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Diagnostic accuracy of laparoscopy following computed tomography (CT) scanning for assessing the resectability with curative intent in pancreatic and periampullary cancer (Review)

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[Diagnostic Test Accuracy Review]

Diagnostic accuracy of laparoscopy following computed tomography (CT) scanning for assessing the resectability with curative intent in pancreatic and periampullary cancer

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ABSTRACT

Background

Surgical resection is the only potentially curative treatment for pancreatic and periampullary cancer. A considerable proportion of patients undergo unnecessary laparotomy because of underestimation of the extent of the cancer on computed tomography (CT) scanning. Laparoscopy can detect metastases not visualised on CT scanning, enabling better assessment of the spread of cancer (staging of cancer). This is an update to a previous Cochrane Review published in 2013 evaluating the role of diagnostic laparoscopy in assessing the resectability with curative intent in people with pancreatic and periampullary cancer.

Objectives

To determine the diagnostic accuracy of diagnostic laparoscopy performed as an add-on test to CT scanning in the assessment of curative resectability in pancreatic and periampullary cancer.

Search methods

We searched the Cochrane Central Register of Controlled Trials (CENTRAL), MEDLINE via PubMed, EMBASE via OvidSP (from inception to 15 May 2016), and Science Citation Index Expanded (from 1980 to 15 May 2016).

Selection criteria

We included diagnostic accuracy studies of diagnostic laparoscopy in people with potentially resectable pancreatic and periampullary cancer on CT scan, where confirmation of liver or peritoneal involvement was by histopathological examination of suspicious (liver or peritoneal) lesions obtained at diagnostic laparoscopy or laparotomy. We accepted any criteria of resectability used in the studies. We included studies irrespective of language, publication status, or study design (prospective or retrospective). We excluded case-control studies.

Data collection and analysis

Two review authors independently performed data extraction and quality assessment using the QUADAS-2 tool. The specificity of diagnostic laparoscopy in all studies was 1 because there were no false positives since laparoscopy and the reference standard are one and the same if histological examination after diagnostic laparoscopy is positive. The sensitivities were therefore meta-analysed using a



univariate random-effects logistic regression model. The probability of unresectability in people who had a negative laparoscopy (post-test probability for people with a negative test result) was calculated using the median probability of unresectability (pre-test probability) from the included studies, and the negative likelihood ratio derived from the model (specificity of 1 assumed). The difference between the pre-test and post-test probabilities gave the overall added value of diagnostic laparoscopy compared to the standard practice of CT scan staging alone.

Main results

We included 16 studies with a total of 1146 participants in the meta-analysis. Only one study including 52 participants had a low risk of bias and low applicability concern in the patient selection domain. The median pre-test probability of unresectable disease after CT scanning across studies was 41.4% (that is 41 out of 100 participants who had resectable cancer after CT scan were found to have unresectable disease on laparotomy). The summary sensitivity of diagnostic laparoscopy was 64.4% (95% confidence interval (CI) 50.1% to 76.6%). Assuming a pre-test probability of 41.4%, the post-test probability of unresectable disease for participants with a negative test result was 0.20 (95% CI 0.15 to 0.27). This indicates that if a person is said to have resectable disease after diagnostic laparoscopy and CT scan, there is a 20% probability that their cancer will be unresectable compared to a 41% probability for those receiving CT alone.

A subgroup analysis of people with pancreatic cancer gave a summary sensitivity of 67.9% (95% CI 41.1% to 86.5%). The post-test probability of unresectable disease after being considered resectable on both CT and diagnostic laparoscopy was 18% compared to 40.0% for those receiving CT alone.

Authors' conclusions

Diagnostic laparoscopy may decrease the rate of unnecessary laparotomy in people with pancreatic and periampullary cancer found to have resectable disease on CT scan. On average, using diagnostic laparoscopy with biopsy and histopathological confirmation of suspicious lesions prior to laparotomy would avoid 21 unnecessary laparotomies in 100 people in whom resection of cancer with curative intent is planned.

PLAIN LANGUAGE SUMMARY

What is the diagnostic accuracy of laparoscopic staging following a CT scan for assessing whether pancreatic and periampullary cancer is resectable?

Background

The pancreas is an organ situated in the abdomen close to the junction of the stomach and small bowel. It secretes digestive juices which are necessary for the digestion of all food materials. The digestive juices secreted in the pancreas drain into the upper part of the small bowel via the pancreatic duct. The bile duct is a tube which drains bile from the liver and gallbladder. The pancreatic and bile ducts share a common path just before they drain into the small bowel. This area is called the periampullary region. Surgical removal is the only potentially curative treatment for cancers arising from the pancreatic and periampullary regions. A considerable proportion of patients undergo unnecessary major open abdominal exploratory operation (laparotomy) because their CT scan has underestimated the spread of cancer. If during the major open operation the cancer is found to have spread within the abdomen, patients are referred for alternate treatments such as chemotherapy, which do not cure the cancer but may improve survival.

This major open abdominal operation can be avoided if the spread of cancer within the abdomen is known, called 'staging' the cancer. The minimum test used for staging is usually the computed tomography (CT) scan. However, CT scan can understage the cancer, that is it can underestimate the spread of cancer. Laparoscopy, a procedure whereby a small telescope is inserted inside the abdomen through a small (keyhole) surgical incision, can detect spread not identified on CT scanning. Different studies report different accuracy of laparoscopy in assessing whether the cancer can be removed. Our aim therefore was to find out the average diagnostic accuracy of laparoscopy for staging pancreatic and periampullary cancers considered to be removable after a CT scan. This review is an update of our previous review.

A glossary of terms is provided in Appendix 1.

Study characteristics

We performed a thorough literature search to identify studies published up to 15 May 2016. We identified 16 studies reporting information on 1146 people with pancreatic or periampullary cancers which were considered to be eligible for potentially curative surgery based on CT scan staging. These studies evaluated diagnostic laparoscopy and compared results of the procedure with the eventual diagnosis by the surgeon that the cancer was not resectable during major abdominal operation or examination under microscope.

Quality of evidence

All of the studies were of unclear or low methodological quality in one or more aspects, which may undermine the validity of our findings.

Key results



Of those people with what CT suggests seems to be a potentially surgically curable cancer, the percentage in whom more extensive cancer was found on further staging with diagnostic laparoscopy or laparotomy ranged between 17% and 82% across studies. The median percentage of people in whom cancer spread was not detected by CT scan was 41%. Adding staging laparoscopy to CT scan might decrease the number of people with unremovable disease undergoing unnecessary major operations to 20% compared to those who undergo unnecessary major operation after CT scan alone (41%). This means that using diagnostic laparoscopy could halve the rate of unnecessary major open operations in people undergoing major surgery for potentially surgically curable pancreatic cancer.



SUMMARY OF FINDINGS

Summary of findings 1. Diagnostic laparoscopy

Population	Males and females aged 15 to 87 years with potentially re	esectable pancreatic or periampullary carcinoma			
•	on computed tomography (CT) scanning				
Setting	Surgical centres in the USA, Germany, the UK, Japan, Isra	ael, and the Netherlands			
Index test	Diagnostic laparoscopy with histologic confirmation				
Reference standard	Paraffin section histology on diagnostic laparoscopy or lasectability on laparotomy	aparotomy or surgeon's judgement of unre-			
	True positive: Suspicious lesion on diagnostic laparoscopexamination of biopsy obtained during diagnostic laparo				
	False positive: This is not possible since laparotomy will of the suspicious lesion on diagnostic laparoscopy show:				
	False negative: No evidence of unresectability by diagnos on laparotomy	stic laparoscopy but evidence of unresectability			
	True negative: No evidence of unresectability by diagnos	tic laparoscopy and laparotomy			
Number of studies	16 studies				
Summary sensitivity	64.4% (95% confidence interval 50.1% to 76.6%)				
Consistent results	No				
Uncertainty (overall risk of bias)	High				
Other limitations	Different definitions of unresectability because studies u parotomy when biopsy confirmation was not possible	sed surgeon's judgement of unresectability on la-			
Pre-test probability from included studies ¹	Post-test probability of unresectable disease for patients with a negative test result (95% confidence interval) ²	Percentage of patients for whom unneces- sary laparotomy can be avoided ³			
Minimum = 17.4	7.0 (4.9 to 9.8)	10.4			
Lower quartile = 34.7	15.9 (11.4 to 21.6)	18.8			
Median = 41.4	20.1 (14.7 to 26.8) 21.3				
Upper quartile = 62.7	37.4 (29.0 to 46.6) 25.3				
Maximum = 81.8	61.5 (52.3 to 70.0)	20.3			
Interpretation	At pre-test probabilities of 17%, 41%, and 82%, adding derative staging of pancreatic cancer avoids 10, 21, and 20 tomies performed for curative resection purposes. These and maximum values obtained from the included studies	Ounnecessary laparotomies out of 100 laparo- e pre-test probabilities are the minimum, middle,			

¹Probability of someone having unresectable disease at laparotomy after CT indicated that the disease is resectable.

²Probability of someone having unresectable disease after the CT and diagnostic laparoscopy indicated that the disease is resectable.



³Calculated as the difference between the post-test probability and the pre-test probability. All probabilities are reported in the table as percentages.



BACKGROUND

Periampullary cancer develops near the ampulla of Vater (National Cancer Institute 2011a). This includes cancer of the head and neck of the pancreas, cancer of the distal end of the bile duct, cancer of the ampulla of Vater, and cancer of the second part of the duodenum. Pancreaticoduodenectomy is the main treatment for cancers arising in the head of the pancreas, ampulla, and second part of the duodenum. Surgical resection is generally considered to be the only cure for pancreatic cancer. However, only 15% to 20% of people with pancreatic cancers undergo potentially curative resection (Conlon 1996; Engelken 2003; Michelassi 1989; Shahrudin 1997; Smith 2008). In all other people, the cancers are not resected because of infiltration of local structures, disseminated disease, or because the person is deemed unfit to undergo major surgery. Computed tomography (CT scan) is generally used for staging pancreatic and periampullary cancers (National Cancer Institute 2011b). Despite undergoing routine CT scanning to stage the disease (Mayo 2009), a substantial proportion of patients (approximately 10% to 25%) undergo unnecessary laparotomy (opening the abdomen using a large incision) with lack of curative resectability identified only during the laparotomy (Lillemoe 1999; Mayo 2009). Laparoscopy can be used to detect metastatic disease in people with periampullary cancer.

Target condition being diagnosed

Inability to perform curative resectability of pancreatic and periampullary cancer ('unresectable' cancers)

Index test(s)

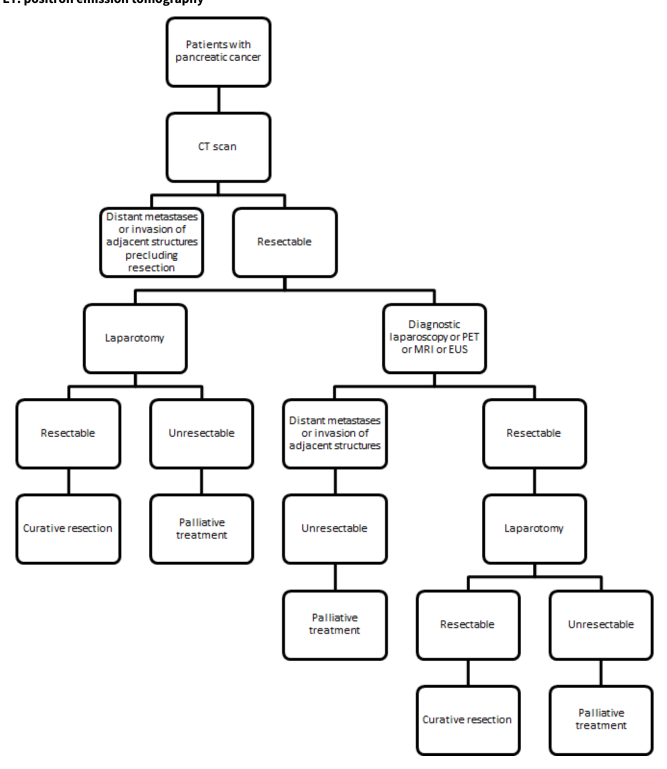
Diagnostic laparoscopy involves the use of a laparoscope (a telescope inserted into the abdominal cavity through a keyhole incision) to visualise and explore the abdominal organs. Also known as staging laparoscopy, it is used following initial staging by CT scanning. Any spread of cancer to the liver, peritoneum, or adjacent structures can be visualised during diagnostic laparoscopy. A biopsy of the suspicious lesion can be performed, and the biopsy specimen can be examined under the microscope to confirm that the suspicious lesion is spread of cancer.

Clinical pathway

No standard algorithm is currently available for assessing the resectability of pancreatic and periampullary cancers, with clinicians following their own algorithms based on either their clinical experience or education. Almost all current algorithms include a CT scan as one of the tests (National Cancer Institute 2011b). CT may be the only test performed before laparotomy. Other tests such as diagnostic laparoscopy, positron emission tomography (PET) scanning, magnetic resonance imaging (MRI), or endoscopic ultrasound (EUS) may be used in addition to CT scan to assess resectability. The possible clinical pathway in the staging of pancreatic cancers is shown in Figure 1. Another review is assessing the accuracy of these various tests and CT scanning (Gurusamy 2015).



Figure 1. Clinical pathway. EUS: endoscopic ultrasound MRI: magnetic resonance imaging PET: positron emission tomography



Prior test(s)

The minimum prior test should be CT, and the cancer should be resectable with curative intent on the basis of the CT scan to be included in this review. Other tests such as PET scanning, MRI, or EUS might be used in addition to CT scanning to assess resectability

prior to diagnostic laparoscopy. We included participants in this review irrespective of whether they underwent these other tests prior to diagnostic laparoscopy.



Role of index test(s)

Diagnostic laparoscopy can be considered as an add-on test to the CT scan prior to laparotomy done with the intention of performing a potentially curative resection.

Alternative test(s)

Other tests such as PET scanning, laparoscopic ultrasound, or EUS may be used as alternative tests to diagnostic laparoscopy in people considered to have CT resectable pancreatic and periampullary cancer. As mentioned earlier, PET scanning and EUS may also be used prior to diagnostic laparoscopy. Laparoscopic ultrasound may be used in combination with diagnostic laparoscopy, and the strategy for determining test positivity of the combination may be either test positive or both tests positive.

Rationale

Diagnostic laparoscopy allows internal visualisation of the abdomen and can detect any peritoneal spread of the cancer or the involvement of any adjacent structures. A biopsy and histopathological examination of any suspicious lesion can be performed and an unnecessary laparotomy to attempt curative resection avoided. If this add-on test can identify unresectable cancers without laparotomy, it might decrease the costs and morbidity associated with unnecessary laparotomy. This is an update to an earlier Cochrane Review assessing the resectability with curative intent in pancreatic and periampullary cancer published in 2013 (Allen 2013).

OBJECTIVES

To determine the diagnostic accuracy of diagnostic laparoscopy performed as an add-on test to CT scanning in the assessment of curative resectability in pancreatic and periampullary cancer.

Secondary objectives

We planned to explore the following sources of heterogeneity.

- 1. Studies at low risk of bias versus those at unclear or high risk of bias based on methodological quality assessment using the QUADAS-2 tool (Whiting 2011).
- 2. Full-text publications versus abstracts (this can inform about publication bias since there may be an association between the results of the study and the study reaching full publication status) (Eloubeidi 2001).
- 3. Prospective studies versus retrospective studies.
- 4. Proportion of participants with pancreatic cancer, ampullary cancer, and bile duct cancers (although classified as periampullary cancers, each has a different prognosis) (Klempnauer 1995). The additional value of diagnostic laparoscopy may be different because of the extent of spread in these different types of periampullary cancers.
- 5. Procedures performed under the same anaesthetic versus procedures performed under a different anaesthetic (there are likely to be differences in the histopathological examinations since the former procedure is associated with frozen section biopsy, while the latter procedure is likely to be associated with paraffin section). Paraffin section is considered to be the gold standard in identifying cancer. Frozen sections can be associated with false-negative results (Yeo 2002). However, frozen section

- results are always confirmed by paraffin section histological examinations.
- 6. Different definitions for resectable cancer on laparotomy. Different surgeons may consider cancer unresectable differently, i.e. they will have different criteria for unresectability on laparotomy (other than the consensus criteria for resectability). For example, one surgeon may judge that the cancer is unresectable on laparotomy because of the involvement of the vessel and consider the reference standard to be positive. This will result in a false-negative result for laparoscopy. Another surgeon may judge the same cancer to be resectable despite the involvement of the vessel and proceed with resection. The reference standard will be negative in this situation, resulting in a true-negative result for laparoscopy. This might have an intrinsic threshold effect.
- Additional pre-tests performed (besides CT scan). This can alter the pre-test probability of unresectability and can help in the assessment of the additional value of diagnostic laparoscopy under various situations.

METHODS

Criteria for considering studies for this review

Types of studies

We included studies that evaluated the accuracy of diagnostic laparoscopy in the appropriate patient population (see below) irrespective of language or publication status, or whether data were collected prospectively or retrospectively. However, we excluded case reports which did not provide sufficient diagnostic test accuracy data. We also excluded case-control studies, which are prone to bias (Whiting 2011).

Participants

People about to undergo curative resection for pancreatic and periampullary cancer with no contraindications (such as metastatic disease) for curative resection on CT scan, and who were anaesthetically fit to undergo major surgery.

Index tests

We included only diagnostic laparoscopy in which histopathological confirmation of metastatic spread was obtained on a paraffin section.

Target conditions

The target conditions were unresectable pancreatic and periampullary cancers, that is diagnostic laparoscopy was considered to be a positive test if the pancreatic or periampullary cancer was unresectable. In these cancers it is not possible to perform curative resectability. There are no uniform criteria for resectability of pancreatic and periampullary cancer. Consensus exists for the definition of borderline resectable cancers (Abrams 2009). Therefore, where there is less tissue involvement than in a borderline resectable cancer, the tumour can be considered as resectable. We accepted any criteria of resectability used by the study authors and acknowledge that this could potentially create a threshold effect. In general, the cancer would not be resected if liver or peritoneal metastases were noted, or if the cancer had invaded important adjacent blood vessels that are beyond the criteria for borderline resectable cancers, for example greater than 180° involvement of the superior mesenteric artery.



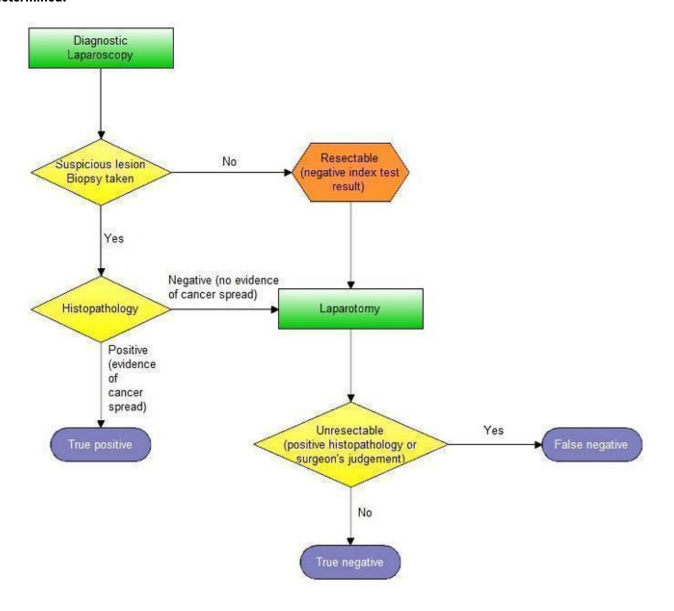
Reference standards

Confirmation of liver or peritoneal involvement histopathological examination of suspicious (liver or peritoneal) lesions obtained at diagnostic laparoscopy or laparotomy. We accepted only paraffin section histology as the reference standard. In clinical practice, depending on the urgency of the results, a frozen section biopsy may be done to obtain immediate results. However, this is always confirmed by subsequent paraffin section histology (which can take several days) because frozen section biopsy is not as reliable as paraffin section histology. We also accepted the surgeon's judgement of unresectability at laparotomy when biopsy confirmation was not possible. For example, if the tumour has invaded the adjacent blood vessels the surgeon may not resect the tumour because of the danger posed by resecting part of a large blood vessel, and so biopsy confirmation cannot be obtained.

Diagnostic laparoscopy results versus reference standard results

A schematic diagram of the results of diagnostic laparoscopy against those of histopathology or laparotomy is shown in Figure 2. Positive histopathology of a biopsy taken during diagnostic laparoscopy confirms the presence of cancer (true positive). Thus, the index test and the reference standard are one and the same if there is positive histopathology after laparoscopy. As a result, false positives are not possible, and there is no sampling error associated with specificity because it is by definition equal to 1. If the histopathology is negative, the surgeon will perform a laparotomy. The cancer may be resectable with curative intent (true negative) or may not be resectable with curative intent (false negative) based on histopathological confirmation or the surgeon's judgement of unresectability on laparotomy if biopsy confirmation cannot be obtained.

Figure 2. Schematic diagram indicating how true-positive, false-negative, and true-negative test results were determined.





Search methods for identification of studies

We included all studies irrespective of language of publication and publication status. We obtained translations of any non-English articles.

Electronic searches

We searched the following databases until 15 May 2016.

- Cochrane Central Register of Controlled Trials (CENTRAL) in the Cochrane Library (Issue 5, 2016) (Appendix 2).
- 2. MEDLINE via PubMed (January 1946 to May 2016) (Appendix 3).
- 3. EMBASE via OvidSP (January 1947 to May 2016) (Appendix 4).
- Science Citation Index Expanded (January 1980 to May 2016) (Appendix 5).

Searching other resources

We searched the references of the included studies to identify additional studies. We also searched for articles related to the included studies by performing the 'related search' function in MEDLINE (PubMed) and EMBASE (OvidSP) and a 'citing reference' search (by searching the articles which cited the included articles) in Science Citation Index Expanded and EMBASE (OvidSP) (Sampson 2008).

Data collection and analysis

Selection of studies

Two review authors (VA and KG or AK) independently searched the references to identify relevant studies. We obtained the full texts for references considered relevant by at least one of the review authors. Two review authors screened the full-text papers against the inclusion criteria. Any differences in study selection were arbitrated by BRD.

Data extraction and management

Two review authors independently extracted the following data from each included study, resolving any differences by discussion with BRD.

· First author.

- Year of publication.
- Study design (prospective or retrospective; cross-sectional studies or randomised clinical trials).
- Inclusion and exclusion criteria for individual studies.
- Total number of participants.
- · Number of females.
- Average age of the participants.
- Type of cancer (i.e. head and neck of pancreas, body and tail of pancreas, ampullary cancers, cancer of the lower end of the bile duct).
- Criteria for unresectability at diagnostic laparoscopy (index test) and at laparotomy (reference standard).
- Preoperative tests carried out prior to diagnostic laparoscopy.
- Description of the index test.
- · Reference standard.
- Number of true positives, true negatives, and false negatives.
- Complications of diagnostic laparoscopy.

The unit of analysis was the participant, meaning that if multiple metastases were found in a participant with a negative index test, the number of false negatives was considered to be one. This is because it is the presence rather than the number of metastases which is important in determining the curative resectability of patients. We considered participants with uninterpretable diagnostic laparoscopy results (no matter the reason given for lack of interpretation) as negative for the test since in clinical practice laparotomy would be carried out on these patients. However, we included such participants in the analysis only if the results of laparotomy were available. We sought further information from study authors if necessary.

Assessment of methodological quality

Two review authors (VA and KG) independently assessed study quality using the QUADAS-2 assessment tool (Whiting 2011). Any differences were resolved by BRD. The criteria used to classify the different studies are shown in Table 1. We considered studies which were classified as 'low risk of bias' and 'low concern' in all the domains as having high methodological quality.



Figure 3. Risk of bias and applicability concerns summary: review authors' judgements about each domain for each included study.

	Risk of Bias		Applicability Con			cerns			
	Patient Selection	Index Test	Reference Standard	Flow and Timing	Patient Selection	Index Test	Reference Standard		
Ahmed 2006	•	•	?		•	•	•		
Arnold 1999	•	•	?	•	•	•	•		
Arnold 2001a	?	•	?	?	•	•	•		
Beenen 2014		•	?	•		•	•		
Brooks 2002	?	•	?	?	•	•	•		
Contreras 2009	•	•	?		•	•	•		
Fernandez-Castillo 1995	•	•	?		•	•	•		
John 1995	?	•	?	?	•	•	•		
Kishiwada 2002	•	•	?	•	•	•	•		
Lavy 2012	•	•	?	?	•	•	•		
Menack 2001	?	•	?	?	•	•	•		
Merchant 1998	•	•	?	?	•	•	•		
Reddy 1999	•	•	?		•	•	•		
Reed 1997	?	•	?	?	•	•	•		
Shah 2008	•	•	?		•	•	•		
Warshaw 1986	?	•	?		•	•	•		
- High	?	Uncle	ear		•	Low			



Statistical analysis and data synthesis

The index test used was diagnostic laparoscopy with biopsy and histopathological confirmation. For the reason mentioned earlier, false positives were not possible. We therefore performed meta-analysis of only sensitivities by using a univariate randomeffects logistic regression model. The analysis was done using the NLMIXED procedure in SAS version 9.2 (SAS Institute Inc, Cary, North Carolina, USA) (Appendix 6). We used the ESTIMATE statement in NLMIXED to obtain the negative likelihood ratio by using a function of the estimated summary sensitivity and a specificity of 1. The median pre-test probability of unresectability was calculated from the pre-test probabilities of the included studies. We calculated the proportion of participants classified as having resectable disease by CT scanning and diagnostic laparoscopy who were actually found to be unresectable at laparotomy (post-test probability) using the median pre-test probability and the negative likelihood ratio (see Appendix 7 for details). The difference in the unresectability proportions (post-test probability minus pre-test probability) gave the overall added value of diagnostic laparoscopy compared to the standard practice of CT scan staging alone.

Investigations of heterogeneity

We planned to explore heterogeneity by using the different sources of heterogeneity as covariate(s) in the regression model. However,

this was not possible because the information was either not available or was the same in all the studies.

Sensitivity analyses

We did not plan any sensitivity analyses.

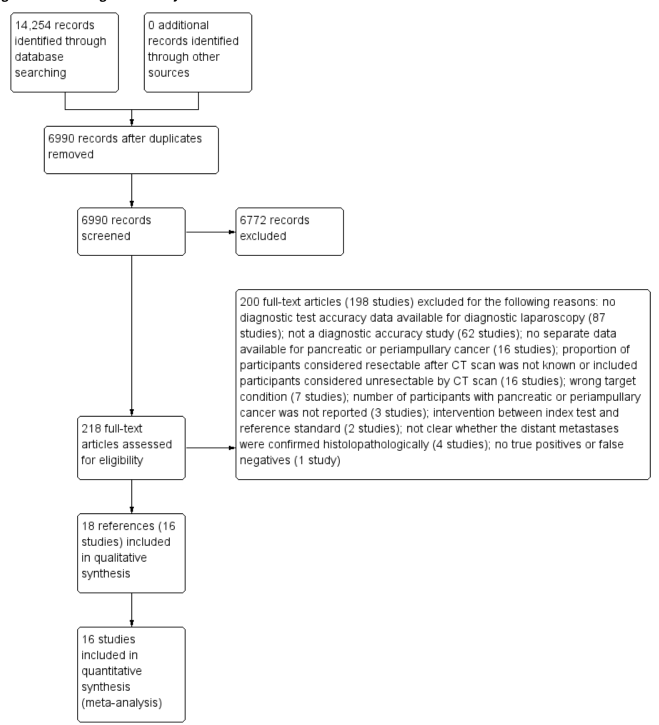
RESULTS

Results of the search

We identified a total of 14,254 references through the electronic searches of the Cochrane Upper Gastrointestinal and Pancreatic Diseases Group Controlled Trials Register and CENTRAL (n = 191), MEDLINE (n = 5228), EMBASE (n = 4460), and Science Citation Index (n = 4375). Figure 4 shows the flow of references through the selection process. We excluded 7264 duplicates and clearly irrelevant references through reading the abstracts. We retrieved 213 references for further assessment. We identified no references through scanning reference lists of the identified studies. Of the 213 references, we excluded 194 for the reasons listed in the Characteristics of excluded studies table. In one study (Hashimoto 2015), all 11 participants who underwent diagnostic laparoscopy and laparotomy had resectable pancreatic cancers. There were therefore no true positives and false negatives for estimation of sensitivity, and we excluded this study from the review. We included 18 references of 16 studies.



Figure 4. Flow diagram of study selection.

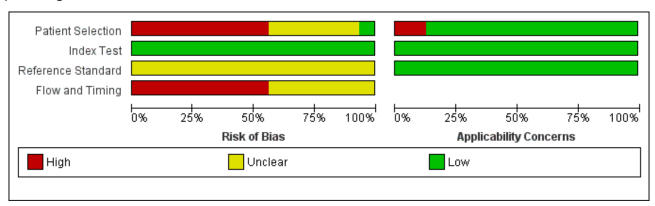


Methodological quality of included studies

The methodological quality of the included studies is shown in the Characteristics of included studies table, Figure 5, and Figure 3.



Figure 5. Risk of bias and applicability concerns graph: review authors' judgements about each domain presented as percentages across included studies.



There was a high risk of bias regarding the selection of participants in most studies (Ahmed 2006; Arnold 1999; Arnold 2001a; Beenen 2014; Brooks 2002; Contreras 2009; John 1995; Kishiwada 2002; Lavy 2012; Menack 2001; Merchant 1998; Reddy 1999; Reed 1997; Shah 2008; Warshaw 1986). This was because the studies did not explicitly state whether a consecutive or random sample of patients was recruited or whether they had made inappropriate exclusions. Only one study had low risk of bias and low applicability concerns regarding the selection of participants (Fernandez-Castillo 1995).

There were no risk of bias issues or concerns regarding applicability of the index test in any of the studies, as was anticipated (Table 1).

As anticipated, it proved impossible to determine whether an appropriate reference standard was used. This is because even in the presence of predefined criteria for unresectability, it may not be ethical to biopsy and confirm that the tumour has invaded the blood vessels because of the risk of major bleeding. Thus it was not possible to determine whether the cancer was truly unresectable. None of the studies reported whether the margins of the resected lesions were clear of cancer. It was therefore not possible to determine whether the cancer was truly resectable with curative intent.

None of the studies reported the time interval between diagnostic laparoscopy and laparotomy. In addition, many studies had excluded some patients inappropriately. All of the studies were therefore at unclear or high risk of bias in the flow and timing domain.

Findings

All of the included studies assessed pancreatic or periampullary cancer. The 16 included studies involved a total of 1146 participants (Data and analyses). The age of participants in the included studies ranged between 15 and 87 years. Studies that provided demographic details of participants reported roughly equal numbers of males and females. Seven studies included only people with pancreatic cancer (Ahmed 2006; Arnold 2001a; Contreras 2009;

Fernandez-Castillo 1995; Kishiwada 2002; Lavy 2012; Warshaw 1986), and two studies included only people with periampullary malignancies (Beenen 2014; Brooks 2002). The remaining studies did not provide information regarding the specific type of cancer they considered.

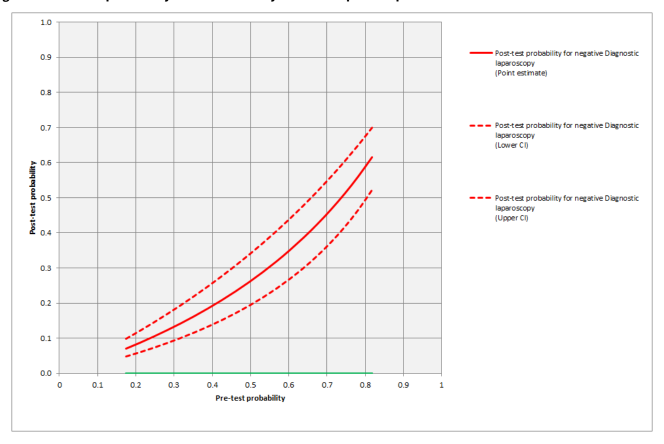
The details of the CT scan; other tests the participants underwent in addition to the CT scan; probability of CT resectable disease identified as unresectable by diagnostic laparoscopy or laparotomy (pre-test probability); reasons for CT resectable disease identified as unresectable by diagnostic laparoscopy; probability of CT and diagnostic laparoscopy resectable disease identified as unresectable at laparotomy (post-test probability); and the reasons for CT and diagnostic laparoscopy resectable disease identified as unresectable at laparotomy are all shown in Table 2.

The pre-test probability of unresectability (due to distant metastases or local infiltration) after CT scanning ranged from 17.4% to 82% in the included studies. The median pre-test probability was 41.4%, meaning that a person that was said to be resectable on CT scanning still had a 41.4% chance that their cancer would be unresectable. Visual inspection of the data in Table 2 did not suggest a relationship between the type of CT scan (such as helical CT or multi-detector row CT, with or without a pancreatic protocol) or date of publication and the pre-test probability of unresectable disease.

The summary estimate of sensitivity was 64.4% (95% confidence interval (CI) 50.1 to 76.6), and the summary negative likelihood ratio was 0.36 (95% CI 0.24 to 0.52). Using the median pretest probability of unresectable disease of 0.414, the post-test probability of unresectable disease for participants with a negative test result was 0.20 (95% CI 0.15 to 0.27). This means that if a person is said to have resectable disease after diagnostic laparoscopy (and a CT scan), there is a 20% chance that their cancer will be unresectable. The post-test probability of unresectable disease is shown at different pre-test probabilities of unresectable disease in Figure 6.



Figure 6. Post-test probability of unresectability for various pre-test probabilities.



None of the studies reported any complications related to diagnostic laparoscopy. In some instances diagnostic laparoscopy provided an inconclusive result, that is it was unclear whether the participant had resectable or unresectable disease. Eight studies reported drop-out rates of: 37.3% (Ahmed 2006), 29.8% (Arnold 1999), 36.1% (Beenen 2014), 67.5% (Contreras 2009), 4.4% (Fernandez-Castillo 1995), 10.6% (Merchant 1998), 1.0% (Reddy 1999), and 61.2% (Shah 2008). In four of these studies the participants underwent laparotomy directly (Ahmed 2006; Beenen 2014; Contreras 2009; Shah 2008), and there was no indication of the selection criteria used for participants who had diagnostic laparoscopy. The other studies did not report drop-out rates.

A subgroup analysis of studies that included only participants with pancreatic cancer gave a summary sensitivity of 67.9% (95% CI 41.1% to 86.5%). The summary negative likelihood ratio was 0.32 (95% CI 0.15 to 0.68). The median pre-test probability of unresectability was 40.0% in this subgroup of studies. Using this pre-test probability, the post-test probability of unresectable disease after negative diagnostic laparoscopy was 0.18 (95% CI 0.31 to 0.92).

We also performed a post hoc meta-regression of studies published before and after the year 2000, to test whether the sensitivity of diagnostic laparoscopy was different in the last decade, because major technological innovations in CT scans such as helical CT scans and multi-slice CT scans became widely available in the last decade. The likelihood ratio test comparing the model with and without this covariate gave a P value of 1.0, indicating no evidence

of a statistically significant difference in sensitivity between studies published before or after the year 2000.

We found an inconsistency in one study between the results reported in the main text of the study and a flow diagram which summarised the results (Kishiwada 2002). In our previous review we investigated the effect of this inconsistency by conducting a sensitivity analysis, which showed no change in the estimates of the summary sensitivity and the confidence intervals (Allen 2013). In another sensitivity analysis, we imputed missing data as false-negative results (that is diagnostic laparoscopy incorrectly classified unresectable disease as resectable in all the missing participants) (Allen 2013). We have not presented the results of the first sensitivity analysis in this update since only participant was misclassified, and the impact on results was negligible. We did not perform the second sensitivity analysis since the reasons for not performing diagnostic laparoscopy were not reported, and it is unlikely that all the participants in diagnostic laparoscopy would have false-negative results.

DISCUSSION

Summary of main results

We have summarised the results in Summary of findings 1. The addition of diagnostic laparoscopy to CT scanning decreases the probability of unresectable disease from 41% to 20%. This means that for every 100 patients who receive a CT scan followed by diagnostic laparoscopy, 21 patients (41 minus 20) will avoid major laparotomy compared to with CT scanning alone. Although this review included studies which were more than



10 years old, with improvements in CT scanning possible over this period, the probability of unresectability was high (63.2%) even after multi-detector row CT using a pancreatic protocol (Table 2). Diagnostic laparoscopy can either be performed as a separate procedure or immediately prior to major laparotomy as part of a larger procedure. These two different approaches have distinct advantages and disadvantages. The advantages of performing diagnostic laparoscopy as part of a larger procedure are that the patient needs only one hospital admission and one general anaesthetic. However, if the patient is diagnosed as having unresectable disease at laparoscopy and the subsequent laparotomy is then cancelled, it means that operation theatre time is wasted. It is also not possible to use paraffin section, the gold standard test, to confirm a histological diagnosis of cancer if diagnostic laparoscopy is undertaken as part of a larger procedure. If laparoscopy is performed as a separate diagnostic procedure, the patient must undergo the burden of two separate hospital admissions and anaesthetics, but no operation theatre time will be wasted if they are found to have unresectable disease. The time delay between the two separate procedures also allows the use of paraffin sections.

We found no complications related to diagnostic laparoscopy in this systematic review, however the literature reports an injury rate of 0.23% involving major blood vessels or the bowel (Azevedo 2009). This indicates that diagnostic laparoscopy should only be performed by appropriately trained healthcare professionals with expertise in the conduct of diagnostic laparoscopy and biopsy during diagnostic laparoscopy.

Strengths and weaknesses of the review

A strength of this review is that we placed no restrictions on the language of publication and conducted a comprehensive search. We avoided the use of search filters and undertook additional searches to find related articles. We also performed a citation search. We therefore minimised the risk of missing relevant studies. Little is known about the mechanisms of publication bias for diagnostic accuracy studies, and so it is not possible to estimate the impact of unpublished studies on our findings. Nevertheless, the studies included in this systematic review are likely to be the majority of studies that provide evidence on this topic. Another strength of this review is that we used a recommended approach for meta-analysis.

Our review has some weaknesses. Firstly, our findings are based on studies with low methodological quality, and there was considerable between-study heterogeneity. There were betweenstudy differences in the conduct and interpretation of diagnostic laparoscopy (in terms of what constitutes a suspicious lesion) and differences in the assessment of resectability on laparotomy. Despite the observed differences in the conduct and interpretation of diagnostic laparoscopy, the procedure appeared to decrease the number of unnecessary laparotomies in 15 of the 16 included studies. With regards to methodological quality, the presence of selection bias may raise doubts about the applicability of our findings in clinical practice. Secondly, determination of unresectability on laparotomy relies on the judgement of individual surgeons, which may not have been appropriate in some of the studies. This could have caused an error in the estimation of diagnostic accuracy. Thirdly, an inappropriate delay between diagnostic laparoscopy and laparotomy can result in patients who had previously resectable cancer developing unresectable cancer because of local or distant spread. This will underestimate the accuracy of diagnostic laparoscopy. Fourthly, inappropriate exclusion of patients is likely to result in an error in the estimation of diagnostic accuracy if the excluded patients had low likelihood of unresectability or high likelihood of unresectability. We performed a sensitivity analysis imputing the results according to the worst-case scenario, that is as false negatives. As mentioned earlier, indeterminate results at diagnostic laparoscopy will result in the patients undergoing laparotomy.

We were able to identify one previous systematic review on this topic (Chang 2009). Despite the inclusion of studies in which histopathological confirmation of suspicious lesions was not obtained, and the lack of meta-analysis on the diagnostic accuracy of diagnostic laparoscopy, the authors of the review suggested that diagnostic laparoscopy decreases unnecessary laparotomy by 4% to 36% and that diagnostic laparoscopy has a role in staging pancreatic cancer (Chang 2009). We agree broadly with the conclusions of the authors of the identified systematic review (Chang 2009).

Applicability of findings to the review question

This review is only applicable to people with pancreatic and periampullary cancer who have had a CT scan which demonstrated resectable disease prior to diagnostic laparoscopy. This review is also applicable only when the interval between diagnostic laparoscopy and laparotomy is sufficient to obtain histopathology results but not too long for the cancer to spread. Diagnostic laparoscopy appears to be beneficial in avoiding unnecessary laparotomies, and the morbidity associated with diagnostic laparoscopy is low. Cost-effectiveness needs to be formally assessed to inform clinical and policy decision making in statefunded health care.

AUTHORS' CONCLUSIONS

Implications for practice

Although the methodological quality of the evidence was limited, diagnostic laparoscopy appears to be useful in decreasing the proportion of people with pancreatic and periampullary cancer that were found to have resectable disease on CT scanning who will undergo unnecessary laparotomy.

Implications for research

- Well-designed diagnostic test accuracy studies are needed to reliably estimate the accuracy of diagnostic laparoscopy. Comparison with positron emission tomography (PET) scanning, endoscopic ultrasound (EUS), and laparoscopic ultrasound may further demonstrate the value of diagnostic laparoscopy in staging pancreatic and periampullary cancers.
- 2. The conclusion of this study needs regular review as the quality of CT scanning improves, and diagnostic laparoscopy should be compared with other tests for staging pancreatic and periampullary cancers.
- 3. Cost-effectiveness studies should be undertaken to determine whether diagnostic laparoscopy should be routinely performed in state-funded clinical practice.



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CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Ahmed 2006 Study characteristics			
Patient sampling	Sample size: 37		
r atient sampling	Females: Not stated		
	Age: Not stated		
Patient characteristics and setting	Patients with potentially cinoma (after CT scan)	resectable, histologically	confirmed pancreatic adenocar-
	Setting: Surgical centre i	n the USA	
Index tests	Diagnostic laparoscopy		
	sectable if laparoscopic e	xamination revealed peri oh node involvement, or to	dered locally advanced and unre- toneal or liver metastasis, coeliac umour invasion or encasement of
Target condition and reference standard(s)	Target condition: Unrese	ctability	
			no evidence of metastases on la- nation of spread for patients with
	sectable if laparoscopic e	xamination revealed perion node involvement, or to	dered locally advanced and unre- toneal or liver metastasis, coeliac umour invasion or encasement of
Flow and timing	Number of indeterminate able: Not stated	es for whom the results of	reference standard were avail-
	Number of patients who	were excluded from the ar	nalysis: 22 (37.3%)
Comparative			
Notes			
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			



Ahmed 2006 (Continued)				
Was a consecutive or random sample of patients enrolled?	Unclear			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate exclusions?	No			
		High	Low	
DOMAIN 2: Index Test All tests				
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes			
		Low	Low	
DOMAIN 3: Reference Standard				
Is the reference standards likely to correctly classify the target condition?	Unclear			
Were the reference standard results interpreted without knowledge of the results of the index tests?	No			
		Unclear	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval between index test and reference standard?	Unclear			
Did all patients receive the same reference standard?	No			
Were all patients included in the analysis?	No			
		High		

Arnold 1999

Study characteristics	
Patient sampling	Sample size: 33
	Females: Not stated
	Age: Not stated
Patient characteristics and setting	Patients with potentially resectable pancreatic adenocarcinoma (after CT scan)
	Setting: Germany (setting not clear)
Index tests	Diagnostic laparoscopy



Arnold 1999 (Continued)			
	Criteria for positive diaş metastases	gnosis: Biopsies of le	sions suspicious of
Target condition and reference standard(s)	Target condition: Unres	sectability	
	Reference standard: La metastases on laparoso mation of spread for pa	copy; biopsy with his	tolopathological confir-
	Criteria for positive dia	gnosis: Not stated	
Flow and timing	Number of indeterminates for whom the results of reference standard were available: Not stated		
	Number of patients who	o were excluded from	n the analysis: 14 (29.8%)
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?	No		
		High	Low
DOMAIN 2: Index Test All tests			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
		Low	Low
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Unclear		
Were the reference standard results interpreted without knowledge of the results of the index tests?	No		
		Unclear	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Unclear		
Did all patients receive the same reference standard?	No		



Arnold 1999 (Continued)

Were all patients included in the analysis?

No

High

Arnold 2001a

Study characteristics					
Patient sampling	Sample size: 61 Females: Not stated				
	Age: Not stated				
Patient characteristics and setting	Patients with potentially resectable pancreatic adenocarcinoma (after CT scan)				
	Setting: Germany (setting not clear)				
Index tests	Diagnostic laparoscopy				
	Criteria for positive diagnosis: Biopsies of lesions suspicious of metastases				
Target condition and reference standard(s)	Target condition: Unresectability				
	Reference standard: Laparotomy for patients with no evidence of metastases on laparoscopy; biopsy with histolopathological confirmation of spread for patients with suspected metastases				
	Criteria for positive diagnosis: Not stated				
Flow and timing	Number of indeterminates for whom the results of reference standard were available: Not stated				
	Number of patients who were excluded from the analysis: Not stated				
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				

Unclear

Unclear

Low

Did the study avoid inappropriate exclusions?



Arnold 2001a (Continued)

DOMAIN 2: Index Test All tests

Were the index test results interpreted without knowledge of Yes the results of the reference standard?

	Low	Low
DOMAIN 3: Reference Standard		

Is the reference standards likely to correctly classify the target condition?

Were the reference standard results interpreted without

knowledge of the results of the index tests?

No

DOMAIN 4: Flow and Timing

Was there an appropriate interval between index test and reference standard?

Did all patients receive the same reference standard?

No

Were all patients included in the analysis?

Unclear

Unclear

Beenen 2014

Study characteristics	
Study Characteristics	
Patient sampling	Sample size: 131
	Females: Not stated
	Age: Not stated
Patient characteristics and setting	Patients with CT and ultrasound resectable periampullary cancer
	Setting: Secondary/tertiary care, the Netherlands
Index tests	Diagnostic laparoscopy
	Criteria for positive diagnosis: Biopsy confirmation of suspicious lesions
Target condition and reference standard(s)	Target condition: Unresectability
	Reference standard: Laparotomy
	Criteria for positive diagnosis: Locally advanced pancreatic cancer or metastatic pancreatic cancer
Flow and timing	Number of indeterminates for whom the results of reference standard were available: 0



eenen 2014 (Continued)	Number of patients who were excluded from the analysis: 74 (36.1%)			
Comparative				
Notes				
Methodological quality				
Item	Authors' judge- ment	Risk of bias	Applicability con cerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	No			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate exclusions?	Unclear			
		High	High	
DOMAIN 2: Index Test All tests				
Were the index test results interpreted without knowledge of the results of the reference standard?	No			
		Low	Low	
DOMAIN 3: Reference Standard			,	
Is the reference standards likely to correctly classify the target condition?	Unclear			
Were the reference standard results interpreted without knowledge of the results of the index tests?	No			
		Unclear	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval between index test and reference standard?	Unclear			
Did all patients receive the same reference standard?	Yes			
Were all patients included in the analysis?	No			
		High		
rooks 2002				
Study characteristics				
Patient sampling Sample size: 14-	4			



Brooks 2002 (Continued)			
	Females: Not stated		
	Age: Not stated		
Patient characteristics and setting	Patients with potentially cancer	resectable periampullary	carcinoma other than pancreatic
	Setting: Surgical centre in	the USA	
Index tests	Diagnostic laparoscopy		
	paroscopy or laparotomy	if they were found to have stant nodal involvement,	ed unresectable at diagnostic la- ve histologically proved peritoneal arterial involvement, or local ex-
Target condition and reference standard(s)	Target condition: Unrese	ctability	
			no evidence of metastases on la- mation of spread for patients with
	paroscopy or laparotomy	if they were found to have stant nodal involvement,	ed unresectable at diagnostic la- ve histologically proven peritoneal arterial involvement, or local ex-
Flow and timing	Number of indeterminate able: 10 (6.9%)	s for whom the results of	reference standard were avail-
	Number of patients who	were excluded from the a	nalysis: Not stated
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		
		Unclear	Low
DOMAIN 2: Index Test All tests			
Were the index test results interpreted with-	Yes		
out knowledge of the results of the reference standard?			



Brooks 2002 (Continued)

DOMAIN	3: Reference	Standard
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Is the reference standards likely to correctly classify the target condition?

Unclear

Were the reference standard results interpreted without knowledge of the results of the index tests?

No

		Unclear	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval between index test and reference standard?	Unclear			
Did all patients receive the same reference standard?	No			
Were all patients included in the analysis?	Unclear			
		Unclear		

Contreras 2009

Study characteristics	
Patient sampling	Sample size: 25
	Females: 12 (32.5%)
	Age: 68 years
Patient characteristics and setting	Patients with potentially resectable pancreatic adenocarcinoma (after CT scan)
	Setting: Surgical referral centre in the USA
Index tests	Diagnostic laparoscopy
	Criteria for positive diagnosis: Biopsies of lesions suspicious of metastases
Target condition and reference standard(s)	Target condition: Unresectability
	Reference standard: Laparotomy for patients with no evidence of metastases on laparoscopy; biopsy with histolopathological confirmation of spread for patients with suspected metastases
	Criteria for positive diagnosis: Not stated
Flow and timing	Number of indeterminates for whom the results of reference standard were available: Not stated
	Number of patients who were excluded from the analysis: 52 (67.5%)



Contreras 2009 (Continued)

Notes

Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability con cerns
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?	No		
		High	Low
DOMAIN 2: Index Test All tests			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
		Low	Low
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Unclear		
Were the reference standard results interpreted without knowledge of the results of the index tests?	No		
		Unclear	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Unclear		
Did all patients receive the same reference standard?	No		
Were all patients included in the analysis?	No		
		High	
Fernandez-Castillo 1995			
Study characteristics			
Patient sampling	Sample size: 109		
	Females: Not stated		
	Age: Not stated		



ernandez-Castillo 1995 (Continued)			
Patient characteristics and setting	Patients with potential CT scan) without gastri		atic adenocarcinoma (on
	Setting: Surgical centre	in the USA	
Index tests	Diagnostic laparoscopy	,	
	Criteria for positive diaş tases	gnosis: Biopsies of le	esions suspicious of metas-
Target condition and reference standard(s)	Target condition: Unres	sectability	
	Reference standard: La metastases on laparoso tion of spread for patie	copy; biopsy with his	stolopathological confirma
	Criteria for positive dia	gnosis: Not stated	
Flow and timing	Number of indetermina were available: not stat		sults of reference standard
	Number of patients wh	o were excluded fror	n the analysis: 5 (4.2%)
Comparative			
Notes			
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability con- cerns
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		
		High	Low
DOMAIN 2: Index Test All tests			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
		Low	Low
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Unclear		
Were the reference standard results interpreted without knowledge of the results of the index tests?	No		
		Unclear	Low



Fernandez-Castillo 1995 (Continued)

DOMAIN 4: Flow and Timing

Was there an appropriate interval between index test and reference standard?	Unclear
Did all patients receive the same reference standard?	No
Were all patients included in the analysis?	No
	High

John 1995

Study characteristics			
Patient sampling	Sample size: 40		
	Females: 22 (100%)		
	Age: 59 years		
Patient characteristics and setting	Patients with potentially resectable pancreatic or periampullary carcinoma		
	Setting: Tertiary referral centre in the UK		
Index tests	Diagnostic laparoscopy		
	Criteria for positive diagnosis: Biopsies of lesions suspicious of metastases		
Target condition and reference standard(s)	Target condition: Unresectability		
	Reference standard: Laparotomy for patients with no evidence of metastases on laparoscopy; biopsy with histolopathological confirmation of spread for patients with suspected metastases		
	Criteria for positive diagnosis: The criteria used to define primary tumour advancement and locoregional unresectability were as follows:		
	1. tumour size of 5 cm or greater;		
	2. extrapancreatic invasion of adjacent tissues (i.e. duodenum, stomach, common bile duct, retroperitoneum); and		
	3. occlusion or stenosis of the portal or superior mesenteric veins, or major branches of the coeliac trunk or superior mesenteric artery.		
Flow and timing	Number of indeterminates for whom the results of reference standard were available: Not stated		
	Number of patients who were excluded from the analysis: Not stated		
Comparative			
Notes			
Methodological quality			
Item	Authors' judgement Risk of bias Applicability concerns		



J	ol	hn	1995	(Continued)
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DOMAIN	1: Pa	tient Se	lection
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Was a consecutive or random sample of patients Unclear enrolled? Was a case-control design avoided? Yes Did the study avoid inappropriate exclusions? Unclear

Unclear Low

DOMAIN 2: Index Test All tests

Were the index test results interpreted without knowledge of the results of the reference standard?

Yes

Low Low

DOMAIN 3: Reference Standard

Is the reference standards likely to correctly classify the target condition?

Unclear

Were the reference standard results interpreted without knowledge of the results of the index tests?

No

Unclear Low

DOMAIN 4: Flow and Timing

Was there an appropriate interval between index Unclear test and reference standard?

Did all patients receive the same reference standard?

No

Were all patients included in the analysis?

Unclear

Unclear

Kishiwada 2002

Study characteristics

Patient sampling Sample size: 16 Females: Not stated Age: Not stated Patient characteristics and setting Patients with potentially resectable pancreatic cancer (only patients with tumours more than 2 cm in diameter were subject to diagnostic la-

paroscopy)



Kishiwada 2002 (Continued)	Setting: Surgical centre	in Japan		
Index tests	Diagnostic laparoscopy			
	Criteria for positive diag	gnosis: Biopsies of les	sions suspicious of metas-	
Target condition and reference standard(s)	Target condition: Unres	sectability		
	Reference standard: La metastases on laparosc tion of spread for patier	copy; biopsy with hist	olopathological confirma-	
	Criteria for positive diag	gnosis: Not stated		
Flow and timing	Number of indetermina were available: Not stat		ults of reference standard	
	Number of patients who	o were excluded from	the analysis: Not stated	
Comparative				
Notes				
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability con- cerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	No			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate exclusions?	Unclear			
		High	High	
DOMAIN 2: Index Test All tests				
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes			
		Low	Low	
DOMAIN 3: Reference Standard				
Is the reference standards likely to correctly classify the target condition?	Unclear			
Were the reference standard results interpreted without knowledge of the results of the index tests?	No			
		Unclear	Low	
DOMAIN 4: Flow and Timing				



(ishiwada 2002 (Continued)				
Was there an appropriate interval between index test and reference standard?	Unclear			
Did all patients receive the same reference standard?	No			
Were all patients included in the analysis?	No			
	High			
avy 2012				
Study characteristics				
Patient sampling	Sample size: 52			
	Females: Not stated			
	Age: Not stated			
Patient characteristics and setting	Patients with potentially resectable pancreatic adenocarcinoma (after CT scan and EUS)			
	Setting: Surgical centre in Israel			
Index tests	Diagnostic laparoscopy			
	Criteria for positive diagnosis: Biopsies of lesions suspicious of metastases			
Target condition and reference standard(s)	Target condition: Unresectability			
	Reference standard: Laparotomy for patients with no evidence of metastases on laparoscopy; biopsy with histopathological confirmation of spread for patients with suspected metastases			
	Criteria for positive diagnosis: Not stated			
Flow and timing	Number of indeterminates for whom the results of reference standard were available: Not stated			
	Number of patients who were excluded from the analysis: Not stated			
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			



Lavy 2012 (Co	ntinued)
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Did t	he study	y avoid inap	propriate exc	:lusions?	Yes
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Did the study avoid inappropriate exclusions?	Yes			
		Low	Low	
DOMAIN 2: Index Test All tests				
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes			
		Low	Low	
DOMAIN 3: Reference Standard				
Is the reference standards likely to correctly classify the target condition?	Unclear			
Were the reference standard results interpreted without knowledge of the results of the index tests?	No			
		Unclear	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval between index test and reference standard?	Unclear			
Did all patients receive the same reference standard?	No			
Were all patients included in the analysis?	Unclear			
		Unclear		

Menack 2001

Study characteristics	
Patient sampling	Sample size: 27
	Females: 10 (100%)
	Age: 66 years
Patient characteristics and setting	Patients with potentially resectable pancreatic or periampullary cancer (after CT scan)
	Setting: Surgical centre in the USA
Index tests	Diagnostic laparoscopy
	Criteria for positive diagnosis: Biopsies of lesions suspicious of metastases
Target condition and reference standard(s)	Target condition: Unresectability
	Reference standard: Laparotomy for patients with no evidence of metas- tases on laparoscopy; biopsy with histolopathological confirmation of spread for patients with suspected metastases



Menack 2001 (Continued)				
	Criteria for positive diagnosis: Patients were considered unresectable if they had histologically proven metastatic disease or carcinomatosis			
Flow and timing	Number of indeterminates for whom the results of reference standar were available: Not stated			
	Number of patients who	were excluded from	the analysis: Not stated	
Comparative				
Notes				
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability con- cerns	
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			
		Unclear	Low	
DOMAIN 2: Index Test All tests				
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes			
		Low	Low	
DOMAIN 3: Reference Standard				
Is the reference standards likely to correctly classify the target condition?	Unclear			
Were the reference standard results interpreted without knowledge of the results of the index tests?	No			
		Unclear	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval between index test and reference standard?	Unclear			
Did all patients receive the same reference standard?	No			
Were all patients included in the analysis?	Unclear			
		Unclear		



Merc	hant	1998
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Study characteristics				
Patient sampling	Sample size: 303			
	Females: Not stated			
	Age: Not stated			
Patient characteristics and setting	Patients with potentially resectable pancreatic or periampullary cancer (after CT scan)			
	Setting: Surgical centre in the USA			
Index tests	Diagnostic laparoscopy			
	Criteria for positive diagnosis: Biopsies of lesions suspicious of metastases			
Target condition and reference standard(s)	Target condition: Unresectability			
	Reference standard: Laparotomy for patients with no evidence of metastases on laparoscopy; biopsy with histolopathological confirmation of spread for patients with suspected metastases			
	Criteria for positive diagnosis: Unresectable if one or more of the following were confirmed histopathologically:			
	1. hepatic, serosal/peritoneal, or omental metastases;			
	2. extrapancreatic extension of tumour (i.e. mesocolic involvement);			
	3. celiac or high portal nodal involvement by tumour; and			
	4. invasion or encasement of the coeliac axis, hepatic artery, or superior mesenteric artery.			
Flow and timing	Number of indeterminates for whom the results of reference standard were available: Not stated			
	Number of patients who were excluded from the analysis: 36 (10.6%)			
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			

High

No

Low

Did the study avoid inappropriate exclusions?



Merchant 1998 (Continued)

DOMAIN 2: Index Test All tests

Were the index test results interpreted without knowledge of the results of the reference standard?

Yes

		Low	Low
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Unclear		
Were the reference standard results interpreted without knowledge of the results of the index tests?	No		

		Unclear	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval between index test and reference standard?	Unclear			
Did all patients receive the same reference standard?	No			
Were all patients included in the analysis?	Unclear			
		Unclear		

Reddy 1999

Study characteristics	
Patient sampling	Sample size: 98
	Females: 47 (49%)
	Age: 65 years
Patient characteristics and setting	Patients with potentially resectable pancreatic cancer (on CT scan)
	Setting: Surgical centre in the USA
Index tests	Diagnostic laparoscopy
	Criteria for positive diagnosis: Biopsies of lesions suspicious of metastases
Target condition and reference standard(s)	Target condition: Unresectability
	Reference standard: Laparotomy for patients with no evidence of metastases on laparoscopy; biopsy with histolopathological confirmation of spread for patients with suspected metastases



Reddy 1999 (Continued)	Criteria for positive dia	gnosis: Not stated	
Flow and timing	Number of indeterminates for whom the results of reference standard were available: Not stated		
	Number of patients wh	o were excluded from	m the analysis: 1 (1%)
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?	No		
		High	Low
DOMAIN 2: Index Test All tests			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
		Low	Low
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Unclear		
Were the reference standard results interpreted without knowledge of the results of the index tests?	No		
		Unclear	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Unclear		
Did all patients receive the same reference standard?	No		
Were all patients included in the analysis?	No		
		High	



Study characteristics			
Patient sampling	Sample size: 11		
	Females: Not stated		
	Age: Not stated		
Patient characteristics and setting	Patients with potentially resectable pancreatic cancer (on		
	Setting: Surgical centre	in the USA	
Index tests	Diagnostic laparoscopy		
	Criteria for positive diag metastases	gnosis: Biopsies of le	sions suspicious of
Target condition and reference standard(s)	Target condition: Unres	ectability	
	Reference standard: La metastases on laparosc mation of spread for pa	opy; biopsy with his	tolopathological confir-
	Criteria for positive diag	nosis: Not stated	
Flow and timing	Number of indeterminates for whom the results of reference standard were available: Not stated		
	Number of patients who	were excluded fror	n the analysis: Not stated
Comparative			
Notes			
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability con- cerns
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		
		Unclear	Low
DOMAIN 2: Index Test All tests			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
		Low	Low



Reed 1997	(Continued)
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Is the reference standards likely to correctly classify the target condition?

Unclear

Were the reference standard results interpreted without knowledge of the results of the index tests?

Nο

		Unclear	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval between index test and reference standard?	Unclear			
Did all patients receive the same reference standard?	No			
Were all patients included in the analysis?	Unclear			
		Unclear		

Shah 2008

Study characteristics	
Patient sampling	Sample size: 19
	Females: Not stated
	Age: Not stated
Patient characteristics and setting	Patients with potentially resectable pancreatic cancer (on CT scan)
	Setting: Surgical centre in the USA
Index tests	Diagnostic laparoscopy
	Criteria for positive diagnosis: Biopsies of lesions suspicious of metastases
Target condition and reference standard(s)	Target condition: Unresectability
	Reference standard: Laparotomy for patients with no evidence of metastases on laparoscopy; biopsy with histolopathological confirmation of spread for patients with suspected metastases
	Criteria for positive diagnosis: Not stated
Flow and timing	Number of indeterminates for whom the results of reference standard were available: Not stated
	Number of patients who were excluded from the analysis: 30 (61.2%)
Comparative	
Notes	

Methodological quality



Shah 2008 (Continued)

Item	Authors' judgement	Risk of bias	Applicability con- cerns
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		
		High	Low
DOMAIN 2: Index Test All tests			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
		Low	Low
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Unclear		
Were the reference standard results interpreted without knowledge of the results of the index tests?	No		
		Unclear	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Unclear		
Did all patients receive the same reference standard?	No		
Were all patients included in the analysis?	No		
		High	

Warshaw 1986

Maisliam 1300	
Study characteristics	
Patient sampling	Sample size: 40
	Females: Not stated
	Age: Not stated
Patient characteristics and setting	Patients with potentially resectable pancreatic adenocarcinoma (after CT scan)
	Setting: Surgical centre in the USA



Diagnostic laparoscopy		
Criteria for positive diag metastases	gnosis: Biopsies of les	sions suspicious of
Target condition: Unres	ectability	
metastases on laparoso	opy; biopsy with hist	colopathological confir-
Criteria for positive dia	gnosis: Not stated	
		ults of reference stan-
Number of patients who	o were excluded from	the analysis: Not stated
Authors' judgement	Risk of bias	Applicability con- cerns
Unclear		
Yes		
Unclear		
	Unclear	Low
Yes		
	Low	Low
Unclear		
No		
	Unclear	Low
Unclear		
	Criteria for positive diag metastases Target condition: Unrest Reference standard: La metastases on laparosc mation of spread for patients of indeterminated dard were available: No Number of patients who Number of Patients Number	Target condition: Unresectability Reference standard: Laparotomy for patient metastases on laparoscopy; biopsy with hist mation of spread for patients with suspected. Criteria for positive diagnosis: Not stated Number of indeterminates for whom the rest dard were available: Not stated Number of patients who were excluded from Authors' judgement Risk of bias Unclear Yes Unclear Yes Low Unclear No Unclear



Warshaw 1986 (Continued)	
Did all patients receive the same reference standard?	No
Were all patients included in the analysis?	No
	High

CT: computed tomography EUS: endoscopic ultrasound

Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion
Abdalla 2003	Insufficient diagnostic test accuracy data available for diagnostic laparoscopy
Adisa 2014	No separate data available for pancreatic or periampullary cancers
Alexakis 2015	No diagnostic test accuracy data available for diagnostic laparoscopy
Altieri 1982	Wrong target condition
Andren-Sandberg 1998	Includes participants who were considered to be unresectable by CT scan
Arnold 2001	Not a diagnostic accuracy study
Atanov 1972	No separate data available for pancreatic or periampullary cancers
Awad 1997	Includes participants who were considered to be unresectable by CT scan
Baghbanian 2013	Not clear whether histopathological confirmation of metastasis was obtained
Baghbanian 2014	Not clear whether histopathological confirmation of metastasis was obtained
Balcom 2000	Not a diagnostic accuracy study
Barabino 2011	No diagnostic test accuracy data available for diagnostic laparoscopy
Barrat 1998	No separate data available for pancreatic or periampullary cancers
Barreiro 2002	Not a diagnostic accuracy study
Barthet 2007	Not a diagnostic accuracy study
Baumgarten 1984	No diagnostic test accuracy data available for diagnostic laparoscopy
Beger 1997	Not a diagnostic accuracy study
Belagyi 2000	Not a diagnostic accuracy study
Bemelman 1995	No diagnostic test accuracy data available for diagnostic laparoscopy
Bohmig 2001	Not a diagnostic accuracy study
Borbath 2005	No diagnostic test accuracy data available for diagnostic laparoscopy



Study	Reason for exclusion
Boselli 2000	No diagnostic test accuracy data available for diagnostic laparoscopy
Bottger 1998	No diagnostic test accuracy data available for diagnostic laparoscopy
Boyce 1992	Not a diagnostic accuracy study
Caldironi 1996	The proportion of participants who were considered to be resectable after CT scan is not known
Callery 1997	No separate data available for pancreatic or periampullary carcinoma
Callery 2009	Not a diagnostic accuracy study
Camacho 2005	Not a diagnostic accuracy study
Carmichael 1995	Not a diagnostic accuracy study
Carpenter 1996	Not a diagnostic accuracy study
Catheline 1998	No diagnostic test accuracy data available for diagnostic laparoscopy
Catheline 1999	No diagnostic test accuracy data available for diagnostic laparoscopy
Chambon 1995	No diagnostic test accuracy data available for diagnostic laparoscopy
Champault 1996	No diagnostic test accuracy data available for diagnostic laparoscopy
Champault 1997	No diagnostic test accuracy data available for diagnostic laparoscopy
Charukhchyan 1998	No diagnostic test accuracy data available for diagnostic laparoscopy
Cipollone 2012	No diagnostic test accuracy data available for diagnostic laparoscopy
Conlon 1997	The number of participants with pancreatic or periampullary cancers is not stated
Conlon 1999	Not a diagnostic accuracy study
Conlon 2002	Not a diagnostic accuracy study
Connor 2004	Not a diagnostic accuracy study
Croome 2009	Insufficient diagnostic test accuracy data available for diagnostic laparoscopy
Croome 2010	Insufficient diagnostic test accuracy data available for diagnostic laparoscopy
Cuesta 1993	No diagnostic test accuracy data available for diagnostic laparoscopy
Cuschieri 1978	No diagnostic test accuracy data available for diagnostic laparoscopy
Cuschieri 1988	The proportion of participants who were considered to be resectable after CT scan is not known
D'Angelica 2003	Wrong target condition
Dadan 1980	Insufficient diagnostic test accuracy data available for diagnostic laparoscopy
Doran 2004	No diagnostic test accuracy data available for diagnostic laparoscopy



Study	Reason for exclusion
Doucas 2007	No diagnostic test accuracy data available for diagnostic laparoscopy
Duffy 2008	Not a diagnostic accuracy study
Durup Scheel-Hincke 1999	No diagnostic test accuracy data available for diagnostic laparoscopy
Eigler 1999	Not a diagnostic accuracy study
Ellsmere 2005	No diagnostic test accuracy data available for diagnostic laparoscopy
Enestvedt 2008	Includes participants who were considered to be unresectable by CT scan
Fernandez-del Castillo 1994	Not a diagnostic accuracy study
Fernandez-del Castillo 1998	Not a diagnostic accuracy study
Ferrone 2006	No diagnostic test accuracy data available for diagnostic laparoscopy
Feussner 2000	No separate data available for pancreatic or periampullary cancer
Fevery 1985	No separate data available for pancreatic or periampullary cancer
Fockens 1993	Not a diagnostic accuracy study
Friess 1997	No diagnostic test accuracy data available for diagnostic laparoscopy
Friess 1998	No separate data available for pancreatic or periampullary cancer
Fristrup 2006	No diagnostic test accuracy data available for diagnostic laparoscopy
Fukumoto 1989	No separate data available for pancreatic or periampullary cancer
Garcea 2012	No diagnostic test accuracy data available for diagnostic laparoscopy
Garofalo 2009	No diagnostic test accuracy data available for diagnostic laparoscopy
Gouma 1996	No diagnostic test accuracy data available for diagnostic laparoscopy
Gouma 1999	Not a diagnostic accuracy study
Gouma 2002	Not a diagnostic accuracy study
Hann 1997	No diagnostic test accuracy data available for diagnostic laparoscopy
Hashimoto 2015	In this study, all 11 participants who underwent diagnostic laparoscopy and laparotomy had resectable pancreatic cancers. There were therefore no true positives and false negatives for estimation of sensitivity, and this study was excluded
Healthcare 1999	Not a diagnostic accuracy study
Heger 2008	Not a diagnostic accuracy study
Hernandezguio 1965	Not a diagnostic accuracy study
Herrera 2003	No diagnostic test accuracy data available for diagnostic laparoscopy



Study	Reason for exclusion
Hidalgo 2004	Not a diagnostic accuracy study
Hohenberger 2000	Not a diagnostic accuracy study
Holzman 1997	No diagnostic test accuracy data available for diagnostic laparoscopy
Hunerbein 1999	Not a diagnostic accuracy study
Hunerbein 2001	No diagnostic test accuracy data available for diagnostic laparoscopy
Ialongo 2010	Not a diagnostic accuracy study
Ialongo 2015	Not a diagnostic accuracy study
ldo 1982	No diagnostic test accuracy data available for diagnostic laparoscopy
Ihse 1984	Not a diagnostic accuracy study
Ishida 1983	No diagnostic test accuracy data available for diagnostic laparoscopy
Ishida 1984	Wrong target condition
Ivanov 1989	No diagnostic test accuracy data available for diagnostic laparoscopy
Jackowski 1997	No diagnostic test accuracy data available for diagnostic laparoscopy
Jakobs 1999	Not a diagnostic accuracy study
Jarnagin 2000	Wrong target condition
Jayakrishnan 2015	Not a diagnostic accuracy study
Jerby 1998	Not a diagnostic accuracy study
Jimenez 2000	Not a diagnostic accuracy study
Jimenez 2000a	No diagnostic test accuracy data available for diagnostic laparoscopy
John 1999	No diagnostic test accuracy data available for diagnostic laparoscopy
Juzkow 1996	Not a diagnostic accuracy study
Kadar 1997	No diagnostic test accuracy data available for diagnostic laparoscopy
Kanazawa 1983	No separate data available for pancreatic or periampullary cancer
Kaplan 1979	Not a diagnostic accuracy study
Karachristos 2005	Intervention between index test and reference standard
Kellokumpu 1996	Not a diagnostic accuracy study
Kelly 2009	No diagnostic test accuracy data available for diagnostic laparoscopy
Khamdanov 1983	Not a diagnostic accuracy study



Study	Reason for exclusion
Kiyonaga 1982	Wrong target condition
Klingler 2000	No diagnostic test accuracy data available for diagnostic laparoscopy
Krahenbuhl 1997	Not a diagnostic accuracy study
Krustev 1998	No diagnostic test accuracy data available for diagnostic laparoscopy
Kubyshkin 2000	No diagnostic test accuracy data available for diagnostic laparoscopy
Kuster 1967	No diagnostic test accuracy data available for diagnostic laparoscopy
Kwon 2002	No diagnostic test accuracy data available for diagnostic laparoscopy
Lavonius 2001	Includes participants who were considered to be unresectable by CT scan
Lightdale 1992	Not a diagnostic accuracy study
Liu 2004	Not a diagnostic accuracy study
Long 2005	Not a diagnostic accuracy study
Luque-de Leon 1998	No diagnostic test accuracy data available for diagnostic laparoscopy
Luque-de Leon 1999	No diagnostic test accuracy data available for diagnostic laparoscopy
Macutkiewicz 2009	No diagnostic test accuracy data available for diagnostic laparoscopy
Madsen 1994	No separate data available for pancreatic or periampullary cancer
Madsen 1994a	No separate data available for pancreatic or periampullary cancer
Maire 2004	No diagnostic test accuracy data available for diagnostic laparoscopy
Maithel 2008	No diagnostic test accuracy data available for diagnostic laparoscopy
Meduri 1994	The proportion of participants who were considered to be resectable after CT scan is not known
Metcalfe 2003	Not a diagnostic accuracy study
Meyer 1973	No diagnostic test accuracy data available for diagnostic laparoscopy
Misra 2012	No diagnostic test accuracy data available for diagnostic laparoscopy
Molnar 2010	The proportion of patients who were considered to be resectable after CT scan is not known
Morak 2009	No diagnostic test accuracy data available for diagnostic laparoscopy
Morganti 2005	No diagnostic test accuracy data available for diagnostic laparoscopy
Mortensen 1996	No diagnostic test accuracy data available for diagnostic laparoscopy
Muniraj 2013	Not a diagnostic accuracy study
Muntean 2009	No diagnostic test accuracy data available for diagnostic laparoscopy



Study	Reason for exclusion
Munteanu 2010	No diagnostic test accuracy data available for diagnostic laparoscopy
Murugiah 1993	The proportion of participants who were considered to be resectable after CT scan is not known
Nagy 1999	Not a diagnostic accuracy study
Nieveen 1996	No diagnostic test accuracy data available for diagnostic laparoscopy
Nieveen 1997	No diagnostic test accuracy data available for diagnostic laparoscopy
Nieveen 1998	No diagnostic test accuracy data available for diagnostic laparoscopy
Nieveen 1999	No diagnostic test accuracy data available for diagnostic laparoscopy
Nieveen 2000	No diagnostic test accuracy data available for diagnostic laparoscopy
Nieveen 2003	No diagnostic test accuracy data available for diagnostic laparoscopy
Nieveen 2003a	No diagnostic test accuracy data available for diagnostic laparoscopy
Occelli 1999	No diagnostic test accuracy data available for diagnostic laparoscopy
Palanivelu 2001	Not a diagnostic accuracy study
Parks 2000	Not a diagnostic accuracy study
Pedrazzoli 1994	No diagnostic test accuracy data available for diagnostic laparoscopy
Pelton 1998	Insufficient diagnostic test accuracy data available for diagnostic laparoscopy
Pietrabissa 1996	No diagnostic test accuracy data available for diagnostic laparoscopy
Pietrabissa 1996a	No diagnostic test accuracy data available for diagnostic laparoscopy
Pietrabissa 1999	Includes participants who were considered to be unresectable by CT scan
Pisters 2001	Not a diagnostic accuracy study
Potkonjak 1974	No diagnostic test accuracy data available for diagnostic laparoscopy
Ramshaw 1999	Not a diagnostic accuracy study
Ribero 1994	No diagnostic test accuracy data available for diagnostic laparoscopy
Rodgers 2003	No separate data available for pancreatic or periampullary cancer
Rothlin 1996	Not a diagnostic accuracy study
Rumstadt 1997	No diagnostic test accuracy data available for diagnostic laparoscopy
Rumstadt 1997a	No diagnostic test accuracy data available for diagnostic laparoscopy
Saeian 1999	Not a diagnostic accuracy study
Sand 1996	No separate data available for pancreatic or periampullary cancer



Study	Reason for exclusion				
Santoro 2012	No information on whether the distant metastases were confirmed histologically as metastases				
Sato 1985	Not a diagnostic accuracy study				
Satoi 2011	No diagnostic test accuracy data available for diagnostic laparoscopy				
Schachter 1999	Wrong target condition				
Schmidt 1997	No diagnostic test accuracy data available for diagnostic laparoscopy				
Schmied 2000	Not a diagnostic accuracy study				
Schmielau 1997	Not a diagnostic accuracy study				
Schneider 2003	The proportion of participants who were considered to be resectable after CT scan is not known				
Schnelldorfer 2014	Not clear whether histopathological confirmation of metastasis was obtained				
Schrenk 1994	Number of participants with pancreatic or periampullary cancer was not reported				
Schrenk 1995	No diagnostic test accuracy data available for diagnostic laparoscopy				
Schwab 1996	Includes participants with unresectable cancers on CT scan				
Sperlongano 2005	Not a diagnostic accuracy study				
Sperlongano 2006	Not a diagnostic accuracy study				
Tang 2001	No separate data available for pancreatic or periampullary cancer				
Tapper 2011	No diagnostic test accuracy data available for diagnostic laparoscopy				
Taylor 2001	No diagnostic test accuracy data available for diagnostic laparoscopy				
Terrosu 2000	Number of participants with pancreatic or periampullary cancer was not reported				
Thomson 2006	No diagnostic test accuracy data available for diagnostic laparoscopy				
Tilleman 2004	Not a diagnostic accuracy study				
Tilleman 2004a	No diagnostic test accuracy data available for diagnostic laparoscopy				
Toughrai 2013	Not a diagnostic accuracy study				
van Delden 1996	No diagnostic test accuracy data available for diagnostic laparoscopy				
van Dijkum 1997	The proportion of participants who were considered to be resectable after CT scan is not known				
Velanovich 1998	No separate data available for pancreatic or periampullary cancer				
Velanovich 2004	No diagnostic test accuracy data available for diagnostic laparoscopy				
Velasco 2000	The proportion of participants who were considered to be resectable after CT scan is not known				
Vollmer 2002	Includes participants who were considered to be unresectable by CT scan				



Study	Reason for exclusion	
Warshaw 1990	Not a diagnostic accuracy study	
Warshaw 1990a	Includes participants who were considered to be unresectable by CT scan	
Watanabe 1993	No diagnostic test accuracy data available for diagnostic laparoscopy	
Weiner 1995	No separate data available for pancreatic or periampullary cancer	
White 2001	Intervention between index test and reference standard	
White 2004	Not a diagnostic accuracy study	
White 2008	Wrong target condition	
Wilson 2010	Not a diagnostic accuracy study	
Yoshida 2002	No diagnostic test accuracy data available for diagnostic laparoscopy	
Zhao 2003	No diagnostic test accuracy data available for diagnostic laparoscopy	

CT: computed tomography

DATA

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

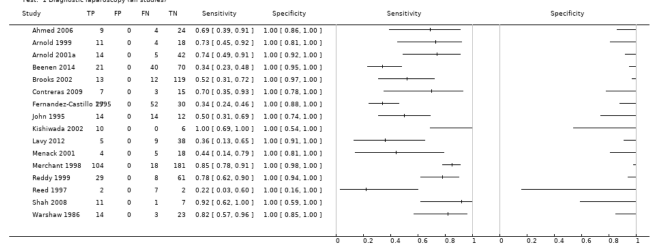
Table Tests. Data tables by test

Test	No. of studies	No. of participants
1 Diagnostic laparoscopy (all studies)	16	1146
2 Diagnostic laparoscopy (pancreatic cancer only)	7	340



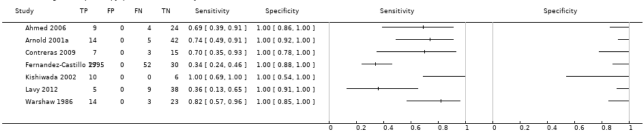
Test 1. Diagnostic laparoscopy (all studies).

Review: Diagnostic accuracy of laparoscopy following computed tomography (CT) scanning for assessing the resectability with curative intent in pancreatic and periampullary cancer Test: 1 Diagnostic laparoscopy (all studies)



Test 2. Diagnostic laparoscopy (pancreatic cancer only).

Review: Diagnostic accuracy of laparoscopy following computed tomography (CT) scanning for assessing the resectability with curative intent in pancreatic and periampullary cancer Test: 2 Diagnostic laparoscopy (pancreatic cancer only)



ADDITIONAL TABLES

Table 1. QUADAS-2 classification

Domain 1: Patient se- lection	Patient sampling	Patients with pancreatic and periampullary cancer considered eligible for surgical resection following a CT scan
	Was a consecutive or random sample of patients enrolled?	Yes: If a consecutive sample or a random sample of patients with pancreatic and periampullary cancer eligible for surgical resection after CT scan was included in the study No: If a consecutive sample or a random sample of patients with pancreatic and periampullary cancer eligible for surgical resection after CT scan was not included in the study Unclear: If this information was not available
	Was a case-control design avoided?	Yes: If a cohort of patients about to undergo surgical resection were studied No: If patients who underwent unsuccessful laparotomy (cases) were compared with patients who underwent successful surgical resection (controls). Such studies were excluded Unclear: We anticipated that we would be able to determine whether the design was case-control



Table 1. QUADAS-2 cla	ssification (Continued)	
		As anticipated, we were able to determine the study design and were able to exclude all case-control studies. So, all studies included in this review were classified as 'yes' for this item
	Did the study avoid in- appropriate exclusions?	Yes: If all patients with pancreatic and periampullary cancer eligible for surgical resection were included No: If the study excluded patients based on high probability of resectability (for example, small tumours) Unclear: If this information was not available
	Could the selection of patients have introduced bias?	Low risk of bias: If 'yes' classification for all the above 3 questions; high risk of bias: if 'no' classification for any of the above 3 questions; unclear risk of bias: if 'unclear' classification for any of the above 3 questions but without a 'no' classification for any of the above 3 questions
	Patient characteristics and setting	Yes: We included only patients with pancreatic and periampullary cancer who were considered eligible for surgical resection following a CT scan. So, we anticipated all the included studies to be classified as 'yes' No: We excluded studies where patients were considered unsuitable for surgery after a CT scan. So, we did use this classification Unclear: We excluded studies in which it was not clear whether the patients had undergone CT scan following which they were still considered suitable for surgical resection
	Are there concerns that the included patients and setting do not match the review question?	Considering the inclusion criteria of this review, we anticipated that all of the included studies would be classified as 'low concern'. However, this was not the case, as shown in Figure 3
Domain 2: Index test	Index test(s)	Diagnostic laparoscopy with histologic confirmation of metastases
	Were the index test results interpreted without knowledge of the results of the reference standard?	The index test would always be conducted and interpreted before the reference standard. So, this classification was always 'yes'
	If a threshold was used, was it prespecified?	Not applicable
	Could the conduct or interpretation of the index test have introduced bias?	We anticipated classifying all studies as 'low risk of bias' because diagnostic laparoscopy indicates that structures within the abdomen were inspected, diagnostic laparoscopy would be conducted and interpreted before reference standard, and because we excluded any studies without histological confirmation of the metastatic spread
		As anticipated, all of the studies were classified as 'low risk of bias' for this domain
	Are there concerns that the index test, its con- duct, or interpretation	Considering the inclusion criteria for this review, we anticipated that all of the included studies will be classified as 'low concern'
	differ from the review question?	As anticipated, all of the studies were classified as 'low concern' for this domain
Domain 3: Target condition and reference standard	Target condition and reference standard(s)	Unresectability. The reasons for unresectability include involvement of adjacent structures or distant metastases. There is currently no universal criteria for unresectability. Consensus exists for the definition of borderline resectable



Table 1. QUADAS-2 classification (Continued)

cancers (Abrams 2009). Therefore where there is less tissue involvement than in a borderline resectable cancer, the tumour can be considered as resectable Positive reference standard: Confirmation of liver or peritoneal involvement by histopathological examination of suspicious (liver or peritoneal) lesions (irrespective of how the tissues were obtained for histopathological examination). We accepted only paraffin section histology as the reference standard. We also accepted the surgeon's judgement of unresectability on laparotomy when biopsy confirmation was not possible (e.g. the surgeon may not resect the tumour if it invaded the adjacent blood vessels but will not obtain a biopsy confirmation of this because of the danger posed by resecting a part of a large blood vessel)

Negative reference standard: Cancer was fully resected, i.e. clear resection margins on histology

Is the reference standard likely to correctly classify the target condition? Yes: If histological confirmation of distant spread or local infiltration of adjacent structures making the cancer unresectable was obtained. The report on the resection margins showed clearly that the cancer was completely resected. We did not anticipate that any studies would meet these criteria because of the danger that biopsy of infiltration of adjacent structures poses No: If resection margins were not clear of cancer

Unclear: If surgeon's judgement was used to assess unresectability or if the information about the resection margins was not available. We anticipated that most studies would be classified as 'unclear' because surgeon's judgement is generally used as a criterion for unresectability in clinical practice

As anticipated, all of the studies were classified as 'unclear' for this item

Were the reference standard results interpreted without knowledge of the results of the index tests? It is not possible to perform the reference standard without knowledge of the results of the index test. However, only patients with suspicious lesions on laparoscopy undergo biopsy, and only patients with negative laparoscopy would undergo laparotomy. The results of the index test are unlikely to influence the results of the reference standard. All studies were classified as 'no' for this question

Could the reference standard, its conduct, or its interpretation have introduced bias? Risk of bias was determined as 'low' if the answer to the first question was 'yes', 'high' if the answer to the first question was 'no', and 'unclear' if the answer to the first question was 'unclear'

Are there concerns that the target condition as defined by the reference standard does not match the question? Considering the inclusion criteria for this review, we anticipated that all of the included studies would be classified as 'low concern'

As anticipated, all of the studies were classified as 'low concern' for this domain

Domain 4: Flow and timing

Flow and timing

The cancer may progress if there is long time interval between diagnostic laparoscopy and laparotomy. So, we chose an arbitrary time interval of 2 months as an acceptable time interval between diagnostic laparoscopy and laparotomy

Was there an appropriate interval between index test and reference standard?

Yes: If the time interval between diagnostic laparoscopy and laparotomy was less than 2 months

No: If the time interval between diagnostic laparoscopy and laparotomy was more than 2 months

Unclear: If the time interval between diagnostic laparoscopy and laparotomy was unclear

Did all patients receive the same reference standard? Yes: If all of the patients received the same reference standard (we anticipated that all the studies would be classified as 'yes')

No: If different patients received different reference standards



Table 1. QUADAS-2 classification (Continued)

Unclear: If this information was not clear

Were all patients included in the analysis?

Yes: If all of the patients were included in the analysis irrespective of whether the results were uninterpretable

No: If some patients were excluded from the analysis because of uninterpretable results

Unclear: If this information was not clear

Could the patient flow have introduced bias?

Low risk of bias: if 'yes' classification for all of the above 3 questions; high risk of bias: if 'no' classification for any of the above 3 questions; unclear risk of bias: if 'unclear' classification for any of the above 3 questions but without a 'no' classification for any of the above 3 questions

CT: computed tomography

Table 2. Prior testing and unresectability

Study name	Type of CT scan	Prior testing in addition to CT scan	Probabili- ty of CT re- sectable disease identified as unre- sectable by diag- nostic la- paroscopy or laparo- tomy	Number of participants (N) and reasons for CT resectable disease identified as unresectable by diagnostic laparoscopy	Probabil- ity of CT and diag- nostic la- paroscopy resectable disease identified as unre- sectable at laparoto- my	Number of participants (N) and reasons for CT and diagnostic la- paroscopy resectable disease identified as unresectable at laparo- tomy
			(Pre-test probabili- ty)		(Post-test probabili- ty of neg- ative diag- nostic la- paroscopy)	
Ahmed 2006	Helical CT		35.1	N = 9	14.3	N = 4
2006	scan			Liver metastases = 6		Metastatic disease = 2
				Peritoneal metastases = 1		Locally advanced disease (1 coeliac artery lymph
				Peritoneal and liver metastases = 2		node, 1 mesenteric vas- cular involvement) = 2
Arnold	No further		45.5	N = 11	18.2	N = 4
1999	informa- tion on CT	underwent en- doscopy and ul-		Liver metastases = 6		Liver metastases = 2
	scan was available	trasound. Some participants un- derwent EUS,		Peritoneal metastasis = 1		Peritoneal metastases = 1
		proportion un- clear		Peritoneal and liver metastases = 3		Liver and peritoneal metastases = 1
				Peritoneal and omental metastases = 1		



Arnold No further	Endoscopy, ul-	31.1	N = 14	10.6	N = 5			
	informa- tion on CT	trasound, and MRI. Proportion		Liver metastases = 8		Liver metastases = 3		
	scan was available	of participants who received each modality is		Peritoneal metastases = 2		Peritoneal metastases = 2		
		unclear		Liver and peritoneal metastases = 4		Metastases in the omentum and mesocolon = 2		
						Some had spread to more than 1 location		
Beenen	No further	All participants	46.6	N = 21	36.3	N = 40		
2014	informa- tion on CT scan was available	underwent ab- dominal ultra- sound and ERCP		Reasons for unre- sectability not stated		Reasons for unresectabil- ity not stated		
Brooks	Contrast	85% of partici-	17.4	N = 13	9.2	N = 10		
2002	enhanced, thin slice	pants underwent ERCP		Liver metastases = 6		Liver metastases = 3		
				Peritoneal metastases		Vascular invasion = 3		
				= 5 Other metastatic dis- ease = 2		Peritoneal metastases = 1		
						Local extension = 1		
						Benign disease = 2		
Contreras		tocol CT some partici-	col CT some partici- pants, propor- tion unclear	40.0	N = 7	16.7	N = 3	
2009	protocol CT scan				Aortocaval node disease = 1			
				Peritoneal metastases = 2		Liver metastases = 1		
				Gross regional lym- phadenopathy = 1		Coeliac node disease = 1		
Fernan-	Further de-	None described	None described	None described	72.4	N = 27	63.4	N = 87
dez-Castillo 1995	tails not known			Liver metastases = 11		Vascular invasion at sub-		
				Peritoneal metastases = 3		sequent angiography and did not undergo la- parotomy = 42		
				Omental metastases = 2		Peritoneal disease at la-		
				Metastases in more than 1 site = 11		parotomy = 2 Reasons for unresectability at laparotomy not stated = 43		
John 1995	Con-	Various scanning	70.0	N = 14	53.8	N = 14		
	trast-en- hanced dy-	trast-en- techniques used.		Liver metastases = 10		Metastatic disease = 2		
n	namic CT scan	namic CT	amic CT and proportion		Peritoneal metastases = 8		Locally advanced and metastatic disease = 1	



unic 2. FIII	or testing and	d unresectability (Conunuea)	Hilar lymph node involvement = 2 Some had spread to more than 1 location		Locoregional spread = 11
Kishiwada 2002	Helical CT scan	All participants received ultra- sound	62.5	Reasons for unre- sectability not stated	0	Reasons for unresectabil- ity at laparotomy not stated
Lavy 2012	No further	All participants	26.9	Peritoneal metastases	19.1	N = 9
	informa- tion on CT	received EUS		= 5		Metastatic disease = 2
	scan was available					Locally advanced cancer = 7
Menack	Con- trast-en-	Transabdominal	33.3	Reasons for unre-	21.7	N = 5
2001	hanced CT	ultrasound, EUS, and ERCP per-		sectability not stated		Portal vein occlusion = 1
thin slices of pancreas	formed in some participants, proportion unclear				Metastatic disease in the lymph nodes or liver on laparoscopic ultrasound and biopsy = 2	
					Portal vein encasement = 1	
					Locally advanced disease at laparotomy = 1	
Merchant	•	40.3	N = 104	9.0	N = 18	
1998				Liver metastases = 48		Liver metastases = 6
				Extrapancreatic spread = 41		Extrapancreatic disease = 3
				Nodal spread = 20		Positive nodal disease =
				Vascular invasion = 37		3 Vascular invasion = 2
				Some had spread to more than 1 location		Benign disease = 4
Daddy 1000	From the end of o	None described	27.0		11.6	-
Reddy 1999	Further de- tails not	None described	37.8	N = 29	11.6	N = 6
known			Liver metastases = 23		Liver metastases = 4	
			Liver and peritoneal metastases = 3		Peripancreatic lymph node involvement = 2	
				Hepatic, peritoneal, and mesenteric metastases = 1		
				Mesenteric involvement = 2		
Reed 1997	Further de-	None described	81.8	Reasons for unre-	77.8	N = 7
	tails not known			sectability not stated		Local tumour spread = 5



Table 2. Pri	or testing and	d unresectability (Continued)			Omental spread = 1 Unclear = 1
Shah 2008	Multi-de- tector row CT using pancreatic protocol	None described	63.2	N = 9 Metastases = 6 Locally advanced disease = 3	12.5	Liver metastasis = 1
Warshaw 1986	Further de- tails not known	All participants received chest roentgenography, transhepatic cholangiography, or ERCP and abdominal ultrasound. Some received coeliac and superior mesenteric angiography	42.5	N = 14 Liver metastases = 6 Parietal peritoneal metastases = 7 Omental metastatic disease = 1	11.5	Liver metastases = 3

CT: computed tomography DL: diagnostic laparoscopy

ERCP: endoscopic retrograde cholangio-pancreatography

EUS: endoscopic ultrasound MRI: magnetic resonance imaging

All probabilities in the table are reported as percentages.

APPENDICES

Appendix 1. Glossary of terms

Index test: The diagnostic test being evaluated. In this review the index test is diagnostic laparoscopy after CT scanning

QUADAS: A tool for assessing the methodological quality of diagnostic accuracy studies in terms of risk of bias and applicability to the review question. The assessment parameters are described in more detail in the main text of the review

Reference standard: The test that is accepted as the best available to classify the target condition correctly in a particular setting. In this review the reference standard is biopsy with histopathological confirmation after diagnostic laparoscopy or laparotomy, or the surgeon's judgement of unresectability at laparotomy when biopsy confirmation was not possible

Sensitivity: Proportion of diseased individuals correctly identified as having the disease by the index test i.e. True positives/(True positives + False negatives)

Specificity: Proportion of disease-free individuals correctly identified as being disease-free by the index test i.e. True negatives/(False positives + True negatives)

Target condition: The disease or condition to be diagnosed. In this review the target condition is unresectable pancreatic and periampullary cancer

Appendix 2. Cochrane Register of Diagnostic Test Accuracy Studies and CENTRAL search strategy

#1 ((ampulla near/2 vater*) or ampullovateric or (papilla near/2 vater*) or periampulla* OR peri-ampulla* OR choledoch* or bile duct* or biliary or cholangio* or gall duct or duoden* or small bowel or small intestin* or enter* or pancrea*)
#2 (carcin* or cancer* or neoplas* or tumour* or tumor* or cyst* or growth* or adenocarcin* or malign*)
#3 (#1 AND #2)



#4 (pancreatect* OR pancreaticojejunost* OR pancreaticogastros* OR pancreaticoduodenect* OR duodenopancreatectom*) #5 (#3 OR #4)

#6 (laparoscop* or peritoneoscop* or celioscop* or coelioscop*)
#7 (#5 AND #6)

Appendix 3. MEDLINE search strategy

(((((ampulla vateri[tiab] OR "Ampulla of Vater" [Mesh] OR ampullovateric[tiab] OR papilla vateri[tiab] OR vater papilla[tiab] OR vater ampulla[tiab] OR peri-ampull*[tiab] OR periampull*[tiab] OR choledoch*[tiab] OR alcholedoch*[tiab] OR bile duct*[tiab] OR biliary[tiab] OR cholangio*[tiab] OR gall duct[tiab] OR duodenum[tiab] OR duodenal[tiab] OR duoden*[tiab] OR small bowel[tiab] OR small instestin*[tiab] OR enteral[tiab] OR enteric[tiab] OR enter*[tiab] OR pancreatic[tiab] OR pancreato*[tiab] OR pancreato*[tiab] OR carcinomas[tiab] OR tumor*[tiab] OR tumor*[tiab] OR cyst*[tiab] OR cyst*[tiab] OR cyst*[tiab] OR growth*[tiab] OR adenocarcin*[tiab] OR malignant[tiab] OR malignancy[tiab])) OR "Duodenal Neoplasms"[Mesh] OR "Pancreatic Neoplasms"[Mesh] OR "Common Bile Duct Neoplasms"[Mesh]) AND (surger*[tiab] OR operat*[tiab] OR resection*[tiab] OR surgical*[tiab] OR Surgical Procedures, Operative[MeSH] OR General Surgery[MeSH])) OR (pancreatect*[tiab] OR pancreaticojejunost*[tiab] OR pancreaticojejunostomy[MeSH] OR Pancreaticoduodenect*[tiab] OR duodenopancreatectom*[tiab] OR pancreaticoduodenectomy[MeSH])) OR coelioscop*[tiab] OR coelioscop*[tiab] OR "Laparoscopy"[Mesh])

Appendix 4. EMBASE search strategy

1 ((ampulla vateri or ampullovateric or papilla vateri or vater papilla or vater ampulla or periampull* or periampull* or choledoch* or alcholedoch* or bile duct* or biliary or cholangio* or gall duct or duoden* or small bowel or small intestin* or enter* or pancrea*) and (carcin* or cancer* or neoplas* or tumour* or tumor* or cyst* or growth* or adenocarcin* or malign*)).ti,ab.

2 exp duodenum cancer/ or Vater papilla tumor/ or exp pancreas cancer/ or exp bile duct tumor/

31 or 2

4 (surger* or surgical* or operat* or resection*). ti,ab.

5 exp Surgery/

64 or 5

73 and 6

8 (pancreatect* OR pancreaticojejunost* OR pancreaticogastros* OR pancreaticoduodenect* OR duodenopancreatectom*). ti, ab.

9 exp pancreas surgery/

107 or 8 or 9

11 (laparoscop* or peritoneoscop* or celioscop* or coelioscop*). ti,ab.

12 laparoscopy/ or laparoscopic surgery/

13 11 or 12

14 10 and 13

Appendix 5. Science Citation Index search strategy

#1 TS=(((ampulla vateri or ampullovateric or papilla vateri or vater papilla or vater ampulla or periampull* or peri-ampull* or choledoch* or alcholedoch* or bile duct* or biliary or cholangio* or gall duct or duoden* or small bowel or small intestin* or enter* or pancrea*) and (carcin* or cancer* or neoplas* or tumour* or tumor* or cyst* or growth* or adenocarcin* or malign*)))

#2 TS=(operat* OR surger* OR surgical* OR resection*)

#3 #1 AND #2

#4 TS=(pancreatect* OR pancreaticojejunost* OR pancreaticogastros* OR pancreaticoduodenect* OR duodenopancreatectom*)

#5 #3 OR #4

#6 TS=(laparoscop* or peritoneoscop* or celioscop* or coelioscop*)

#7 #5 AND #6

Appendix 6. SAS code for analysis

data DiagnosticTestMetaAnalysis; input Study_id TP FP FN TN; datalines;

190424

2110418

3 14 0 5 42

4 21 0 40 70



```
5 13 0 12 119
670315
7 27 0 52 30
8 14 0 14 12
910006
1050938
1140518
12 104 0 18 181
13 29 0 8 61
142072
15 11 0 1 7
16 14 0 3 23
run;
/* Modify the dataset for the analysis */
data dt;
set DiagnosticTestMetaAnalysis;
sens=1; spec=0; true=tp; n=tp+fn; output;
sens=0; spec=1; true=tn; n=tn+fp; output;
/* Ensure that both records for a study are clustered together */
proc sort data=dt;
by study_id;
run;
ods output ParameterEstimates=pet4 FitStatistics=fitt4 additionalestimates=addest4;
/* Run random effects logistic regression model for sensitivity only*/
proc nlmixed data=dt tech=quanew lis=5 qpoints=10;
parms msens=2 s2usens=0;
logitp=(msens+usens)*sens;
p = exp(logitp)/(1+exp(logitp));
model true ~ binomial(n,p);
random usens ~ normal([0],[s2usens]) subject=study_id out=randeffs;
/* logLR based on spec=1 */
estimate 'logLR-' log((1-(exp(msens)/(1+exp(msens)))));
/* Obtain summary sens and spec from the model 4 */
data summary4;
set pet4;
if parameter = 'msens' then name = 'Sensitivity';
if parameter = 'msens' then summary=100 * exp(estimate)/(1 + exp(estimate));
if parameter = 'msens' then summlower=100 * exp(lower)/(1 + exp(lower));
if parameter = 'msens' then summupper=100 *exp(upper)/(1 + exp(upper));
output;
run;
/* Obtain summary LR- */
data summaryLR;
set addest4;
summary=exp(estimate);
summlower=exp(lower);
summupper=exp(upper);
```



output; run;

Appendix 7. Calculation of post-test probability of unresectable disease for patients with a negative test result

The post-test probability of unresectable disease for patients with a negative test result can be calculated from the pre-test probability of unresectable disease and the negative likelihood ratio. The calculation using the median pre-test probability from the included studies, as an example, is shown below.

Pre-test probability = 0.414

Pre-test odds = Pre-test probability/(1 - Pre-test probability) = 0.414/0.586 = 0.706

Post-test odds of negative test = Post-test odds * negative likelihood ratio = 0.706 * negative likelihood ratio

Post-test probability of unresectable disease for patients with a negative test result = Post-test odds/(1 + Post-test odds)

WHAT'S NEW

Date	Event	Description
2 June 2016	New search has been performed	Searches were updated. One new study was added and the data re-analysed.
2 June 2016	New citation required but conclusions have not changed	The conclusions remain unchanged.

HISTORY

Protocol first published: Issue 10, 2011 Review first published: Issue 11, 2013

Date	Event	Description
28 August 2014	Amended	Review republished solely to include the plain language summary.

CONTRIBUTIONS OF AUTHORS

VB Allen selected studies for inclusion, extracted the data, and wrote the draft of the review. KS Gurusamy wrote the protocol, selected studies for inclusion, and extracted the data and critically commented on the review. Y Takwoingi helped in the statistical analysis and critically commented on the review. A Kalia selected the studies for inclusion and extracted the data for some of the studies. BR Davidson critically commented on the review.

DECLARATIONS OF INTEREST

VB Allen: None. KS Gurusamy: None. Y Takwoingi: None. A Kalia: None. BR Davidson: None.

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• University College London, UK.

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External sources

· None, Other.

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

The QUADAS tool was replaced by the QUADAS-2 tool.

The software used for meta-analysis was different from the one stated in the protocol.

The median pre-test probability rather than the pre-test probability calculated by a meta-analysis of proportions was used to calculate the post-test probability.

INDEX TERMS

Medical Subject Headings (MeSH)

*Ampulla of Vater; *Unnecessary Procedures; Common Bile Duct Neoplasms [diagnostic imaging] [pathology] [*surgery]; Laparoscopy [*methods]; Laparotomy [*statistics & numerical data]; Neoplasm Staging [methods]; Pancreatic Neoplasms [diagnostic imaging] [pathology] [*surgery]; Randomized Controlled Trials as Topic; Tomography, X-Ray Computed

MeSH check words

Humans