Unsedated transnasal endoscopy: A Canadian experience in daily practice

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BACKGROUND: Endoscopie oesophagastroduodénoscopique (EGD) est le plus souvent pratiqué examen diagnostique. Elle est couramment exécuté sous sédation consciente en Amérique du Nord. Une forte proportion de la morbidité et de la mortalité associées à l'EGD est liée à l'hypoxie attributable à la sédation consciente. La sédation est également liée à une augmentation des coûts, à l'absence du travail et à la nécessité que le patient se fasse accompagner à la maison après l'intervention. L'endoscopie transnasale comporte des avantages, tels que l'absence de sédation et une diminution de la surveillance du patient, du temps des infirmières et des dépenses par rapport à l'EGD par voie orale classique.

OBJECTIVES: Évaluer la faisabilité et l'acceptabilité de l'EGD transnasale sans sédation dans la pratique quotidienne.

METHODS: Les patients devant subir une EGD ont eu le choix entre une EGD transnasale sans sédation et une EGD par voie orale sous sédation. Les patients qui choisissaient l'EGD transnasale sans sédation subissaient l'intervention en cabinet par un gastroentérologue principal expérimenté dans les EGD transnasales. Toutes les interventions étaient effectuées à l'aide d'un esophagastroduodénoscope de petit calibre. Tous les patients ont été interrogés à l'aide d'un questionnaire sur la satisfaction des patients et ont été invités à attribuer des scores précis sur la sensation d'étouffement, le mal de gorge, le malaise nasal et le malaise abdominal. Toutes les variables ont été évaluées au moyen d'indices de 0 à 10 correspondant au degré le plus grave de chaque variable. Les complications ont également été enregistrées.

RESULTS: Entre mars 2002 et août 2003, 231 patients ont subi une EGD transnasale. Les patients avaient un âge médian de 57 ans (15 à 87 ans). Il a été possible de procéder à un examen complet chez 98% des patients. Les patients ont déclaré un fort degré d'acceptabilité (indice moyen de 6.6, fourchette de 1 à 10) et un faible degré de sensation d'étouffement (indice moyen de 1.8, fourchette de 0 à 10), de malaise nasal (indice moyen de 1.7, fourchette de 0 à 10), de malaise abdominal (indice moyen de 1.1, fourchette de 0 à 10). Les complications ont été rapportées de la façon suivante: 6% d'épistaxis, 12% de sinusite et 5% d'amélioration. Ces patients ont également déclaré des étourdissements (n=12, 5%) et un écoulement de mucus (n=2, 0.9%). Certains patients ont également déclaré des brûlures abdominales (n=12, 5%) et une sensation d'étouffement (n=2, 0.9%). Lorsqu'on a posé la question, 185 patients (88%) ont affirmé que l'EGD transnasale sans sédation est bien tolérée, facile et sûre. On peut l'effectuer en consultations externes, au moyen d'une anesthésie topique. Le faible taux de complications, la forte satisfaction des patients et les économies potentielles font de l'endoscopie transnasale une solution de rechange intéressante par rapport à l'EGD classique pour dépister les patients atteints d'une maladie oesophagastroduodénale.

CONCLUSIONS: L'endoscopie transnasale sans sédation : Une expérience canadienne dans la pratique quotidienne.
Esophagogastroduodenoscopy (EGD) is the most frequently performed diagnostic procedure for upper gastrointestinal disorders. It is a relatively safe procedure, but complications do occur, with overall morbidity and mortality rates of 0.13% and 0.004%, respectively (1). The procedure is routinely performed under conscious sedation in North America. Although studies with conventional endoscopes have suggested that EGD with sedation is more comfortable than without sedation (2,3), a significant proportion of morbidity and mortality is related to hypoxia due to conscious sedation, with elderly patients and individuals with underlying cardiopulmonary disease at a greater risk (4-6). In addition, the use of sedation is associated with an increase in costs due to the greater need for patient monitoring, experienced nursing staff, recovery rooms and drugs (7). It also results in an increased loss of work time on the day of endoscopy (8) and the need for the patient to be accompanied home after the procedure.

In Asia, Latin America and in some of the European countries, unsedated EGD is the routine practice (9,10). With the development of small-calibre endoscopes, several studies have explored the feasibility of unsedated EGD with ultra-thin endoscopes via both per oral and transnasal routes. The transnasal route for upper endoscopy causes less gagging and is better tolerated overall than the per oral route. Despite the literature indicating that transnasal endoscopy has many advantages over sedated per oral EGD, it is not often used in Canadian gastroenterology. This may be due to concern over patient selection in the reported trials and the expectations that Canadian patients have to receive sedation when they undergo EGD. We report a Canadian single-centre experience of unsedated transnasal EGD and a review of the literature. The objective of the present study was to assess the feasibility and acceptability of unsedated transnasal EGD in daily practice.

METHODS

Patients due to undergo EGD were given a choice of undergoing unsedated transnasal EGD within the same month or per oral EGD with sedation in six to nine months. Patients who chose unsedated transnasal EGD had the procedure performed in the office by a senior gastroenterologist with experience in transnasal EGD.

All procedures were performed using a Pentax small-calibre esophagogastroduodenoscope (Pentax FG 16V, Pentax Precision Instrument Co, USA). It had an insertion tube outer diameter of 5.3 mm and a 2 mm biopsy channel. Topical anesthesia was induced in the nasal cavity by spraying a solution of 5% lidocaine. No sedation was used. A nurse was present to assist in biopsies and oral suction. The endoscope was inserted through the nose and the upper esophageal sphincter under direct vision, and the esophagus, stomach and duodenum down to the second part were examined as usual. Biopsies were performed when clinically indicated.

All patients were contacted within seven days by a research assistant who was not present at the time of the procedures, and were surveyed using a patient satisfaction questionnaire. They were asked whether they would accept the same procedure in the future if necessary. Patients who had undergone conventional EGD with conscious sedation in the past were asked which procedure they preferred. All patients were also asked to give specific scores in terms of choking sensation, sore throat, nasal discomfort and abdominal discomfort. All variables were assessed by scores between 0 and 10, with 10 being the most severe degree of choking, sore throat, nasal discomfort and abdominal discomfort. Any complications were also recorded.

RESULTS

Between March 2002 and August 2003, 231 patients underwent transnasal EGD. The median age of the patients was 57 years (range 15 to 87 years) and 100 patients were women. The indications for the procedure were dyspepsia (n=115), reflux symptoms (n=52), anemia (n=36), dysphagia (n=10), family history of gastric cancer (n=5), nausea and vomiting (n=4), weight loss (n=3), radiological abnormality (n=3), postsurgical screening (n=2) and postpolypectomy evaluation (n=1) (Table 1). Eighty-seven patients (38%) were found to have no abnormalities. However, other abnormalities found in descending frequency included hiatus hernias, esophagitis, Schatzki's rings, pangastritis, antritis, gastric erosions, gastric polyps, nodules, Cameron lesions, peptic ulcer disease, esophageal and gastric cancers, gastric antral vascular ectasia and bezoars.

Transnasal EGD failed in five of the 231 patients (2%) due to difficulties in intubating the nasal passage. Complete examinations, including the second part of the duodenum and biopsy sampling where indicated, were possible in all other patients once the nasal tract was passed. Patients reported a high degree of acceptability (mean score 6.6, range 1 to 10) and low degrees of choking sensation (mean score 1.8, range 0 to 10), nasal discomfort (mean score 1.7, range 0 to 10), sore throat (mean score 0.8, range 0 to 9) and abdominal discomfort (mean score 1.1, range 0 to 10) (Table 2). The only complications reported by the patients were epistaxis (n=2, 0.9%) and sinusitis (n=1, 0.4%). Some patients also reported transient light-headedness (n=12, 5%) and mucous discharge (n=2, 0.9%). When asked, 185 patients (88%) stated that they were willing to undergo the same procedure in the future if medically indicated. Of the 84 patients who had conventional EGD under conscious sedation in the past, 52 patients (62%) preferred transnasal EGD without sedation.

DISCUSSION

With the development of small-calibre endoscopes, unsedated transnasal EGD is a potentially cost-effective option in investigating patients with symptoms warranting endoscopic
examination of the upper gastrointestinal tract. Two studies (8,11) comparing transnasal EGD using a small-calibre endoscope to per oral EGD using a standard endoscope have shown that the transnasal route of intubation for unsedated upper endoscopy is better tolerated overall than the per oral route. Compared with per oral intubation, transnasal intubation is thought to result in less gagging and retching due to the absence of contact between the endoscope and the base of the tongue, thereby reducing the risk of aspiration. The added advantages of transnasal intubation also include the reduced risk of patients biting the tube and the the ability of patients to speak during the procedure, which may reduce their anxiety levels (12,13). In the present study, significant numbers of patients who in the past had conventional EGD with conscious sedation preferred unsedated transnasal EGD with an ultra-thin endoscope (62%). Furthermore, 88% of the patients stated that they were willing to undergo the same procedure in the future if medically indicated. These results are consistent with the high level of patient tolerability and acceptability of transnasal EGD in the literature. Dumortier et al (14) reported that 982 of 1033 patients (95.2%) who underwent transnasal EGD with 5.3 mm and 5.9 mm endoscopes would agree to undergo the procedure again if necessary.

Several studies have compared unsedated transnasal EGD and unsedated per oral EGD using small-calibre endoscopes. Zaman et al (15) suggested from their prospective study that the per oral route may be more preferable with a 5.9 mm endoscope. However, Press et al (13), in their prospective study of 150 patients, found that patients undergoing transnasal EGD with a 5.9 mm endoscope required less sedation than per oral EGD using the same size endoscope. Some studies suggest that the diameter of the endoscope might play an important role in patient tolerance and in the success of transnasal intubation. In the present study, a 5.3 mm outer diameter fibre optic endoscope was used, and the failure rate of transnasal EGD due to a narrow nasal tract was only 2%. Craig et al (16), who compared a 5.9 mm video endoscope to a 5.3 mm fiberoptic endoscope, found that failure was significantly more frequent with the larger endoscope (19.5% versus 4.6%). Similar findings were noted by Dumortier et al (14) (16.7% versus 4.7%). Some of the other predictive factors for poorly tolerated unsedated endoscopy described in the literature include young age, female sex, high anxiety levels and low body mass index (12,13).

Complications of transnasal EGD are rare. Epistaxis is the most common complication but it is usually mild and resolves spontaneously. Less than 1% of the patients in the present study had epistaxis. The rate of epistaxis ranges from 0% to 22.6% in the literature, and appears to occur more often with a 5.9 mm video endoscope than with a 5.3 mm fibre optic endoscope (14-16). Only a single incidence of upper esophageal perforation has been reported (15).

One major issue with transnasal EGD is the decreased ability of physicians using the small-calibre endoscopes to carry out satisfactory examination because of the technical difficulty due to greater flexibility and limited optical capabilities. The development of the small-calibre video endoscopes has provided an improved image compared with the older small-calibre fibre optic endoscopes, although the resolution is inferior to that of modern high-resolution endoscopes. In our experience, image quality is somewhat impaired with small-calibre endoscopes of both generations when secretions and bubbles are present due to poorer suction, lavage and air suflation. However, several studies have reported that the rates of completion of the examination to the second part of the duodenum are high (13,15), as was shown in the present study. The histological results with the small-calibre endoscopes using pediatric biopsy forceps (cup diameter of 1.8 mm compared with the 2.2 mm cup diameter of conventional forceps), through their 2 mm channels have been reported in some studies to be of good quality (13,16) and even comparable to that of standard endoscopy (12). Saeian et al (17) showed in their prospective study that specimens obtained from conventional forceps and pediatric forceps were equivalent for providing documentation of Helicobacter pylori eradication with a rapid urease test. Whether the specimens provided by pediatric biopsy forceps are adequate in providing a diagnosis with equivalent accuracy to those obtained by the standard forceps needs to be further investigated.

Unsedated transnasal EGD can also be used in patients in whom per oral intubation is not possible (eg, tongue cancer and jaw fracture). Cirrhotic patients are at particular risk for developing prolonged hepatic encephalopathy following sedation. Unsedated transnasal EGD has been used in this patient group and was shown to be accurate in detecting and grading esophageal varices (18). Although transnasal EGD is not routinely used for therapeutic endoscopy, unsedated transnasal endoscopic placement of percutaneous endoscopic gastrostomy has already been reported (19,20).

Although transnasal EGD appears to be a safe and effective alternative to conventional diagnostic EGD, it should be avoided in patients with coagulopathy and a history of nasal trauma or surgery. Conventional EGD is also more appropriate if there is a possible need for a therapeutic procedure or large biopsies.

CONCLUSIONS
Transnasal EGD is generally well tolerated, feasible and safe. It can be performed with topical anesthesia, without intravenous sedation in an outpatient setting. The low complication rate, high patient satisfaction and potential cost savings make transnasal endoscopy an attractive alternative to conventional EGD to screen patients for upper gastrointestinal tract diseases.

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