Adherence to guidelines: A national audit of the management of acute upper gastrointestinal bleeding. The REASON registry.

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OBJECTIVES: To assess process of care in nonvariceal upper gastrointestinal bleeding (NVUGIB) using a national cohort, and to identify predictors of adherence to ‘best practice’ standards.

METHODS: Consecutive charts of patients hospitalized for acute upper gastrointestinal bleeding across 21 Canadian hospitals were reviewed. Data regarding initial presentation, endoscopic management and outcomes were collected. Results were compared with ‘best practice’ using established guidelines on NVUGIB. Adherence was quantified and independent predictors were evaluated using multivariable analysis.

RESULTS: Over all, 2020 patients (89.4% NVUGIB, variceal in 10.6%) were included (mean ± SD age 66.3±16.4 years; 38.4% female). Endoscopy was performed in 1612 patients: 1533 with NVUGIB had endoscopic lesions (63.1% ulcers; high-risk stigmata in 47.8%). Early endoscopy was performed in 65.6% and an assistant was present in 83.5%. Only 64.5% of patients with high-risk stigmata received endoscopic hemostasis; 9.8% of patients exhibiting low-risk stigmata also did. Intravenous proton pump inhibitor was administered after endoscopic hemostasis in 95.7%. Rebleeding and mortality rates were 10.5% and 9.4%, respectively. Multivariable analysis revealed that low American Society of Anesthesiologists score patients had fewer assistants present during endoscopy (OR 0.63 [95% CI 0.48 to 0.83]), a hemoglobin level <70 g/L predicted inappropriate high-dose intravenous proton pump inhibitor use in patients with low-risk stigmata, and endoscopies performed during regular hours were associated with longer delays from presentation (OR 0.33 [95% CI 0.24 to 0.47]).

CONCLUSION: There was variability between the process of care and ‘best practice’ in NVUGIB. Certain patient and situational characteristics may influence guideline adherence. Dissemination initiatives must identify and focus on such considerations to improve quality of care.

Key Words: Gastrointestinal hemorrhage; Guideline adherence; Peptic ulcer hemorrhage; Quality of health care

Upper gastrointestinal bleeding (UGIB) is associated with significant morbidity and mortality. The incidence of acute nonvariceal UGIB (NVUGIB) ranges from 50 to 150 cases per 100,000 adults per year (1), while mortality varies from 2.5% to 10% (1,2). A large bleeding registry from 1999 to 2002 (3) revealed significant practice variations in the management of NVUGIB across Canadian hospitals (4). Several consensus recommendations have since been developed, setting well-defined standards (5-7). However, wide practice variations among physicians continue to be reported (8). Such variability calls to assess the quality of care delivered in NVUGIB. The latter extends beyond the measurement of outcomes such as rebleeding and mortality, and includes the evaluation of the structure and delivery of care through process-based measures (9,10). Optimally, the process of care should be consistent with evidence-based ‘best practice’ (8).

The Registry of patients undergoing Endoscopic and/or Acid Suppression therapy and an Outcome analysis for upper gastrointestinal bleeding (REASON) recorded the ‘real-life’ hospital and physician practice managing acute UGIB across Canadian hospitals. The process of care was compared with ‘best practice’ recommendations available at the time, namely international NVUGIB consensus guidelines (5) published one year before the national cohort study. We present Canadian nationwide data regarding the structure, process and results.

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outcomes of care as measures of quality of care (10) in light of ‘best practice’ guidelines and assess the impact of the latter.

METHODS

REASON initiative and data collection

A retrospective chart review of unselected hospitalized patients with a diagnosis of UGIB was performed in 21 Canadian hospitals. One hundred charts per institution were reviewed, starting from January 2004 until a total of 2000 cases was reached. Data collection was performed using an electronic system. Data were audited for the first 10 patients at each site, and randomly thereafter in 10% to ensure standardization.

Patient population

All hospitalized patients at least 18 years of age with a primary or secondary discharge diagnosis of NVUGIB and variceal UGIB were identified using International Classification of Diseases, Ninth and 10th Revision (ICD-9 and ICD-10) codes via each hospital’s electronic record database. Only hospitalized patients were included. Patients discharged from the emergency room or those transferred from another institution were excluded. Individuals already hospitalized for another reason, but who developed UGIB during their stay, were included. The performance of endoscopy was not an inclusion criterion.

Study variables

Structure and process of care: initial presentation and endoscopic management: Baseline data with patient demographics (age, sex, relevant medical history and medications) and institutional information (staffing and facility) were obtained. A description of the initial bleeding event was collected for all patients. The performance of endoscopy (if any), in addition to timing, endoscopic findings and hemostatic event was collected for all patients. The performance of endoscopy (if any), in addition to timing, endoscopic findings and hemostatic event was collected for all patients. The performance of endoscopy was not an inclusion criterion.

Outcomes: Rebleeding, recourse to angiography and surgery, length of stay and mortality were reported. Rebleeding was defined as overt hematemesis; passage of fresh blood per rectum; fall in hemoglobin concentration >20 g/L within any 24 h period after the first 24 h; shock (systolic blood pressure of ≤90 mmHg or a heart rate of ≥110 beats/min) in the presence of continuing melaena; or fresh blood in the stomach and/or duodenum at repeat endoscopy when further bleeding was suspected (3,11,12). All outcome data were collected for a 30-day period following the initial bleeding event.

Best practice: adherence to guidelines and its possible predictors: Adherence to guidelines was determined using the most recent guidelines available at the time of data collection (5), which featured 20 recommendations pertaining to the management of NVUGIB. Each recommendation was reviewed in light of available data from REASON, and adherence rates were calculated using ratios and percentages, except when data were insufficient or missing from the registry. Data were compared with a similar cross-sectional study preceding the publication of the guidelines (4).

For selected recommendations, possible independent predictors of guideline adherence were assessed, including the patient’s age (>60, 60 to 80 or >80 years of age), sex, pre-endoscopic Rockall score (0 to 2, or >2) and Blatchford scores (0 to 2, 3 to 11, or >11), American Society of Anesthesiologists (ASA) classification (1 to 2, 3, or 4 to 5), presenting hemoglobin level (<70 g/L) and time of endoscopy (regular hours defined as Monday to Friday from 08:00 to 17:00 versus after hours).

Data analysis

Categorical data were expressed as proportions and 95% CIs, and continuous data as means ± SD. Interquartile range (IQR) and median were also used when appropriate. Univariable analysis was performed using χ² or Wilcoxon-ranking test. Independent predictors of guideline adherence were identified using the best fitting multivariable logistic regression models that were clinically relevant; P<0.05 was considered to be statistically significant. All statistical analyses were performed using SAS version 9.2 (SAS Institute Inc, USA).

Ethics

Informed consent from individual patients was not deemed necessary because the study was considered to be an audit at all sites. The institutional review boards from all participating centres gave approval for the REASON data collection and analysis.

RESULTS

Structure of care: institutional and endoscopic facility data

Across the 21 hospitals, the mean number of endoscopists on staff was 11 per hospital (median nine; range three to 26). After-hours trained endoscopic support staff were available in 17 of 21 (81%) hospitals, of which 13 of 17 (76%) were available 24 h/day. Assistants were present to assist endoscopy in 83.5% of cases. Fewer than one-half (10 of 21 [48%]) of the hospitals had explicit written guidelines on intravenous proton pump inhibitor (IV PPI) use; a standardized critical care path for the management of patients with suspected UGIB existed in a similar proportion of institutions (11 of 21 [52%]).

Clips were accessible in all hospitals, followed by injection (20 of 21 [95%]); other devices included gold probe (17 of 21 [81%]), argon plasma coagulation (15 of 21 [71%]), heater probe (15 of 21 [71%]), and bipolar or multipo lar electrocoagulation (13 of 21 [62%]).

Study population

Data from 2020 patients with UGIB were collected between January 2004 and May 2005. A total of 1805 (89.4%) patients presented with NVUGIB and 215 (10.6%) with variceal bleeding. The mean (± SD) age was 66.3±16.4 years, with a male predominance (1245 of 2020 [61.6%]). Nearly one-fifth (356 of 2020 [17.6%]) of all patients experienced the bleeding episode while already hospitalized for another reason. The mean Blatchford score was 9.1±4.1 (minimum = 0; maximum = 23), while the pre-endoscopic mean Rockall scores were 2.0±1.5 (minimum = 0; maximum = 7). The overall ASA score distribution was 80.2% ASA score 1 to 3, 19.8% ASA score 4 and 5.

Process of care

Initial management: Twenty-two percent (448 of 2020) of patients underwent nasogastric tube lavage; 38.4% (172 of 448) revealed coffee-ground material, 27.5% (123 of 448) bright red blood and 4.0% (18 of 448) bile. Only five of the 21 (24%) hospitals routinely obtained nasogastric tube aspirates for the majority of their patients. The mean presenting hemoglobin level was 98±27.8 g/L. Sixty-nine percent of patients received a blood transfusion, at a median time of 8.7 h (IQR 3.9 h to 24.0 h) from presentation, with the vast majority (74.9%) occurring in the first 24 h. Among those transfused, 16.8% had an initial hemoglobin level ≤70 g/L, 39.2% between 71 g/L and 100 g/L and 44.0% a level >100 g/L.

Overall, 68.2% (1113 of 1632) of patients presenting with acute NVUGIB received pre-endoscopic IV PPIs. These were administered at high dose with an 80 mg bolus, followed by 8 mg/h infusion in 54.5% (698 of 1281). Prokinetic use was not recorded.

Time to endoscopy: The median time to endoscopy was 17.7 h (IQR 6.1 h to 29.4 h). Patients with suspected variceal bleeding underwent endoscopy significantly earlier than individuals with suspected non-variceal bleeding, with a median time to endoscopy of 12.4 h (IQR 4.5 h to 25.0 h) compared with 18.3 h (IQR 6.3 h to 30.0 h)
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for nonvariceal bleeding (P=0.0038). Overall, 65.6% of patients underwent endoscopy within 24 h. The distributions of time to endoscopy are shown in Figures 1A and 1B.

**Endoscopic findings:** Findings at endoscopy were recorded for both variceal and nonvariceal bleeding, although details pertaining to variceal bleeding will not be discussed further. A total of 89.4% (1805 of 2020) of patients were diagnosed with NVUGIB (Figure 2).

The etiologies of NVUGIB were distributed as follows: ulcers (967 of 1533 [63.1%]), erosions (341 of 1533 [22.2%]), esophagitis (313 of 1533 [20.4%]), Mallory-Weiss tear (113 of 1533 [7.4%]), malignancy (40 of 1533 [2.6%]), Dieulafoy lesion (35 of 1533 [2.3%]), gastric antral vascular ectasia syndrome (35 of 1533 [2.3%]) and esophageal strictures (34 of 1533 [2.2%]). More than one endoscopic diagnosis was possible for the same patient. Figure 3 illustrates the distribution of stigmata of recent hemorrhage in patients with nonvariceal bleeding, including ulcers and nonulcer lesions. While several diagnoses could be identified for the same patient, only the stigma from the lesion most likely to be the source of bleeding was recorded. Detailed documentation of endoscopic findings, more specifically, bleeding stigmata and description were unavailable in 30.1% (135 of 448) of cases. Excluding patients with no documentation of stigmata of recent bleeding, 71% had HRS.

**Endoscopic therapy:** Nearly two-thirds (64.5%) of patients with HRS from both ulcer and nonulcer bleeding received endoscopic hemostasis. Moreover, 9.8% of patients LRS were also treated with endoscopy (Table 1). Epinephrine injection was used as monotherapy in 23.8% of patients with high-risk ulcers and 20.7% of all high-risk lesions. The most common endoscopic modalities used for the management of HRS lesions were injection monotherapy (32.8%), combination therapy of injection and thermal (32.3%), followed by thermal monotherapy (14.5%) and clips (alone or in combination) (14.3%). A routine (preplanned) second-look endoscopy was performed in 9.0% (145 of 1612) of patients, while repeat endoscopy due to suspected rebleeding was performed in 56.7% (97 of 171).

**Pharmacotherapy:** High-dose IV PPI (80 mg bolus followed by continuous infusion) was administered to 95.7% of patients with HRS after successful endoscopic hemostasis. Overall, 91.5% of patients with LRS received high-dose IV PPI after endoscopy, irrespective of endoscopic therapy.

**H pylori testing and treatment:** H pylori status was available on presentation for 10.8% of patients (7.5% positive, 3.3% negative). Thirty-six percent of patients with NVUGIB were further tested for H pylori during the course of their hospitalization, with eradication therapy initiated in 9.2% who tested positive. No data regarding the confirmation of eradication were available.

**Outcomes:** Rebleeding occurred in 10.5% (190 of 1805) of patients, with an overall mortality rate of 9.4% (169 of 1805). A small proportion...
of patients (47 of 1805 [2.5%]) required surgery, while 1.2% (22 of 1805) required interventional radiology to control their bleeding. Of the patients who developed UGIB while hospitalized for another reason (n=356), rates of rebleeding, need for surgery and mortality were 14.4%, 1.4% and 21.9%, respectively. Length of hospital stay was, on average, six days. Patients with NVUGIB who were admitted to the intensive care unit were discharged to the floor after a median of 5.8±10.1 days.

Adherence to guidelines: The data from REASON were applicable to 13 of the 23 consensus recommendations (5) (Table 2). Comparing current adherence rates with data with a quantitative analysis of appropriateness in Canadian practice patterns performed before guidelines publication (4), higher adherence was noted in 67% (eight of 12) of the recommendations (Figure 4). Two recommendations showed arithmeticly decreased adherence rates (two of 12 [17%]), namely the performance of early endoscopy within 24 h and endoscopic hemostatic therapy in HRS. Comparison was not performed for one recommendation due to the lack of presguideline values.

Predictors of adherence: Predictors of recommended use of endoscopic resources: Multivariable analysis revealed that the presence of qualified assistants during endoscopy occurred more frequently when endoscopy was performed during regular hours (OR 2.9 [95% CI 2.21 to 3.81]). In contrast, patients with low ASA classification (1 to 2) more often underwent endoscopy without assistance from support staff (OR 0.63 [95% CI 0.48 to 0.83]) (Table 3). Furthermore, the targeted delay of <24 h from presentation for early endoscopy was less likely to be met when an endoscopy was performed during regular hours (OR 0.33 [95% CI 0.23 to 0.47]). High Blatchford scores of 3 to 11 predicted the performance of endoscopic therapy in patients with HRS (OR 1.67 [95% CI 1.03 to 2.70]). No such predictors were identified to influence rates of endoscopic hemostasis in patients with LRS.

Predictors of recommended use of pharmacotherapy: The administration of high-dose IV PPI, which was not otherwise indicated in patients with LRS, was noted more frequently in patients with a presenting hemoglobin level <70 g/L (OR 6.84 [95% CI 1.97 to 23.76]). This practice was less often encountered in patients who were on antithrombotic therapy (OR 0.39 [95% CI 0.20 to 0.79]). No significant factors impacting on the use of high-dose acid suppressant were identified among patients with HRS. Significant predictors of adherence to the different recommendations and their respective ORs and 95% CIs are listed in Table 3.

DISCUSSION

The assessment of quality of care in the management of patients with NVUGIB extends beyond the sole measurement of clinical outcomes, especially in view of high-level evidence now available to guide best practice. Process-based measures (9) were applied to REASON, a retrospective multicentre national registry of 2020 patients, to evaluate the quality of care in NVUGIB in Canada. Compared with data collected before the publication of consensus guidelines (4,5), REASON demonstrates an improved adherence across one-half of the recommendations, the median adherence remains low (65.6%). While the process of care is a dynamic exchange of events (8) and may preclude rigid application of guideline recommendations, adherence to guidelines remains an important surrogate of quality measurement. Such data for NVUGIB are scarce, with some reports suggesting marked differences in practice across countries (13) and suboptimal uptake of national guidelines (14). Predictors of adherence are poorly defined in the literature, even though some have been suggested as a result of a multifaceted randomized educational intervention for guideline dissemination (15). Inferential testing performed as part of the REASON data analysis shed some light on possible discrepancies between observed practice and existing guidelines. These are discussed below.

Areas where practice can be further improved include the implementation of a protocol to manage UGIB and access to adequate endoscopy assistance, even out-of-hours (recommendations 1 and 2). Interestingly, a low ASA class was associated with the absence of an assistant during endoscopy, independent of the time to endoscopy, suggesting a relaxing of standards for cases perceived to be less urgent or more stable.

Attention should be drawn to the use of appropriate resuscitation (recommendation 3) and transfusion thresholds because they impact...
### TABLE 2
**Recommendation statements with associated adherence rates after guideline publication**

<table>
<thead>
<tr>
<th>Recommendation statement (5)</th>
<th>Adherence, % (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Hospitals should develop institution-specific protocols for multidisciplinary management, which should include access to an endoscopist with training in endoscopic hemostasis</td>
<td>52 (32.4–71.7)</td>
</tr>
<tr>
<td>2. Support staff trained to assist in endoscopy should be available for urgent endoscopy</td>
<td>83.5 (59.4–92.9)</td>
</tr>
<tr>
<td>3. Immediate evaluation and appropriate resuscitation are critical to proper management</td>
<td>N/A (–)</td>
</tr>
<tr>
<td>4. In selected patients, the placement of a nasogastric tube can be considered because the findings may have prognostic value</td>
<td>22 (20.4–24.0)</td>
</tr>
<tr>
<td>5A* Clinical (nonendoscopic) stratification of patients into low- and high-risk categories for rebleeding and mortality is important for proper management. Available prognostic scales may be used to assist in decision making</td>
<td>N/A (–)</td>
</tr>
<tr>
<td>5B* Early stratification of patients into low- and high-risk categories for rebleeding and mortality, based on clinical and endoscopic criteria, is important for proper management. Available prognostic scales may be used to assist in decision making</td>
<td>N/A (–)</td>
</tr>
<tr>
<td>6* Early endoscopy (within the first 24 h) with risk classification by clinical and endoscopic criteria allows for safe and prompt discharge of patients classified as low risk; improves patient outcomes for patients classified as high risk and reduces resource use for patients classified as either low or high risk</td>
<td>65.6 (62.7–68.4)</td>
</tr>
<tr>
<td>7A A finding of low-risk endoscopic stigmata (a clean-based ulcer or nonprominent pigmented dot in ulcer bed) is not an indication for endoscopic hemostatic therapy</td>
<td>90.2 (85.3–93.0)</td>
</tr>
<tr>
<td>7B* Endoscopic hemostatic therapy is indicated for a patient with a clot in an ulcer bed, including targeted irrigation in an attempt at dislodgment, with appropriate treatment of the underlying lesion</td>
<td>N/A (–)</td>
</tr>
<tr>
<td>7C A finding of high-risk endoscopic stigmata (active bleeding, nonbleeding visible vessel and/or adherent clot) in an indication for immediate endoscopic hemostatic therapy</td>
<td>64.5 (60.6–68.1)</td>
</tr>
<tr>
<td>8* No single solution for endoscopic injection therapy is superior to another for hemostasis</td>
<td>N/A (–)</td>
</tr>
<tr>
<td>9 No single method of endoscopic thermal coaptive therapy is superior to another</td>
<td>N/A (–)</td>
</tr>
<tr>
<td>10* Monotherapy, with injection or thermal coagulation, is an effective endoscopic hemostatic technique for high-risk stigmata; however, the combination is superior to either treatment alone</td>
<td>N/A (–)</td>
</tr>
<tr>
<td>11* The placement of clips is a promising endoscopic hemostatic therapy for high-risk stigmata</td>
<td>14.3 (11.2–18.1)</td>
</tr>
<tr>
<td>12† Routine second-look endoscopy is not recommended</td>
<td>91.0 (89.5–92.3)</td>
</tr>
<tr>
<td>13 In case of rebleeding, a second attempt at endoscopic therapy is generally recommended</td>
<td>56.7 (49.2–63.9)</td>
</tr>
<tr>
<td>14 Surgical consultation should be sought for patients who have failed endoscopic therapy</td>
<td>N/A (–)</td>
</tr>
<tr>
<td>15 H2-receptor antagonists are not recommended in the management of patients with acute upper gastrointestinal bleeding</td>
<td>95.1 (94.1–96.0)</td>
</tr>
<tr>
<td>16 Somatostatin and octreotide are not recommended in the routine management of patients with acute nonvariceal upper gastrointestinal bleeding</td>
<td>100 (100)</td>
</tr>
<tr>
<td>17† An intravenous bolus followed by continuous proton pump inhibitor infusion is effective in decreasing rebleeding in patients who have undergone successful endoscopic therapy</td>
<td>95.7 (93.2–97.4)</td>
</tr>
<tr>
<td>18* In patients awaiting endoscopy, empirical therapy with a high-dose proton pump inhibitor should be considered</td>
<td>54.4 (51.8–57.2)</td>
</tr>
<tr>
<td>19 Patients considered at low risk for rebleeding after endoscopy can be fed within 24 h</td>
<td>N/A (–)</td>
</tr>
<tr>
<td>20* Patients with upper gastrointestinal bleeding should be tested for Helicobacter pylori and receive eradication therapy if infection is present</td>
<td>N/A (–)</td>
</tr>
</tbody>
</table>

*Recommendations that have been revised in the 2010 guidelines (7); †Statement updated from 2003 guidelines, but no change in the actual recommendation in the 2010 guidelines (7). N/A Not applicable

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### Figure 4
**Adherence to recommendations comparing practice before and after guidelines publication.** *Guidelines in which collected data were insufficient to assess adherence are not included; **Data before guidelines publication were not available for recommendation 16. REASON Registry of patients undergoing Endoscopic and/or Acid Suppression therapy and an Outcome analysis for upper gastrointestinal bleeding*

### TABLE 3
**Selected predictors of guideline adherence showing statistical significance**

<table>
<thead>
<tr>
<th>Recommendation with associated predictors of adherence</th>
<th>OR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Presence of assistants during endoscopy</td>
<td>0.63 (0.48–0.83)</td>
</tr>
<tr>
<td>Blatchford score 3 to 11</td>
<td>1.67 (1.03–2.70)</td>
</tr>
<tr>
<td>Early endoscopy &lt;24 h</td>
<td>0.33 (0.24–0.47)</td>
</tr>
<tr>
<td>Endoscopic therapy in patients with high-risk stigmata</td>
<td>6.84 (1.97–23.76)</td>
</tr>
<tr>
<td>High-dose IV PPI administration in patients with low-risk stigmata</td>
<td>0.39 (0.20–0.79)</td>
</tr>
</tbody>
</table>

*Regular hours endoscopy defined as Monday to Friday, 08:00 to 17:00. IV PPI Intravenous proton pump inhibitor*
rebleeding and survival rates in UGIB (16). The mean initial hemoglobin value in REASON was 98.0±27.8 g/L, comparable with RUGBE (3) and the recent randomized control trial by Villanueva et al (16). Over the first 24 h, only 16.8% of those who received a transfusion had a presenting hemoglobin level ≤70 g/L. Of note, this proportion includes all patients, regardless of associated cardiovascular morbidity and hemodynamic status, a group excluded, in large part, by inclusion criteria in the large multisite Spanish trial (16).

The administration of IV PPIs before endoscopy is common (54.4%) even though their use is considered to be optimal (recommendation 18) and no longer supported in recent United Kingdom guidelines (17) despite some clinical settings identified as possibly cost effective (18). Overall, 65.6% of patients underwent early endoscopy within the recommended 24 h (recommendation 6). The point estimate is inferior to the 76% reported in RUGBE and European values ranging from 70% to 93% (13), while remaining superior to the 55% value reported in a true United Kingdom national bleeding audit (19). Additionally, longer delays (>24 h) to endoscopy were associated with the endoscopies being performed during regular hours. This suggests that some patients have their endoscopy postponed until regular working hours, even if this results in delays exceeding 24 h, perhaps because of limited out-of-hours support. Consistent with guidelines (20), patients with suspected variceal bleeding nevertheless underwent endoscopy significantly earlier, with a median of 12.4 h compared with 18.3 h for patients with NVUGIB (P=0.0038).

At endoscopy, even though lesions were identified in the great majority of cases after one or multiple endoscopies, a substantial proportion of endoscopy reports (26.9% for ulcers) failed to document the severity of the stigmata of recent bleeding. Such a high value is unsettling because this information is needed for adequate risk stratification into high- and low-risk categories, which in turn is crucial for appropriate endoscopic and pharmacological management (recommendation 5). Failure to record endoscopic stigmata was even more frequent (70.5%) among patients with nonvariceal, nonulcer bleeding, although both groups carry similar risk for unfavourable outcomes (21).

Patients exhibiting LRS do not require endoscopic therapy (recommendation 7A). Nonetheless, 9.8% of patients with LRS received endoscopic hemostasis in REASON, exceeding the 2.6% noted in an Italian prospective survey (22), although significantly inferior to a 24% endoscopic treatment rate for pigmented spots reported in the Clinical Outcomes Research Initiative database (23).

In contradistinction, although endoscopic hemostasis of lesions with HRS has clearly been shown to be associated with significant decreases in rebleeding, need for surgery and mortality (24) (recommendation 7C), adherence to such practice is suboptimal, with rates of 64.5% across all patients with NVUGIB and 69.9% among patients with ulcers. Reported rates of endoscopic treatment in high-risk patients in the literature range from 74% (3) to 85% (22,23). A French multicentre cohort proposes the lack of trained endoscopic assistance and out-of-hours endoscopy as possible factors for such practice (14).

The rate of injection monotherapy in REASON was 20.7%. This practice is not consistent with guidelines that rather suggest the added value of combination therapy (recommendation 10); it is, nonetheless, improved compared with the preguidelines value of 39% (3). Furthermore, IV PPI administration following successful endoscopy reached 95.7%, also superior to the previously reported 70.5% (recommendation 17). The only significant predictor of poor adherence in this regard was a low presenting hemoglobin level, which was associated with the use of high-dose IV PPI even in patients exhibiting LRS.

REASON did not record eradication rates, but reveals that *H pylori* testing (recommendation 20) was performed in only one-third of patients.

The REASON registry informs ‘real life’ practice in the management of UGIB, although it is limited by its retrospective nature. Our sampling excluded patients discharged from the emergency room, although it includes patients already hospitalized who develop a secondary diagnosis of UGIB, a subset at higher risk for complications (25). Missing data inherent to retrospective chart analyses somewhat undermine the accuracy of the calculated guideline adherence rates, although our values remain within ranges reported in the literature. Furthermore, the selected predictors of guideline adherence evaluated in our multivariate analysis may not be exhaustive and should be interpreted as exploratory because objective data regarding such predictors are scarce.

The above observations identify both areas of discrepant clinical care and some possible reasons for poor adherence that may guide future guideline dissemination initiatives. However, the impact of such educational efforts is often only modest. In fact, recent national data obtained from a 48-site cluster randomized clinical trial show that a multifaceted educational intervention over a 12-month period to disseminate UGIB guidelines did not improve adherence (15). The issue of low guideline adherence is longstanding and complex involving attitude, knowledge, contextual and behavioural barriers (26,27). Alternatively, the development of quality indicators using a list of more context-specific detailed clinical scenarios (28) may further help clinicians to reconcile ‘best practice’ in ‘real-life’ settings.

**CONCLUSION**

The process-based measures from the REASON registry reveal variable adherence to ‘best practice’ in NVUGIB in Canada. Proper endoscopic documentation, and adequate use of appropriate pharmacological and endoscopic therapy are key elements that clinicians should target to improve quality of care in NVUGIB. Both patient- and process-related factors may predict guideline adherence, or its lack. More precise characterization of such variables can help identify both clinician and patient subgroups for interventions, enhancing clinical management.

**REFERENCES**


