Topical glyceryl trinitrate relaxes the sphincter of Oddi

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Abstract

**Background/Aim**—Nitric oxide (NO) may be involved in non-adrenergic non-cholinergic (NANC) inhibitory innervation of the sphincter of Oddi (SO). The effects of topical application of glyceryl trinitrate (GTN), a NO donor, upon SO motility were examined.

**Methods**—Nineteen patients undergoing routine SO manometry for investigation of abdominal pain were studied. After routine recording of SO motility, they were randomised into three groups to receive 10 ml of normal saline, 5 mg GTN (0.5 mg/ml) or 10 mg (1 mg/ml) GTN. Drug solutions were infused topically onto papilla via the manometry catheter and recordings were continued for a further 5 minutes.

**Results**—There was no significant change in SO motor variables following application of normal saline. GTN reduced SO tonic and phasic contractions. In four patients, there was complete abolition of all phasic contraction.

**Conclusions**—Local application of GTN inhibits SO motility. This may have application for diagnostic and therapeutic biliary endoscopy.

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Delivery of bile into duodenum is controlled by hepatic bile secretion, gall bladder contraction and the sphincter of Oddi (SO). The hormone cholecystokinin (CCK) is believed to be responsible for postprandial gall bladder contraction and SO relaxation. Recent studies have shown that the effect of CCK upon the SO is mediated through stimulation of the non-adrenergic non-cholinergic (NANC) nerves and nitric oxide (NO) is an important element of this pathway.

There have been few studies on the effect of NO on the function of the SO. Sivka et al localised the presence of NO synthase (NOS) in nerve fibres and bundles of human SO and demonstrated that topical application of S-nitroso-N-acetylcysteine (SNAC), an NO donor, inhibits SO motility. However, SNAC needs to be freshly prepared, is not stable over a prolonged period of time, and is not widely available. The aim of the present study was to examine the effect of glyceryl trinitrate (GTN; Nitrocin, Schwarz Pharma, UK), a form of NO donor that is commercially available, on SO motility.

**Methods**

**PATIENTS**

Patients undergoing routine SO manometry for investigation of upper abdominal pain were examined. The majority of patients underwent manometry for investigation of postcholecystectomy pain with a median duration of 2.1 (SD 1–5) years after surgery. Only two patients still had gall bladder in situ. None of the patients was on regular medication with nitrates or calcium channel blockers. All of the patients had been investigated with normal abdominal ultrasound scans and endoscopic retrograde cholangiopancreatography (ERCP) before manometry.

**SO MANOMETRY**

After an overnight fast, patients were sedated with 5–10 mg intravenous midazolam. A standard triple lumen polyethylene manometric catheter (SOM-21-LEHMAN, Wilson Cook Medical Inc, Winston-Salem, USA) with an external diameter of 1.7 mm and luminal diameter of 0.5 mm, was introduced into the common bile duct via an Olympus JFIT10 duodenoscope. The manometric catheter had a length of 200 cm with three lateral openings of 0.5 mm in diameter at 2 mm intervals. The most distal end of the catheter was marked by six black rings 2 mm apart to facilitate positioning of catheter in relation to the ampulla.

The catheter was perfused continuously with sterile water using a low compliance pneumohydraulic capillary infusion pump (Arndorfer Medical Specialities, Wisconsin, USA) at a flow rate of 0.25 ml/min. This was connected via a transducer to a computerised polygraph (Albyn Medical Version 6-0, UK). The transducer was calibrated before each study and catheter lumen occlusion produced a pressure rise in excess of 250 mm Hg/s.

After recording duodenal pressure, which was taken as the zero reference, the papilla was cannulated and the catheter was withdrawn across the SO in 2 mm increments using the black marks on the catheter as a guide. Recordings were obtained for at least 60 seconds at each station. After two such pullthroughs, the catheter was repositioned so that the distal two channels were recording phasic SO contractions. The most proximal channel was therefore located just outside the papilla. Patients were randomised into three groups. The first group of patients received topical application of 10 ml normal saline; the two other groups received 10 ml of GTN at either 10 mg (1 mg/ml; GTN 10) or 5 mg (diluted with 5 ml of...
Figure 1: Change in basal SO tonic contraction (A), amplitude (B), frequency (C), and duration (D) of phasic contraction with the different infusions.

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<th>Baseline manometric motor variables. Results are expressed as mean (SE)</th>
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CBD=common bile duct.

Results

Nineteen patients were recruited into the study. There were 17 females; the median age was 41 (range 29–61) years. Five patients were randomised to saline injection; two groups of seven patients were each randomised to either 5 mg or 10 mg GTN.

The baseline results of SO motor activity are presented in the Table. Apart from the basal SO pressure, the means of all the motor variables were within normal limits as defined by previous studies using a similar perfusion technique in healthy volunteers. The basal SO pressures are slightly higher than previously reported. This is because some of these patients had SO dysfunction according to previously defined criteria. One patient exhibited a basal SO tonic pressure of 90 mm Hg accompanied by tachyodynia (frequency of 12 contraction per second); she was randomised to saline. The second patient who was randomised to 10 mg GTN also had tachyodynia although tonic SO pressure was normal.

There was a non-significant increase in basal SO tonic pressure, amplitude and frequency of phasic activity after injection of 10 ml normal saline (Fig 1). In contrast, mean tonic SO basal pressure and all the variables of phasic SO activity significantly diminished after injection with 5 mg and 10 mg GTN; specifically, phasic contractions became significantly less frequent, were shorter in duration, and had decreased amplitude. Consequently, the motility indices fell from 312 (35) mm Hg min to 40 (19) mm Hg min in the group who received GTN 5 mg and from 462 (125) to 57 (27) mm Hg min in the higher dose GTN group (Fig 2). These effects were seen both in patients who had normal SO function and in those who had abnormal SO motility.

The duration of GTN inhibition of the SO was approximately 2 minutes. Although duodenal motility was not quantitated, video recording also showed cessation of contractility.

Discussion

We have demonstrated that topical application of GTN but not normal saline significantly inhibited SO tonic and phasic activity. Doses of 5 and 10 mg GTN had similar efficacy with a duration of approximately 2 minutes in both cases.

In the human gastrointestinal tract, non-adrenergic non-cholinergic (NANC) innervation is important in nerve mediated relaxation and membrane hyperpolarisation and present evidence indicates that NO is a NANC neurotransmitter. NO is synthesised from L-arginine by the enzyme nitric oxide synthase (NOS); NO then activates soluble guanylate cyclase, catalysing formation of cyclic GMP
Topical glyceryl trinitrate relaxes the sphincter of Oddi which is an inhibitor of smooth muscle contraction. Identification of NOS in a subpopulation of myenteric nerves supports the physiological role of NO in the regulation of intestinal motility. Complex neurogenic and hormonal mechanisms are involved in the control of SO muscle tone and motility. Ingestion of a meal and injection of CCK produce a decrease in SO basal pressure and amplitude and reduce the frequency of phasic contractions, thereby facilitating bile flow into the duodenum. The inhibitory effect of CCK is believed to be mediated through NANC inhibitory nerves. NO has been demonstrated to control SO motility in animals and the presence of NOS has been confirmed in the SO neurons of rabbits. In humans, NOS has been localised to nerve bundles and fibres by NADPH diaphorase immunohistochemical staining and topical application of SNAP inhibits SO motility.

In humans, sublingual GTN was found to be effective in improving post-cholecystectomy pain associated with SO dysfunction although side effects limited its therapeutic potential. Staritz et al demonstrated that sublingual GTN lowered SO basal pressure and contraction amplitude during SO manometry. In contrast with our findings, SO frequency was not inhibited but this difference could be attributed to drug dosage and mode of administration. The same investigators subsequently showed that common bile duct stones between 6 and 12 mm in diameter could be removed from an intact papilla after GTN induced SO relaxation. Sublingual GTN spray could facilitate papillotomy during ERCP. We speculate that topical application of GTN could be of clinical application during ERCP as it inhibits both duodenal and SO motility. When administered orally, sublingual GTN can cause headache and may cause hypotension. Local administration to the papilla was not associated with systemic effects and high volumes can be targeted to the SO. This may have advantages over the administration of systemic drugs. It would be helpful in inhibiting duodenal and SO motility for patients in whom buscopan or glucagon are contraindicated. It could also facilitate extraction of small stones from the biliary tree without resort to sphincterotomy or papillary balloon dilatation. However, its duration of action is relatively short and this may limit its clinical application.

In summary, our present study showed that topical application of GTN, an NO donor, inhibits SO tonic and phasic contraction. This mode of GTN delivery may be of clinical application during ERCP cannulation and stone extraction.