Endoscopy reporting standards

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OBJECTIVES: The Canadian Association of Gastroenterology (CAG) recently published consensus recommendations for safety and quality indicators in digestive endoscopy. The present article focuses specifically on the identification of key elements that should be found in all electronic endoscopy reports detailing recommendations adopted by the CAG consensus group.

METHODS: A committee of nine individuals steered the CAG Safety and Quality Indicators in Endoscopy Consensus Group, which had a total membership of 35 voting individuals with knowledge on the subject relating to endoscopic services. A comprehensive literature search was performed with regard to the key elements that should be found in an electronic endoscopy report. A task force reviewed all published, full-text, adult and human studies in French or English.

RESULTS: Components to be entered into the standardized report include identification of procedure, timing, procedural personnel, patient demographics and history, indication(s) for procedure, comorbidities, type of bowel preparation, consent for the procedure, pre-endoscopic administration of medications, type and dose of sedation used, extent and completeness of examination, quality of bowel preparation, relevant findings and pertinent negatives, adverse events and resulting interventions, patient comfort, diagnoses, endoscopic interventions performed, details of pathology specimens, details of follow-up arrangements, appended pathology report(s) and, when available, management recommendations. Summary information should be provided to the patient or family.

CONCLUSION: Continuous quality improvement should be the responsibility of every endoscopist and endoscopy facility to ensure improved patient care. Appropriate documentation of endoscopic procedures is a critical component of such activities.

Key Words: Colonoscopy/standards; Electronic reporting; Research report/standards; Review

Les normes des rapports d’endoscopie

OBJECTIFS: L’Association canadienne de gastroentérologie (ACG) a récemment publié des recommandations consensuelles sur les indicateurs de sécurité et de qualité en endoscopie digestive. Le présent article s’attarde sur la détermination des principaux éléments qui devraient figurer dans tous les rapports électroniques d’endoscopie détaillant les recommandations adoptées par le groupe consensuel de l’ACG.

MÉTHODOLOGIE: Un comité de neuf personnes a dirigé les indicateurs de sécurité et de qualité du groupe consensuel d’endoscopie de l’ACG, formé d’un total de 35 personnes ayant droit de vote qui avaient des connaissances sur les services d’endoscopie. Ils ont procédé à une analyse bibliographique détaillée des principaux éléments qui devraient figurer dans un rapport d’endoscopie électronique. Un groupe de travail a analysé le texte intégral de toutes les études sur des adultes et des humains publiées en anglais ou en français.

RÉSULTATS: Les éléments à inclure dans le rapport normalisé sont le nom et le montant de l’intervention, le personnel présent, la démographie et l’historique des patients, les indications de l’intervention, les comorbidités, le type de préparation intestinale, le consentement à l’intervention, l’administration de médicaments avant l’endoscopie, le type et la dose de sédatif utilisé, l’étendue et l’exhaustivité de l’examen, la qualité de la préparation intestinale, les observations pertinentes et les résultats négatifs pertinents, les effets indésirables et les interventions en résultant, le confort du patient, les diagnostics, les interventions endoscopiques effectuées, le détail des échantillons pathologiques, le détail des dispositions de suivi, les rapports de pathologie connexes et, si elles sont disponibles, les recommandations de prise en charge. Le résumé de l’information devrait être fourni au patient ou à sa famille.

CONCLUSION: L’amélioration continue de la qualité devrait incomber à chaque endoscopiste et établissement d’endoscopie afin de garantir de meilleurs soins aux patients. La consignation conviviale des interventions endoscopiques est un élément essentiel de ces activités.

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membership of 35 voting individuals with knowledge on the subject relating to endoscopic services. This multidisciplinary group included gastroenterologists, surgeons, gastroenterology nurses, health policy experts and a lawyer. In addition, nine subcommittees were mandated to address specific issues in greater detail, including a group that was tasked with reviewing the literature pertaining to endoscopy reporting standards. The methodology behind the consensus group process has been described in the CAG consensus guidelines on safety and quality indicators in endoscopy (3). The present article will focus specifically on the identification of key elements that should be found in all electronic endoscopy reports and the resulting recommendations adopted by the CAG consensus group.

Literature search
A comprehensive literature search was performed in several databases including Ovid MEDLINE, Embase, the Cochrane Library and ISI Web of Knowledge from 1980 to March 2012. All databases were searched using a validated search string specific to colonoscopy and quality reporting. Recursive searches and cross-referencing were also performed; manual searches for articles were performed after the initial search. All full-text, published, adult and human studies in French or English were included. Details regarding the consensus conference and voting have been reported previously (3).

RESULTS/DISCUSSION
The relevant statement pertaining to the content of the standardized report can be found in the main consensus meeting document reported previously (3). Detailed below are the results of the literature search leading to the justification of the different items to be entered in a standardized report as listed in Table 1.

Components to be entered into the standardized report
Identification of procedure, timing, and procedural personnel: Type of procedure, date and time of procedure and name of endoscopists and assistants are elements that should be documented in each endoscopic report.

Patient demographics and history: There are differences in the incidence rate of adenomas and mortality of CRC based on age and sex. Therefore, they are important risk factors that should be documented in the standardized electronic report to enable meaningful analysis of adenoma detection rates and prevalence estimates (7).

Indication(s) for the procedure: Colonoscopists should be familiar with the appropriate indications for colonoscopy. Screening and surveillance for colonic neoplasia represent the most common indications for a colonoscopy; emphasis is, therefore, made on specific quality-control issues relative to these two indications (8). Statement 3 from the CAG consensus guidelines (3) states that “endoscopic procedures are performed for an appropriate, clearly documented indication, consistent with current, evidence-based guidelines”. The indication for every procedure should be documented in the procedure report, and the indication should be consistent with accepted guidelines to ensure continuous quality improvement. There is evidence that the diagnostic yield of endoscopy is significantly increased when consensus guidelines for appropriate indications are outlined and followed; recent experience also suggests that such practice facilitates appropriate triaging of requests amid already stretched endoscopic resources with the advent of population-based CRC screening (AN Barkun, personal communication). More positive outcomes arise when procedures are performed with appropriate indications. Consensus guidelines provide explicit statements of appropriate indications for endoscopic procedures (9). Unfortunately, studies indicate that 11% to 39% of endoscopic procedures are performed for inappropriate indications (9), and that surveillance endoscopies may be performed at inappropriate intervals or unnecessarily. It appears that a lack of knowledge by the endoscopists about the proper intervals is an important factor. Even those who are fully aware tend to not respect guidelines and perform surveillance colonoscopy sooner than recommended (10). In all cases, a nonstandard indication should be clearly justified in the report. The American Society for Gastrointestinal Endoscopy and the United States Multi-Society Task Force on Colon Cancer have published appropriate indications for colonoscopy (2), as has the CAG, with corresponding suggested target wait times (11). In addition, the appropriateness of screening or surveillance intervals for colonoscopy screening need be assessed according to contemporary guidelines that have recently been updated (12).

Comorbidities: Physical status classifications of The American Society of Anesthesiologists (ASA) have been used by clinicians for more than 50 years to predict perioperative morbidity and mortality. Although this tool has not been validated by studies in digestive endoscopy, the classification is widely accepted as a surrogate of comorbidity (7). The classification category serves to modify the setting and enhance precautions when needed. For example, patients with ASA class 3 or higher are considered to be at high risk for cardiopulmonary events, and endoscopists should consider performing procedures on ASA class 3 patients in a setting that can accommodate adverse events (8).

Type of bowel preparation: The type of bowel preparation should be recorded in each report, and should also address whether a same-day or split-preparation approach was used (13,14).

Consent for the procedure: Documentation of consent is a component that also needs to be entered in the endoscopy report; the CAG consensus indicated in statement 1 that “For a patient to give a physician informed consent to perform an elective endoscopic procedure, the patient must be advised, in a timely fashion, of all relevant information about the procedure, its risks, benefits and alternatives, if any, and be given an opportunity to ask questions that the physician must answer” (3). Lieberman et al (2) further suggest that obtaining consent from the patients is an important component of quality that needs to be documented in the endoscopic report. The risks of endoscopic procedures include bleeding, perforation, infection, adverse events due to sedation, bad diagnosis, missed lesions and other complications. Some have suggested that the informed consent that notifies the patient of all these significant adverse events and the possibility of failure to

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### TABLE 1

Required endoscopy report elements

| Report field                                      | 1. Type of procedure | 2. Date and time of procedure | 3. Name of endoscopist | 4. Name(s) of assistant(s) | 5. Age and sex of patient | 6. Indications(s) for procedure | 7. Comorbidities | 8. Type of bowel preparation | 9. Documentation of consent | 10. Type and dose of sedation used | 11. Other medication and related information | 12. Extent and completeness of examination | 13. Quality of bowel preparation | 14. Relevant findings | 15. Pertinent negatives | 16. Adverse events and resulting interventions | 17. Patient comfort | 18. Diagnoses | 19. Endoscopic interventions performed | 20. Details of pathology specimens | 21. Details of follow-up arrangements | 22. Appended pathology report(s), when available | 23. Management recommendations | 24. Information provided to patient and/or family |
|--------------------------------------------------|----------------------|-----------------------------|------------------------|---------------------------|---------------------------|--------------------------------|----------------|-------------------------------|-----------------------------|----------------------------------|-----------------------------------|----------------------------------|----------------------------------|-----------------------------|----------------|---------------------------------|-----------------------------|---------------------------|-------------------------------|-----------------------------|----------------|---------------------------------|-----------------------------|---------------------------|-------------------------------|-----------------------------|
detect neoplasia in the colon even if present should be documented in the standardized colonoscopic report (8) in addition to a documented description of alternatives, when available.

Pre-endoscopic administration of medications: Any such relevant administration should be recorded; however, with regard to antibiotic prophylaxis, clinically significant infections following a colonoscopic procedure are very rare. Thus, evidence that supports such administration is very limited – in fact, no standard recommendations currently exist for such practice (15,16).

Type and dose of sedation used: The method and intended level of sedation should be recorded in all cases (8). Colonoscopy reports should indicate whether anesthesia or nursing staff participated in the administration of sedation. Objectives and guidelines have been described in detail elsewhere (17). Some of the objectives include assuring the patient’s well-being, providing sedation adequate for the intervention and allowing the patient to recover his/her initial state after the procedure. It has been shown that the use of sedation increases rates of cecal intubation and polyp detection (18). Appropriate sedation also favourably influences both completeness and the risk of acute complications (19). It is reasonable to believe that sedation would render the procedure more tolerable to the patient, in turn improving the potential results of polyp and adenoma detection rates of the procedure, although the practice of sedation and the medications used vary widely across the world.

Extent and completeness of examination: Cecal intubation rates have been reported in previous quality assurance studies (20). Completeness of the colonoscopy is critical to the adequacy of the examination and its benefits with regard to detection of neoplasia (5,21), and perhaps favour the subsequent prevention of cancer as recently suggested (22). From the pertinent quality indicators described previously (3), visualization of the cecum by notation and photodocumentation of landmarks should appear in every report to confirm completion of the procedure. In colonoscopy, cecal intubation is a good indicator of completeness and recognized performance, which is pertinent for future practice evaluations (2,23). Moreover, identification of two of the following key landmarks should also be included in the standardized electronic report: the appendiceal orifice, the ileocecal valve or the cecal strap fold (2). Photography of the landmarks, although subject to interpretation, is also recommended and will be discussed further below. If for some reason the procedure cannot be completed, this must be documented in the report and another colonoscopy should be scheduled.

It has also been recommended that the following times should be recorded: insertion of the endoscope; start of withdrawal from cecum; and complete withdrawal of the endoscope (8). Withdrawal time is the time required to withdraw the colonoscope from cecal intubation to the anus. There exists a significant correlation between withdrawal time and adenoma and polyp detection (although the significance of those detected has been brought into question); thus, sufficient time must be allotted for the removal of the scope (24,25). Withdrawal time has been shown to increase the detection rate of colonic neoplastic lesions with varying reported threshold values. The most influential publication on this topic suggested the endoscopist spend at least 6 min or more withdrawing the scope. Thus, the CAG has set the mean withdrawal time at 6 min under normal conditions (no tissue sampling or staining, and no polypectomy) (2). However, many factors, such as the length of the intestine and bowel preparation quality, can affect withdrawal time, influencing the polyp detection rate (26). Therefore, withdrawal time should be considered as an indirect quality indicator of colonoscopy and, although currently adopted by many workgroups and societies, including the CAG, may not withstand the test of time as a useful, independent quality indicator. The documentation of withdrawal time should be in every report, but is less interpretable and of less use when applied to a procedure requiring biopsies or polypectomy. Reasons for a withdrawal time shorter than recommended should be documented in the standardized electronic report.

Finally, for the procedure to be complete, retroflexion can be considered, although routine rectal retroflexion during colonoscopy has been shown to carry a low yield for the detection of advanced neoplasia (28).

Quality of bowel preparation: The quality of bowel preparation should be recorded in all cases, using a validated scale (or the very least in the context of a population-based screening program) (29,30) because it is a necessary indicator in determining the appropriate interval for the examination (8). Indeed, the quality of bowel preparation is an indicator of quality and performance recognized and relevant for the evaluation of colonoscopic practice (2). Poor bowel preparation is a major reason for an inability to meet preplanned appropriate screening intervals. It can also lead to prolonged cecal intubation and withdrawal time, and a reduced polyp detection rate (2). Good bowel preparation is also associated with a higher proportion of complete colonoscopies (19). Current guidelines established by the United States Multi-Society Task Force for Colonoscopy state that a preparation should be judged adequate if polyps ≥5 mm can be detected (7). Suggested intervals for repeat examinations in case of poor colonic preparations have recently been proposed (12).

Relevant findings and pertinent negatives: Appropriate description of findings is required; a number of descriptor compendia exist (31). In the case of colonic polyps, characteristics to be recorded include polyp number, size and location. This information will permit the subsequent tracking of adenoma detection rates. Studies have shown that polyps and adenoma removal results in a lower than expected incidence of CRC (22). The incidence is lowered to 76% to 90% after polypectomy (32,33). More than 95% of the detected polyps should be removed and sent to pathology. Small hyperplastic-appearing rectal polyps (<5 mm, sessile) do not require removal (34). If in doubt, a biopsy should be obtained to confirm the histology of the polyp and its location adequately described (7,8). Depending on the nature of the findings, where appropriate, widely recognized scales to better standardize reporting should be used (eg, the Forrest score in peptic ulcer bleeding [35], or the Mayo score for inflammatory colitis [36]). Pertinent negatives should also be specified where appropriate.

Adverse events and resulting interventions: In each report, there should be documentation of unplanned interventions during the procedure, if applicable. The record should reflect any intra- and postprocedural complications. Currently, no link exists between postprocedural complications that were not recognized at the time of the procedure and the endoscopic report in most units. Optimally, such a link between databases should be established. Indeed, a system should be set up to report and evaluate these postprocedure complications so that they can be discovered and corrected. The optimal standardized time for capturing such events remains controversial (37). Furthermore, adverse events should be recorded using relevant, standardized descriptions and validated scales (38,39).

Patient comfort: Patient comfort during the procedure should be documented, ideally using validated scales (40).

Diagnoses: The diagnosis should take into consideration all of the available data derived from history, laboratory, radiographies and new endoscopic findings. It should also be performed using standard terminology and validated scales, if appropriate (for example, the Los Angeles classification scoring of esophagitis) (41).

Endoscopic interventions performed: A clear statement of what the endoscopist did during the procedure should be included in the report, again using standard terminology and descriptions. Also, the number and location of the biopsies performed should be recorded.

Details of pathology specimens: The details of the polyps seen and resected should be part of all reports, with a clear description of whether tissue was sent to pathology and what sample is present in each container.

Details of follow-up arrangements: Details about the recommendations for discharge planning and follow-up arrangements should be included with the colonoscopy report and given to the patient. Endoscopists should indicate the expected interval for the next examination, recommended according to contemporary published surveillance guidelines, or if there is reason to deviate from the guidelines, as stated above (2). Whenever biopsies are performed, final recommendations and treatment are made after the pathology results.
are reviewed; however, the endoscopist should ensure that there is a system in place to communicate these final arrangements to both the patient and referring clinician (8).

Appropriate management following endoscopy (in addition to appropriate preparation before the procedure) must also be documented when managing a patient undergoing antithrombotic therapy. Guidelines have been proposed by the American Society for Gastrointestinal Endoscopy (42), the British Society of Gastroenterology (43), the American College of Chest Physicians (44), and the American College of Cardiology/American Heart Association (45). Practitioners need to stay up to date with the introduction of newer medications and recommended optimal times of discontinuation, if indicated, such as new anticoagulants such as dabigatran (46).

Appendix pathology report(s), when available: No specific mechanism or manpower support currently exist in most Canadian units to allow for a reliable and complete link between the databases that generate the endoscopic report and the pathology results (47). An effort must be made to ensure that the pathology results are conveyed in a timely fashion to both endoscopist and referring physician, and that adequate documentation be ensured through providing some form of linkage of endoscopic findings and subsequent histological characterization.

**Management recommendations:** When necessary, the management of antithrombotic prophylaxis should be outlined in the report as discussed above (42).

**Information provided to patient and/or family:** The CAG consensus specified in statement 9, further discussed below, that all patients should be provided with written information regarding the procedure the same day. The information should include procedural findings, follow-up plans and treatment, symptoms to watch for and the next steps that should be taken (3).

### THE ENDOSCOPY REPORT PLATFORM

#### Need for an adapted system

A complete endoscopy report is an essential element of a quality endoscopy service. Previously, narrative reporting was commonly used, but it is often associated with incomplete documentation, variations in the recording of positive findings, pertinent negative findings and other procedural details. An electronic standardized endoscopy report template results in effective communication of procedural findings, and successful practice audit and quality improvement processes that are exceedingly difficult to achieve using narrative or written reports.

This is why the CAG consensus indicated in statement 20: “Endoscopic procedures should be reported in a standardized electronic format, including mandatory reporting fields, to provide full documentation of all necessary clinical and quality measures” (3). A country-wide initiative is currently underway in an attempt to define national synoptic reporting standards in digestive endoscopy (P Rossos, personal communication).

Colonoscopy reporting practices of clinicians are highly variable and often suboptimal. Studies have revealed substantial variation in the completeness of endoscopic text reports and in adherence to the use of standard terms. Inconsistent endoscopy reporting includes differences in disease definition such as the description and quantification of mucosal inflammation in ulcerative colitis (48). Reports also show marked variations in the completion of different report elements. The items that often score most poorly are demographic data, patient history, preparation quality with visualization and procedure interpretation. Other report elements that vary greatly include lesion identification and removal, and sedation practice (49). The use of grading systems is suggested; however, to date, the recommended systems are disease specific and require further assessment before adopting them for use in standardized endoscopy reporting.

It is well established in the literature that a well-structured reporting system leads to improved completeness in endoscopy reports, and this can be achieved with a standardized electronic reporting system. The value of electronic reporting also lies in its ability to establish a method and process of increased standardization facilitating timely audits, benchmarking and data archiving.

**Standardization of electronic reports includes the mandatory reporting of elements.** Electronic data, such as digital transcripts of dictated reports, are preferable to handwritten reports but are still considered inferior to a standardized electronic report.

### Benefits of an electronic report

Comparing handwritten, dictated and computerized reports, Soehkoe et al (50) showed that preparing an endoscopy report using a computer-generated method does not take more time than preparing a report the conventional way, such as with dictation. Electronic reports also offer some advantages over handwritten and dictated reports. To have a complete endoscopic report, clinicians need to capture images of the gastrointestinal (GI) tract, for example, as recommended by the European Society for Gastrointestinal Endoscopy (51). In handwritten and dictated reports, images are printed on photographic paper with ordinary printers, a practice that is expensive. One of the advantages of a computer-generated report is that an unlimited number of endoscopic images can be stored digitally in the electronic database. Real-time comparison with previously captured images is also possible with such a system. Although not the aim of the study, the authors determined that the standardized text in computerized, pre-defined reports also enables statistical analysis of endoscopic findings, whereas the use of free-text handwritten and dictated reports does not do so in a direct way. Therefore, the capacity to store all endoscopic findings, including endoscopic images, in a database offers an additional advantage over handwritten and dictated reports. Many examples of such computer-generated, pre-defined electronic reports are now used globally (52,53), with examples of systems that offer a quick, user-friendly way of report writing; however, none produce a synoptic report. Furthermore, studies have shown that a standardized electronic report system produces superior reports in terms of completeness of the endoscopic report compared with free-text reports that are handwritten or dictated (52).

**Cost of electronic reporting**: Groenen et al (54) assessed the costs of the different ways to generate endoscopic reports, comparing hand-written, dictated and computerized reports. It was concluded that electronic reports were beneficial and cost effective in the long run. Although they require a larger initial acquisition cost compared with other means of reporting, electronic reports gain their cost advantage after five years. The high initial investment is due to the need for hardware, software and linkage to other medical computerized systems. After five years, the cost per report declines below the cost of hand-written and dictated reports. The more conventional ways of reporting require more personnel, increasing the risks of errors and costs. Electronic reporting minimizes material, workspace and personnel costs. Furthermore, the cost of printing and storing images is less in computerized reporting because only storage capacity is required to store the recommended number of images per procedure.

All the aforementioned advantages were taken into account by the CAG consensus group, which recommends the routine use of a standardized electronic reporting system in digestive endoscopy.

**Electronic image capture**

High-quality videendoscopic imaging is currently available to endoscopists to document photographs of specific parts of the GI tract during the course of an endoscopic examination. Imaging can be used to confirm cecal intubation. However, a problem occurs due to the non-specific appearance of cecal landmarks in many individuals (55). Variations in normal cecal anatomy limit the extent to which still imaging can be used as a definite indication of cecal intubation, whereas multiple still photographs can be more convincing. Photographing characteristic features of the GI tract can also render the photographs more conclusive. Rex (56) suggested that the cecal view is best taken when the valve lips are en face or a notch in the
intervention at a different site. Each endoscopy facility should, of the report may also facilitate unscheduled postprocedural medical stress and medication of the procedure have less influence. Finally, a reasons for this might be because the report clarifies the findings, intervention regarding the procedure findings, plans for treatment and follow-up, worriesome symptoms to watch for, and steps to be taken (3).

A study conducted by Spodick et al (6) has shown that providing an endoscopy report to patients after an endoscopic examination diminishes postprocedure anxiety, improves recollection of findings and recommendations, and increases the level of compliance to the given recommendations and follow-up plan (6). Thus, when a patient is discharged, an endoscopy report with the details of the procedure should be provided at that time. Furthermore, information should also be transmitted to any physician providing subsequent care. As recommended by the CAG, the endoscopy report should contain the following information: description of key findings, interventions, recommendations and follow-up plan (3). Measurement of quality indicators in clinical practice can identify areas for quality improvement and permit good monitoring of the clinical practice (4).

**Patient perception and timing of the report**

**Statement 9:** “All patients, on discharge, are given written information regarding the procedure findings, plans for treatment and follow-up, worriesome symptoms to watch for, and steps to be taken” (3).

**Impact of implementing education in reporting**

Lieberman et al (4) conducted a study to determine the quality of colonoscopy reporting in diverse practice settings because, as mentioned above, to improve colonoscopy quality, reports must include key quality indicators that can be monitored. As expected, there is significant variation in the quality of colonoscopy reports across diverse practices. Even with the use of a standardized computer-generated reports, many quality indicators were missing, which could harm quality improvements in the practice of colonoscopy. Hence, implementing education in reporting is crucial for continuous quality improvement in endoscopy. In a study performed in a Canadian institution, various deficits in reporting were outlined and recommendations to improve the appropriate use of an existing reporting system were made, leading to a documented postintervention quality improvement (41). Measurement of quality indicators in clinical practice can identify areas for quality improvement and permit good monitoring of the clinical practice (4).

**CONCLUSION**

Reduction in the variation of quality is of primary importance for endoscopy practice. High-quality endoscopy requires more than a good performance on the part of the endoscopist. From the initial patient interaction with a health care provider, the delivery of high-quality endoscopy services must be provided in an optimal and cost-effective manner following established quality assurance guidelines. Such an approach requires a well-established quality improvement framework, addressing all aspects of endoscopy, such as is outlined in the Global Rating Scale, the Canadian version of which will soon be published (C Dubé, personal communication). Continuous quality improvement should be the responsibility of every endoscopy facility to ensure improved patient care. Quality assurance for endoscopy requires that only competent physicians be given privileges to practice, and that continuous implementation of education in reporting and other areas of endoscopy is performed. Endoscopy units should regularly participate in a continuous process of quality improvement. In May 2010, the scarcity of high-grade evidence relevant to endoscopy service delivery was one of the CAG consensus’ outcomes. Following this conclusion, participants identified important quality indicators. There was a high level of agreement on key features that should be addressed to improve endoscopy quality among the large multidisciplinary group of health care professionals forming the consensus (3). The present article reviews current evidence and expert consensus on these quality indicators to be used in the process of improving quality in endoscopic reporting and patient care. Following the consensus among experts, the present study suggests a framework for a quality improvement structure in endoscopic units based on explicit recommendations that support systematic monitoring, assessment and modifications in endoscopy service delivery through optimal endoscopy reporting.

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