Clinical TNM staging for esophageal, gastric, and colorectal cancers in the era of neoadjuvant therapy: A systematic review of the literature

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Abstract

Aim: Clinical staging is vital for selecting appropriate candidates and designing neoadjuvant treatment strategies for advanced tumors. The aim of this review was to evaluate diagnostic abilities of clinical TNM staging for gastrointestinal, gastrointestinal cancers.

Methods: We conducted a systematic review of recent publications to evaluate the accuracy of diagnostic modalities on gastrointestinal cancers. A systematic literature search was performed in PubMed/MEDLINE using the keywords “TNM staging,” “T4 staging,” “distant metastases,” “esophageal cancer,” “gastric cancer,” and “colorectal cancer,” and the search terms used in Cochrane Reviews between January 2005 to July 2020. Articles focusing on preoperative diagnosis of: (a) depth of invasion; (b) lymph node metastases; and (c) distant metastases were selected.

Results: After a full-text search, a final set of 55 studies (17 esophageal cancer studies, 26 gastric cancer studies, and 12 colorectal cancer studies) were used to evaluate the accuracy of clinical TNM staging. Positron emission tomography–computed tomography (PET-CT) and/or magnetic resonance imaging (MRI) were the best modalities to assess distant metastases. Fat and fiber mode of CT may be useful for T4 staging of esophageal cancer, CT was a partially reliable modality for lymph node staging in gastric cancer, and CT combined with MRI was the most reliable modality for liver metastases from colorectal cancer.

Conclusion: The most reliable diagnostic modality differed among gastrointestinal cancers depending on the type of cancer. Therefore, we propose diagnostic algorithms for clinical staging for each type of cancer.

KEYWORDS

colorectal cancer, distant metastases, esophageal cancer, gastric cancer, lymph node metastases, T4
1 | INTRODUCTION

Despite recent advances in surgical techniques, perioperative care, and multimodal treatment, postoperative recurrence is observed in approximately 40% of esophageal cancers\(^1\) and 20%-30% of gastric\(^2\) and colorectal\(^3\) cancers with advanced tumors.\(^4\) Lymph node metastasis remains crucial for applying adjuvant treatment and predicting oncological outcome. Various studies have shown that such postoperative recurrence was frequently reduced by neoadjuvant chemotherapy (NAC).\(^7\) The present review evaluates the accuracy of preoperative diagnosis in gastrointestinal cancers, including lymph node metastasis.

Potential T4 esophageal cancer should be treated with neoadjuvant chemoradiation therapy to ensure a negative surgical margin for cancer cells. NAC became the standard management for stage II/III esophageal cancer following the results of the JCOG9907 trial.\(^12\) JCOG1002 evaluated NAC for locally advanced gastric cancer with extended lymph node metastasis and/or bulky positive nodes.\(^13\) Two more ongoing trials also evaluating NAC for locally advanced gastric cancer.\(^14\) Distant metastases should be classified as a noncurative factor for surgical approach. Therefore, clinical TNM staging should be accurate, based on high sensitivity and specificity to predict T4 and/or distant metastases. Since definitive chemoradiation therapy showed a similar overall survival to radical surgery for clinical stage I esophageal cancer,\(^16\) an accurate diagnosis of lymph node metastases is also vital to design treatment strategies for potential stage I tumors.

In Western countries, preoperative chemotherapy or chemoradiotherapy is a standard therapeutic strategy for advanced gastric cancer, based on the findings from large-scale randomized clinical trials.\(^17\) While advanced stage gastric cancer is the main target of NAC, 8.3% of pathological T1 patients were included in the surgery alone group,\(^17\) indicating that some early gastric cancer patients underwent unnecessary NAC. This problem may be due to inaccuracy of clinical diagnosis of T and N staging. In Japan, the efficacy of NAC for type 4 and large-sized type 3 was not demonstrated in the JCOG0501 trial.\(^21\) The JCOG1302A trial, which evaluated the accuracy of clinical diagnosis of gastric cancer, was conducted as prospective setting prior to starting the JCOG1509 trial\(^22\) regarding the efficacy of NAC for stage III gastric cancer.\(^23\)

The JCOG1310 trial (PRECIOUS study) is intended to compare preoperative vs postoperative chemotherapy for lower rectal cancer patients with suspected lateral pelvic node metastasis.\(^24\) MRI has been reported to be the most effective tool for the preoperative stage diagnosis of rectal cancer.\(^25\) It remains controversial whether chemotherapy with or without primary tumor resection is effective for patients with incurable stage IV colorectal cancer. The precise detection of distant metastases\(^26\) is vital in order to enroll patients for such a typical randomized study.

Thus, the impact of clinical TNM staging is more important than ever since neoadjuvant therapy for gastrointestinal cancers is becoming established. Therefore, we evaluated the accuracy of clinical TNM staging through multimodal diagnostic tools using a systematic review of recent publications from January 2005 to July 2020. We propose the use of standard diagnostic algorithms for gastrointestinal cancers. The present review aimed to summarize the fundamental information about the accuracy of clinical TNM staging to design future guidelines and clinical protocols for preoperative adjuvant therapy for gastrointestinal cancers.

2 | METHODS

2.1 | Research themes and study selection criteria

The present review focused on esophageal, gastric, and colorectal cancers. An eligible trial was a clinical study which evaluate accuracy of clinical TNM staging based on imaging modalities including computed tomography (CT), magnetic resonance imaging (MRI), and positron emission tomography–computed tomography (PET-CT). Articles including information related to these research themes were searched for independently by H. S., Y. H., TF, S. O., and K. O. using PubMed and MEDLINE between January 2005 and July 2020. In PubMed, the search terms "esophageal cancer," "gastric cancer," "colorectal cancer," and "TNM staging" were used. In MEDLINE, the search terms used in Cochrane Reviews were used (advanced search system, Appendix 1).\(^27\) The relevance of each article was evaluated (by H. S., Y. H., S. O., and T. F.) and categorized as either relevant or irrelevant. Irrelevant articles were excluded from the review.

2.2 | Data extraction

Key messages and information were extracted from each article and organized. The following information from eligible articles was used: authors, title, countries of origin, publication year, total sample size, study design, study period, diagnostic modality, conclusion, and summary statistics (sensitivity, specificity, and number of positive and negative patients) for diagnosis. We focused on two statistical measurements of diagnostic accuracy of the modality: sensitivity (the proportion of positively diagnosed patients with disease) and specificity (the proportion of negatively diagnosed patients without disease).

3 | RESULTS

3.1 | Studies included in this paper

Our systematic search identified 23,126 articles using PubMed and MEDLINE. After a manual search of eligible papers, 3,553 studies (640 esophageal cancer studies, 587 gastric cancer studies, and 2,232 colorectal cancer studies) were considered eligible based on their title and abstract. After a full-text search, a final set of 55 studies (17 esophageal cancer studies, 26 gastric cancer studies, and 12...
colorectal cancer studies) were used to evaluate the accuracy of TNM staging.

3.2 | Esophageal cancer staging

3.2.1 | Diagnosis for T4 invasion

Computed tomography has been used for the majority of diagnostic modalities for T4 ever since Picus et al. and Thompson et al. first reported that CT images were useful to detect T4 invasion of esophageal cancer, with 80% accuracy. Recently, endoscopic ultrasonography (EUS) and MRI have also become standard tools to predict T4 invasion. Six recent studies in five reports were selected to evaluate the diagnostic impact of predicting T4 invasion of esophageal cancer (Table 1A). The sensitivity ranged from 27.3% to 84%, with 69% to 100% specificity. Although the accuracy of EUS was the highest among these diagnostic modalities, CT or MRI are still appropriate modalities in cases of stenosis or obstruction due to the tumor, which make EUS examination impossible. On the other hand, PET/CT had a sensitivity of 32% to 89%, specificity of 70% to 98%, and accuracy of 53% to 82% in detecting T4 invasion of esophageal cancer (Table 1B). M staging was evaluated using computerized tomography and PET/CT, which showed a sensitivity of 42% to 93%, specificity of 50% to 93%, and accuracy of 59% to 87% (Table 1C).

TABLE 1 | The summary of diagnostic modalities for TNM staging in esophageal cancer

| Year | Author [Ref] | Country | Journal | Modality | Number of patients | Sensitivity (%) | Specificity (%) | Accuracy (%) |
|------|--------------|---------|---------|----------|-------------------|----------------|----------------|-------------|
| 2013 | O’Farrell NJ [30] | Ireland | World J Surg | EUS | 222 | 66 | 93 | 71 |
| 2016 | Lin-na Luo [31] | China | Plos One | EUS | 2880 | 84 | 96 | 79 |
| 2018 | Jie Yang [32] | China | Ann Surg Oncol | EUS | 1434 | 27 | 99 | 99 |
| 2018 | Jinrong Qu [33] | China | Eur Radiol | EUS | 43 | 57 | 100 | 68 |
| 2018 | Jinrong Qu [33] | China | Eur Radiol | MRI | 43 | 71 | 100 | 91 |
| 2018 | Yue Zhou [34] | China | World J Gastroenterol | CT | 120 | 84 | 69 | 72 |

B) Summary of diagnostic accuracy for N staging in esophageal cancer.

| Year | Author [Ref] | Country | Journal | Modality | Number of patients | Sensitivity (%) | Specificity (%) | Accuracy (%) |
|------|--------------|---------|---------|----------|-------------------|----------------|----------------|-------------|
| 2008 | van Vliet [37] | The Netherlands | Br J Cancer | EUS | 1841 | 80 | 70 | 75 |
| 2008 | van Vliet [37] | The Netherlands | Br J Cancer | CT | 943 | 50 | 83 | 63 |
| 2008 | van Vliet [37] | The Netherlands | Br J Cancer | PET | 424 | 57 | 85 | 67 |
| 2012 | Li H [38] | China | Eur Radiology. | CT | 205 | 76 | 75 | 76 |
| 2012 | Yano M [39] | Japan | Esophagus | PET/CT | 81 | 32 | 70 | 53 |
| 2014 | Yamada H [40] | Japan | Surgery Today | PET | 258 | 26 | 98 | 82 |
| 2016 | Parry K [41] | The Netherlands | Eur J Surgical Oncology | EUS + CT | 266 | 31 | 84 | 68 |
| 2017 | Foley KG [42] | UK | Clin Radiol. | PET | 112 | 40 | 77 | 55 |
| 2018 | Jeong DY [43] | Korea | Cancer Medicine | EUS | 435 | 90 | 42 | 75 |
| 2018 | Jeong DY [43] | Korea | Cancer Medicine | PET/CT | 435 | 89 | 39 | 73 |
| 2018 | Jiancheng li [44] | China | Rev Assoc MeD Bras | CT | 305 | 55 | 88 | 82 |
| 2018 | Harrington C [45] | United Kingdom | World J Gastrointest Endosc. | PET/CT | 121 | 93 | 50 | 59 |
| 2018 | Liu J [46] | China | Eur Radiol. | CT | 204 | 67 | 92 | 87 |

C) Summary of diagnostic accuracy for M staging in esophageal cancer.

| Year | Author [Ref] | Country | Journal | Modality | Number of patients | Sensitivity (%) | Specificity (%) | Accuracy (%) |
|------|--------------|---------|---------|----------|-------------------|----------------|----------------|-------------|
| 2004 | van Westreeren [47] | The Netherlands | J Clin Oncol. | PET | 452 | 67 | 97 | 86 |
| 2008 | van Vliet [37] | The Netherlands | Br J Cancer | CT | 437 | 52 | 91 | 77 |
| 2008 | van Vliet [37] | The Netherlands | Br J Cancer | PET | 475 | 71 | 93 | 85 |
| 2018 | Lucas Goense [48] | USA | Eur J Nuc Med Mol Imag | PET | 783 | 75 | 94 | 92 |
hand, EUS or MRI was appropriate to determine non-T4 status. The most reliable diagnostic modalities include a combination of EUS and CT. Kobayashi et al analyzed the characteristics of the esophageal motion and esophageal internal target volume margins to assess the differences between clinical T1-T3 and clinical T4 using four-dimensional CT.35 Although the accuracy of EUS was the highest among these diagnostic modalities, CT or MRI were appropriate modalities to detect T4 status. Figure 1 shows the differential diagnosis between T4 and T3 tumors of esophageal cancer after chemoradiation therapy using an image reconstruction method according to the CT value of the tissue histology of enhanced CT, so called fat and fiber mode.36 In this examination, the contrast agent (3 mL/kg body) was administrated and the lesions were scanned with a 50-second delay and a thickness of 1 mm. The fibrotic area induced by the chemoradiation therapy was emphasized as green and the presence of a fibrotic layer between the tumor and the adjacent organs could be interpreted as not T4.

### 3.2.2 | Lymph node staging

A total of 13 studies in 10 manuscripts37-46 evaluated the diagnostic impact of EUS/CT/PET on the nodal involvement in esophageal cancer (Table 1B). Four out of 13 eligible studies used combination diagnosis with either PET/CT or EUC/CT. The sensitivity of nodal involvement ranged from 29% to 94%, and the specificity ranged from 38% to 98% (Table 1B).

**FIGURE 1** Representative CT images for clinical staging in esophageal cancer. The differential diagnosis between T4 and T3 tumors in esophageal cancer by fat and fiber mode. (A) Standard enhanced CT, (B) Fat and fiber mode

### 3.2.3 | Diagnosis for distant metastases

Four studies in three manuscripts37,47,48 evaluated the diagnostic accuracy of imaging to detect distant metastases (Table 1C). The useful modalities were CT and PET. The advantage of CT was its high resolution to detect the lesion with an accuracy of 77%,37 while the advantage of PET was the ability to perform whole-body scanning with highly qualitative contrasted metastatic lesions identified by high glucose uptake with an accuracy of 85% – 92%.46-51 These accuracies were around 10% greater than that of CT; therefore, CT and PET should be used together as morphological and qualitative modalities.

### 3.2.4 | Algorithm of image modalities for clinical staging in esophageal cancer

Based on these findings, we suggest an algorithm of image modalities for clinical staging in esophageal cancer (Figure 2). After routine endoscopic examination to determine the pathology by biopsy and exclude T1 tumors, PET-CT and/or MRI should be performed. Surgical resection should be performed in patients without distant metastasis or T4 invasion, whereas PET-CT should be employed to detect further metastasis in patients with distant metastasis. Chemotherapy with or without surgery or radiation should be selected depending on the involved lesions. Precise assessment by EUS should be performed to select tumors indicated for endoscopic resection when the tumor depth is evaluated as T1. Findlay et al reported “pragmatic staging” of esophageal cancer using decision theory involving selective EUS, PET, and laparoscopy.51 They concluded that EUS was used in 71.8% of patients with T2-T4a disease and that it was moderately accurate for pT1 N0 disease. PET-CT altered management in 23.0% of patients and laparoscopy in 7.1% of patients, including those with T2 and distal esophageal tumors. Furthermore, although EUS provided additional information on T and N categories, its risk outweighed any potential benefits in patients with T2-T4a disease on CT.

### 3.3 | Gastric cancer staging

#### 3.3.1 | T staging

T staging of gastric cancer is evaluated by conventional endoscopy, EUS, CT, and MRI. A total of 10 studies in eight manuscripts21,52-58 evaluated the diagnostic accuracy for T staging (Table 2A). In East Asia, where early gastric cancer is frequently detected, conventional endoscopy is the main modality of T staging. EUS could provide additional diagnostic value for distinction between T1a (m) and T1b (sm). Hwang et al concluded that the accuracy of multidetector-row CT was close to that of EUS and both EUS and multidetector-row CT were useful complementary modalities for the locoregional staging of gastric cancer.56 For advanced gastric cancer, T staging can be performed by CT, and has reported diagnostic accuracies of 82%,53
The reported accuracies from retrospective studies involve potential bias about patient selection. The diagnostic accuracy is likely to be better when higher numbers of T1 cases are included in studies. Fukagawa reported a diagnostic accuracy of 73% in a large-scale prospective study limited to advanced tumor. The diagnostic ability of MRI for T staging is reported to be 78%, which is similar to that of CT.

### 3.3.2 Lymph node staging

The clinical evaluation of lymph node metastases in gastric cancer is performed by either CT or EUS. A total of 13 studies in 10 manuscripts analyzed the diagnostic accuracy for N staging (Table 2B). Pathological N staging is determined by the number of positive nodes. However, accurate diagnosis of the number of positive nodes is challenging, and positive/negative is incorporated into the clinical staging (TNM 8th). In a review that included a high volume of cases of stage T1-4 cancer, the sensitivities and specificities were reported as 83% and 67%, respectively, by Mocelin et al., 77% and 78%, respectively, by Seevaratnam et al., 63% and 67%, respectively, by Wang et al. The incidence of lymph node metastases was higher in advanced tumors than in early tumors. When limited to T1 tumors, the sensitivities and specificities were reported to be 17% and 90%, respectively, by Ahn et al. and 43% and 98%, respectively, by Fujikawa et al. When limited to T2-T4, the specificity and sensitivity were reported to be 63% and 66%, respectively, by Fukagawa et al.

One of the reasons for difficulties in lymph node diagnosis is the diagnostic difficulty for small-sized lymph node metastases. However, this patient chose to be monitored using CT examinations every 6 months without additional surgery for lymph node dissection. The lymph node was found to be clinically metastatic at the No. 6 station. Following this CT finding, the patient underwent distal gastrectomy and one lymph node was found to be pathologically positive for metastasis at the No. 6 station, which was compatible with the CT findings. Looking at these CT images, a tiny lymph node was visible (Figure 3A, B) in the same area, with swollen node visible (Figure 3C). This tiny lymph node may have been positive for metastasis at that time but was not found to be clinically positive due to its small size. This patient underwent distal gastrectomy after this CT finding, and one lymph node was pathologically positive for metastasis at No. 6, the same with the CT finding. Looking back at these CT images, a tiny lymph node was visible (in A and B) at the same area with swollen node in (C). This tiny lymph node was positive for metastasis at that time, which was not clinically positive for its small size.

### 3.3.3 Diagnosis for distant metastases

Peritoneal dissemination is diagnosed using CT, with findings of ascites and multiple mesenteric or omental nodules; however, its diagnostic accuracy is not high (Table 2C). The standard therapeutic strategy for advanced gastric cancer with peritoneal dissemination is systemic chemotherapy without gastrectomy, as determined by the REGATTA trial. The detection of small peritoneal dissemination by staging laparoscopy can avoid unnecessary laparotomy. A total of 10 manuscripts evaluated the diagnostic accuracy of peritoneal metastases by staging laparoscopy (Table 2C). The detection ratio of peritoneal dissemination was found to be 7.8%-36%. In Western countries, the indication of staging laparoscopy is basically resectable advanced gastric cancer diagnosed by PD and routine examination modality as CT, ultrasound, and EUS. In contrast, 46%-53.4%
# TABLE 2
The summary of diagnostic accuracy for TNM staging in gastric cancer

| Year | Author [Ref] | Location | Modality | T          | Number of Patients | Sensitivity (%) | Specificity (%) | Accuracy (%) |
|------|--------------|----------|----------|------------|--------------------|-----------------|-----------------|--------------|
|      |              |          |          |            |                    |                 |                 |              |
|      |              |          |          |            |                    |                 |                 |              |
|      |              |          |          |            |                    |                 |                 |              |
|      |              |          |          |            |                    |                 |                 |              |
| 2008 | Puli [52]    | USA      | EUS      | T1-4       | 1896               | 88/82/90/99     | 60              | NA           |
| 2009 | Pan [53]     | China    | CT       | T3         | 135                | 85              | 81              | 82           |
| 2010 | Hwang [54]   | Korea    | EUS      | T1-4       | 277                | NA              | NA              | 75           |
| 2010 | Hwang [54]   | Korea    | CT       | T1-4       | 277                | NA              | NA              | 77           |
| 2010 | Huang [55]   | China    | MRI      | T1/2 vs. T3/4 | 213               | 93              | 91              | NA           |
| 2011 | Choi [56]    | Korea    | EUS      | T1         | 955                | NA              | NA              | 67           |
| 2011 | Makino [57]  | Japan    | CT       | T1-4       | 616                | NA              | NA              | 91           |
| 2013 | Feng [58]    | China    | EUS      | T1-4       | 610                | NA              | NA              | 77           |
| 2013 | Feng [58]    | China    | CT       | T1-4       | 610                | NA              | NA              | 78           |
| 2017 | Fukagawa [21]| Japan    | CT       | T2 /T3,4   | 1222               | 85              | 49              | 73           |

| Year | Author [Ref] | Location | Modality | T          | Number of Patients | Sensitivity (%) | Specificity (%) | Accuracy (%) |
|------|--------------|----------|----------|------------|--------------------|-----------------|-----------------|--------------|
| 2007 | Bentrem [60] | US       | EUS      | T1-4(T1:30%)| 223               | 75              | 66              | 71           |
| 2009 | Ahn [61]     | Korea    | CT       | T1         | 434                | 17              | 90              | 84           |
| 2010 | Pan [62]     | China    | CT       | T1-4       | 350                | NA              | NA              | 87           |
| 2012 | Seevaratnam  | Canada   | CT       | T1-4       | 2909               | 77              | 78              | 66           |
| 2012 | Seevaratnam  | Canada   | MRI      | T1-4       | 109                | 85              | 75              | 53           |
| 2012 | Seevaratnam  | Canada   | PET      | T1-4       | 422                | 40              | 60              | 98           |
| 2013 | Feng [58]    | China    | EUS      | T1-4       | 610                | NA              | NA              | 49           |
| 2013 | Feng [58]    | China    | CT       | T1-4       | 610                | NA              | NA              | 45           |
| 2013 | Hasegawa [64]| Japan    | CT       | T1-4(T1:60%)| 315               | 46              | 97              | 81           |
| 2014 | Fujikawa [65]| Japan    | CT       | T1         | 761                | 4.3             | 99              | 90           |
| 2015 | Wang [66]    | China    | CT       | T1-4       | 6788               | 67              | 84              | NA           |
| 2015 | Mocellin [67]| Italy    | EUS      | T1-4       | 3573               | 83              | 67              | NA           |
| 2017 | Fukagawa [21]| Japan    | CT       | T2-4       | 1241               | 63              | 66              | 64           |

| Year | Author [Ref] | Location | Number of Patients | Indication of SL | Yield | False negative (%) |
|------|--------------|----------|--------------------|------------------|-------|--------------------|
| 2006 | Sarela [70]  | US       | 657                | Resectable GC & EGJ | 23    | 10% (41/401) (p:56%) |
| 2013 | Munasinghe [71]| UK     | 316                | Resectable GC & EGJ | 23    | 0% (0/183)         |
| 2014 | Ishigami [72] | Japan   | 178                | GC ≥T2           | 43    | NA                 |
| 2015 | Convie [73]  | UK       | 295                | Resectable GC, EGC & EC, ≥T3 or N+ | 21    | NA                 |
| 2015 | Mirza [74]   | UK       | 378                | Resectable GC, EGC & EC, ≥T3 or N+ | 14    | NA                 |
| 2016 | Simon [75]   | France   | 116                | Resectable GC, EGC & EC, ≥T3 or N+ | 13    | NA                 |
| 2016 | Ikoma [76]   | US       | 711                | Resectable GC & EGC (EGJ: 43.2%) | 36    | NA                 |
| 2016 | Hu [77]      | China    | 582                | GC ≥T2           | 26    | NA                 |
| 2017 | Hosogi [78]  | Japan    | 120                | ≥5 cm and/or bulky N | 45    | 5.9% (1/17)       |
| 2018 | Irino [79]   | Japan    | 156                | Large type 3 & type 4, bulky N/PAN, suspicious for P | 47    | 11% (7/66)       |
was reported in Japan because staging laparoscopy is performed for more limited patients who are potentially associated with peritoneal dissemination, including type 4, large-sized type 3 (>8 cm), and high lymph node metastases. The diagnostic accuracy of peritoneal dissemination by staging laparoscopy is not always 100%. The percentage of “false negatives” is reported to be 11%–17% in Japan and 0%–8% in Western countries. The reason for this discrepancy is considered to be the difference in the indication of staging laparoscopy.

### 3.3.4 Algorithm of image modalities for clinical staging in gastric cancer

Figure 4 shows an algorithm of image modalities for clinical staging in gastric cancer. Endoscopy and CT scan should be performed first for pretreatment diagnosis. If there are no findings for distant metastases (cM0) by CT scan, clinical stage is defined by T staging and N staging. In cases of type 4 and large type 3 tumors, staging laparoscopy is recommended for screening peritoneal dissemination and positive cytology that cannot be detected by CT. If distant metastases are diagnosed by CT (cM1), the patient is evaluated as cStage IVb. For liver metastases, enhanced MRI is effective for detecting small metastatic nodules; therefore, correct diagnosis of the number of liver metastases is available. Distant metastases of other sites (including lung, bone, adrenal gland, distant lymph node) should be confirmed by PET.

#### 3.4 Colorectal cancer staging

##### 3.4.1 T staging

The selected papers are summarized in Table 3. For T staging (Table 3A), Chen et al revealed that dual-energy CT showed a high accuracy, with a sensitivity of 90% and specificity of 97%. Komono et al advocated the new criteria using CT–colonography with multiplanar reconstruction to differentiate between T2, T3, and T4a. They focused on new blood vessels produced by tumor angiogenesis at the subserosal layer, designated as “bordering vessels.” They defined the criteria that tumors that do not involve bordering vessels and have a smooth outer border are considered T2, while those with...
3.4.2 Lymph node staging

For lymph node staging (Table 3B), the sensitivity was found to range from 44% to 73%, and the specificity ranged from 41% to 68%.84,95 The accuracies of these studies were around 50%–60%. PET showed a relatively high specificity of 84%, but a sensitivity of only 44%.91 These results highlight the requirement for more reliable modalities. Colon cancer patients were surgically resected regardless of preoperative nodal status, and thus clinical N staging is not essential in these patients. Neoadjuvant therapy is only considered for locally advanced colon cancer.97 For rectal cancer, neoadjuvant chemoradiation therapy is more common against nodal positive cancer in Western countries. Nonetheless, the accuracy of clinical staging has been reported to be medium.94

3.4.3 Diagnosis for distant metastases

The diagnostic accuracy of imaging to detect distant metastases is shown in Table 3C. Oh et al compared the use of MRI and PET-CT to detect liver metastasis.92 The per patient analysis revealed similar specificities and sensitivities between the modalities. On the other hand, the per nodule analysis showed that the sensitivity of PET-CT was 68.7%, which was significantly lower than that of MRI (96.2%). Colagrande et al and Moreno et al also demonstrated a high accuracy of MRI to detect liver metastasis.94,95 Figure 5 illustrates the superiority of MRI to detect small liver metastasis over enhanced CT. CT could only detect a 1-cm sized lesion in the S5 segment, whereas MRI was able to detect the lesions as well as another 4-mm sized lesion on the back side. A comparison of CT and MRI to detect liver metastasis in colorectal cancer was conducted. MRI was able to detect small metastasis of the liver that could not be detected by CT. Georgakopoulos et al revealed that PET-CT was able to detect extrahepatic disease, which was missed by conventional imaging in 50% of patients who were found to have liver metastasis prior to surgery.95 These findings may alter the treatment strategy.

### Table 3
  The summary of diagnostic modalities for TNM staging in colorectal cancer

| Year | Author [Ref] | Country | Journal | Modality | Number of patients | Sensitivity (%) | Specificity (%) | Accuracy (%) |
|------|--------------|---------|---------|----------|--------------------|----------------|----------------|-------------|
| 2014 | Cho SH [82]  | Korea   | Am J Roentgenol | MRI      | 146                | 74             | 87             | 85          |
| 2014 | Chen CY [83] | China   | PLoS One  | CT       | 103                | 90             | 97             | 95          |
| 2017 | So JS [84]   | Korea   | Ann Coloproctol | CT      | 285                | 90             | 68             | 55          |
| 2018 | Malmstrøm ML [85] | Denmark | Int J Colorectal Dis | CT | 615                | 65             | 89             | 49          |
| 2019 | Komono A [86] | Japan   | Int J Colorectal Dis | CT | 172                | 79             | 99             | 97          |
| 2019 | Korsbakke K [87] | Sweden | Acta Radiologica Open | CT | 383                | 28             | 93             | 74          |

| Year | Author [Ref] | Country | Journal | Modality | Number of patients | Sensitivity (%) | Specificity (%) | Accuracy (%) |
|------|--------------|---------|---------|----------|--------------------|----------------|----------------|-------------|
| 2014 | de Vries FE [88] | Netherlands | Eur J Surg Oncol | CT | 106                | 71             | 41             | 54          |
| 2016 | Ogawa S [89] | Japan   | Ann Surg Oncol | MRI    | 449                | 73             | 55             | 64          |
| 2017 | So JS [84]   | Korea   | Ann Coloproctol | CT      | 285                | 72             | 63             | 55          |
| 2017 | Lee JY [90]  | Korea   | Intest Res PET | PET | 220                | 44             | 84             | 67          |
| 2019 | Korsbakke K [87] | Sweden | Acta Radiologica Open | CT | 383                | 55             | 66             | 61          |

| Year | Author [Ref] | Country | Journal | Modality | Number of patients | Sensitivity (%) | Specificity (%) | Accuracy (%) |
|------|--------------|---------|---------|----------|--------------------|----------------|----------------|-------------|
| 2016 | Oh JW [92]   | Korea   | Biomed Res Int PET/CT | PET/CT | 67                | 95             | 100            | 97          |
| 2016 | Oh JW [92]   | Korea   | Biomed Res Int Gd-MRI | Gd-MRI | 67                | 98             | 93             | 98          |
| 2016 | Colagrande S [93] | Italy | Eur J Radiol | MRI | 115                | 97             | 85             | 96          |
| 2017 | Lee JY [90]  | Korea   | Intest Res PET | PET | 220                | 79             | 94             | 93          |
| 2017 | Lee JY [90]  | Korea   | Intest Res CT | CT | 220                | 79             | 87             | 86          |
3.4.4 | Algorithm of image modalities for clinical staging in colorectal cancer

Based on these results, we propose an algorithm of image modalities for clinical staging in colorectal cancer (Figure 6). Colonoscopy, CT, and MRI should preoperatively be performed in colorectal cancer patients. CT–colonography is useful for T staging, and MRI is much more sensitive than CT for the detection of small liver metastasis. Surgical resection should be performed in patients without distant metastasis, whereas PET-CT should be used to detect extrahepatic metastasis in patients with distant metastasis. Chemotherapy with or without surgery or radiation should be selected according to the involved lesions.

A comparison of the best diagnostic accuracy for clinical N staging among esophageal, gastric, and colorectal cancers are shown in Table 4. The common criteria for metastatic nodes were “round shape” and “enhancement” in gastrointestinal cancers. The optimal cutoff size to classify the positive lymph nodes differed according to the type of cancer as follows: 5-10 mm in esophageal nodes,86 8-10 mm in gastric nodes,21 and 4-5 mm in colorectal nodes.88,97 Although the diagnostic accuracy in esophageal cancer was relatively higher than that in gastric and colorectal cancers, the accuracies in all three types of cancer were unsatisfactory.

4 | DISCUSSION

This systematic review of clinical staging of gastrointestinal cancers included 55 articles published between January 2005 and July 2020 that were retrieved from PubMed/MEDLINE. Since the present review examined patient selection for neoadjuvant therapy, the main targets of diagnosis were T2-T4, positive nodes, and distant metastases. Although several systematic reviews have evaluated the performance of clinical staging for gastrointestinal cancers, most focused on just one cancer type. The present review evaluated the diagnostic modalities to detect T2-T4 invasion, nodal involvement, and distant metastases in patients with gastrointestinal cancers based on the studies published during the same period. Favorable diagnostic modality for lymph node metastasis in each type of cancer differed; however, the sensitivities ranged from 60% to 80%. PET-CT was the
best modality to detect distant metastases for esophageal cancer, staging laparoscopy was the best modality for detecting peritoneal metastasis of gastric cancer, and MRI was the best modality for detecting liver metastases of gastric cancer and colorectal cancer.

Detection of lymph node metastases in superficial gastric cancer is to differentiate tumors for ESD indication. Since superficial esophageal cancer is more likely have lymph node metastases than other gastrointestinal cancers,98 precise evaluation of lymph node metastases is essential to determine the indication for ESD. These topics are reviewed elsewhere.99,100

Detection of T4 invasion and distant metastases are the most important critical issues regarding the clinical staging of advanced esophageal cancer. Based on the present systematic review, CT was found to be the best modality to evaluate potential invasion to adjacent organ, while PET-CT or EUS/CT was useful to detect nodal metastases. Since subtotal esophagectomy is one of the most stressful surgical procedures, neither T4 invasion or distant metastases should be detected prior to surgery to differentiate noncurative tumors. Based on selected papers, although the sensitivities for the detection of T4 invasion were not high enough, the specificities were nearly 100%. The majority of suspected T4 cases were treated by chemoradiation rather than surgery; therefore, the number of patients included in the papers were limited. The positive predictive value for distant metastases gradually increased according to the time period of published papers; however, the sensitivities and accuracies remained unsatisfactory, although the resolution of PET images has improved during the last 10 years.101 The identification of patients who may not benefit from potentially curative surgery may be associated with high resolution. However, the present review also demonstrated that the use of PET-CT restaging resulted in a 5% false positive rate, which may introduce unnecessary physical and psychological intervention to the patient via additional testing and anxiety.

Clinical staging after chemoradiation therapy should be essential for esophageal cancer. Among previous reports using various diagnostic modalities, PET-CT may be the best tool for response assessment after neoadjuvant chemoradiotherapy. Stiekema et al reported that maximum standardized uptake value, metabolic tumor volume, and total lesion glycolysis were correlated with the pathologic response.102 Assessment of changes to these parameters may be the best tools for restaging after neoadjuvant therapy.

Important critical issues of clinical staging of gastric cancer include detection of early gastric cancer for ESD indication, T3 invasion and more with lymph node metastases positive for the indication of NAC, and distant metastases. The detailed indication of ESD is defined in the Japanese Gastric Cancer Association guidelines. Mucosal cancer is a basic target for ESD, and clinical distinction between mucosal and submucosal invasion by endoscopic examination is required. The positive predictive value for pT1b (sm) by endoscopic diagnosis was reported to be 63%–89%.54,103-105 and additional diagnostic values by EUS were not demonstrated in some reports.103,106 The diagnostic characteristics of submucosal invasion are not described clearly and diagnostic ESD is performed in some cases. In Western countries, the standard therapeutic strategy for advanced gastric cancer is NAC based on the results of pivotal clinical trials, such as the FLOT trial20 and others.16-18

Surgical outcomes of p stage I/II gastric cancer patients are favorable, and p stage III patients are the main target of NAC. However, p stage I/II patients were included in the NAC group in the FLOT trial due to clinical misdiagnosis. In JCOG1302A,23 the proportion of p stage I patients who were diagnosed as clinical stage III, T3/T4 and N1-3, and T3/T4 were 4.6%, 6.5%, and 12%, respectively. The sensitivities for p stage III patients were 52%, 65%, and 88%, respectively. Based on these findings, the eligibility criteria in JCOG1509 regarding NAC for advanced gastric cancer is defined as "T3/4 and N1-3." An essential consideration of clinical diagnosis of gastric cancer is "How can an accurate diagnosis of T3/4 and N positive be performed?" Difficulties remain concerning the accurate diagnosis of lymph node metastases of gastric cancer patients because lymph node evaluation by size alone has potential limitations.107 A cutoff value of 8 mm is commonly used, but smaller-sized lymph node metastases are frequently seen, especially for poorly differentiated adenocarcinoma. If a smaller cutoff value is defined for metastases, diagnostic "false positives" will be more frequent. Even a diagnosis of node positive/negative is not sufficiently accurate that we can give up the clinical N staging based on the number of metastatic nodes. The clinical diagnosis of peritoneal dissemination is commonly determined by ascites, thickness of omentum, hydrenephrosis, and definite disseminated nodules by CT imaging, but small disseminated nodules cannot be detected by imaging. Staging laparoscopy is recommended prior to surgery for advanced gastric cancer patients with possible peritoneal dissemination (linitis plastica, large-sized tumor, and suspicious findings of dissemination by imaging).

In colorectal cancer, the important factors to consider when selecting candidates for neoadjuvant therapy are tumor depth and distant metastases. In stage II/III, neoadjuvant therapy is uncommon and upfront surgery is the first priority in colorectal cancer.91 Therefore, these findings indicate that N staging is not important. One of the most important clinical features is liver metastases in colorectal cancer. Therefore, MRI should first be performed prior to surgery, which is reported to be better in detecting liver metastasis than PET-CT and CT.92,93,108 Afterwards, CT should be performed to detect lung metastases as well as for T staging. Regarding the diagnostic definition of lymph node metastases in colorectal cancer, Ogawa et al reported a better diagnostic accuracy using a cutoff size of 5 mm compared with 10 mm.89 These cutoff values are relatively smaller than those for gastric cancer.

### TABLE 4
Comparison of the best diagnostic accuracy for clinical N staging between esophageal, gastric, and colorectal cancers

| Reference cutoff size | Esophagus [96] | Gastric [21] | Colorectal [97] |
|-----------------------|----------------|-------------|-----------------|
| Sensitivity           | 67%            | 63%         | 73%             |
| Specificity           | 92%            | 66%         | 55%             |
| Accuracy              | 87%            | 64%         | 64%             |

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Important critical issues of clinical staging of gastric cancer include detection of early gastric cancer for ESD indication, T3 invasion and more with lymph node metastases positive for the indication of NAC, and distant metastases. The detailed indication of ESD is defined in the Japanese Gastric Cancer Association guidelines. Mucosal cancer is a basic target for ESD, and clinical distinction between mucosal and submucosal invasion by endoscopic examination is required. The positive predictive value for pT1b (sm) by endoscopic diagnosis was reported to be 63%–89%.54,103-105 and additional diagnostic values by EUS were not demonstrated in some reports.103,106 The diagnostic characteristics of submucosal invasion are not described clearly and diagnostic ESD is performed in some cases. In Western countries, the standard therapeutic strategy for advanced gastric cancer is NAC based on the results of pivotal clinical trials, such as the FLOT trial20 and others.16-18

Surgical outcomes of p stage I/II gastric cancer patients are favorable, and p stage III patients are the main target of NAC. However, p stage I/II patients were included in the NAC group in the FLOT trial due to clinical misdiagnosis. In JCOG1302A,23 the proportion of p stage I patients who were diagnosed as clinical stage III, T3/T4 and N1-3, and T3/T4 were 4.6%, 6.5%, and 12%, respectively. The sensitivities for p stage III patients were 52%, 65%, and 88%, respectively. Based on these findings, the eligibility criteria in JCOG1509 regarding NAC for advanced gastric cancer is defined as "T3/4 and N1-3." An essential consideration of clinical diagnosis of gastric cancer is "How can an accurate diagnosis of T3/4 and N positive be performed?" Difficulties remain concerning the accurate diagnosis of lymph node metastases of gastric cancer patients because lymph node evaluation by size alone has potential limitations.107 A cutoff value of 8 mm is commonly used, but smaller-sized lymph node metastases are frequently seen, especially for poorly differentiated adenocarcinoma. If a smaller cutoff value is defined for metastases, diagnostic "false positives" will be more frequent. Even a diagnosis of node positive/negative is not sufficiently accurate that we can give up the clinical N staging based on the number of metastatic nodes. The clinical diagnosis of peritoneal dissemination is commonly determined by ascites, thickness of omentum, hydrenephrosis, and definite disseminated nodules by CT imaging, but small disseminated nodules cannot be detected by imaging. Staging laparoscopy is recommended prior to surgery for advanced gastric cancer patients with possible peritoneal dissemination (linitis plastica, large-sized tumor, and suspicious findings of dissemination by imaging).

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At present, the number of positive nodes cannot be diagnosed accurately by imaging. Since the tumor depth is significantly associated with the number of positive nodes, combination diagnosis using tumor depth and clinical positive nodes may be the most reliable clinical diagnosis under the current performance of imaging technology. On the other hand, the accuracy of diagnostic imaging to detect distant metastases from gastrointestinal cancers is becoming more reliable with the use of PET-CT and/or MRI with the latest technologies. So far, we speculated that PET was useful for the esophagus squamous cell carcinoma, but less useful for gastric and colorectal adenocarcinomas. Highly antigenic tumors generally tend to develop swelling of metastatic lymph nodes, whereas low antigenic tumors tend to have smaller metastatic lymph nodes.

The rate of accurate diagnosis of conventional diagnostic imaging was evaluated in patients who underwent radical surgery without preoperative treatment. However, many advanced cancers will become candidates for preoperative treatment. Therefore, it will be necessary to perform diagnostic imaging before and after preoperative chemotherapy to monitor changes in staging and the rate of agreement with postoperative pathological staging. It is not possible to verify whether pretreatment staging was correct in patients undergoing preoperative chemotherapy. However, if the staging by diagnostic imaging after preoperative treatment matches the postoperative pathological staging, it may be possible to ensure the accuracy of the staging prior to treatment. In the future, more accurate pathological therapeutic effects and staging will be required after preoperative treatment. In patients receiving preoperative treatment, difficulties remain in terms of lymph node metastasis diagnosis and the usefulness of PET is predicted to become more important.

In conclusion, our literature review suggests that the recent diagnostic modalities can make precise differential diagnoses for T4, N1, and M1 for gastrointestinal cancers. However, the accuracy is still not sufficient to design preoperative treatment strategies. The most important purpose of clinical staging is to determine whether neoadjuvant therapy should be performed on each patient. Overstaging could occur in some patients without a standard algorithm for clinical staging and may lead to overtreatment. Accurate diagnostic modalities that adhere to a standard algorithm may improve both oncological outcomes and patient quality of life. Since there are only a few large-scale prospective cohort studies in this field, further multi-institutional prospective studies are required.

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APPENDIX 1

MEDLINE SEARCH TERMS

1. exp Tomography, Emission-Computed/
2. exp Tomography, X-Ray Computed/
3. exp Magnetic Resonance Imaging/
4. (computed tomography* or CT or CECT or MDCT or MSCT or magnetic resonance imaging or MRI or emission tomography or PET). mp. [mp = title, abstract, original title, name of substance word, subject heading word, keyword heading word, protocol supplementary concept word, rare disease supplementary concept word, unique identifier]
5. exp Ultrasonography/
6. (ultrasound or ultrasonography* or US or CEUS). mp. [mp = title, abstract, original title, name of substance word, subject heading word, keyword heading word, protocol supplementary concept word, rare disease supplementary concept word, unique identifier]
7. 1 or 2 or 3 or 4
8. 5 or 6
9. exp Colorectal Neoplasms/
10. exp Stomach Neoplasms/
11. exp esophageal neoplasms/
12. 9 or 11 or 12
13. 8 and 12
14. limit 13 to yr="2005 -Current"