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Recommendations for Reporting Tumor Budding in Colorectal Cancer based on the International Tumor Budding Consensus Conference (ITBCC) 2016


1Institute of Pathology, University of Bern, Bern, Switzerland
2Pathology and Laboratory Medicine, Mount Sinai Hospital, University of Toronto, Toronto, Canada
3Division of Molecular and Diagnostic Pathology, Niigata University Graduate School of Medical and Dental Sciences, Niigata, Japan.
4University Institute of Pathology, Lausanne University Medical Center, Lausanne, Switzerland
5Institute of Pathology, Kantonsspital Liestal, Liestal, Switzerland
6Dynacare Laboratory, Brampton, Toronto, ON, Canada
7Pathology Department, Saint-Antoine Hospital, Pierre et Marie Curie University, Paris, France
8Department of Pathology, Copenhagen University Hospital, Herlev, Denmark
9Department of Pathology, University Hospital Erlangen, Erlangen, Germany
10Department of Anatomic Pathology, University of California, San Francisco, USA
11Institute of Pathology, Medical University of Graz, Graz, Austria
12Department of Pathology, Radboud University Medical Centre, Nijmegen, The Netherlands
13Department of Clinical Pathology, Geneva University Hospital, Geneva, Switzerland
14Department of Pathology, University of Helsinki and Helsinki University Hospital, Helsinki, Finland
15Department of Pathology, St. Vincent's University Hospital, Dublin, Ireland
16Divisions of Anatomic Pathology and Mayo Clinic, Rochester, MN, USA
17Department of Surgical Oncology, Tokyo Medical and Dental University, Graduate School of Medical and Dental Sciences, Bunkyo-ku, Tokyo, Japan
18Pathology Department, Hôpital Cochin and Université Paris Descartes Sorbonne Paris Cité, Paris, France
19Department of Surgery, National Defense Medical College, Tokorozawa, Saitama, Japan
20Institute of Pathology, Klinikum Bayreuth, Bayreuth, Germany
21Pathology and Tumour Biology, University of Leeds, St James's University Hospital, Leeds, United Kingdom
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Corresponding author:

Prof. Alessandro Lugli, MD
Institute of Pathology
University of Bern
Murtenstrasse 31
3008 Bern, Switzerland
Ph: +41 31 632 9958
Fax: +41 31 632 4995
e-mail: alessandro.lugli@pathology.unibe.ch
Abstract

Tumor budding is a well-established independent prognostic factor in colorectal cancer but a standardized method for its assessment has been lacking. The primary aim of the International Tumor Budding Consensus Conference (ITBCC) was to reach agreement on an international, evidence-based standardized scoring system for tumor budding in colorectal cancer.

The ITBCC included 9 sessions with presentations, a pre-meeting survey, and an e-book covering the key publications on tumor budding in colorectal cancer. The “Grading of Recommendation Assessment, Development and Evaluation” method was used to determine the strength of recommendations and quality of evidence.

The following 10 statements achieved consensus: Tumor budding is defined as a single tumor cell or a cell cluster consisting of 4 tumor cells or less (22/22, 100%). Tumor budding is an independent predictor of lymph node metastases in pT1 colorectal cancer (23/23, 100%). Tumor budding is an independent predictor of survival in stage II colorectal cancer (23/23, 100%). Tumor budding should be taken into account along with other clinico-pathological features in a multidisciplinary setting (23/23, 100%). Tumor budding is counted on H&E (19/22, 86%). Intra-tumoral budding exists in colorectal cancer and has been shown to be related to lymph node metastasis (22/22, 100%). Tumor budding is assessed in one hotspot (in a field measuring 0.785 mm²) at the invasive front (22/22, 100%). A three-tier system should be used along with the budding count in order to facilitate risk stratification in colorectal cancer (23/23, 100%). Tumor budding and tumor grade are not the same (23/23, 100%). Tumor budding should be included in guidelines/protocols for colorectal cancer reporting (23/23, 100%).

Members of the ITBCC were able reach strong consensus on a single international, evidence-based method for tumor budding assessment and reporting. It is proposed that this method be incorporated into colorectal cancer guidelines/protocols and staging systems.
Introduction

The TNM classification system remains the gold standard for stratification of colorectal cancer patients into prognostic subgroups. Nevertheless, heterogeneity in survival within the same tumor stages points to the need for additional prognostic biomarkers. Tumor budding is defined as single cells or clusters of up to four cells at the invasive margin of colorectal cancer and can be stratified into peritumoral budding (PTB, tumor buds at the tumor front) and intratumoral budding (ITB, tumor buds in the tumor center). PTB can only be assessed in endoscopic or surgical resection specimens, whereas ITB can be assessed in both colorectal cancer biopsies and resection specimens. Both ITB and PTB are morphologic manifestations of epithelial-mesenchymal transition (EMT). A ZEB1, SNAIL1, TWIST1 positive microenvironment is conductive to the tumor budding phenotype. Additionally, tumor buds show loss of the adhesion molecule E-cadherin and express markers of an activated Wnt signaling pathway such as nuclear beta-catenin and APC. Furthermore, tumor buds express matrix metalloproteinases, cyclin D1, VEGF and p16, but do not show increased proliferation as determined by MIB1. Also of interest is that tumor buds may express stem cell markers such as EpCAM and ABCG5. The presence of ABCG5 in buds was associated with a worse prognosis in node-negative colorectal cancer patients.

Tumor budding is an independent adverse prognostic factor in colorectal cancer. It is also associated with a higher TNM stage, high tumor grade, the presence of lymphovascular invasion and consequently with lymph node and distant metastases. In colorectal cancer, tumor budding can potentially be applied as an additional quantitative prognostic factor to facilitate the management of colorectal cancer patients in three clinical scenarios. First, in endoscopically resected pT1 colorectal cancer, tumor budding is associated with an increased risk of lymph node metastases. Therefore, patients with budding may benefit from surgical resection. Second, in stage II colorectal cancer, the presence of tumor budding is an indicator of shorter disease-free survival compared to stage II colorectal cancer with low grade budding, or no budding. Therefore, stage II colorectal cancer patients with high grade tumor budding may be considered for adjuvant therapy. Third, ITB assessed in preoperative biopsies could help select patients who might qualify for neo-adjuvant therapy and could potentially predict tumor regression.

One of the main reasons that tumor budding is not routinely reported is the lack of a standardized scoring system that is simple and reproducible. The selection of the tumor slide, the location of counting, the applied stain (H&E vs immunohistochemistry) and the scoring system (cut-off vs continuous scale) are practical points that need to be clarified and then validated in multiple studies. Nevertheless, tumor budding seems to be a robust biomarker which retains its prognostic value independently of selected scoring systems used in different studies. The primary objective of the International Tumor Budding Consensus Conference (ITBCC), which took place in Bern in April 2016, was to reach agreement on an evidence-based, standardized scoring system for tumor budding to be used in international colorectal cancer guidelines and routine practice.
Material and Methods

Consensus process

The ITBCC was initiated by members of the ITBCC steering committee at the annual meeting of the United States and Canadian Academy of Pathology (USCAP) in March 2015 in Boston. The ITBCC was held in April 27-29 2016 in the Kursaal, Bern, Switzerland, and included participants from eleven countries. The primary objective was to determine whether consensus could be reached on a standardized scoring method for assessing tumor budding. The ITBCC included nine sessions with presentations, a pre-meeting survey, and an e-book covering the most important publications on tumor budding in colorectal cancer.

Steering Committee and participants

Twenty-five participants (22 gastrointestinal pathologists, 2 surgeons and 1 translational researcher) from the United States, Canada, Japan and Europe with expertise in tumor budding were invited to attend the face-to-face meeting in April 28-29 2016. Two participants could not attend the meeting, but participated in the pre-meeting survey. The steering committee, composed of six participants (AL, RK, RR, HD, GC and IZ), organized the meeting in Bern and prepared the pre-meeting survey and the e-book in collaboration with two chairmen (FB, PQ). All 25 participants were voting members and differences in vote numbers are due to participants who were not able to attend the meeting or could only attend on one day.

Pre-meeting survey

The 9-question pre-meeting survey was sent to all the participants before the meeting and results were presented during the sessions (summarized in Table 1).

Sessions

Based on the results of the pre-meeting survey the ITBCC was organized in nine sessions, with preliminary statements serving as a starting point for discussions (see below):

- Session 1: Definitions of tumor budding
  - Statement 1: Tumor is defined as a single cell or a cell cluster.
  - Statement 2: Tumor budding is different from poorly differentiated clusters.

- Session 2: Clinical scenarios and tumor budding
  - Statement 1: Tumor budding is an independent predictor of lymph node metastases in pT1 colorectal cancers.
  - Statement 2: Tumor budding is an independent predictor of survival in stage II colorectal cancer. Statement 3: Tumor budding is an adverse prognostic factor in preoperative biopsies of colorectal cancer.

- Session 3: H&E and immunohistochemistry for the tumor budding score
  - Statement 1: Tumor budding is assessed on H&E provided there are no features that limit its assessment (e.g. peritumoral inflammation, glandular fragmentation).
Statement 2: Immunohistochemistry, as optimal visualization tool, is applied in cases where H&E assessment is limited.

Statement 3: On immunohistochemical stained slides, digital software provides an objective budding count.

- **Session 4: Intratumoral and peritumoral budding**
  - Statement 1: ITB is applied on pre-operative biopsies.
  - Statement 2: The prognostic impact of tumor budding is independent of its location (i.e. ITB versus PTB)
  - Statement 3: All tumor buds (i.e. ITB+PTB) are counted in pT1 colorectal cancers and stage II colorectal cancers).

- **Session 5: Field number and size for the tumor budding score**
  - Statement 1: Tumor budding assessment in pre-operative biopsies and pT1 colorectal cancers is performed using the “hot spot” approach.
  - Statement 2: In surgical resection specimens, the “hot spot” and 10HPF methods are similar in terms of prognostic information (despite superior reproducibility of the 10HPF method) and are both used.
  - Statement 3: The field size definition is independent of the microscope type used.

- **Session 6: Cut-offs and continuous scale for the tumor budding score**
  - Statement 1: A cut-off for high grade tumor budding is used in order to facilitate meaningful risk stratification in colorectal cancer.
  - Statement 2: Upon specific request by the responsible clinician a continuous scale for tumor budding score (which allows more precise risk assessment) is provided.
  - Statement 3: The chosen method is sufficiently reproducible.

- **Session 7: reporting tumor budding**
  - Statement 1: Tumor budding should be a standard element in guidelines/protocols for colorectal cancer reporting.
  - Statement 2: Tumor budding should included in the next TNM classification as an additional prognostic factor equal to L, V or Pn stage.
  - Statement 3: Tumor budding should not be taken into account in the assessment of tumor grade.

- **Session 8 and 9: proceedings, conclusions and further studies**
Sources

A systematic literature search in PubMed for tumor budding in colorectal cancer was performed. An e-book including 38 key publications relevant to specific sessions was circulated to all participants ahead of the meeting: Session 1: definitions of tumor budding\textsuperscript{14,18,23-30}, session 2: clinical scenarios and tumor budding\textsuperscript{2,17-22,31-40}, session 3: H&E and immunohistochemistry for the tumor budding score\textsuperscript{11-13,41-46}, session 4: peritumoral and intratumoral budding\textsuperscript{2,20,21,47}, session 5: field number and size for the tumor budding score\textsuperscript{11,13,18,44-46,48,49}, session 6: cut-offs and continuous scale for the tumor budding score\textsuperscript{22}, session 7: reporting tumor budding\textsuperscript{11,12,18,44-46,49}.

Grading of Recommendations Assessment, Development and Evaluation (GRADE)

The quality of evidence and the strength of recommendations were based on the GRADE System\textsuperscript{50-58}:

level of evidence:

High: further research is very unlikely to change our confidence in the estimate of effect.

Moderate: further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low: further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low: any estimate of effect is very uncertain.

Strength of recommendation:

Strong: for intervention: desirable effects outweigh undesirable effects; against intervention: undesirable effects outweigh desirable effects.

Weak: for intervention: desirable effects probably outweigh undesirable effects; against intervention: undesirable effects probably outweigh desirable effects).
Results and discussion

Recommendations

Statement 1: Tumor budding is defined as a single tumor cell or a cell cluster of up to 4 tumor cells. GRADE: Strong recommendation, Vote: 22/22 (100%), quality of evidence high

Evidence and decision: Most outcome data are based on studies defining tumor budding as single tumor cells or clusters of up to 4 tumor cells, although a cut-off of 5 cells has also been used\textsuperscript{1,5}. The prognostic power of tumor budding is not affected by the number of cells that defines a cluster\textsuperscript{11,13-15,18}. Since 4 tumor cells is the most widely used cut-off for tumor budding, and since this cut-off distinguishes tumor budding from the novel histopathological parameter "poorly differentiated cluster (PDC)" which is defined as 5 or more cells\textsuperscript{27,59}, the ITBCC group agreed on a cut-off of up to 4 cells to define tumor budding (Figure 1).

Statement 2: Tumor budding is an independent predictor of lymph node metastasis in pT1 colorectal cancer GRADE: Strong recommendation, Vote: 23/23 (100%), quality of evidence high

Evidence and decision: In pT1 colorectal cancer, there is need for reliable predictors of local lymph node metastasis as their presence may help select candidates for radical surgery following endoscopic resection. Numerous studies as well as four meta-analyses have shown tumor budding to be a strong and independent predictor of local lymph node metastases in pT1 colorectal cancer\textsuperscript{15-17,31-33,60-68}. The ITBCC group therefore strongly recommends the reporting of tumor budding in pT1 colorectal cancer along with other histopathologic predictors of lymph node metastasis such as poor differentiation, lymphovascular invasion and depth/level of submucosal invasion\textsuperscript{16-18}.

Statement 3: Tumor budding is an independent predictor of survival in stage II colorectal cancer GRADE: Strong recommendation, Vote: 23/23 (100%), quality of evidence high

Evidence and decision: The UICC/AJCC TNM staging system remains the gold standard for prognostication in colorectal cancer and strongly influences adjuvant therapy decisions. Patients with stage III colorectal cancer are generally offered adjuvant chemotherapy, while those with stage II are not unless other high risk features are present (i.e. tumor perforation, lymphovascular invasion, serosal involvement [pT4a], poor tumor differentiation in microsatellite stable tumors, close/indeterminate/positive margins, perineural invasion, and low lymph node yield). However some stage II colorectal cancer patients show worse survival than stage III colorectal cancer patients who receive adjuvant chemotherapy\textsuperscript{69-71}. Numerous studies and meta-analysis have shown tumor budding to be an independent predictor of recurrence and survival in stage II colorectal cancer\textsuperscript{11,12,35-37,39,43,48,49}, with outcomes similar to stage III colorectal cancer\textsuperscript{38,72,73}. Based on these data the ITBCC group strongly recommends that tumor budding be included among the high risk factors reported in stage II colorectal cancer.
Statement 4: Tumor budding should be taken into account along with other clinico-pathological factors in a multidisciplinary setting GRADE: Strong recommendation, Vote: 23/23 (100%), quality of evidence high

Evidence and decision: In addition to the TNM classification, prognostic factors such as tumor grade (G), histological subtype, vascular invasion (V), perineural invasion (Pn), and margin status are routinely reported, in accordance with current guidelines and protocols69-71. Molecular biomarkers include microsatellite, KRAS mutation and BRAF mutation status74. Based on available evidence, the ITBCC group considers the prognostic value of tumor budding to be at least equivalent to that of V, G and Pn status, and therefore recommends that tumor budding (Bd) should be taken into account along with these and other clinicopathologic factors in the risk assessment of colorectal cancer.

Statement 5: Tumor budding is counted on H&E GRADE: Strong recommendation, Vote: 19/22 (86%), quality of evidence moderate

Evidence and decision: Outcome data for tumor budding are largely based on studies using H&E assessment. Relatively few studies have evaluated tumor budding by immunohistochemistry (IHC)13,14,18,45,75. Meta-analyses suggest that the prognostic power of tumor budding assessed on H&E and IHC do not differ materially11,13-15,18. Some studies have found IHC to be superior to H&E with regard to reproducibility and inter-observer agreement14,18,45, while others have not41,48. The ITBCC group recommends that tumor budding should be evaluated on H&E, since the vast majority outcome data are based on H&E assessment (particularly in pT1 colorectal cancer). In addition, the cost-effectiveness of H&E allows tumor budding to be assessed worldwide. This recommendation may change as more data on IHC assessment become available. It should be noted that tumor buds may be obscured by a peritumoral inflammatory infiltrate, making their identification difficult on H&E. In addition, tumor buds may, on occasion, be difficult to distinguish from reactive stromal cells. In such scenarios, pan-keratin IHC allows for better visualization of tumor buds, although it may also stain apoptotic bodies and cellular debris, which should not be counted as buds. While the final tumor bud count is performed on H&E, IHC can be helpful in challenging cases (i.e. glandular fragmentation, strong peritumoral inflammation) to confirm that the cells being counted are indeed tumor buds.

Statement 6: Intratumoral budding in colorectal cancer has been shown to be related to lymph node metastasis GRADE: Strong recommendation, Vote: 22/22 (100%), quality of evidence low

Evidence and decision: In contrast to conventional or so-called “peritumoral budding” (PTB) which is seen at the invasive front, intratumoral budding (ITB) refers to budding within the main tumor body. Although the terms ITB and PTB were introduced in 20112, ITB was first reported in 1989 in a series in rectal cancer biopsies and found to be associated with lymph node metastases23. More recently, ITB in pre-operative biopsies has been shown to correlate with high grade PTB, lymph node metastases and tumor regression grade in the corresponding colorectal cancer resection specimens20-22. Although ITB may prove to be a promising biomarker in the pre-operative management of colorectal cancer patients,
there is insufficient evidence to support its routine reporting at this time. The ITBCC group therefore recommends further research in this area before reporting of ITB is implemented in routine practice.

Statement 7: Tumor budding is assessed in one hotspot (in a field measuring 0.785 mm²) at the invasive front GRADE: Strong recommendation, Vote: 22/22 (100%), quality of evidence moderate

Evidence and decision: Standardization is crucial for the implementation of any biomarker in clinical practice. Several different scoring systems for tumor budding are currently in use as highlighted by recent reviews, meta-analyses and the ITBCC pre-meeting survey. Although the prognostic significance of tumor budding is largely independent of the scoring system used, the presence of a single international standard is critical for future clinical trials and diagnostic practice. To ensure standardization of field size, the ITBCC group recommends reporting by area (i.e. mm²) rather than objective lens (e.g. 20x) since the field of vision varies widely between different microscopes. The field area selected by the ITBCC group is 0.785mm², which corresponds to the field area used by Ueno et al. and adopted by the Japanese Society for Cancer of the Colon and Rectum (20x objective lens with a 20mm eyepiece field number diameter). A conversion table has been developed to normalize bud counts to 0.785mm² for microscopes with ocular lenses associated with different fields of vision (Figure 2).

Statement 8: For tumor budding assessment in colorectal cancer, the hotspot method is recommended GRADE: Strong recommendation, Vote: 22/22 (100%), quality of evidence moderate

Evidence and decision: Most studies have performed tumor bud counts in a single field with the highest density of tumor buds (“hotspot” method), while others have used multiple fields (e.g. “5 high power field” and “10 high power field” methods). Counting across multiple fields has the advantage of being more representative of the entire invasive front, and there is also some evidence of improved inter-observer agreement using this approach. On the other hand, counting multiple fields may “dilute” the final (mean) tumor bud count in cases with focally many tumor buds. The “hotspot” method therefore better reflects the maximal extent of tumor budding at the invasive front. The ITBCC group recommends the use of the “hotspot” method, since this is the method used in the vast majority of outcome based studies, and inter-observer agreement using this method is quite acceptable. However, to ensure that the field with the highest tumor budding is selected, it is recommended that 10 separate fields (20x objective) along the invasive front are scanned prior to counting of tumor buds in the single selected “hotspot”.

Statement 9: A three-tier system should be used along with the budding count in order to facilitate risk stratification in colorectal cancer GRADE: Strong recommendation, Vote: 23/23 (100%), quality of evidence moderate

Evidence and decision: For risk stratification based on tumor bud counts, most studies have used numerical cut-offs (including two-tier and three-tier systems), while a few studies have used a continuous
scale to predict probability of recurrence\textsuperscript{11-16,18}. Since tumor budding behaves as a continuous variable, a continuous scale provides more precise risk stratification than does a numerical cut-off\textsuperscript{22}. Nonetheless, cut-offs are more practical in the clinical setting and there is insufficient evidence to support the use of a continuous scale for tumor budding in clinical decision making. The ITBCC group recommends the use of a three tier system as used by Japanese Society for Cancer of the Colon and Rectum\textsuperscript{77}:

- 0-4 buds – low budding (Bd1)
- 5-9 buds – intermediate budding (Bd2)
- 10 or more buds – high budding (Bd3)

This system allows for risk stratification of both pT1 colorectal cancer and stage II colorectal cancer. In pT1 colorectal cancer, Bd2 and Bd3 are associated with an increased risk of lymph node metastasis\textsuperscript{32,61,63,65,68,78-81} whereas in stage II colorectal cancer, Bd3 is associated with an increased risk of recurrence and mortality\textsuperscript{15,16,31,37,67,82-85}. The ITBCC group recommends that, in addition to the Bd category, the absolute bud count is provided (e.g. Bd3 [count 17]). This avoids loss of information that may occur when applying a cut-off to borderline cases. For example, a bud count of 9 (Bd2) may be biologically similar to a bud count of 10 (Bd3), but falls into a different risk category. As indicated in statement 4, it is important that tumor budding is taken into account along with other clinico-pathological features in a multidisciplinary setting. Histological examples of Budding 1, 2 and 3 are represented in Figure 3.

**Statement 10:** Tumor budding should be included in guidelines/protocols for colorectal cancer reporting

GRADE: Strong recommendation, Vote: 23/23 (100%), quality of evidence high

Evidence and decision: Tumor budding is a well-established, independent prognostic factor in colorectal cancer with the potential to impact clinical decision making in pT1 and stage II disease (see Statements 2 and 3\textsuperscript{11-16,18}). The standardized, international, evidence-based method for tumor budding assessment agreed on at the ITBCC (Figure 4), provides a basis for future reporting of tumor budding in routine practice. The ITBCC group therefore recommends that tumor budding should be included in guidelines and protocols for the pathology reporting of colorectal cancer.

**Statement 11:** Tumor budding and tumor grade are not the same

GRADE: Strong recommendation, Vote: 23/23 (100%), quality of evidence high

Evidence and decision: Tumor budding is defined by the presence of single cells or clusters of up to four cells at the invasive front (see Statement 1), whereas tumor grade is defined by the proportion of tumor demonstrating gland formation. In multivariate analyses, the prognostic effect of tumor budding is independent of tumor grade and growth pattern\textsuperscript{1,11-16,18}. Therefore, the ITBCC group considers tumor budding to be different from tumor grade, and to have prognostic value that is independent of tumor grade.
All the statements and the corresponding GRADE recommendation and evidence are summarized in table 2.

**Additional practical aspects for tumor budding in colorectal cancer**

In some histological subtypes of colorectal cancer (e.g. mucinous, signet-ring cell, medullary and micropapillary), the assessment of tumor budding should be performed with caution. In mucinous and signet ring cell carcinomas, tumor buds suspended in pools of mucin should not be counted. In medullary carcinomas, discohesion or separation of tumor cells secondary to inflammation may be impossible to distinguish from true tumor buds. In micropapillary carcinoma, care should be taken not to include poorly differentiated clusters in the tumor bud count (see statement 1). Finally, glandular fragmentation secondary to heavy (often neutrophil rich) inflammation may be difficult to distinguish from tumor budding. In cases where an accurate tumor bud count cannot be performed, $B_d$ can be reported as “cannot be assessed” with an explanatory note. In rectal cancer resections after neo-adjuvant therapy tumor budding should not be reported since there are insufficient data regarding its prognostic significance.

**Conclusion and future perspectives**

Tumor budding is a well-established prognostic factor with the potential to refine clinical management decisions in patients with colorectal cancer. Consensus on a standardized, evidence-based method for tumor budding assessment at the ITBCC paves the way for future reporting of tumor budding in routine practice. The ITBCC is not intended to be an end-point, but rather a foundation for further multi-centre collaborations and clinical trials to prospectively validate and further refine the proposed ITBCC method. The ITBCC recommends that tumor budding should be included in future colorectal cancer guidelines and protocols.

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Figure legend

Figure 1: Example of tumor budding (A) which is defined as single tumor cells or tumor cell clusters at up to 4 cells. Example of poorly differentiated clusters (B) which are defined as 5 tumor cells or more.

Figure 2: Conversion table to adjust and standardize the tumor bud count for different microscope types.

Figure 3: Examples of different tumor budding grades (hotspot, 0.785mm2) at the invasive front of colorectal cancer based on the ITBCC 2016. A: Bd 1 (low), B: Bd 2 (intermediate) and C: Bd 3 (high).

Figure 4: Procedure proposed by the ITBCC 2016 for reporting tumor budding in colorectal cancer in daily diagnostic practice.