




Measuring depth of invasion of submucosa – invasive adenocarcinoma in oesophageal endoscopic specimens: how good are we?☆

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Measuring depth of invasion of submucosa – invasive adenocarcinoma in oesophageal endoscopic specimens: how good are we?

Aims: Emerging data support that submucosa-invasive (pT1b) esophageal adenocarcinomas are cured via endoscopic resection, provided that invasion measures $\leq 500 \mu\text{m}$, they lack other histological features predictive of nodal metastasis and have negative margins. Hence, pathologists' measurement of the depth of submucosal invasion in endoscopic resections may dictate further management (i.e. endoscopic follow-up versus oesophagectomy). In this study, we assessed the inter-observer agreement in measuring the depth of submucosal invasion in oesophageal endoscopic resections.

Methods and results: Six subspecialised gastrointestinal (GI) pathologists from five academic centres independently measured the depth of submucosal invasion in μm from the deepest muscularis mucosae on 37 oesophageal endoscopic resection slides (round 1 scoring). A consensus meeting with a systematic approach for measuring and discussion of pitfalls was undertaken and

remeasuring (round 2 scoring) was conducted. Inter-observer agreement was assessed by the intraclass correlation coefficient (ICC) and Cohen's kappa statistics. A lack of agreement was seen among the six reviewers with a poor ICC for both rounds: 1 [0.40, 95% confidence interval (CI) = 0.26–0.56] and 2 (0.49, 95% CI = 0.34–0.63). When measurements were categorised as $<$ or $> 500 \mu\text{m}$, the overall agreement among the six reviewers was only fair for both rounds: 1 (kappa = 0.37, 95% CI = 0.22–0.53) and 2 (kappa = 0.29, 95% CI = 0.12–0.46).

Conclusions: Our study shows a lack of agreement among gastrointestinal pathologists in measuring the depth of submucosal invasion in oesophageal endoscopic resections despite formulating a consensus approach for scoring. If important management decisions continue to be based upon this parameter, more reproducible and concrete guidelines are needed.

Keywords: adenocarcinoma, depth, invasion, oesophagus

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Introduction

Endoscopic management is now considered the standard of care for patients with Barrett's dysplasia and early oesophageal adenocarcinoma.^{1–6} Recent data support that submucosa-invasive (pT1b) oesophageal adenocarcinomas are cured via endoscopic resection when the following criteria are met: submucosal invasion ≤ 500 μm , lack of other histological features predictive of nodal metastasis (i.e. poor tumour differentiation and lymphovascular invasion) and clear margins.^{1,4,5,7} In these cases of 'low-risk adenocarcinoma', endoscopic management is a valid alternative to oesophagectomy, given the significant morbidity associated with the latter.^{1,4,5,7–12} Hence, the pathologists' measurement of depth of submucosal invasion in oesophageal endoscopic resections may be vital for guiding further management (i.e. endoscopic follow-up versus oesophagectomy). In this study, we assessed the interobserver agreement among gastrointestinal (GI) pathologists in measuring the depth of submucosal invasion in oesophageal endoscopic resection specimens.

Methods

The study was approved by institutional review board of all the participating authors' institutions. Six subspecialised GI pathologists from five different academic centres cumulatively collected and then independently reviewed haematoxylin and eosin (H&E)-stained slides from oesophageal endoscopic resections with submucosal invasive (pT1b) adenocarcinoma. Endoscopic resections included slides from endoscopic submucosal dissection specimens and endoscopic mucosal resection specimens. Five of the six pathologists had at least 6 years of clinical sign-out experience, while one pathologist had 1 year of clinical sign-out experience. For the purpose of this study, the slides were reviewed solely for the depth of submucosal invasion in μm (no other histopathological parameters were assessed). All study slides had some discernible muscularis mucosae identified. Improperly orientated specimens, including tangentially sectioned specimens, were excluded.

Each pathologist individually selected an area on the slide for measuring the maximal depth of submucosal invasion using their best judgement, similar to what they would do in routine clinical practice. The depth of submucosal invasion was measured for each case via computer-captured photomicrograph of the H&E slide.

The pathologists used their available measuring software [Cell Sens for five reviewers and Nikon Imaging Software (NIS) elements for one reviewer] for their Olympus or Nikon cameras that were properly calibrated for each objective. All the pathologists were instructed to take one image using their preferred objective that allowed them to capture and best demonstrate the entire tumour depth in their field of view. The measurement was performed independently by each pathologist as per the recommended guidelines, i.e. measurement in μm from the deepest aspect of the muscularis mucosae (MM) to the deepest extent of submucosal invasion.^{4,5} This was designated as round 1 scoring. Three months after round 1 scoring, an online consensus meeting via Zoom (Zoom Video Communications, San Jose, CA, USA) between all the pathologists was undertaken. Selected ($n = 7$) cases were re-assessed together, and a consensus approach to measure the depth of invasion and potential pitfalls were discussed among the authors based upon the review of recent literature as well as the authors' own personal practices and experience. These selected cases included four cases with disagreement when measurements were categorised at 500 μm (i.e. the same case scored as >500 μm and <500 μm by different pathologists), two cases which were diagnosed as intramucosal carcinoma (IMC) by some pathologists and as submucosal invasion by the others and one case which was measured as >500 μm by all pathologists. The consensus approach and pitfalls are tabulated in Table 1 and shown in Figure 1. Following the consensus meeting, remeasurements were performed by all the pathologists independently on all the cases on the same H&E slides, using the consensus approach developed in the meeting. This was designated as round 2 scoring. A photograph displaying the measurement was taken by each reviewer for every case scored in round 2, again using their preferred objective. The pathologists were blinded from each other's results for both rounds of scoring.

STATISTICAL ANALYSIS

Interobserver agreement was assessed by the intraclass correlation coefficient (ICC) for continuous variables and by Cohen's kappa statistic for categorical variables. When conducting a reliability study, it is recommended to obtain at least 30 heterogeneous samples and involve at least three raters whenever possible. Under such conditions, ICC values less than 0.5 are indicative of poor reliability, values of 0.5–0.75 indicate moderate reliability, values of 0.75–0.9

Table 1. Consensus approach formulated for the re-measuring (round 2 scoring) of oesophageal endoscopic resection specimens

1. Confirm the diagnosis of submucosal invasive (pT1b) oesophageal adenocarcinoma on the endoscopic resection specimen. Stromal desmoplasia, tumour adjacent to thick-walled submucosal vessels and/or oesophageal ducts and/or submucosal glands are the best indicators of submucosal invasion (Figure 1A)
2. Look at all the fragments on the slides. Choose the fragment with the deepest extent of submucosal invasion by eyeballing and/or by measuring if there is confusion as to which fragment shows the greatest depth of invasion
3. Be wary of muscularis mucosae (MM) abnormalities that are commonly seen in endoscopic resections (Figure 1A). Depth of invasion in the submucosa should be measured in μm from the outermost extent of the outer (deeper) MM to the deepest point of invasion
4. Deepest layers of MM are often destroyed by the invading tumour. In these cases, try to approximate the deepest extent of MM by drawing an imaginary line from the base of the existing deepest MM (which is usually comprised of a thicker muscle bundle, rather than wisps of smooth muscle) at the shoulder(s) of the invasive carcinoma. Use this approximated (imaginary) line as the deepest point of MM (Figure 1A)
5. Draw a vertical line from visible or approximated (imaginary) deepest MM to the deepest extent of tumour in the submucosa and measure
6. If neoplastic glands do not extend beyond the bottom aspect of the imaginary MM compared to that of the adjacent intact MM, they are best classified as intramucosal carcinoma (pT1a)
PITFALLS
1. Smooth muscle walls of thick-walled submucosal blood vessels (in tangential sections) can mimic the wisps of MM (Figure 1B)
2. Wisps of muscle when identified in the submucosal tissue can cause problems with depth of invasion (Figure 1C). A better indicator is to always look at the shoulder of intact deepest MM adjacent to the invasive tumour to gain the best estimate of the location of deepest MM
3. Diathermy artefact of the submucosal tissue can mimic MM (Figure 1D)
4. Dilated, large-calibre vessels are seen in lamina propria within duplicated MM. Do not confuse them with submucosal vessels, which are thick-walled (Figure 1D)

indicate good reliability and values greater than 0.90 indicate excellent reliability.¹³ Cohen suggested that the kappa results should be interpreted as follows: values ≤ 0 as indicating no agreement and 0.01–0.20 as none to slight, 0.21–0.40 as fair, 0.41–0.60 as

moderate, 0.61–0.80 as substantial and 0.81–1.00 as almost perfect agreement.¹⁴

Results

The six participating pathologists cumulatively collected 37 H&E-stained slides from 34 oesophageal endoscopic specimens originally diagnosed as submucosa-invasive (pT1b) adenocarcinoma by the contributing pathologist. For the sake of numbers, different slide(s) of the same lesion from the same patient were accepted as being separate measurable test cases (six slides from three endoscopic specimens; one slide each from 31 endoscopic specimens).

ROUND 1 SCORING

The scoring results for all six reviewers for round 1 are tabulated in Table 2. Of the 37 cases, 28 were scored as demonstrating submucosal invasion by all six reviewers; nine were scored as submucosal invasion by four or five reviewers and as IMC by the remaining one or two reviewers (Figure S1). The lack of agreement among the six reviewers can be seen by the variation around the line of agreement in each plot (Figure 2). The overall estimate of agreement (ICC) among the six reviewers in measuring the depth of invasion was moderate [0.64, 95% confidence interval (CI) = 0.51–0.76]; however, this agreement was skewed higher due to a single outlying scoring observation (with great agreement; all measured $>2000 \mu\text{m}$) that was identified along the line of agreement, seen in the plots (Figure 2). When this outlying observation was excluded the ICC was poor (0.48, 95% CI = 0.33–0.63). When the measurements were categorised as either $>$ or $\leq 500 \mu\text{m}$, one case (3%) was measured as $\leq 500 \mu\text{m}$ (including IMC by two reviewers) and 14 cases (38%) were measured $>500 \mu\text{m}$ by all reviewers (Figure 3). There was disagreement in 22 cases (59%), i.e. a case measured variably as $>500 \mu\text{m}$ and $\leq 500 \mu\text{m}$ by six reviewers (Figure 4). The percentage of measurements scored as $> 500 \mu\text{m}$ varied among the reviewers from 49 to 90% (Table 2). The overall agreement was only fair (Cohen's kappa 0.37, 95% CI = 0.22–0.53) among the reviewers when measurements were categorised at $500 \mu\text{m}$.

ROUND 2 SCORING

The rescoring results for all six reviewers' post-consensus meeting are shown in Table 2. Of the 37

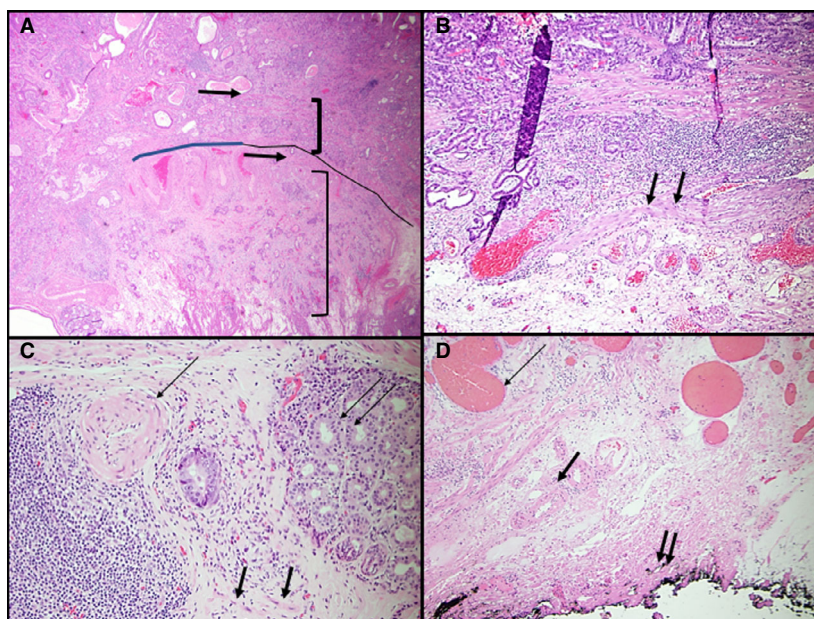


Figure 1. Approach and pitfalls discussed during consensus meeting. **A**, Submucosal thick-walled vessels and stromal desmoplasia confirm submucosal invasion (lower thin bracket). Duplication of muscularis mucosae (MM) is seen (arrows), and the carcinoma within the duplicated muscle represents intramucosal carcinoma (upper thick bracket). The black thin curved line represents the deepest aspect of visible MM and the blue thick line represents the imaginary (approximated) site of the deepest MM destroyed by the tumour. **B**, Smooth muscle wall of the thick-walled submucosal blood vessels (in tangential sections) can mimic the wisps of MM (black arrows). **C**, Wisps of muscle identified in the submucosal tissue (highlighted by thick black arrows) can cause problems with staging and scoring depth of submucosal invasion. This carcinoma is clearly submucosal, given the thick-walled submucosal blood vessel (single long thin arrow) and the presence of submucosal oesophageal glands (double long thin arrows). **D**, Diathermy artefact of the submucosal tissue can mimic MM (double arrows). Dilated, large-calibre vessels can be seen in lamina propria and cause confusion with submucosa (long thin arrow); however, they are not thick-walled as submucosal vessels (short thick black arrow).

cases, 34 were scored as submucosal invasion by all six reviewers; three were scored as submucosal invasion by three or more reviewers and as IMC by the remaining reviewers (Figure S2). The lack of agreement among the measurements between reviewers can be seen again by the variation around the line of agreement in each plot (Figure 2). The overall ICC among the six reviewers remained moderate (0.60, 95% CI = 0.46–0.72). There still was an outlying observation (with great agreement; all measured > 2000 μm) (Figure 2), and when the furthest outlying observation along the line of agreement was removed the ICC indicated poor agreement (0.49, 95% CI = 0.34–0.63). The overall estimate of agreement among the six reviewers when measurements were categorised at 500 μm indicated only fair agreement (0.29, 95% CI = 0.12–0.46). The percentage of measurements scored as >500 μm varied from 65 to 95% among the reviewers. When the measurements were categorised at 500 μm there was disagreement in 17 cases (46%) (Figure 5), while in the remaining 20 cases all reviewers scored as >500 μm (54%) (Figure 6).

AGREEMENT BETWEEN EACH PATHOLOGIST'S SCORING FOR ROUNDS 1 AND 2

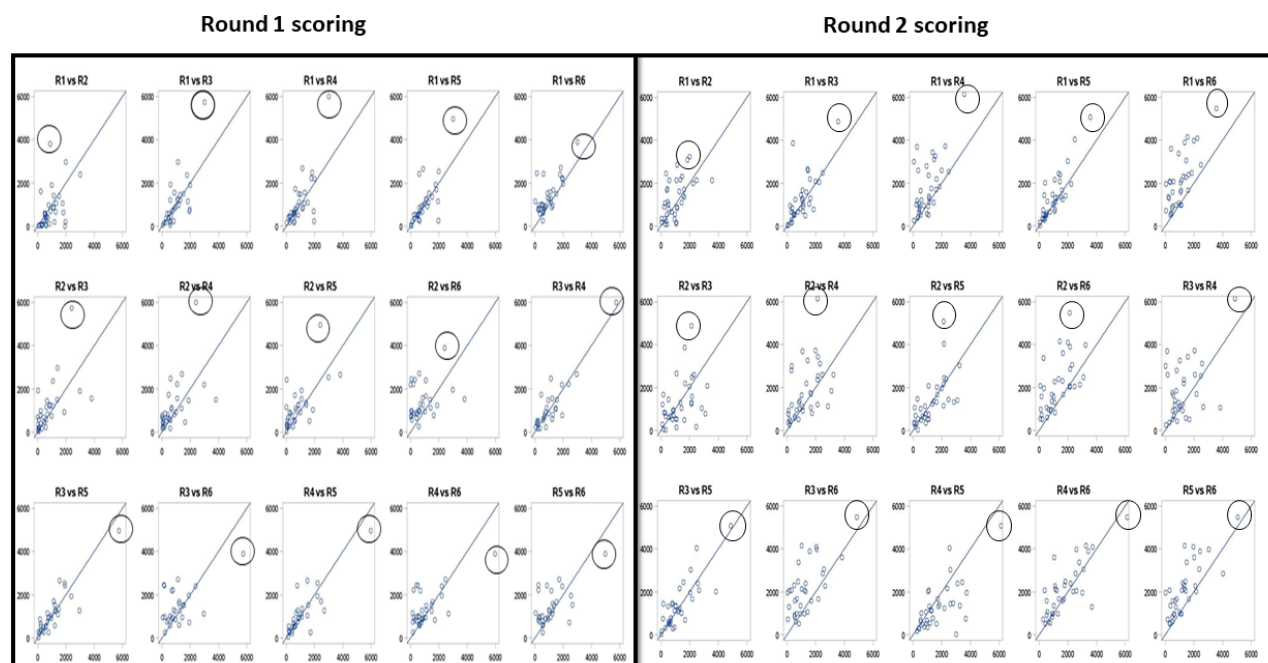
The agreement for each pathologist between their rounds 1 and 2 measurements ranged from 0.45 (poor; reviewer 2) to 0.87 (good, reviewer 1); both these reviewers had at least 6 years of clinical experience. The agreements for other reviewers were 0.67 (reviewers 3 and 4), 0.84 (reviewer 5) and 0.48 (reviewer 6). Of note, as round 2 measurements were undertaken after discussion and establishment of a 'consensus' approach, this was not considered as true 'intra-observer reliability'.

Discussion

Many recent papers have emphasised that the curative status of an endoscopic resection depends upon the adequacy of resection and risk of nodal metastases. The latter cannot be evaluated upon histological examination of endoscopic resection specimens; however, certain histopathological features are

Table 2. Results of the depth of submucosal invasion measurements in oesophageal endoscopic resection specimens among the six reviewers for rounds 1 and 2 scoring

Reviewer (R) no.	Round 1			Round 2		
	Number of cases diagnosed as intramucosal adenocarcinoma	Number of cases scored as submucosal invasion $\leq 500 \mu\text{m}$	Number of cases scored as submucosal invasion $>500 \mu\text{m}$	Number of cases diagnosed as intramucosal adenocarcinoma	Number of cases scored as submucosal invasion $\leq 500 \mu\text{m}$	Number of cases scored as submucosal invasion $>500 \mu\text{m}$
R1	0 (0%)	9 (24%)	28 (76%)	0 (0%)	13 (35%)	24 (65%)
R2	2 (6%)	17 (46%)	18 (49%)	0 (0%)	9 (24%)	28 (76%)
R3	5 (14%)	10 (27%)	22 (59%)	2 (5%)	6 (16%)	29 (78%)
R4	3 (8%)	9 (24%)	25 (68%)	0 (0%)	3 (8%)	34 (92%)
R5	3 (8%)	9 (24%)	25 (68%)	1 (3%)	7 (19%)	29 (78%)
R6	3 (8%)	1 (3%)	33 (90%)	2 (5%)	0 (0%)	35 (95%)
Total reads	16	55	151	5	38	179

**Figure 2.** Matrix of scatterplots showing the lack of agreement (i.e. variation around the line of agreement in each plot) among the six reviewers for the measurements of depth of submucosal invasion in oesophageal endoscopic resections for both round 1 (left panel) and round 2 (right panel). The circle highlights the single outlying observation, which was close to the line of agreement in each plot.

predictive of lymph node metastases, i.e. depth of invasion, poor differentiation and lymphovascular invasion.^{4,5,15,16} Although recently published studies suggest that tumour budding may be an independent prognostic factor in resected oesophageal adenocarcinoma, unlike similar colonic specimens,¹⁷ currently

there are no strong guidelines to report tumour budding in oesophageal endoscopic resections.^{18,19}

The current guidelines support that oesophageal adenocarcinoma invading into the upper one-third of the submucosa with the absence of other negative histological risk factors has a low risk for lymph node

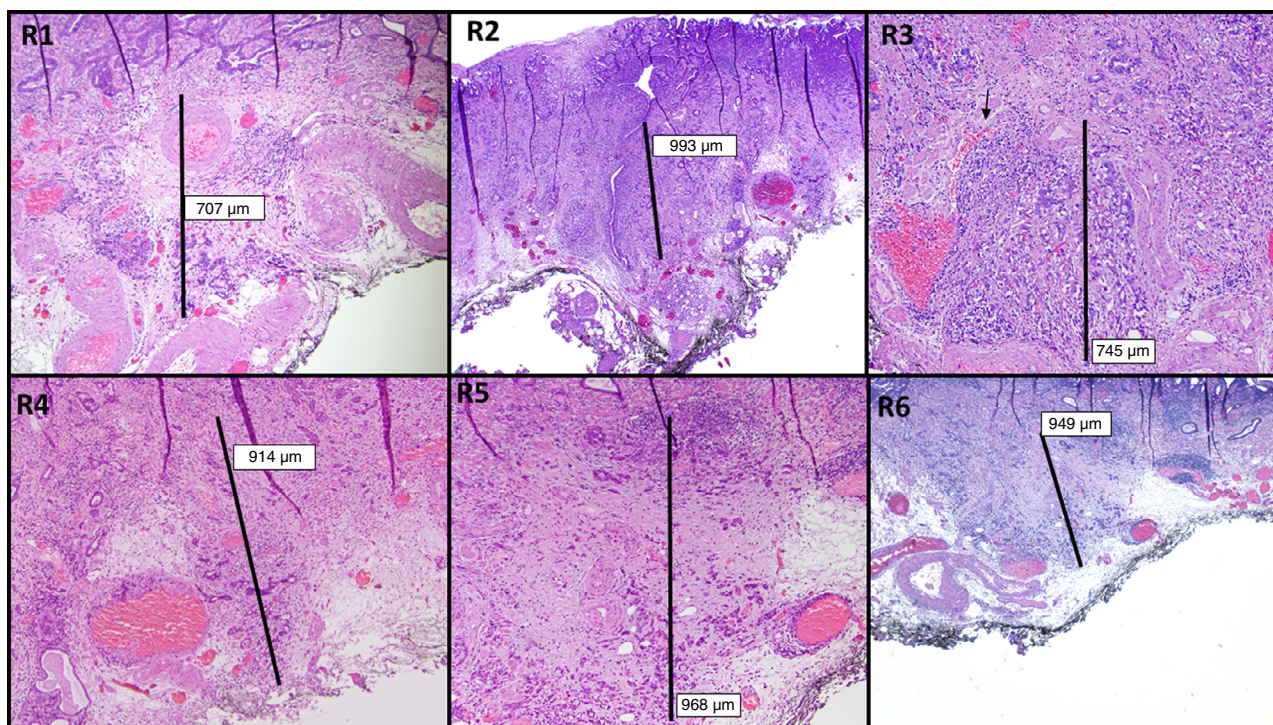


Figure 3. Example of a case in round 1, where all six reviewers measured >500 µm; however, measurements varied from 707 to 993 µm. [Colour figure can be viewed at wileyonlinelibrary.com]

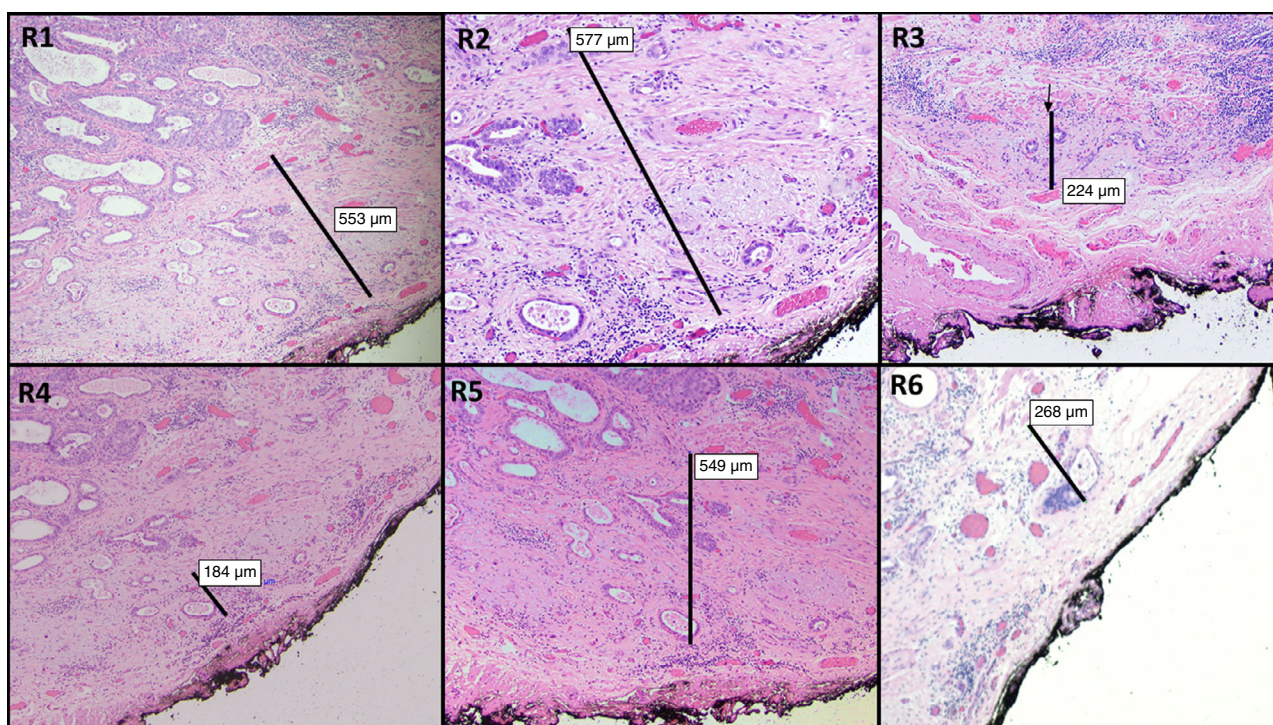


Figure 4. Example of a case with disagreement in round 1 among the six reviewers when the measurements were categorized at 500 µm. Three reviewers (R1, R2, and R5) measured >500 µm and three reviewers (R3, R4 and R6) measured <500 µm. [Colour figure can be viewed at wileyonlinelibrary.com]

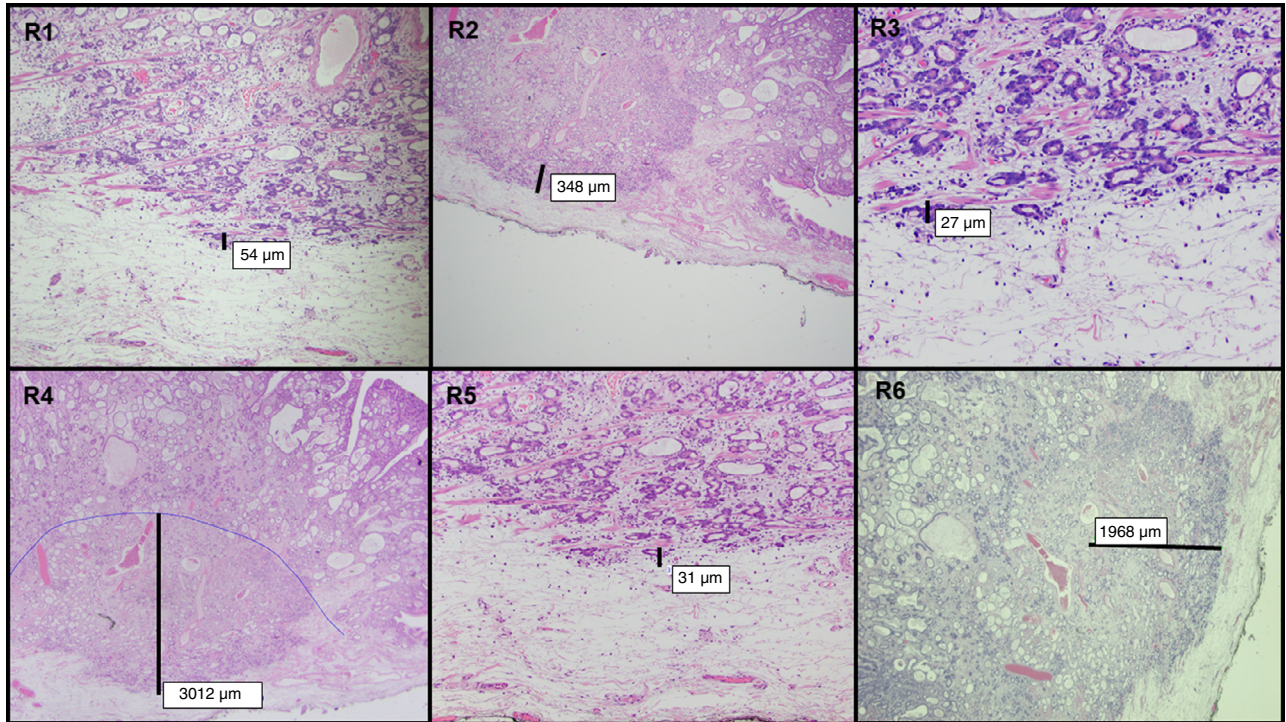


Figure 5. Example of a case with disagreement in round 2 among the six reviewers when the measurements were categorised at 500 µm. Four reviewers (R1, R2, R3 and R5) measured <500 µm and two reviewers (R4 and R6) measured >500 µm. The blue curved line in R4 represents the reviewer's assessment of the approximated (imaginary) muscularis mucosae. [Colour figure can be viewed at wileyonlinelibrary.com]

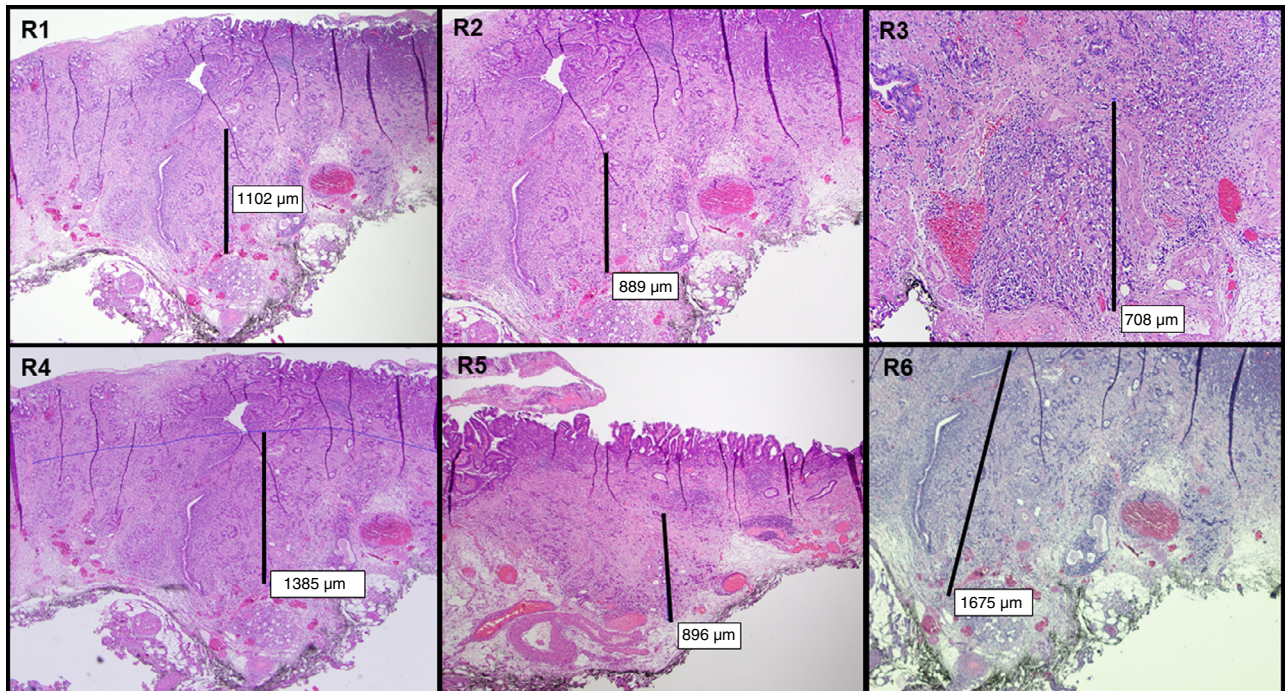


Figure 6. Round 2 measurements (after the consensus meeting) for the same case as Figure 3. All six reviewers still measured >500 µm; however, measurements varied from 708 to 1675 µm. [Colour figure can be viewed at wileyonlinelibrary.com]

metastasis and is regarded as low-risk adenocarcinoma.^{7–11} The pragmatic classification of dividing submucosa into three equal thirds (sm1, superficial one-third submucosa; sm2, intermediate one-third submucosa; sm3, outer one-third submucosa) is considered reliable in oesophagectomy specimens wherein the whole submucosa can be visualised and muscularis propria is present as a landmark for subdivision of submucosa; however, this classification scheme can be challenging and inaccurate in endoscopic resections. Therefore, the Paris endoscopic classification of superficial neoplasms recommends measurement of submucosal invasion in μm as an alternative, wherein 500 μm corresponds approximately to sm1.^{20–22} Recent endoscopic data report a lower risk of nodal metastases (0–2%) when submucosal invasion is $\leq 500 \mu\text{m}$ and there are no other associated negative histological risk factors.^{4,5,7,10,22–25} Given these recent data, measuring depth of invasion within the submucosa has become an increasingly reported histopathological parameter in oesophageal endoscopic resections. However, if significant treatment decisions are to be based upon this parameter it needs to be accurate and reproducible amongst pathologists. Our study shows a lack of agreement among GI pathologists in assessing the depth of submucosal invasion in oesophagus endoscopic resections, even after developing a consensus approach for measurement and discussing potential pitfalls. Our results are similar to a recent study wherein the authors showed a substantial discordance among three GI pathologists for the assessment of depth of submucosal invasion depth in pT1b oesophageal adenocarcinoma. The authors concluded that the discordance may potentially lead to a false estimation of risk of nodal metastases, with serious implications for further therapy.²⁶ The same study found that there was good to excellent agreement between pathologists for the histological assessment of differentiation grade and lymphovascular invasion in pT1b adenocarcinoma. Of note, their study included both oesophageal endoscopic resection specimens ($n = 35$) and oesophagectomy specimens ($n = 50$). Additionally, only one round of scoring was conducted and no consensus guidelines were discussed.

Given the crucial cut-off of 500 μm , wherein close surveillance is recommended for $\leq 500 \mu\text{m}$ invasion and additional oesophagectomy with lymphadenectomy is recommended for invasion depth of $> 500 \mu\text{m}$,^{7,27–33} the above results raise concern and question the validity of using this guideline for clinical decision-making purposes. Although the guideline for measurement (i.e. measure from deepest aspect of

MM to deepest extent of submucosal invasion) may appear straightforward, it can be difficult in routine clinical sign-out. The substantial discordance may be explained by subjective interpretation of the deepest layer of MM and deepest invasion, together with several other practical difficulties associated with the measurement of submucosal invasion. Even in properly fixed and well-orientated specimens, the deepest layers of MM may be destroyed by the invading tumour, leaving no discernible MM present above the deepest extent of invasion.

In these cases, one approach used is to approximate the deepest extent of MM by drawing an imaginary line from the base of the existing deepest MM (which is usually comprised of a thicker muscle bundle, rather than wisps of smooth muscle) at the shoulder(s) of the invasive carcinoma and using this line as the deepest point of MM. Even this approach can be extremely challenging, given the well-known MM abnormalities reported to be present in more than 90% of the Barrett's resections.^{34–36} These include distortion and duplication of MM and the presence of a discontinuous or hypertrophic MM, which further adds to the wide subjective assessment of the deepest layer of MM, as seen in our study. These abnormalities of the MM, together with variability in method of measurement (whether from the deepest visible MM or from an imaginary line of MM) have also been postulated to be a cause of disagreement in other sites within the GI tract.^{37,38} We also found that the pathologists were not uniform in selecting the area with deepest invasion on the same slide, adding to the discordance in measurements among pathologists. Lastly, tissue artefacts during fixation and processing together with specimen curling may make it difficult to assess the focus of deepest invasion as well as the orientation plane, further adding to subjectivity in scoring.

Despite the numerous challenges that lead to poor reproducibility, even after a consensus meeting, we identified several important pitfalls which pathologists should be aware of. First, the space between the duplicated muscle layers may resemble submucosa. We found it helpful to compare the wall thickness of blood vessels in this space to those in areas that were clearly submucosal in order to more readily distinguish lamina propria from submucosa, given that submucosal vessels are thick-walled. Secondly, we found that fibromuscularisation of the submucosa produced by tumoral infiltration can mimic MM, particularly against a background of diathermy artefact; and thirdly, that the muscular layer of thick wall submucosal vessels can simulate wisps of MM when

sectioned tangentially. Of note, prior studies have shown that distinction of IMC from pT1b tumours itself is less than perfect, with an overall kappa value of 0.71 and a kappa of 0.76 between the GI pathologists even in oesophagectomy specimens.^{39,40} In our study, there were a few endoscopic resection cases diagnosed as submucosal invasion by most reviewers, but as IMC by some.

A strength of our study is that we performed two rounds of measurements: the first round simulated authors' routine clinical practice and the second round was after discussion of a consensus approach and pitfalls. Despite that, there remained a wide variation in scoring among pathologists, emphasising that measuring depth of submucosal invasion remains challenging in clinical practice. Also, oesophagectomy specimens were not included in our study in an attempt to mimic the real-life decision process on endoscopic specimens. A limitation of the study was that these endoscopic resection specimens were collected, fixed and processed at five different institutions with possibly different protocols. However, given that all pathologists reviewed all study slides, we believe that this would not affect the interobserver agreement for scoring. Also, this simulates real-world clinical practice, as even many tertiary academic centres in the United States do not pin endoscopic specimens before fixation and frequently evaluate slides prepared at other institutions. A recent study compared the pinning and non-pinning methods and failed to demonstrate a statistically significant difference for clinically relevant histopathological parameters, including depth of tumour infiltration.⁴¹ Also, as the authors of this study typically measure the depth of invasion on H&E-stained slides in routine clinical sign-out practice, ancillary stains were not performed in this study to simulate the routine sign-out decision-making process. Of note, the agreement for scoring depth of invasion did not improve by the use of immunostains in a recent study.²⁶ We also did not attempt to look at interobserver agreement of other histological features typically reported in endoscopic resections, given that studies have already documented good to excellent agreement between pathologists for the histological assessment of differentiation grade and lymphovascular invasion in pT1b adenocarcinoma.²⁶

To conclude, our study shows a lack of agreement among GI pathologists in measuring the depth of submucosal invasion in oesophageal endoscopic resections performed for pT1b adenocarcinoma. If important clinical management decisions continue to be based upon this parameter, the recommendations

need to be revisited with more reproducible and concrete guidelines.

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Conflicts of interest

None.

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Supporting Information

Additional Supporting Information may be found in the online version of this article:

Figure S1. Example of a case in Round 1 wherein measurements of submucosal invasion varied from 227 microns to 1629 microns among five reviewers and one diagnosed intramucosal adenocarcinoma. (R-reviewer; R1, R4- Hematoxylin & eosin x 100; R2, R5- Hematoxylin & eosin x 40x, R6- Hematoxylin & eosin x 200).

Figure S2. Example of a case in Round 2 wherein measurements of submucosal invasion varied from 269 microns to 1190 microns among five reviewers and one diagnosed intramucosal adenocarcinoma. (R-reviewer; R1- Hematoxylin & eosin X100; R2- Hematoxylin & eosin X20; R3, R5- Hematoxylin & eosin x 200x; R4- Hematoxylin & eosin x 40).