The COlorectal NEoplasia Endoscopic Classification to Choose the Treatment classification for identification of large laterally spreading lesions lacking submucosal carcinomas: A prospective study of 663 lesions

Clementine Brule 1 | Mathieu Pioche 2 | Jeremie Alouys 1 | Jerome Rivory 2 | Sophie Geyl 1 | Romain Legros 1 | Florian Rostain 2 | Martin Dahan 1 | Hugo Lepetit 1 | Denis Sautereau 1 | Thierry Ponchon 2 | Emilie Auditeau 3 | Jeremie Jacques 3

1 Department of Endoscopy and Gastroenterology, Dupuytren University Hospital, Limoges, France
2 Department of Endoscopy and Gastroenterology, Edouard Herriot Hospital, Hospices Civils de Lyon, Lyon, France
3 Department of Epidemiology, Dupuytren University Hospital, Limoges, France

Correspondence
Clementine Brule, Department of Endoscopy and Gastroenterology, Dupuytren University Hospital, Limoges, France.
Email: brule.clementine@gmail.com

Abstract

Introduction: Optical diagnosis is necessary when selecting the resection modality for large superficial colorectal lesions. The COlorectal NEoplasia Endoscopic Classification to Choose the Treatment (COlNECCT) encompasses overt (irregular pit or vascular pattern) and covert (macroscopic features) signs of carcinoma in an all-in-one classification using validated criteria. The COlNECCT IIC subtype corresponds to adenomas with a high risk of superficial carcinoma that should be resected en bloc with free margins.

Methods: This prospective multicentre study investigated the diagnostic accuracy of the COlNECCT classification for predicting submucosal invasion in colorectal lesions >20 mm. Optical diagnosis before en bloc resection by endoscopic submucosal dissection (ESD) was compared with the final histological diagnosis. Diagnostic accuracy for the COlNECCT IIC subtype was compared with literature-validated features of concern considered to be risk factors for submucosal invasion (non-granular large spreading tumour [NG LST], macronodule >1 cm, SANO IIIA area, and Paris 0-IIC area).

Results: Six hundred 63 lesions removed by ESD were assessed. The en bloc, R0, and curative resection rates were respectively 96%, 85%, and 81%. The COlNECCT classification had a sensitivity (Se) of 100%, specificity (Sp) of 26.2%, positive predictive value of 11.6%, and negative predictive value (NPV) of 100% for predicting at least submucosal adenocarcinoma. The sensitivity of COlNECCT IIC (100%) to predict submucosal cancer was superior to all other criteria evaluated. COlorectal NEoplasia Endoscopic Classification to Choose the Treatment IIC lesions constituted 11.5% of all submucosal carcinomas.
Conclusion: The CONECCCT classification, which combines covert and overt signs of carcinoma, identifies with very perfect sensitivity (Se 100%, NPV 100%) the 30% of low-risk adenomas in large laterally spreading lesions treatable by piecemeal endoscopic mucosal resection or ESD according to expertise without undertreatment. However, the low specificity of CONECCCT leads to a large number of potentially not indicated ESDs for suspected high-risk lesions.

KEYWORDS
colonoscopy, EMR, endoscopy, ESD, optical diagnosis, submucosal carcinoma

INTRODUCTION

Real-time optical diagnosis of polyps enables the prediction of histology and selection of an optimal therapeutic approach. Several classification systems have been developed for colorectal polyps. The macroscopic aspect is included in the Paris$^1$ and LST$^2$ classifications, via white light imaging. An invasive carcinoma can be present despite the absence of an irregular pattern$^3$ in non-granular Laterally spreading tumour (LST) (NG LST) or under a macronodule or an area of depression (Paris O-IIc). Closed analysis (chromoendoscopy with/without magnification) yields useful details, including the pit pattern of the Kudo classification$^4$ and the vascular patterns of the Sano,$^5$ NBI International Colorectal Endoscopic (NICE),$^6$ and JNET classifications.$^7,8$ A national evaluation confirmed the poor knowledge of the usual classifications, and many benign lesions are wrongly treated by surgery.$^7$ Therefore, physicians from the research and development committee of the Société Française d’Endoscopie Digestive (SFED) wanted to simplify optical diagnosis by combining macroscopic, pit, and vascular patterns with the validated criteria used in the Paris, LST, NICE, Kudo, JNET, and Sano classifications based on the latest European guidelines.$^9$ This CONECCCT for COlorectal Neoplasia Endoscopic Classification to Choose the Treatmen (Figure 1) classification was first validated in a teaching program,$^10$ then validated by experts for optical diagnosis. The CONECCCT classification had higher inter-observer agreement and reproducibility than other classifications.$^{11}$ The European guidelines,$^{12–15}$ state that it is important to differentiate between lesions with a very low risk of undetected carcinoma, which should be resected by low-risk endoscopic procedures, and lesions with a high risk of at least superficial carcinoma, which should be resected en bloc with free margins. En bloc resection allows correct pathological examination and avoids underestimation of the invasion depth. The CONECCCT IIC subtype corresponds to lesions at high risk of submucosal carcinoma based on macroscopic aspects—such as NG LST or granular LST (G-LST)—with a macronodule > 10 mm or a depressed area (Paris O-IIc) and because of an irregular mucosal/vascular pattern (Kudo Vi and Sano’s IIIA; also type 2B of the JNET classification).

Here, we evaluated the diagnostic accuracy of the CONECCCT classification (optical diagnosis) compared to pathological examination, in particular for predicting at least submucosal carcinoma in large laterally spreading colorectal lesions (>20 mm) removed by en bloc ESD.

MATERIALS AND METHODS

Study design

This prospective study included lesions (>20 mm) that were consecutively treated by ESD in two referral centres. The study was approved by the Institutional review board and the Ethics Committee of Limoges University Hospital (n° 87RI20-0021_FECCo; NCT04592003). All patients were informed of their right to object to use of their data, which were collected and managed using Microsoft Excel (Microsoft).
Endoscopists and equipment

Four experienced operators performed optical diagnosis of lesions before resection by ESD. All operators had at least 5 years of experience in characterisation and were trainers in a teaching program for endoscopic characterisation. Before starting the study, each operator had experience comprising 50–100 colorectal ESDs in humans and animal models. The endoscopy suites were equipped with Olympus Exera III CV 190 units (190-generation colonoscopes) or Fujifilm 700 Eluxeo units (760-generation colonoscopes). All lesions were assessed by white light imaging and virtual chromoendoscopy without magnification.

Participants

We included all patients who were referred for endoscopic resection of a large lesion (>20 mm), regardless of its characteristics. In the participating centres, beginning in January 2016, we performed ESD for lesions >20 mm, for both benign lesions and lesions at risk of superficial cancer; this approach followed Japanese guidelines.

ESD allows endoscopic en bloc resection with negative margins (R0 resection). This facilitates pathological examination and prevents local recurrence, although it is associated with a higher risk of complications when performed by inexperienced endoscopists.

The exclusion criteria were adenocarcinoma associated with chronic inflammatory bowel disease, a recurrent or residual lesion after EMR, or prior piecemeal resection that created histological uncertainties and/or rendered characterisation incomplete because one or more of the LST, Paris, CONECT, KUDO, SANO, and JNET classifications could not be applied.

Lesions associated with obvious deeply invasive adenocarcinomas (CONECT III and SANO IIIB lesions) were analysed separately.

Lesion characterisation and data collection

All lesions were cleaned and subjected to both an overview and a close-up examination under white light imaging. We performed virtual chromoendoscopy without magnification under optimal illumination conditions. All lesions were described in real-time using the Paris, LST, Sano, and CONECT classifications. Patient data (age, sex), the type of endoscope used, and complete lesional data (size, location, type, presence of a macronodule >10 mm in diameter, Paris morphological analysis, Sano vascular pattern analysis, Kudo pit pattern analysis, and CONECT type) were recorded immediately after the procedure by the operators performing the ESD.

Diagnostic gold standard

Diagnosis employed the CONECT classification. A presumptive diagnosis was made in real-time (before resection). The lesions were pinned on corks and sent to the pathologists for histological diagnosis (the gold standard) using 2 mm sections. The pathologists received information concerning lesion morphology, location, and size; they were not informed of the optical diagnoses. Histological diagnoses...
were performed within 10 days of resection, in line with the revised Vienna classification. Serrated polyps were analysed using the criteria of the World Health Organization. Each report included the histological diagnosis, resection margin details, the extent of malignant differentiation, the invasion depth, and any evidence of lymphatic/vascular emboli or budding. Submucosal invasion was measured as described by the Japanese guidelines commencing from the lower edge of the muscularis mucosa. When the muscularis mucosa could not be identified, the depth of submucosal invasion was measured commencing from the mucosal layer. Lesions lacking deep invasion included those histologically diagnosed as hyperplastic polyps, sessile serrated lesions without dysplasia, low-grade dysplasia (LGD, Vienna 3), carcinoma in situ (pTIS, Vienna 4), or adenocarcinomas with submucosal invasion ≤1000 μm (Vienna 5, pT1a). Deep invasion was diagnosed if the submucosal invasion depth was >1000 μm (Vienna 5, pT1b) or if the lesion invaded the muscularis propria (≥pT2).

Definitions

Laterally spreading tumours were defined in accordance with the Japanese classification as lesions >10 mm that were wider than high. NG-LSTs featured smooth (non-granular) surfaces and were further divided into slightly elevated LSTs (0-IIa NG-LST) or pseudo-depressed (0-IIc NG-LST) LSTs. G-LSTs featured granular surfaces that were otherwise homogenous (GH-LSTs) or exhibited macro-nodules (granular mixed-type LSTs [GM-LSTs]). A protruding lesion was >2 cm and was more high than wide (Appendix A).

Outcomes

The principal outcome was the diagnostic accuracy of the CONECCCT classification in terms of predicting invasive adenocarcinoma of the submucosa (or deeper) compared to histology.

**FIGURE 2** Flow chart
The secondary outcomes were comparisons with other risk factors considered independently (macronodule >1 cm, Paris 0-IIc area, NG-LST, and SANO IIIA), comparisons of the endoscopes used (Olympus/NBI or Fujifilm/Blue laser imaging [BLI]), and the development of risk tables.

### Sample size and statistical analysis

Statistical analyses were performed with the aid of Stata ver. Twelve software (Stata Corporation, College Station). A p-value <0.05 was considered to indicate statistical significance. All analyses followed the standards for Reporting Diagnostic accuracy studies recommendations of 2015. Quantitative variables are described as means (±standard deviations, minima and maxima). Qualitative data are presented as counts with percentages. Diagnostic accuracy parameters (Se, Sp, positive predictive value (PPV), and NPV) are shown in percentages with 95% confidence intervals. The Pearson chi-squared test was used to compare the diagnostic accuracies of CONECCT IIC and SANO IIIA (compared to histology) in terms of predicting invasive adenocarcinomas in the submucosa or deeper. The number of subjects required was based on the hypothesis that CONECCT IIc would exhibit a Se of 84% when predicting adenocarcinomas involving the submucosa or deeper layers. Assuming a significance level of 0.05 and a precision of 3%, the number of subjects required was 574. The diagnostic accuracy of CONECCT and other classifications was compared by calculating areas under the receiver operating characteristic curves with the aid of the Pearson chi-squared test, as was the comparison of CONECCT diagnostic accuracies when different endoscopes were used. Multivariate analysis employed a descending, stepwise, logistic regression model. The explanatory variables were those yielding p-values <0.20 on univariate analysis.

### RESULTS

#### Flow chart

Between January 2016 and December 2019, 993 colorectal lesions were removed en-bloc using ESD; 330 lesions met the exclusion criteria. We thus assessed 663 lesions of 623 patients, including 467 colonic and 196 rectal lesions (Figure 2).
Cohort characteristics

The mean large diameter was 57.9 mm; 80.2% of lesions were >40 mm and 56% of lesions were >50 mm. Of all lesions, 43.7% (290) were in the recto sigmoid region, 8.5% (56) were in the left side colon, 6.8% (45) were in the transverse colon, and 40.9% (271) were in various regions of the right side colon. Morphologically, 12.8% (85) were protruded lesions, 66.4% (440) were G-LST lesions (26% GH-LST, 40.4% GM-LST), and 19% (126) were NG-LST lesions (11.9% flat and 7.1% pseudo-depressed). Features of concern included macronodules >1cm (53.4% of the lesions [354]), depressed areas (Paris 0-IIc; 13.9%), and SANO IIIA or Kudo VI areas (40% [265]). Lesions exhibiting submucosal invasion risk factors (macronodules >1cm, SANO IIIA areas, or NG-LST or depressed areas [0-IIc]) were classified as CONECCT IIC in 76% of cases (504).

The en bloc resection rate was 96%, the R0 resection rate was 85%, and the curative resection rate was 81%. Only 0.9% of the patients required surgery for a complication (Table 1).

Histological analysis

Histology revealed sessile serrated lesions in 12 cases, LGD (Vienna 3) in 216 cases (32.6%), pTis (V4) in 376 cases (57%), adenocarcinomas with submucosal invasion ≤1000 µm (Vienna 5-pT1a) in 29 cases (4.4%), adenocarcinomas with submucosal invasion >1000 µm (Vienna 5-pT1b) in 26 cases (3.9%), and ≥pT2 adenocarcinomas in four cases (Tables 1 and 2).

Primary outcome

The CONECCT IIC classification exhibited a sensitivity (Se) of 100.0%, a specificity (Sp) of 26.2%, a PPV of 11.9%, and a NPV of 100.0% (95% confidence interval 0.59–0.64) for predicting invasion of the submucosa or beyond. The accuracy was 32.6%. Of CONECCT IIA lesions lacking features of concern, none exhibited invasion of the submucosa or beyond (Table 3).

Performance of endoscopic predictive factors in terms of predicting submucosal carcinoma

The CONECCT Iic classification detected submucosal invasion significantly more sensitively (100%) than the other risk factors (Se values 66.1% for a Sano IIIA area, 76.3% for a macronodule >1 cm; and 27.1% for a depressed area [Paris 0-II]; all p < 0.05) (Table 3).

Prevalence and predictive factors of submucosal invasion

Of GH-LST tumours (without macronodules), 1.2% contained adenocarcinomas exhibiting submucosal invasion; the corresponding values were 11.9% for GM-LST tumours (with macronodules >1 cm), 5.1% for flat NG-LST tumours, 14.9% for pseudo-depressed NG-LST tumours, and 16.5% for protruding lesions. The submucosal
carcinoma rates were 17.3% for lesions with depressed (Paris 0-IIC) areas, 14.7% for lesions with SANO IIIA areas, and 8.7% for NG lesions. All GH-LST tumours with submucosal invasion exhibited depressed areas (Paris 0-IIC) (Table 2).

NG-LST tumours with submucosal invasion showed a SANO IIIA vascular pattern. Even in cases of LST-NG-PD, no submucosal cancer was found in the absence of SANO IIIA area (Table 4).

All G-LST tumours with submucosal invasion had a macroscopic morphologic risk factor of carcinoma (macronodule or depressed area); vascular abnormality (SANO IIIA) did not influence this risk of submucosal carcinoma (Table 4).

### Differences between the colon and rectosigmoid region

The rate of submucosal cancer was two fold higher in the rectum than in the colon, irrespective of the morphological risk factor:

- Macronodules (16.6% in the rectosigmoid region vs. 7.7% elsewhere; p = 0.013).
- SANO IIIA areas (20.9 vs. 8.8%; p = 0.003).
- CONECCCT IIc ranking (16 vs. 7.7%; p = 0.006; Appendix B).

### Endoscope model

The diagnostic accuracies of all predictive factors did not vary according to the endoscope used (Fujifilm/BLI or Olympus/NBI) (Appendix C).

### Submucosal cancer prediction based on the number of endoscopic predictive factors

When the number of features of concern increased (thus an NG-LST tumour, macronodule >1 cm in diameter, depressed area [Paris 0-IIC], and SANO IIIA area) the percentages of adenocarcinomas exhibiting submucosal invasion or beyond also increased: no feature of concern (n = 154) 0% submucosal invasion; one feature (n = 236) 8.5%; two features (n = 218) 11.9%; and three features (n = 55) 23.6% (p < 0.0001).

### Prediction of adenocarcinomas exhibiting deep submucosal invasion

Thirteen lesions subjected to ESD were finally classified as CONECCCT III and analysed separately. All were either resected via ESD or underwent surgery. Seven (53.8%) exhibited adenocarcinomas with submucosal invasion beyond 1000 μm, four exhibited superficial submucosal cancers (30.8%), and two (15.4%) exhibited intramucosal carcinomas. The Se, Sp, PPV, and NPV of the CONECCCT III classification in terms of predicting deep submucosal invasion were respectively 18.9%, 99.1%, 53.8%, and 95.5%.

### DISCUSSION

Endoscopic characterisation is recommended prior to resection of any superficial lesion. An effort to predict the final histology is necessary when selecting the optimal resection modality (piecemeal endoscopic mucosal resection [PM-EMR], ESD, or surgery) for large spreading lesions. The existing classifications (Kudo, SANO, and JNET) are based principally on overt carcinoma signs; they have been validated principally on lesions <2 cm using an NBI endoscope under magnification. For lesions >2 cm, the risk of covert carcinoma is related to macroscopic features (NG-LST status, macronodule >1 cm in diameter, O-IIc depressed area), but this has not been integrated into the classifications. The CONECCCT classification combines the macroscopic aspect with the pit/vascular pattern and...
allows any lesion to be described using any endoscope with a virtual chromoendoscopy function.

The CONECCIIc subtype combines overt signs of carcinoma validated by Japanese classifications (Sano IIIA, Kudo Vi, and JNET IIB) with covert signs significantly associated with macroscopic aspects (NG-LST status, a macronodule >1 cm in diameter, and a depressed area). The combination of these risk factors was essential because it is difficult to analyze the whole glandular and vascular patterns of large lesions (>2 cm).

The sensitivity of CONECCII (100%) for predicting submucosal cancer was superior to all other criteria; all classifications, including the CONECCII, had a very low PPV and accuracy, which led to difficulty identify submucosal cancer. Such lesions may always be curatively treated via PM-EMR; there is no need for ESD. However, these lesions represent only 25% of lesions referred for endoscopic resection. NG-LST with a regular pit and vascular pattern could be considered low-risk according to our results; they could be curatively removed by PM-EMR from a carcinologic perspective. This result increases the proportion of low-risk lesions to 30%.

Indeed, all granular lesions with submucosal cancer had a macroscopic morphologic risk factor—macronodule or depressed area or both (Figure 3). This is unsurprising because cancer can develop in the deepest part of the nodule without any superficial visible abnormalities.

Moreover, PM-EMR is often challenging in NG-LST or GM-LST with large macronodules because of frequent fibrosis. ESD may have a technical advantage for resection of such lesions.

A CONECCII status was associated with a non-negligible risk of adenocarcinoma with submucosal invasion. For example, Sano IIIA area, depressed area, or macronodule >1 cm in diameter was associated with a >10% risk of submucosal invasive cancers, of which 50% were superficial and curatively treated via ESD. Using the CONECCII status, no adenocarcinoma with submucosal invasion was missed, but only 5.8% of lesions with shallow submucosal invasion benefitted from ESD. In contrast, 6% of CONECCII lesions were not curatively resected via ESD; further surgery was necessary. However, ESD could treat curatively 47% of CONECCII lesions with small SANO IIIB areas, demonstrating the potential for overestimation by endoscopic characterisation. New criteria are needed to differentiate deep and shallow submucosal tumours and propose endoscopic or surgical resections. However, because primary ESD followed by salvage surgery

### Table 4

| LST G | MACRONODULE? |
|-------|--------------|
| +     | -            |
| 30.8% | 9.1%         |
| 16.7% | 12.5%        |
| 15.9% | 0%           |
| 7.1%  | 0%           |

| LST NG | SANO IIIA? |
|--------|-------------|
| +      | -           |
| 20%    | 0%          |
| -      | 8.5%        |

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has no carcinologic impact, it is difficult to propose primary surgery even for CONECCCT IIC lesions with three features of concern, because 80% of them are curatively treated by ESD.

Choosing a classification with high sensitivity decreases the rate of undertreated lesions (T1 with PM-EMR) but leads to a high number of ESDs. PM-EMR would have been a curative option for 322 adenomas (63.9% of CONECCCT IIC lesions) and an acceptable option for 128 intramucosal adenocarcinomas (25.4%). Several authors consider these ESDs to be unnecessary from a carcinologic perspective. However, ESD decreases the recurrence rate, as well as the need for uncomfortable and costly control colonoscopy. Indeed, a recent retrospective study of more than 500 patients from Korea reported that the 3-year cumulative total costs of patients treated via ESD or PM-EMR were similar. ESD was more expensive in year 1, but such patients required fewer control colonoscopies than the EMR group, despite the low recurrence rate (7.5%) in the EMR group. New ESD devices and strategies must be evaluated in comparative medico-economic trials (e.g., the ongoing French trial comparing ESD to PM-EMR [National Clinical Trials 03962868]).

Increasing specificity and PPV would improve the CONECCCT classification accuracy. In terms of the increasing risk of submucosal invasion as the number of features of concern increases, further studies are needed to define the minimum number of such features to trigger selection of ESD instead of PM-EMR. When three such factors were present, approximately 25% of lesions exhibited submucosal invasion; we suggest that piecemeal resection is unacceptable in such cases.

Although we believe that this approach should not be considered as overtreatment for a benign lesion, the low specificity of CONECCCT leads to a large number of potentially not indicated ESDs for suspected high-risk lesions. In fact, classification directly depends on the endoscopic resection technique used. If PM-EMR is used, a classification with high specificity to predict submucosal cancer is needed (e.g., the O-IIc criteria) to select ESD for these lesions. If ESD is used, as in the present study and our previous study, a classification with a high sensitivity (e.g., CONECCCT) is favourable because it prevents undertreatment.

The ESD results are very good, as in the present study: 96% en bloc resection rate, 85% R0 resection rate, and 49 min median procedure time for 57 mm lesions. The perforation rate was high in the present study, identical to the rates in the largest Asian studies for lesions >5 cm, which can be explained by the greater proportion of large lesions in the present study (10.6% in the study by Hong et al. and 8% in the study by Saito et al. for lesions >50 mm). Only 0.9% of patients had surgery for a complication.

Surgery is overtreatment in all cases, PM-EMR is undertreatment if there is a risk of submucosal cancer, and ESD with poor technical and carcinologic results is overtreatment of benign lesions. Moreover, the prevalence (pre-test likelihood) of submucosal cancer
in large LST is low (10%), thus statistically justifying a classification with high sensitivity.

Finally, at a time when extensive criteria are being evoked, our strategy allows adapted treatment by ESD of nine supplementary intermediate-risk lesions (submucosal infiltration 1000–2000 μm without budding or lymphovascular involvement).

The risk of submucosal invasion by lesions in the rectosigmoid region was two fold greater than the risk of lesions in the proximal colon, irrespective of the lesional features, as reported previously.

This higher distal risk is poorly understood but should be considered. ESD should be prioritised earlier for distal lesions.

The strengths of our study are its real-world design and the high numbers of lesions treated. This is one of the few studies to consider only lesions resected en bloc (ESD), reducing the risk of missing pathological information compared to studies based on PM-EMR. All lesions referred for endoscopic resection that lacked any sign of a deep submucosal feature were included, whatever their size; there was no selection bias. Moreover, this study is the first to compare two scopes from different suppliers with similar accuracies when applying the CONECCCT classification.

The first limitation of this study was the high endoscopic expertise of the four operators. The results should be confirmed by other teams. Moreover, because this was a real-time optical diagnosis study, inter- and intra-observer agreement could not be evaluated. However, we recently published a study based on pictures for which the inter- and intra-observer agreement were significantly higher than the agreements of other classifications.

The non-use of magnification was also a limitation of this study. However, therapeutic colonoscopes dedicated to ESD do not have a zoom function. Moreover, the combination of chromoendoscopy and zoom increases diagnostic accuracy for superficial submucosal adenocarcinoma and deep submucosal carcinoma, but it does not affect diagnostic accuracy for low- and high-risk lesions. Importantly, the use of endoscopes lacking a zoom function could be considered a strength that enables our results to be extrapolated throughout the West.

In conclusion, our optical diagnosis strategy (Figure 3), the CONECCCT classification, combines covert and overt signs of carcinoma to identify—with very high sensitivity (Se 100%, NPV 100%)—the 30% of low-risk adenomas in large laterally spreading lesions treatable by PM-EMR or ESD according to expertise without undertreatment (PM-EMR for T1 cancer). For the remaining 70% of lesions, ESD prevents unnecessary surgery for superficial sm-cancer, whereas PM-EMR prevents ESD for benign lesions at the cost of more surgery for carcinologic reasons and more colonoscopies to treat recurrences. A comparative trial incorporating technical and medico-economic endpoints is needed to clarify the optimal treatment modality in this situation; patients should be informed of the results and included in the decision-making process.

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CONFLICT OF INTERESTS

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DATA AVAILABILITY STATEMENT

Research data are not shared.

ORCID

Clementine Brule  https://orcid.org/0000-0002-1711-1845

REFERENCES

1. Participants in the Paris Workshop. The Paris endoscopic classification of superficial neoplastic lesions: esophagus, stomach, and colon. Gastrointest Endosc. 2003;58:S3–S43.
2. Oka S, Tanaka S, Kanao H, Oba S, Chayama K. Therapeutic strategy for colorectal laterally spreading tumor. Dig Endosc. 2009;21:S43–6.
3. Yamada M, Saito Y, Sakamoto T, Nakajima T, Kusuma R, Parra-Blanco A, et al. Endoscopic predictors of deep submucosal invasion in colorectal laterally spreading tumors. Endoscopy. 2016;48:456–64.
4. Kudo S, Rubino C, Teixeira C, Kashida H, Kagure E, Pit pattern in colorectal neoplasia: endoscopic magnifying view. Endoscopy. 2004;33:67–73.
5. Uraoka T, Saito Y, Ikematsu H, Yamamoto K, Sano Y. SANO’S capitular pattern classification for narrow-band imaging OF early colorectal lesions: SANO’S classification OF NBI. Dig Endosc. 2011;23:112–15.
6. Hayashi N, Tanaka S, Hewett DG, Kaltenbach TR, Sano Y, Ponchon T, et al. Endoscopic prediction of deep submucosal invasive carcinoma: validation of the Narrow-Band Imaging International Colorectal Endoscopic (NICE) classification. Gastrointest Endosc. 2013;78:625–32.
7. Sumimoto K, Tanaka S, Shigita K, Hirano D, Tamaru Y, Ninomiya Y, et al. Clinical impact and characteristics of the narrow-band imaging magnifying endoscopic classification of colorectal tumors proposed by the Japan NBI Expert Team. Gastrointest Endosc. 2017;85:816–21.
8. Sano Y, Tanaka S, Kudo S, Saito S, Matsuda T, Wada Y, et al. Narrow-band imaging (NBI) magnifying endoscopic classification of colorectal tumors proposed by the Japan NBI Expert Team: Japan NBI Expert Team classification. Dig Endosc. 2016;28:526–32.
9. Le Roy F, Manfredi S, Hamonic S, Piette C, Bouguen G, Riou F, et al. Frequency of and risk factors for the surgical resection of nonmalignant colorectal polyps: a population-based study. Endoscopy. 2015;48:263–70.
10. Pimentel-Nunes P, Dinis-Ribeiro M, Ponchon T, Repici A, Vieth M, De Ceglie A, et al. Endoscopic submucosal dissection: European society of gastrointestinal endoscopy (ESGE) guideline. Endoscopy. 2015;47:829–54.
11. Fabritius M, Gonzalez J-M, Béçq A, Dray X, Coron E, Brenet-Defour L, et al. A simplified table using validated diagnostic criteria is effective to improve characterization of colorectal polyps: the CONECCCT teaching program. Endosc Int Open. 2019;07:E1197–206.
12. Bonniaud P, Jacques J, Gonzalez JM, Dray X, Coron E, Leblanc S, et al. Endoscopic characterisation of colorectal neoplasia with the different published classifications: comparative study involving Conecct classification. Endosc Int Open. 2022 Jan 14;10(1) E145–E153 https://doi.org/10.1055/a-1613-5328
13. Pimentel-Nunes P, Pioche M, Albenzio E, Barr F, Delprez P, Ebibio A, et al. Curriculum for endoscopic submucosal dissection training in europe: European society of gastrointestinal endoscopy (ESGE) position statement. Endoscopy. 2019;51:980–92.
14. Ferlitsch M, Moss A, Hassan C, Bhandari P, Dumonceau J-M, Pas- 
patis G, et al. Colorectal polypectomy and endoscopic mucosal 
resection (EMR): European society of gastrointestinal endoscopy 
(ESGE) clinical guideline. Endoscopy. 2017;49:270–97.
15. Kamiński M, Hassan C, Bisschops R, Pohl J, Pellisé M, Dekker E, et al. 
Advanced imaging for detection and differentiation of colorectal 
neoplasia: European Society of Gastrointestinal Endoscopy (ESGE) 
Guideline. Endoscopy. 2014;46:435–57.
16. Tanaka S, Kashida H, Saito Y, Yahagi N, Yamano H, Saito S, et al. 
JGES guidelines for colorectal endoscopic submucosal dissection/ 
endoscopic mucosal resection. Dig Endosc. 2015;27:417–34.
17. Farhat S, Chaussade S, Ponchon T, Coumaros D, Charachon A, 
Barrioz T, et al. Endoscopic submucosal dissection in a European 
setting. A multi-institutional report of a technique in development. 
Endoscopy. 2011;43:664–70.
18. Schlemper RJ, Kato Y, Stolte M. Diagnostic criteria for gastrointes-
tinal carcinomas in Japan and Western countries: proposal for a new 
classification system of gastrointestinal epithelial neoplasia. J Gas-
troenterol Hepatol. 2000;15:G49–57.
19. Rex DK, Ahnen DJ, Baron JA, Batts KP, Burke CA, Burt RW, et al. 
Serrated lesions of the colorectum: review and recommendations 
from an expert panel. Am J Gastroenterol. 2012;107:1315–29.
20. Bossuyt PM, Reitsma JB, Bruns DE, Gatsonis CA, Glasziou PP, Irwig 
L, et al. StARD 2015: an updated list of essential items for reporting 
diagnostic accuracy studies. Radiol. 2015;277:826–32.
21. Sidhu M, Tate D, Desomer L, Brown G, Hourigan L, Lee E, et al. The 
size, morphology, site, and access score predicts critical outcomes of 
endoscopic mucosal resection in the colon. Endoscopy. 2018; 
50:684–92.
22. Ham NS, Kim J, Oh EH, Hwang SW, Park SH, Yang D-H, et al. Cost of 
endoscopic submucosal dissection versus endoscopic piecemeal 
mucosal resection in the colorectum. Dig Dis Sci. 2020;65:969–77.
23. Bordillon P, Pioche M, Wallenhorst T, Rivory J, Legros R, Albouys J, 
et al. Double-clip traction for colonic endoscopic submucosal 
dissection: a multicenter study of 599 consecutive cases (with video). 
Gastrointest Endosc. 2021;94:333–43.
24. Jacques J, Charissoux A, Bordillon P, Legros R, Rivory J, Hervieu V, 
et al. High proficiency of colonic endoscopic submucosal dissection 
in Europe thanks to countertraction strategy using a double clip and 
rubber band. Endosc Int Open. 2019;07:E1166–74.
25. Hong SN, Byeon JS, Lee B-I, Yang D-H, Kim J, Cho KB, et al. Pre-
diction model and risk score for perforation in patients undergoing 
colorectal endoscopic submucosal dissection. Gastrointest Endosc. 
2016;84:98–108.
26. Saito Y, Yamada M, So E, Abe S, Sakamoto T, Nakajima T, et al. 
Colorectal endoscopic submucosal dissection: technical advantages 
compared to endoscopic mucosal resection and minimally invasive 
surgery. Dig Endosc. 2014;26(Suppl 1):52–61.
27. Iwatate M, Sano Y, Tanaka S, Kudo Se, Saito S, Matsuda T, 
et al. Validation study for development of the Japan NBI Expert 
Team classification of colorectal lesions. Dig Endosc. 2018;30: 
642–51.

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classification for identification of large laterally spreading 
lesions lacking submucosal carcinomas: a prospective study of 
663 lesions. United European Gastroenterol J. 2022; 
10(1):80–92. https://doi.org/10.1002/ueg2.12194
**APPENDIX A**

**Definitions of LST**

| O-Ia | PROTRUDING LESION | GH-LST | MN-LST | Flat NG-LST | PD NG-LST |
|------|-------------------|--------|--------|-------------|-----------|

**APPENDIX B** Differences between the colon and rectosigmoid region

|                      | Recto-sigmoid (n = 291) | Colon (n = 372) | p value |
|----------------------|-------------------------|-----------------|---------|
| Size (mean +/- SD)   | 61.48 +/- 29.17         | 54.50 +/- 22.84 | 0.0007  |
| Type of lesion (n, %)|                         |                 |         |
| Protruding           | 54 (18.6%)              | 31 (8.3%)       | <0.0001 |
| Sm invasion          | 10 (18.5%)              | 4 (12.9%)       |         |
| G LST                | 208 (71.7%)             | 231 (62.1%)     |         |
| Sm invasion          | 25 (12%)                | 9 (3.9%)        |         |
| NG LST (n, %)        |                         |                 |         |
| Yes                  | 28 (9.7%)               | 98 (26.3%)      | <0.0001 |
| Sm invasion          | 4 (14.3%)               | 7 (7.1%)        | 0.23    |
| Macronodule (n, %)   |                         |                 |         |
| Yes                  | 199 (68.6%)             | 155 (41.7%)     | <0.0001 |
| Sm invasion          | 33 (16.6%)              | 12 (7.7%)       | 0.0013  |
| Depressed area (O-IIc) (n, %) |         |                 |         |
| Yes                  | 37 (12.8%)              | 55 (14.8%)      | 0.455   |
| Sm invasion          | 9 (24.3%)               | 7 (12.7%)       | 0.15    |
| SANO IIIA area (n, %) |                         |                 |         |
| Yes                  | 129 (44.5%)             | 136 (36.6%)     | 0.039   |
| Sm invasion          | 27 (20.9%)              | 12 (8.8%)       | 0.003   |

(Continues)
APPENDIX B (Continued)

|                      | Recto-sigmoid (n = 291) | Colon (n = 372) | p value |
|----------------------|-------------------------|----------------|---------|
| CONECCT Iic (n, %)   |                         |                |         |
| Yes                  | 244 (84.1%)             | 260 (69.9%)    | <0.0001 |
| Sm invasion          | 39 (16%)                | 20 (7.7%)      | 0.006   |
| Adenocarcinoma > V4 (n, %) |                 |                |         |
| Yes                  | 119 (41%)               | 77 (20.7%)     | <0.0001 |
| Adenocarcinoma > V5 (n, %) |                 |                |         |
| Yes                  | 39 (13.5%)              | 20 (5.4%)      | <0.0001 |

APPENDIX C Comparison of the endoscopes used in terms of predicting invasion of the submucosa or beyond

|                      | FUJIFILM 700 | OLYMPUS 190 | p value |
|----------------------|--------------|-------------|---------|
| Se (%)               |              |             |         |
| SANO IIIA            | 64           | 80          | 0.574   |
| CONECCT Iic          | 100          | 100         | 1       |
| Spe (%)              |              |             |         |
| SANO IIIA            | 62.5         | 58.7        | 0.641   |
| CONECCT Iic          | 25.3         | 28.9        | 0.588   |
| PPV (%)              |              |             |         |
| SANO IIIA            | 14           | 13.8        | 0.988   |
| CONECCT Iic          | 11.4         | 10.4        | 0.941   |
| NPV (%)              |              |             |         |
| SANO IIIA            | 94.8         | 97.3        | 0.668   |
| CONECCT Iic          | 100          | 100         | 1       |

Abbreviation: CONECCT, COlorectal NEoplasia Endoscopic Classification to Choose the Treatment.