant or specialist gynec onc pathologist? | Unclear | | | |
| | | | | Unclear |
| DOMAIN 3: Reference Standard | | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | | |
| | | | | Low |
| DOMAIN 4: Flow and Timing | | | | |
| Did all patients receive a reference standard? | Yes | | | |
| Did all patients receive the same reference standard? | Yes | | | |
| Were un-interpretable/intermediate test results reported? | Yes | | | |
| Were withdrawals from the study explained? | Yes | | | |
| | | | | |

Toneva 2012

| Study characteristics | | | |
|--|--|---------------------|-------------------------------|
| Patient sampling | Design: Retrospective Setting: UK Accrual dates: Oct 2005 - Sept 2008 No participants: 67 No assessed: 66 Inclusion criteria: FS in 67 cases (29.7%), 1 excluded due to missing data Included previous histological diagnosis: unclear | | |
| Patient characteristics and setting | Unclear inclusion criteria. UK. | | |
| Index tests | FS. 3-5 sections. | | |
| Target condition and reference standard(s) | Malignant or borderline on PS. | | |
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | - | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Yes | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Yes | | |
| | | | Low |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |

| | | | | |
|--|---------|--|--|------------|
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Yes | | | |
| | | | | Low |
| DOMAIN 3: Reference Standard | | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | | |
| | | | | Low |
| DOMAIN 4: Flow and Timing | | | | |
| Did all patients receive a reference standard? | Yes | | | |
| Did all patients receive the same reference standard? | Yes | | | |
| Were un-interpretable/intermediate test results reported? | Yes | | | |
| Were withdrawals from the study explained? | Yes | | | |

Torres 1998

| | |
|------------------------------|---|
| Study characteristics | |
| Patient sampling | Design: Retrospective Setting: Colombia Accrual dates: Jan1994-Dec 1997 No participants: 199 No assessed: 199- (73 excluded, 3 deferred diagnoses)=123 Inclusion criteria: Mass Included previous histological diagnosis: unclear |

Torres 1998 (Continued)

| | | | |
|--|---------------------------|---------------------|-------------------------------|
| Patient characteristics and setting | Pelvic masses. Columbia. | | |
| Index tests | FS. No details given. | | |
| Target condition and reference standard(s) | Malignnat on PS. | | |
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | - | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Unclear | | |
| Was a case-control design avoided? | Unclear | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| | | | Unclear |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Unclear | | |
| | | | Unclear |
| DOMAIN 3: Reference Standard | | | |

Torres 1998 (Continued)

| | | | |
|--|---------|--|--|
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | |
| Low | | | |
| DOMAIN 4: Flow and Timing | | | |
| Did all patients receive a reference standard? | Yes | | |
| Did all patients receive the same reference standard? | Yes | | |
| Were un-interpretable/intermediate test results reported? | Yes | | |
| Were withdrawals from the study explained? | Yes | | |

Twaalfhoven 1991

Study characteristics

| | |
|--|--|
| Patient sampling | Design: Retrospective Setting: Netherlands Accrual dates: Jan 1984-Jan 1990 No participants: 176 No assessed: 176 inc. 11 deferred Inclusion criteria: Included 27 ovarian biopsies and 149 ovaries. Included previous histological diagnosis: unclear |
| Patient characteristics and setting | Ovarian biopsies and ovaries submitted for FS. Netherlands. |
| Index tests | FS. No details given. |
| Target condition and reference standard(s) | Malignant or borderline on PS. |
| Flow and timing | FS before PS. |

Twaalfhoven 1991 (Continued)

| | | | |
|--|--|---------------------|-------------------------------|
| Comparative | | | |
| Notes | 11 Deferred (1 benign, 4 borderline, 6 malignant). | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Unclear | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| | | | Unclear |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Unclear | | |
| | | | Unclear |
| DOMAIN 3: Reference Standard | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | |
| | | | Low |

Twaalfhoven 1991 (Continued)

| DOMAIN 4: Flow and Timing | | | |
|---|---------|--|--|
| Did all patients receive a reference standard? | Yes | | |
| Did all patients receive the same reference standard? | Yes | | |
| Were un-interpretable/intermediate test results reported? | Yes | | |
| Were withdrawals from the study explained? | Unclear | | |
| | | | |

Wakahara 2001

| Study characteristics | | | |
|--|--|--------------|------------------------|
| Patient sampling | Design: Prospective Setting: Japan Accrual dates: 1994 - 1999 No participants: 292 No assessed: 187 Inclusion criteria: None given Included previous histological diagnosis: unclear | | |
| Patient characteristics and setting | Unclear inclusion criteria. Japan. | | |
| Index tests | FS. Single pathologist reported all FS and PS | | |
| Target condition and reference standard(s) | Malignant or borderline on PS. | | |
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | Principal aim to assess performance of US / tumour markers in differentiating malignant from benign adnexal masses | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |

Wakahara 2001 (Continued)

| DOMAIN 1: Patient Selection | | | |
|--|---------|--|----------------|
| Was a consecutive or random sample of patients enrolled? | Unclear | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| | | | Unclear |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Unclear | | |
| | | | Low |
| DOMAIN 3: Reference Standard | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | |
| | | | Low |
| DOMAIN 4: Flow and Timing | | | |
| Did all patients receive a reference standard? | Yes | | |
| Did all patients receive the same reference standard? | Yes | | |

Wakahara 2001 (Continued)

| | | | |
|---|---------|--|--|
| Were un-interpretable/intermediate test results reported? | No | | |
| Were withdrawals from the study explained? | Unclear | | |
| | | | |

Wang 1998

| Study characteristics | | | |
|--|---|--------------|------------------------|
| Patient sampling | Design: Retrospective Setting: Taiwan Accrual dates: Jan 1991 to June 1996 No participants: 299 No assessed: 299 Inclusion criteria: Unclear if all were pelvic masses. Also reported FS in lymphatic, uterine and other tissue samples. Included previous histological diagnosis: nr | | |
| Patient characteristics and setting | Unclear if hospital setting. Pelvic masses. Taiwan. | | |
| Index tests | FS. Between 1 and several FS were performed each case. Different grades of pathologist were employed. Most were general surgical pathologists (not gynaecological oncology pathologists.) | | |
| Target condition and reference standard(s) | Benign, borderline and malignant ovarian diagnosis on PS. | | |
| Flow and timing | FS before PS. Interpretation of ovaries varied in number of sections submitted for FS. Same pathologist reported both the FS and PS in most cases. Interpretation bias | | |
| Comparative | | | |
| Notes | Also reported FS in lymphatic, uterine and other tissue samples | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Yes | | |

Wang 1998 (Continued)

| | | | |
|--|---------|--|-------------|
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| | | | High |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | No | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | No | | |
| | | | High |
| DOMAIN 3: Reference Standard | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | No | | |
| | | | Low |
| DOMAIN 4: Flow and Timing | | | |
| Did all patients receive a reference standard? | Yes | | |
| Did all patients receive the same reference standard? | Yes | | |
| Were un-interpretable/intermediate test results reported? | Yes | | |

Wang 1998 (Continued)

| | | | |
|--|-----|--|--|
| Were withdrawals from the study explained? | Yes | | |
| | | | |

Wasinghon 2008

| Study characteristics | | | |
|--|---|---------------------|-------------------------------|
| Patient sampling | Design: Retrospective Setting: Thailand Accrual dates: Jan 2002 - Dec 2006 No participants: 376 No assessed: 376 Inclusion criteria: Consecutive ovarian tumours undergoing surgery where FS and PS performed Included previous histological diagnosis: unclear | | |
| Patient characteristics and setting | Consecutive ovarian masses. Thailand. | | |
| Index tests | FS 1-2 slides, reported by 5 pathologists (expertise unclear) | | |
| Target condition and reference standard(s) | Malignant or borderline on PS. | | |
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | No deferred cases. | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Yes | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |

Wasinghon 2008 (Continued)

| | | | | |
|--|---------|--|--|------------|
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | | |
| | | | | Low |
| DOMAIN 2: Index Test All tests | | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | | |
| Were the index tests Interpreted by consultant or specialist gyn-onc pathologist? | Unclear | | | |
| | | | | Low |
| DOMAIN 3: Reference Standard | | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | | |
| | | | | Low |
| DOMAIN 4: Flow and Timing | | | | |
| Did all patients receive a reference standard? | Yes | | | |
| Did all patients receive the same reference standard? | Yes | | | |
| Were un-interpretable/intermediate test results reported? | Yes | | | |
| Were withdrawals from the study explained? | Yes | | | |
| | | | | |

Wootipoom 2006

| Study characteristics | | | |
|--|---|---------------------|-------------------------------|
| Patient sampling | Design: Retrospective Setting: Thailand Accrual dates: May 1999 to Oct 2004 No participants: 229 No assessed: 213 Inclusion criteria: Excluded 16 deferred diagnoses. Included previous histological diagnosis: unclear | | |
| Patient characteristics and setting | Unclear inclusion criteria. Thailand. | | |
| Index tests | FS. No details given. | | |
| Target condition and reference standard(s) | Malignant or borderline on PS. | | |
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | 16 deferred diagnoses on FS excluded. | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Unclear | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| | | | Unclear |
| DOMAIN 2: Index Test All tests | | | |

Wootipoom 2006 (Continued)

| | | | |
|--|---------|--|--|
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Unclear | | |
| Unclear | | | |
| DOMAIN 3: Reference Standard | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | |
| Low | | | |
| DOMAIN 4: Flow and Timing | | | |
| Did all patients receive a reference standard? | Yes | | |
| Did all patients receive the same reference standard? | Yes | | |
| Were un-interpretable/intermediate test results reported? | No | | |
| Were withdrawals from the study explained? | Yes | | |

Yarandi 2008

| | |
|------------------------------|---|
| Study characteristics | |
| Patient sampling | Design: Retrospective Setting: Iran Accrual dates: Jan 2004 - Aug 2006 No participants: 106 No assessed: 106 Inclusion criteria: All women with ovarian tumours who had a FS diagnosis. Excluded patients with |

Yarandi 2008 (Continued)

| | | | |
|--|--|---------------------|-------------------------------|
| | neo-adjuvant chemo/radiotherapy. Included previous histological diagnosis: unclear | | |
| Patient characteristics and setting | Ovarian masses. Iran. | | |
| Index tests | 2 - 5 samples for FS, samples for PS taken 'from 1cm over maximum tumour diameter'. Single gynaec-pathologist reported all FS and PS | | |
| Target condition and reference standard(s) | Malignant or borderline on PS. | | |
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | - | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Unclear | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| | | | Low |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests interpreted by consultant or specialist gynaec pathologist? | Yes | | |

Yarandi 2008 (Continued)

| | | | | | |
|--|---------|--|--|--|------------|
| | | | | | Low |
| DOMAIN 3: Reference Standard | | | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | | | |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear | | | | |
| | | | | | Low |
| DOMAIN 4: Flow and Timing | | | | | |
| Did all patients receive a reference standard? | Yes | | | | |
| Did all patients receive the same reference standard? | Yes | | | | |
| Were un-interpretable/intermediate test results reported? | No | | | | |
| Were withdrawals from the study explained? | Yes | | | | |
| | | | | | |

Yeo 1998

| | |
|-------------------------------------|---|
| Study characteristics | |
| Patient sampling | Design: Retrospective Setting: China Accrual dates: Jan 1990 to Dec 1995 No participants: 316 No assessed: 316 Inclusion criteria: Pelvic masses. Included previous histological diagnosis: unclear |
| Patient characteristics and setting | Pelvic masses. Hospital, unknown if tertiary. China. |
| Index tests | FS. 2 sections taken at FS in 85% cases. Experienced pathologist. Same pathologist reported PS and FS |

Yeo 1998 (Continued)

| | | | |
|--|--|---------------------|-------------------------------|
| Target condition and reference standard(s) | Malignant or borderline on PS. PS performed by pathologist who interpreted FS. Third party quality assurance performed by a third author | | |
| Flow and timing | FS before PS. | | |
| Comparative | | | |
| Notes | - | | |
| Methodological quality | | | |
| Item | Authors' judgement | Risk of bias | Applicability concerns |
| DOMAIN 1: Patient Selection | | | |
| Was a consecutive or random sample of patients enrolled? | Yes | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | Unclear | | |
| Was the sample representative of patients in practice (90% stage I/II with RMI>200)? | Unclear | | |
| | | | Unclear |
| DOMAIN 2: Index Test All tests | | | |
| Were the index tests interpreted without knowledge of the reference standard? | Yes | | |
| Were the index tests Interpreted by consultant or specialist gynec pathologist? | Yes | | |
| | | | Low |
| DOMAIN 3: Reference Standard | | | |
| Is the reference standards likely to correctly classify the target condition? | Yes | | |

Yeo 1998 (Continued)

| | | | | |
|--|-----|--|--|--|
| Were the reference standard results interpreted without knowledge of the results of the index tests? | No | | | |
| Low | | | | |
| DOMAIN 4: Flow and Timing | | | | |
| Did all patients receive a reference standard? | Yes | | | |
| Did all patients receive the same reference standard? | Yes | | | |
| Were un-interpretable/intermediate test results reported? | No | | | |
| Were withdrawals from the study explained? | No | | | |

FS: frozen section; PS: paraffin section.

Characteristics of excluded studies [ordered by study ID]

| Study | Reason for exclusion |
|---------------------|--|
| Abbasi 2010 | Conference abstract. Retrospective study of 105 patients comparing value of intraoperative cytology and FS. Unable to construct 3 x 3 table from data |
| Abdel-Hady 2012 | Not a DTA study. A study of fertility-conserving surgery for ovarian tumours in children and young adults 6-20 years of age. Although frozen section was performed, evaluation of its accuracy was not part of the study |
| Abe 2013 | Retrospective review of accuracy of FS and imprint cytology in 23 ovarian germ cell tumours |
| Ahmad 2008 | Retrospective study of all FS analyses performed at an institution in Pakistan during 2006 (N = 356). Cohort was not limited to women with ovarian tumours, who comprised only 9% of the sample |
| Alvarez Santin 2011 | Retrospective study of intraoperative consultations of ovarian neoplastic and non-neoplastic lesions (N = 337). Intraoperative diagnoses based on macroscopic exam, FS, imprint cytology or smears, and cyto-histological correlation. Intraoperative diagnoses compared with final histologic diagnoses |

(Continued)

| | |
|----------------------|--|
| Anastasiadis 2002 | Retrospective study of PW and imprint cytology for 52 patients undergoing primary surgery for ovarian cancer |
| Aslam 2010 | Retrospective study of FS analyses compared to preoperative ultrasound and final paraffin section diagnosis. Investigators selected the first 400 women with malignant ovarian tumours and 400 with benign tumours between August 2000 and March 2007. Cohort was not limited to early ovarian cancer and comprised a significant proportion of metastatic tumours (56%). Sensitivity, specificity, PPV and NPV calculated but unable to construct 3 x 3 table |
| Atallah 2004 | Commentary on management of ovarian masses at laparoscopy, including role of FS. No comparison of FS to PS |
| Basaran 2014 | Only borderline cases |
| Bensaid 2006 | Retrospective study of 313 patients to assess performance of laparoscopy +/- FS to identify malignancy. FS performed in 111 (35%) patients. Results compared to final histology |
| Brun 2008 | Retrospective study of patients with epithelial ovarian cancer only |
| Canis 1997 | Laparoscopic versus laparotomy for management of pelvic masses |
| Chapron 1998 | Review of laparoscopic management of pelvic masses. FS reserved for 26 pelvic masses with CA125 within range of 4-76. Not representative of population being studied |
| Cheung 1992 | Retrospective review of all ovarian masses sent for FS, including those to determine extent of metastases and bilaterality of tumour; unable to construct 3 x 3 table from data for FS sent for suspicious pelvic masses |
| Cingillioglu 2011 | Conference abstract. Retrospective study of borderline tumours diagnosed at FS, PS or both in a single unit in Turkey 2000-2011 |
| Coffey 2005 | Review of role of intraoperative consultation, no data provided |
| Da Cunha Bastos 1983 | Included obviously malignant masses. Expertise of pathologists unclear. Unable to extract data for 3 x 3 table |
| Dede 2005 | Use of frozen section laparoscopically for predicted benign masses |
| Dottino 1999 | Prospective study of 160 women undergoing laparoscopic evaluation with FS for adnexal masses. Large masses above umbilicus excluded. No comparison with PS, only discordant cases reported |
| Fain-Kahn 2009 | Ovarian cryopreservation amongst young women undergoing surgery for borderline ovarian tumours |
| Freitag 2004 | Retrospective review of management of 38 patients with borderline ovarian tumours. Not comparing FS to PS |
| Ganesan 2013 | Survey of UK practice of FS in gynaecological oncology |

(Continued)

| | |
|--------------------|--|
| Garg 2011 | Conference abstract. Retrospective study of 166 patients with borderline ovarian tumours only. FS and PS results compared. Unable to construct 3 x 3 table from data |
| Geomini 2005 | Meta-analysis |
| Geomini 2009 | Survey on women's attitudes towards frozen section diagnosis |
| Ghaemmaghami 2008 | Retrospective study of 150 women undergoing laparotomy for adnexal masses in Iran. 143 had FS. Unable to construct 3 x 3 table from data |
| Gocku 2013 | Conference abstract only. Retrospective review of 113 tumours diagnosed on either PS or FS |
| Gol 2003 | All of the data adds up to 221 women, but authors report 222 women. Table 2 data does not add up and does not match data in text |
| Gultekin 2011 | Retrospective study of 82 patients with borderline tumours only. FS and PS results compared |
| Gupta 2013 | Conference abstract. Retrospective study of 52 patients with borderline tumours only. FS and PS results compared, unable to construct 3 x 3 table |
| Guzel 2011 | Prospective study of postmenopausal and women of reproductive age (N = 80) with predicted benign adnexal masses. FS in 75% of cases but no data on accuracy |
| Guzin 2013 | Conference abstract. Retrospective review of 40 borderline tumours diagnosed on either FS or PS |
| Harmon 2011 | Conference abstract. Retrospective study of 100 consecutive ovarian mucinous tumours that underwent FS. FS and PS results compared |
| Hua 2005 | Full article in Chinese |
| Ismiil 2009 | Retrospective review 731 FS from all gynaecological operations performed, 29 performed for ovarian cyst and 591 from ovary/tube. 257 of these were performed by general gynaecologist or surgeon. This is not representative of the population being studied |
| Ivanov 2005 | Full article in Bulgarian |
| Jaafar 2005 | Review of frozen section concepts. Not original research |
| Kato 2011 | Review of histopathological features of FS for 40 clear cell carcinomas and 30 serous ovarian tumours |
| Kay, kç öğ lu 2000 | Retrospective study of 33 patients with borderline tumours only. FS and PS results compared |
| Khunamornpong 2003 | Prospective study of 131 ovarian masses submitted for scrape cytology and not intraoperative frozen section analysis |
| Kim 2009a | Retrospective study of 101 patients with borderline tumours only. FS and PS results compared |

(Continued)

| | |
|-------------------|---|
| Kim 2009b | Retrospective study of 209 patients with borderline tumours only. FS (182 cases) and PS results compared |
| Kim 2013 | Conference abstract only. Retrospective review of 179 borderline tumours diagnosed on FS |
| Konopacka 2012 | Prospective observational study of 131 patients undergoing laparoscopic surgery for adnexal masses. FS performed in 87 cases. Unable to construct 3 x 3 table from data |
| Kumpulainen 2007 | Prospective study of 65 patients with borderline tumours only to assess staging/treatment and outcomes in different hospital settings. FS in half of cases, results compared to PS |
| Kushima 2013 | 3 case reports on usefulness of intraoperative cytology |
| Leng 2006 | Retrospective review of benign pelvic masses managed with laparoscopy |
| Li 2009 | Full article in Chinese |
| Lin 1993 | Retrospective review of 80 women undergoing laparotomy for pelvic mass. FS in 48 cases with discussion of discordant cases but no data provided |
| Liu 2010 | Prospective review of diagnostic accuracy of haptoglobin level in ovarian cyst fluid for intraoperative triage of epithelial ovarian cancers |
| Marana 2005 | Prospective study of FS of adnexal masses at laparoscopy for ultrasonographically non-suspicious adnexal mass |
| Maruoka 2003 | Full article in Japanese |
| Medeiros 2005 | Quantitative systematic review of diagnostic accuracy of FS, including 14 studies |
| Mendilcioglu 2002 | Retrospective study of 61 patients undergoing laparoscopy for adnexal masses, aiming to assess the safety of laparoscopic approach. FS performed in only 8 (13%) of cases |
| Menzin 1995 | Retrospective review of 48 patients with FS diagnosis borderline tumour. 2 patients were stage II, 10 patients were stage III. This is not representative of the population being studied |
| Michael 1996 | Comparison of cytology and frozen section. No comparison to paraffin section |
| Moodley 2005 | Commentary on frozen section. Not original research |
| Morotti 2011 | Conference abstract. Retrospective review of 98 borderline tumours diagnosed by FS, PS or both. Unable to construct 3 x 3 table from data |
| Nasfi 2012 | Retrospective study of 79 ovarian mucinous tumours that underwent FS. FS and PS results compared |
| Nevin 2010 | Letter in response to Warwick 2009 . |
| Obiakor 1991 | Retrospective review of 311 FS classified as benign or malignant. Unable to construct 3 x 3 table from data |

(Continued)

| | |
|-----------------------|--|
| Ozdamar 2006 | Retrospective review of all FS analysed in a pathology laboratory 2001-2005. No details provided for ovarian masses alone |
| Parker 2011 | Conference abstract only. Review of 831 frozen sections interpreted by general or specialist gynaecological pathologists |
| Pongsuvareeyakul 2012 | Retrospective study of mucinous tumours only |
| Puga 2011 | Conference abstract. Retrospective study of 67 patients with borderline tumours only. Unable to construct 3 x 3 table from data |
| Quan 2004 | 31 patients with stage IV breast cancer with either adnexal mass or undergoing therapeutic BSO. Not representative of population being studied in this review |
| Saglam 2006 | Letter to editor discussing 4 discordant cases of a total 174 FS performed during 2002 in a single unit |
| Sakurai 2004 | Full article in Japanese |
| Salman 2013 | Conference abstract only. 745 pelvic masses undergoing FS |
| Scurry 1989 | Retrospective review of 203 FS from all gynaecological operations, including 73 ovarian. No report that these were suspicious masses. Authors agree that many FS were performed on grossly benign appearing cysts. Not representative of population being studied in this review |
| Seckin 2011 | Retrospective study of females 25 years or younger undergoing laparoscopic surgery for presumed benign ovarian cysts |
| Shahid 2012 | Reports role of intraoperative cytology not frozen section. |
| Shih 2011 | Retrospective study of 120 patients with borderline tumours diagnosed at FS |
| Slavutin 1979 | Retrospective study of 55 patients with serous ovarian tumours. FS and PS reviewed by 2 pathologists for study and compared with original results |
| Song 2011 | Retrospective study of 354 patients with borderline tumours only. FS and PS results compared |
| Souka 1990 | Retrospective review of combined use of imprint cytology and FS to evaluate 50 pelvic masses at laparotomy. Borderline tumours at PS were grouped together with malignant. Unable to construct 3 x 3 table |
| Spann 1994 | Report on role of FS and gross inspection combined. Unable to extract data for FS alone. May not be representative of study population as 88% of intraoperative consultations were benign diagnoses |
| Springel 2009 | Retrospective study of FS intraoperative consultations reported as epithelial ovarian tumours |
| Stewart 2005 | Retrospective study of 914 patients in Australia looking at accuracy of FS to determine primary from metastatic disease, 1999-2003. 32 patients known to have extra-ovarian disease at time of FS. Patient selection bias therefore high. FS omentum and lymph node included |

(Continued)

| | |
|------------------|--|
| Stewart 2008 | Results for clear cell carcinoma were assessed separately and compared with a similar number with serous and endometrial cancer |
| Stewart 2010 | 402 cases where cytology was compared to frozen section. No comparison to paraffin section made |
| Storms 2012 | Retrospective review of 73 ovarian mucinous tumours |
| Takemoto 2014 | Retrospective review of benign masses diagnosed at FS at laparoscopy. Not representative of study population |
| Tempfer 2007 | Borderline tumours only |
| Twigg 2012 | Letter in response to Cross 2012 |
| Uguz 2005 | Prospective study of 62 women having FNAC of ovarian masses and not intraoperative frozen section analysis |
| Ulrich 2000 | Retrospective analysis of FS results for 226 adnexal masses. Excluded simple masses on USS and suspicious masses that required conversion to laparotomy. Heavy selection bias in that masses not considered to need laparotomy were excluded. As a result, 202 of 211 studied women had benign disease on PS |
| Usubutun 1998 | Retrospective review of 360 ovarian masses with FS. 12 deferred cases. Unable to construct 3 x 3 table from data |
| Vemavarapu 2014 | Conference abstract only. Retrospective review of 73 pelvic masses submitted for FS |
| Vijayakumar 2013 | Prospective study of intraoperative imprint cytology in 50 patients with suspected ovarian malignancy |
| Warwick 2009 | Retrospective study to determine optimal management strategy for women with suspected stage I ovarian cancer. No data on FS accuracy |
| Wingo 2006 | Retrospective study of 32 patients with borderline (low malignant potential) tumours only |
| Zhang 1993 | Full article in Chinese |

FNAC: fine needle aspiration cytology; **FS:** frozen section; **NPV:** negative predictive value; **PPV:** positive predictive value; **PS:** paraffin section; **PW:** peritoneal washing; **USS:** ultrasound scan.

DATA

Presented below are all the data for all of the tests entered into the review.

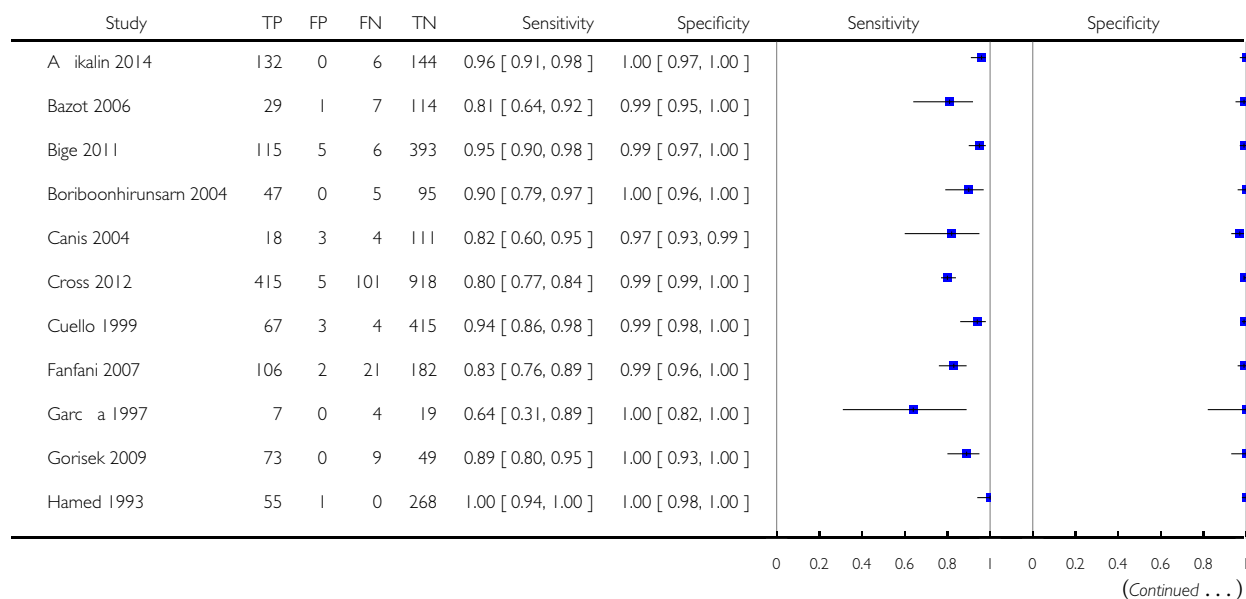
Tests. Data tables by test

| Test | No. of studies | No. of participants |
|---|----------------|---------------------|
| 1 Frozen section: Threshold Malignancy vs Borderline or Benign | 38 | 11181 |
| 2 Frozen section: Threshold Malignancy or Borderline vs Benign | 38 | 11181 |
| 3 Frozen section: Threshold Malignancy vs Borderline or Benign when FS indicated Mal or BOT | 38 | 3953 |

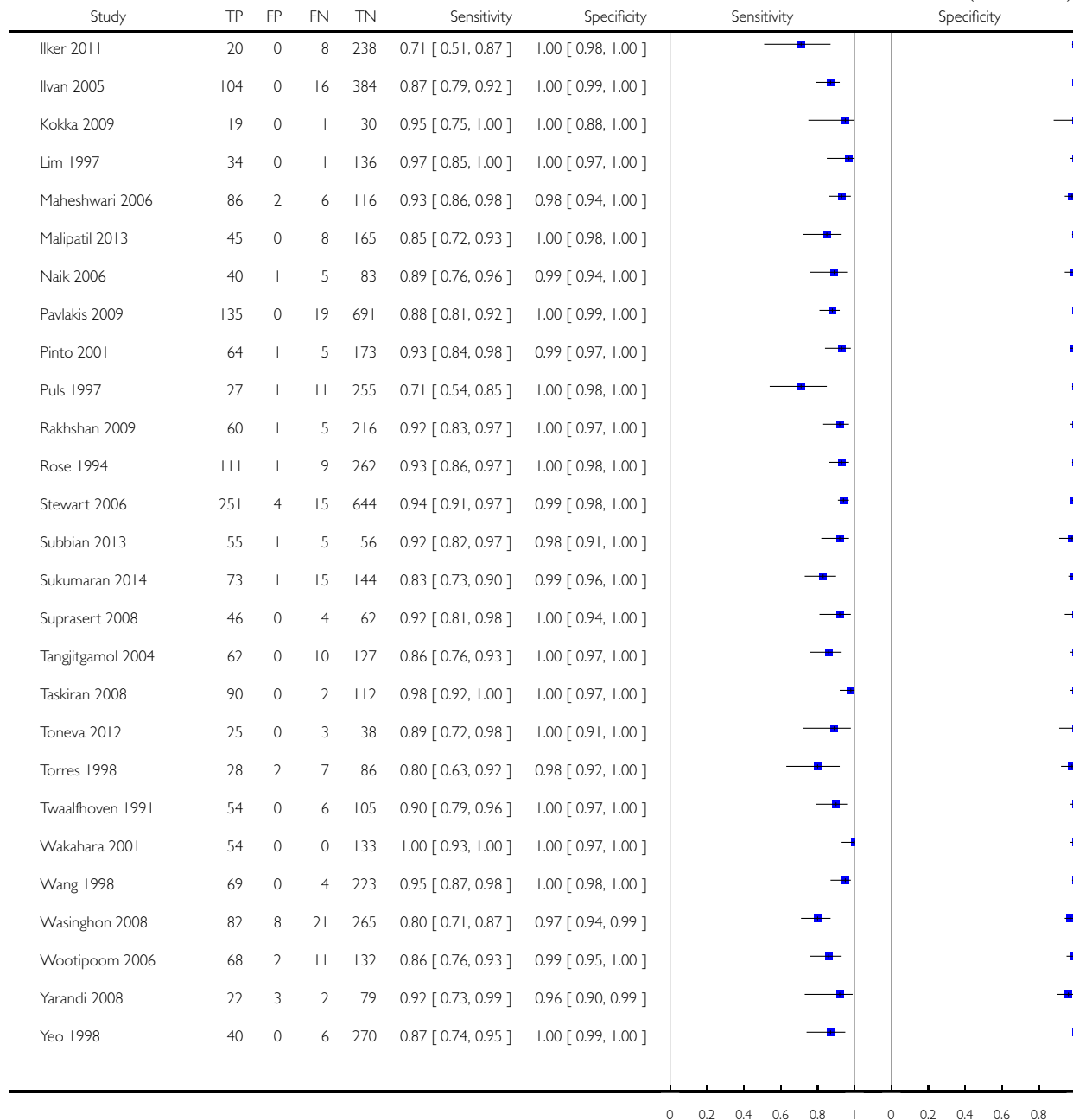
Test 1. Frozen section: Threshold Malignancy vs Borderline or Benign.

Review: Intraoperative frozen section analysis for the diagnosis of early stage ovarian cancer in suspicious pelvic masses

Test: 1 Frozen section: Threshold Malignancy vs Borderline or Benign



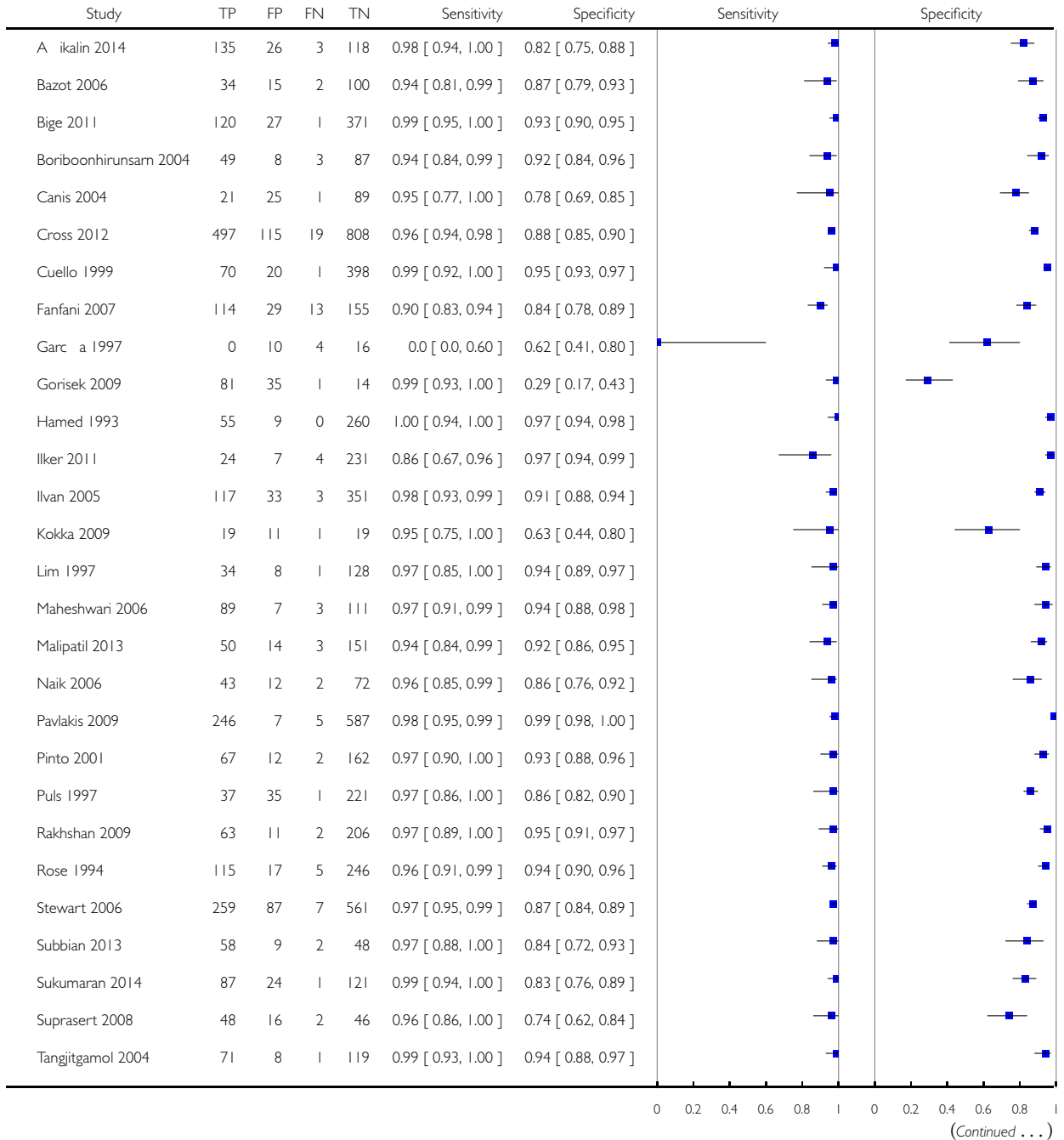
(... Continued)

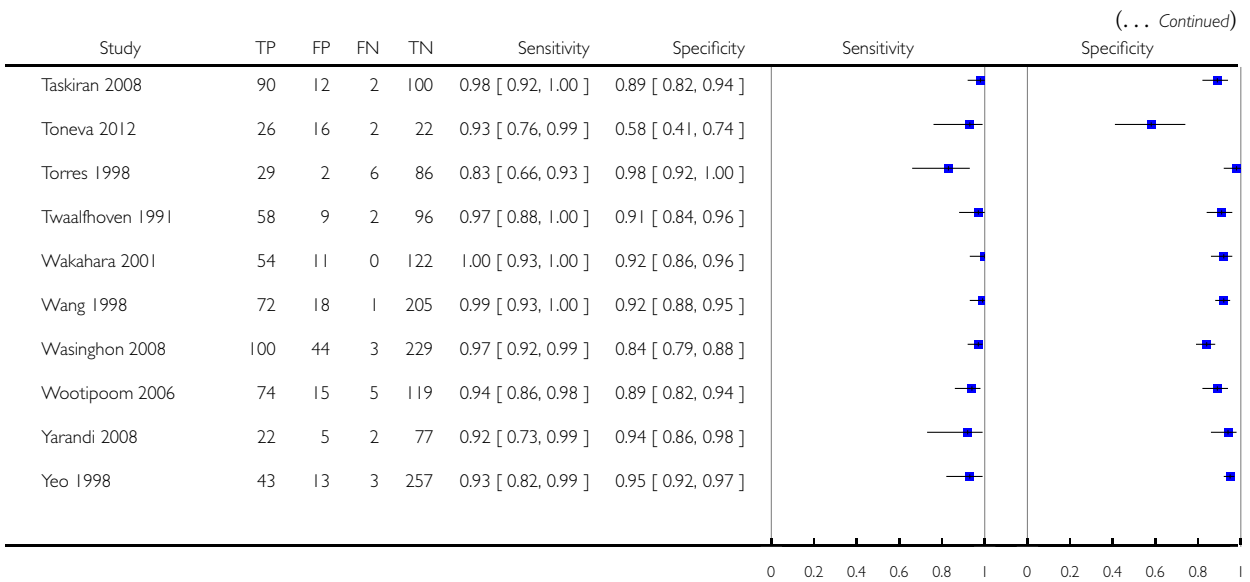


Test 2. Frozen section: Threshold Malignancy or Borderline vs Benign.

Review: Intraoperative frozen section analysis for the diagnosis of early stage ovarian cancer in suspicious pelvic masses

Test: 2 Frozen section: Threshold Malignancy or Borderline vs Benign

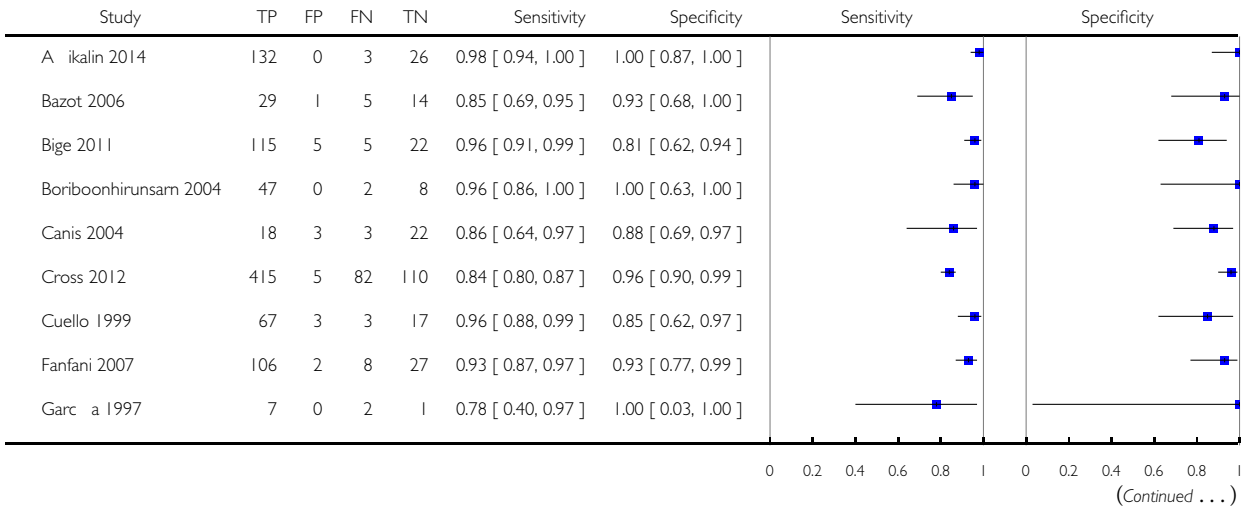




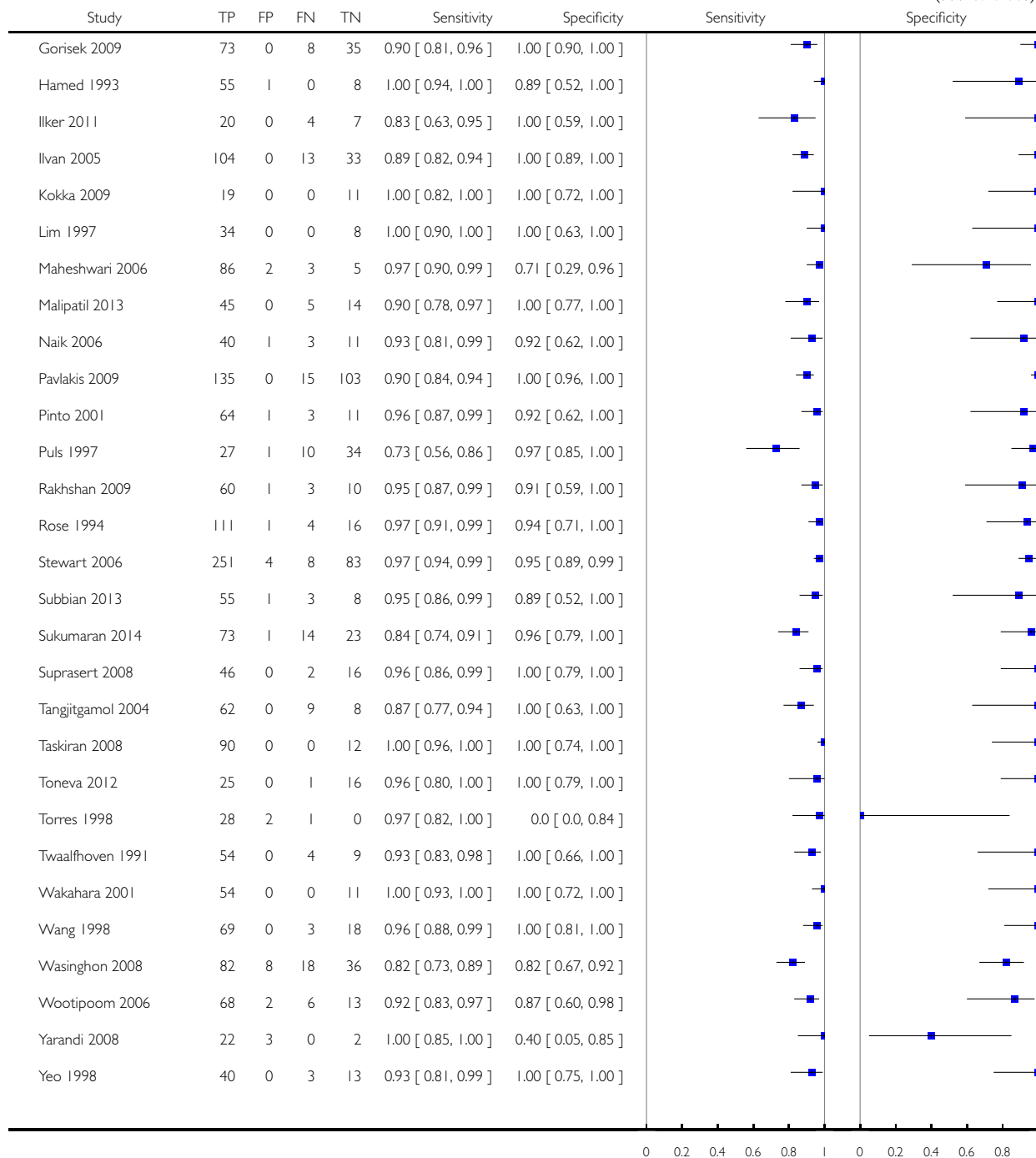
Test 3. Frozen section: Threshold Malignancy vs Borderline or Benign when FS indicated Mal or BOT.

Review: Intraoperative frozen section analysis for the diagnosis of early stage ovarian cancer in suspicious pelvic masses

Test: 3 Frozen section: Threshold Malignancy vs Borderline or Benign when FS indicated Mal or BOT



(... Continued)



ADDITIONAL TABLES

Table 1. Accuracy of frozen section malignant results to identify women with malignancy

| Frozen section | Paraffin section positive test | Paraffin section negative test | |
|----------------|--------------------------------|--------------------------------|----------------|
| | Malignant | Borderline | Benign |
| Malignant | True positive | False positive | False positive |
| Borderline | False negative | True negative | True negative |
| Benign | False negative | True negative | True negative |

Table 2. Accuracy of frozen section malignant or borderline results to identify women with malignancy

| Frozen section | Paraffin section positive test | Paraffin section negative test | |
|----------------|--------------------------------|--------------------------------|--------------------------|
| | Malignant | <i>Borderline</i> | Benign |
| Malignant | True positive | False positive | <i>FP</i> False positive |
| Borderline | True positive | False positive | False positive |
| Benign | False negative | True negative | True negative |

APPENDICES

Appendix I. International Federation of Gynecology and Obstetrics (FIGO) staging of ovarian cancer

Stage I. Stage I consists of tumour limited to the ovaries or fallopian tubes.

- Stage IA includes the following: tumour limited to one ovary (capsule intact) or fallopian tube. No tumour on the external surface of the ovary or fallopian tube. No malignant cells in ascites or peritoneal washings
- Stage IB includes the following: tumour limited to both ovaries (capsules intact) or fallopian tubes. No tumour on the external surface of the ovaries or fallopian tubes. No malignant cells in ascites or peritoneal washings
- Stage IC includes tumour limited to one or both ovaries or fallopian tubes, with any of the following: Stage IC1: Surgical spill. Stage IC2: Capsule ruptured before surgery, or tumour on ovarian or fallopian tube surface. Stage IC3: Malignant cells in the ascites or peritoneal washings

Stage II. In stage II tumour involves one or both ovaries or fallopian tubes, with pelvic extension (below pelvic brim) or primary peritoneal cancer.

- Stage IIA: Extension, implants or both on at least one of the following: uterus, ovaries and fallopian tubes.
- Stage IIB: Extension to other pelvic intraperitoneal tissues

Stage III . In stage III, tumour involves one or both ovaries or fallopian tubes, or primary peritoneal cancer, with cytologically or histologically confirmed spread to the peritoneum outside the pelvis and/or metastasis to the retroperitoneal lymph nodes.

- Stage IIIA includes the following: Stage IIIA1: Positive (cytologically or histologically proven) retroperitoneal lymph nodes only. Stage IIIA1(i) Metastasis up to 10 mm in greatest dimension. Stage IIIA1(ii) Metastasis more than 10 mm in greatest dimension. Stage IIIA2: Microscopic extrapelvic (above the pelvic brim) peritoneal involvement with or without positive retroperitoneal lymph nodes

- Stage IIIB involves macroscopic peritoneal metastasis beyond the pelvis up to 2 cm in greatest dimension, with or without metastasis to the retroperitoneal lymph nodes.

- Stage IIIC involves macroscopic peritoneal metastasis beyond the pelvis more than 2 cm in greatest dimension, with or without metastasis to the retroperitoneal lymph nodes. Stage IIIC includes extension of tumour to the capsule of liver and spleen without parenchymal involvement of either organ.

Stage IV . Stage IV consists of distant metastasis, excluding peritoneal metastases, and includes the following:

- Stage IVA: pleural effusion with positive cytology.
- Stage IVB: parenchymal metastases and metastases to extra-abdominal organs (including inguinal lymph nodes and lymph nodes outside of the abdominal cavity)

Appendix 2. MEDLINE (Ovid) search strategy

1. exp Ovarian Neoplasms/
2. (ovar* adj5 (cancer* or tumor* or tumour* or adenocarcinoma* or carcinosarcoma* or cystadenocarcinoma* or carcinoma* or malignan* or neoplas* or carcinogen* or teratoma* or metasta* or mass or masses)).tw,ot.
3. (thecoma* or luteoma*).tw,ot.
4. 1 or 2 or 3
5. Frozen Sections/
6. (FS or FSA or IFS or IFSA).tw,ot.
7. (frozen or quick) adj5 section*.tw,ot.
8. ((intraoperative or intra-operative) adj5 (consultation* or histolog* or diagnos* or patholog*)).tw,ot.
9. (cryosection* or cryogenic*).tw,ot.
10. (fresh or frozen) adj5 tissue*).tw,ot.
11. 5 or 6 or 7 or 8 or 9 or 10
12. 4 and 11
13. exp animals/ not humans.sh.
14. 12 not 13

key: tw=textword, ot=original title

CONTRIBUTIONS OF AUTHORS

- Guarantor of the review: RN
- Conceiving the idea: RN, AP, PC
- Designing and coordinating the review: NR, RN
- Data collection for the review; designing search strategies; undertaking searches; screening search results: TL, NR, JH
- Organising retrieval of papers: NR, AB
- Screening retrieved papers against inclusion criteria: NR, AP
- Appraising quality of papers: NR, AB, RS
- Extracting data from papers: NR, AB, CF, RS, SM
- Providing additional data about papers: NR
- Obtaining and screening data on unpublished studies: NR, AP

- Data management of the review: NR
- Entering data into RevMan: NR, SM, RS
- Analysis and interpretation of data: SM, NR, RS
- Providing a methodological perspective; providing a clinical perspective; providing a policy perspective; providing a consumer perspective: RN, NR, AP, PC
- Writing the review: NR, SM
- Providing general advice on the review: RN, NR, AP, PC, KG
- Securing funding for the review: NR, RN

DECLARATIONS OF INTEREST

PC, AP and RN were authors in a study that met the inclusion criteria in the review.

NR: none known.

AB: none known.

SM: Received payment for methodology work on review.

RS: none known.

CF: none known.

KG: none known.

SOURCES OF SUPPORT

Internal sources

- No sources of support supplied

External sources

- Department of Health, UK.
NHS Cochrane Collaboration programme Grant Scheme CPG-10/4001/12

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

We clarified that we used sensitivity and specificity in our primary analysis to assess accuracy.

We clarified that our objectives included assessment for the following two thresholds for frozen section.

1. Test positive is cancer.
2. Test positive is cancer and borderline.

The reference standard test threshold for all analyses is test positive cancer and test negative borderline or benign.

The secondary objective of the protocol (renamed secondary objective #1 in the review) could not be addressed due to lack of data in included studies. We included an additional analysis, Secondary objective #2, which was the closest substitute to secondary objective #1, which could be addressed.

There was insufficient data to examine heterogeneity except for pathologist reader experience. We did not assess reporting bias, based on recommendations in the *Cochrane Handbook for Diagnostic Test Accuracy Reviews*.