Colonoscopy quality across Europe: a report of the European Colonoscopy Quality Investigation (ECQI) Group

Authors
Cristiano Spada1,2, Anastasios Koulaouzidis1, Cesare Hassan4, Pedro Amaro5, Anurag Agrawal8, Lene Brink7, Wolfgang Fischbach8, Matthias Hüniger9, Rodrigo Jover10, Urpo Kinnunen11, Akiko Ono12, Árpád Patai13, Silvia Pecere14, Lucio Petruzzelli14, Jürgen F. Riemann15, Bharat Amlani16, Harry Staines17, Ann L. Stringer18, Ervin Toth19, Giulio Antonelli4,20,21, Lorenzo Fuccio22

Institutions
1 Digestive Endoscopy Unit and Gastroenterology, Fondazione Poliambulanza, Brescia, Italy
2 Digestive Endoscopy Unit, Università Cattolica del Sacro Cuore, Rome, Italy
3 Pomeranian Medical University in Szczecin-Department of Social Medicine and Public Health, Faculty of Health Sciences, Szczecin, Zachodniopomorskie, Poland
4 Digestive Endoscopy, Nuovo Regina Margherita Hospital, Rome, Italy
5 Gastroenterology Department, Centro Hospitalar e Universitário de Coimbra, Coimbra, Portugal
6 Doncaster Royal Infirmary, Doncaster, UK
7 Herlev and Gentofte Hospital, Copenhagen University, Gastro Unit, Division of Endoscopy, Herlev, Denmark
8 Gastroenterologie und Innere Medizin, Aschaffenburg, Germany
9 Private Practice for Internal Medicine, Würzburg, Germany
10 Hospital General Universitario de Alicante – Instituto de Investigación Sanitaria ISABIAL – Servicio de Medicina Digestiva, Alicante, Spain
11 Tampere University Hospital-Gastroenterology, Tampere, Finland
12 Hospital Clínico Universitario Virgen de la Arrixaca-Gastroenterology, El Palmar, Murcia, Spain
13 Markusovszky University Teaching Hospital-Gastroenterology, Szombathely, Hungary
14 Digestive Endoscopy Unit, Fondazione Policlinico Universitario A. Gemelli IRCCS, Rome, Italy
15 LebensBlicke Foundation, Ludwigshafen, Germany
16 Norgine Ltd-Medical Affairs, Harefield, UK
17 Sigma Statistical Services Ltd, Saint Andrews, UK
18 ECQI Secretariat, Buckinghamshire, UK
19 Skåne University Hospital, Lund University, Department of Gastroenterology, Malmö, Sweden
20 Department of Anatomical, Histological, Forensic Medicine and Orthopedics Sciences, “Sapienza” University of Rome, Rome, Italy
21 Gastroenterology and Digestive Endoscopy Unit, Ospedale dei Castelli Hospital, Ariccia, Rome, Italy
22 Gastroenterology Unit, Department of Medical and Surgical Sciences, S. Orsola-Malpighi Hospital, Bologna, Italy

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Georg Thieme Verlag KG, Rüdigerstraße 14, 70469 Stuttgart, Germany
Corresponding author
Cristiano Spada, Fondazione Poliambulanza – Digestive Endoscopy Unit and Gastroenterology, Via L. Bissolati 57, 25124 Brescia, Italy
Fax: +390303518221
cristianospada@gmail.com

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ABSTRACT

Background and study aims The European Colonoscopy Quality Investigation (ECQI) Group comprises expert colonoscopists and investigators with the aim of raising colonoscopy standards. We assessed the levels of monitoring and achievement of European Society of Gastrointestinal Endoscopy (ESGE) performance measures (PMs) across Europe using responses to the ECQI questionnaires.
**Methods** The questionnaire comprises three forms: institution and practitioner questionnaires are completed once; a procedure questionnaire is completed on multiple occasions for individual total colonoscopies. ESGE PMs were approximated as closely as possible from the data collected via the procedure questionnaire. Procedure data could provide rate of adequate bowel preparation, cecal intubation rate (CIR), withdrawal time, polyp detection rate (PDR), and tattooing resection sites.

**Results** We evaluated ECQI questionnaire data collected between June 2016 and April 2018, comprising 91 practitioner and 52 institution questionnaires. A total of 6445 completed procedure forms were received. Institution and practitioner responses indicate that routine recording of PMs is not widespread: adenoma detection rate (ADR) is routinely recorded in 29.5% of institutions and by 34% of practitioners; PDR by 42% and 47%, CIR by 62% and 64%, bowel preparation quality by 56% and 76%, respectively.

Procedure data showed a rate of adequate bowel preparation of 84.2%, CIR 73.4%, PDR 40.5%, mean withdrawal time 7.8 minutes and 12.2% of procedures with possible removal of a non-pedunculated lesion ≥ 20 mm reporting tattooing.

**Conclusions** Our findings clearly show areas in need of quality improvement and the importance of promoting quality monitoring throughout the colonoscopy procedure.

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**Introduction**

Colonoscopy has been shown to greatly reduce colorectal cancer (CRC) incidence and mortality as it allows for both identification of early neoplasia and removal of precancerous lesions [1,2]. While adenoma detection rate (ADR) is considered a primary quality indicator [3,4], it is dependent on other quality measures, such as cecal intubation rate (CIR), withdrawal time, and quality of bowel preparation [5].

There is considerable variability in the quality of colonoscopy [6] with a three- to six-fold variation in ADR among endoscopists [7,8]. Given the substantial impact of CRC on patients and healthcare systems [9,10], and that screening can be effective provided the services are of high quality [11], it is clearly important to ensure that colonoscopy is delivered to a high standard across the endoscopy community.

The European Society of Gastrointestinal Endoscopy (ESGE) has published both performance measures (PMs) for lower gastrointestinal endoscopy [12] and PMs for the endoscopy service as a whole [13], providing to all stakeholders (patients and their advocacy groups; service leaders; staff, including endoscopists; professional societies; payers and regulators) recommendations on the necessary parameters needed to meet the requirements of the ESGE quality improvement initiatives. These measures include those related to leadership, organisation, and service delivery, as well as those associated with the patient journey, and comprise recommendations for a minimum and target standard for endoscopy services to achieve. A crucial aspect of the guidelines is periodical monitoring of PMs both on individual practitioners, as well as at institutional service levels.

The European Colonoscopy Quality Investigation (ECQI) Group comprises expert colonoscopists and investigators with the aim of raising colonoscopy standards across Europe. ECQI does not wish to create any specific quality criteria, but rather document how the recent ESGE guidelines are implemented in daily practice and assess the quality of colonoscopy practice in Europe. We aimed to assess the levels of monitoring and achievement of ESGE PMs across Europe using responses to the ECQI questionnaires.

Methods

At the inaugural meeting of the ECQI Group in 2013, the Group chose to develop a clinical practice questionnaire to enable colonoscopists to evaluate current practice. The online questionnaire was based on the ESGE position statement on quality in screening colonoscopy published in 2012 [14]. An iterative process was used to hone the questionnaire ensuring that the time to complete the form was not too onerous. It was validated in November 2014 and May 2015 during two pilot phases, via a collaborative approach to ensure pertinent information was being recorded and data on 1861 patient procedures were collected [15,16]. The questionnaire comprises three forms: institution (18 questions) and practitioner questionnaires (12 questions) are each completed once, recording routine practice at respective levels; a procedure questionnaire (34 questions) is completed on multiple occasions for individual total colonoscopies (see Supplementary Material).

Participation was open to all Europe-based colonoscopists via web-based registration at the ECQI Group website. Awareness of the questionnaire came from abstracts, posters, presentations at national and international congresses and individual communications from ECQI Group members. Interested participants applied via the ECQI Group website or to the ECQI Group Secretariat. Following verification, log-in access to the web-based questionnaire site was provided by email.

Calculation of performance measures

ESGE PMs [12] were approximated as closely as possible from the data collected via the procedure questionnaire. We determined that our questionnaires could provide approximations for rate of adequate bowel preparation, CIR, withdrawal time, polyp detection rate (PDR), and tattooing resection sites (Table 1). For tattooing, we were unable to include polyps with suspicious macroscopic features regardless of size, as we were limited in the ability to determine the presence of suspicious macroscopic features due to questionnaire design, so this measure only includes procedures with a non-pedunculated polyp ≥ 20 mm. We were also unable to determine from ques-
tionnaire responses whether tattooing was performed on the resection site; we were able to determine whether it was performed during the same procedure as an endoscopic intervention that could have resulted in polyp removal. We also provide an indication of polyp removal rate for procedures with a polyp > 5 mm, as our questionnaire was unable to determine the polyp retrieval rate for histopathology examination.

A score of ≥6 on the Boston Bowel Preparation Scale (BBPS) was used to define adequate bowel preparation [17]. In procedures with data missing for one segment, if it could not be determined that the BBPS was definitely either ≥6 or <6, procedures were classified as missing, along with all other procedures with more than one segment missing data. To calculate CIR, only those procedures reporting the cecum as the intended endpoint were included, which excluded some procedures with terminal ileum/neo-terminal ileum as the intended endpoint, because given the questionnaire design, we could not determine whether the cecum was photo-documented in these procedures.

Diagnostic and screening procedures were determined using the reason that was provided for performing them. Our questionnaire had no method of collecting histopathological data so ADR could not be calculated. Polyp detection was regarded as positive if either a polyp or a polypectomy was reported. Age at the date of the procedure was derived assuming the date of birth was June 30 (to preserve anonymity, only the patient’s year of birth was recorded).

Calculation of mean withdrawal time was restricted to those procedures (screening or diagnostic) in which the cecum was the intended endpoint, the endpoint was reached, and no endoscopic intervention was reported. Procedures with a definite non-pedunculated lesion ≥20 mm were identified when in any segment reporting a polyp ≥20 mm, only non-pedunculated classifications were recorded for that segment. This may have excluded some non-pedunculated lesions ≥20 mm, as it excluded procedures in which both pedunculated and non-pedunculated lesions were reported in a segment with a polyp ≥20 mm. We only included procedures in which an endoscopic intervention that could have removed the polyp was reported, i.e. endoscopic mucosal resection, endoscopic submucosal dissection, polypectomy (complete or incomplete) or biopsy.

Results

We evaluated ECQI questionnaire data collected between June 2, 2016 and April 30, 2018, comprising 91 completed practitioner questionnaires and 52 completed institution questionnaires from 12 European countries. A total of 6445 completed procedure forms were received from 25 academic hospitals (2270/6445, 35.2%), 14 hospitals (1235/6445, 19.1%), eight private institutions (2657/6445, 41.2%), three group practices (160/6445, 2.5%), and one other (123/6445, 1.9%). Results are summarized in Table 2 and Table 3.

Pre-procedure

A reason for colonoscopy was provided for 6413 of 6445 procedures (99.5%). These were classified as: diagnostic (3182/6445, 49.3%), screening (1274/6445, 19.8%), follow-up (1837/6445; 28.5%), previous unsuccessful procedure (99/6445, 1.5%), and other (21/6445, 0.3%). Screening was classified as “due to familial risk” in 29.7% (378/1274) of procedures, “following a positive screening test” in 39.2% (499/1274) and “without a pre-screening test” in 30.9% (394/1274), and one other.

The collected responses showed that scale-based bowel cleansing quality was reported as “routinely recorded” by 56% of institutions and “routinely used” by 76% of practitioners. From the procedure data, 84.2% (5427/6445) of procedures reported an adequate bowel cleansing (data missing for 209 procedures, 3.2%).

Completeness of procedure

CIR was reported as “routinely recorded” by 64% of practitioners and by 62% of institutions. Procedure data showed that the cecum was the intended endpoint in 69.4% (4473/6445) of

<table>
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<th>Denominator</th>
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<td>Procedures that report reaching the cecum (documented in written form and by photo/video)</td>
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<td>Withdrawal time</td>
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<td>Procedures with an endoscopic intervention</td>
<td>All procedures with a polyp &gt; 5 mm reported</td>
</tr>
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</table>

BBPS, Boston Bowel Preparation Scale.

1 This is not an ESGE performance measure, however, within the restrictions of data provided, it provides an indication of whether polyp retrieval is attempted, although the rate of success cannot be determined.
procedures (ileum 28.1%, anastomosis 1.3%, data unavailable 1.2%). For those colonoscopies in which the cecum was the intended endpoint, 94.7% reported reaching the cecum but only 77.5% (3281/4234) of those stated endpoint photo-documentation.

Identification of pathology
ADR was reported as “routinely recorded” by only 34% of practitioners and in 29% of institutions. PDR was “routinely recorded” by 47% of practitioners and in 42% of institutions. Retraction time was “routinely recorded” by 60% of practitioners.

At least one polyp was detected in 40.5% (1363/3365) of qualifying procedures. Withdrawal time was assessed in the 1150 qualifying procedures providing data, the overall mean (± SD) withdrawal time was 7.8 ± 3.1 minutes, the median withdrawal time was 7 minutes.

Management of pathology
The proportion of practitioners reporting “routinely recording” polyp removal rate was 44% and the polyp retrieval rate was 37%. Routine use of a polyp classification scale was reported by 77% of practitioners and 54% routinely placed tattoos following polyp removal based on guidelines.

In the 1294 procedures where a polyp >5 mm was reported, 89.3% (1156/1294) reported an endoscopic intervention (Table 4). In procedures in which a non-pedunculated lesion ≥20 mm could be definitively identified and an endoscopic intervention to remove the polyp reported, 12.2% (17/139) reported tattooing.

Complications, patient experience and post-procedure
Patient satisfaction was recorded in 25% of institutions, during-procedure complications were reported to be “routinely recorded” in 83%, but post-procedure complications by only 56%. Quality guidelines were reported to be “routinely followed” in 69% of institutions.
Discussion

In this study, we sought to evaluate the adoption of colonoscopy PMs across Europe. In 2017, the ESGE published PMs for lower gastrointestinal endoscopy [12], recommending that endoscopy services across Europe should adopt a list of key and minor PMs for objective assessment and evaluation in daily practice at both center and individual endoscopist level. Several key performance indicators have been established for adoption to achieve consistently high-quality endoscopic practice. We analysed a sample of procedures conducted across Europe, between June 2016 and April 2018, spanning a period before and after publication of the PMs, to evaluate the baseline achievement of standards, as defined by the ESGE. The analysis was performed at institution, practitioner and procedure levels. A set of variables listed in the ESGE lower gastrointestinal endoscopy PM document was considered. Interestingly, although some of the PMs seem to be relatively commonly assessed, documentation of other relevant PMs is far from routine.

Scale-based bowel cleansing quality was reported as routinely recorded in only 56% of institutions, and an adequate level was achieved in 84.2% of procedures, slightly below the ≥90% minimum standard as recommended by ESGE [12]. The quality of bowel preparation is crucial for the overall efficacy of colonoscopy, with a suboptimal ADR and CIR related to an inadequate cleansing level and a higher risk of interval cancer. [18, 19] In addition, a suboptimal cleansing level results in further costs and organizational issues since colonoscopy needs to be rescheduled or patients may be referred for alternative tests [20, 21].

CIR was reported as routinely recorded in only 62% of institutions. In addition, using the ESGE definition of CIR, which requires both written and photo-documentation, only 73.4% of procedures met requirements, which is short of the ≥90% minimum standard. However, when considering just written documentation, 94.7% reported reaching the cecum, almost reaching the ≥95% target standard.

Overall, PDR can be considered as a surrogate for ADR and is easier to monitor, because it is automatically collated by endoscopists and institutions while generating procedure reports and/or billing codes, making it more practical to measure than ADR, even if less robust. Although our data show that PDR is more commonly recorded, it seems even this parameter falls short of the recommended standard, with only a minority of practitioners (47%) and institutions (42%) routinely recording the quality measure. When looking at the procedure forms, in terms of PDR, at least one polyp was detected in 40.5% of qualifying procedures, being borderline with the ≥40% ESGE minimum standard of screening and diagnostic colonoscopies performed in those aged 50 years or older. Initiatives such as education, adequate training, creating awareness, feedback, and colonoscopy quality benchmarking have been shown to contribute to improvement in these parameters [22–25].

Retraction time is recorded by only 60% of practitioners. Procedure data indicate the mean withdrawal time was 7.8 ± 3.1 minutes, which reached the minimum standard (i.e. mean 6 minutes) defined by Kaminski et al [12]. Although we did not directly measure ADR, our data show that it was routinely recorded by only 34% of practitioners and in 29% of institutions. When considering the role of ADR as a universal key quality indicator, this is quite disappointing and might be one of the limiting factors for ADR underperformers.

Measurement of complication rate only partially entered routine practice: the collected responses showed that although during-procedure complications are usually recorded by the majority of institutions (83%), almost one of two institutions (56%) do not record post-procedure complications. This is comparable to the results by Adler et al [26]. This substantial under-recording probably reflects difficulty in monitoring patients after the procedure and the lack of availability of methods that allow the identification of a late complication.

Patient feedback, to enhance patient experience and colonoscopy quality, is important, and should be routinely monitored with adequate feedback mechanisms in place [12, 13]. However, only a minority of institutions (25%) record patient satisfaction and this merits further evaluation. Such underperformance in terms of recording of patient experience could be related to cultural issues (at least in some countries) as well as to logistic limitations related to the collection of patient feedback.

An important strength of this study is its size, both in terms of the number of colonoscopies analyzed and the Europe-wide coverage of the survey. Many colonoscopy quality studies are either single-center or restricted to a small number of endoscopists. However, we accept that both the present study and the questionnaires have some limitations. The current findings are not based on consecutive reporting and a selection bias for those procedures recorded cannot be ruled out. Nevertheless, it reflects real-world data and can provide an efficient method to monitor colonoscopy quality measures both at an institutional and endoscopist level, with the aim to support initiatives and improve clinical practice standards. It identifies the quality measures that are adhered to, and how effective they can be in driving standards. Another important limitation of the present study is the self-selection of endoscopists across Europe for participation in the survey rather than random selection. It is debatable whether this self-selection bias might have selected a subgroup of endoscopists not representative of the general endoscopist population, leading to better results than in the general population of endoscopists. The same limitations, however, apply to all measures of voluntary quality control.

It is noteworthy that the publication of the ESGE PMs occurred after this version of the questionnaires was compiled; therefore, there are some areas in which the ECQI measures do not exactly match those specified by the ESGE [12].

In general, looking at the picture coming from the present study, we should admit that quality measures for colonoscopy are far from being routinely recorded in clinical practice. Performance measurement is the first step in a process aimed at improving quality in colonoscopy. Further steps include the identification of underperformers, of the barriers that need to be addressed, and subsequent reevaluation after corrective interventions have taken place. Measurement of performance parameters is the prerequisite without which concrete im-
Improvement actions cannot be developed. Initiatives such as education, creating awareness, and training should be implemented to contribute to the overall improvement of colonoscopy. The final goal should be to improve quality, reducing gaps between clinical practice and evidence.

Conclusions
In conclusion, our findings clearly show areas in need of quality improvement and the importance of promoting quality monitoring throughout the colonoscopy procedure. They also underscore the necessity of regularly recording individual quality parameters to measure daily performance against well-established recommendations and evaluate their wider dissemination and adoption.

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Competing interests
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