Hereditary proctalgia fugax and constipation: report of a second family

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Abstract
A second family with hereditary proctalgia fugax and internal anal sphincter hypertrophy associated with constipation is described. Anorectal ultrasonography, manometry, and sensory tests were conducted in two symptomatic and one asymptomatic subjects within the same family and further clinical information was obtained from other family members. The inheritance would correspond to an autosomal dominant condition with incomplete penetration, presenting after the second decade of life. Physiological studies showed deep, ultraslow waves and an absence of internal anal sphincter relaxation on rectal distension in the two most severely affected family members, suggesting the possibility of a neuropathic origin. Both of these patients had an abnormally high blood pressure. After treatment with a sustained release formulation of the calcium antagonist, nifedipine, their blood pressure returned to normal, anal tone was reduced, and the frequency and intensity of anal pain was suppressed. These together improved the quality of the patients’ sleep, which had previously been very troubled because of night time attacks of anal pain.

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Keywords: hereditary proctalgia fugax, internal anal sphincter hypertrophy.

Proctalgia fugax is an unusual condition characterised by sudden, severe transient attacks of anorectal pain without evidence of anorectal disease. Attacks can occur at any time, but are most common at night. Proctalgia fugax is more common in men than women. It begins in early adult life and stops spontaneously in late middle life.\(^1\) The origin of the pain is still a subject of some controversy. Hurst\(^2\) suggested that sphincter spasm might be the cause, and Douthwaite\(^3\) later ascribed it to spasmmodic contractions of all the muscles of the pelvic floor. Other uncontrolled studies have noted that patients with proctalgia fugax exhibit excessive contractility of the sigmoid colon\(^4\) and a high incidence of symptoms of irritable bowel syndrome,\(^5,6\) although the episodes of pain were not accompanied by an acute bowel disturbance. The abnormally high prevalence of psychiatric disturbance\(^7\) in patients with this condition has suggested that it is a functional disorder of the anal sphincter that can be aggravated by psychological factors.

Kamm \textit{et al}\(^9\) recently described several members of a family with proctalgia fugax and constipation associated with hypertrophy and hyperemia of the internal anal sphincter. We recently discovered a second family living in the Sheffield area with similar symptoms and laboratory (physiological) findings. This report confirms the description of the first family and presents some new clinical, physiological, and therapeutic data that help to explain the pathophysiology of this syndrome and may indicate its management.

Methods

Patients

Case 1 (PB)
This 60 year old man was referred with moderate constipation and anal pain. His constipation had been present since childhood but has become more troublesome in the past 15 years. His stools were hard in consistency and required considerable effort to evacuate, but he had treated himself successfully with lactulose and bulking agents. His episodes of anal pain occurred particularly at night, lasted about 10 minutes, and caused him to wake up five to six times per night.

Physical examination showed high blood pressure (185/120 mm Hg) and a thickened anal sphincter, which was so tight that we could not insert a simulated stoo. Sigmoidoscopy findings were normal, but a barium enema showed a megacanth.

Case 2 (NC)
A 66 year old woman, a sibling of case 1 (PB), was first referred to our unit in 1988 with a seven month history of cramping anal pain and moderate constipation that had started shortly after an episode of viral gastroenteritis. At that time, physical examination showed appreciable perineal descent, a tight anal sphincter, and faecoloma in the rectum. Barium enema showed a megacanth with multiple diverticula in the descending and sigmoid colon. Her general health was good except for raised blood pressure (160/110 mm Hg).

During a one and a half year period of follow up, her anal pain was not relieved by quinine, tranquilisers, or antispasmodics, but her moderate constipation was improved by combination of lactulose, fibre supplement, and suppository. The patient did not make any other contact with us until we learned that she was related to case 1 (PB). Her anal pain was
still present, but of lower intensity than previously and her bowel habit was regular on a combination of lactulose, bulk laxatives, and suppositories.

Case 3 (HB)
This 32 year old woman was the daughter of case 1 (PB) and had no complaints except that her stools were hard and pelletty and often required much effort to evacuate. She had not experienced anal pain. Her past medical history showed a seronegative rheumatological disease for which she was receiving intra-articular steroid injections. Physical examination was unremarkable. Her blood pressure was normal (120/80 mm Hg).

Other family members
Other family members were not available or did not wish to be studied. However, their medical history, and whether they had or have the same symptoms were known by the above patients, from whose information the family tree was constructed (Fig 1). Additional information was gained from telephone conversations with affected family members.

Apart from describing her own symptoms RS gave information about her mother, EL, who had had a colostomy to remove a distal rectal mass, which was thought to be malignant, but was later identified histologically as hypertrophied muscle (nearly 40 years ago). The operation did not improve the pain. In all subjects, the anal pain had begun between the second and sixth decade of life.

METHODS

Anorectal function tests
With the subjects lying in the left lateral position with the hips flexed, a manometric probe, consisting of a polyvinyl 7 lumen tube with an external diameter (OD) of 4 mm was inserted into the rectum. When correctly positioned, manometric side holes were situated in the anal canal at approximately 0-5, 1-0, 2-0, and 4-5 cm from the anal margin. A highly compliant, thin walled balloon, constructed from a 6 cm cylinder of unstretched condom (Durex Dry; LRC Products Ltd, London, England), was tied to the probe at a normal position of 5–11 cm from the anal verge. The side holes were perfused with water at a rate of 0-2 ml/min by a low compliance, pressurised perfusion system (Arndorfer Medical Specialities Inc, Wisconsin, USA), and pressures were measured by transducers (PDCR 75, Druck Ltd, Groby, Leics, UK), situated in each perfusion line and connected through amplifiers to a multichannel chart recorder (Lectromed MTPX, Ormed Ltd, Welwyn Garden City, UK). The pressure within the balloon was measured by a water filled, non-perfused catheter connected to a transducer.

The electrical activity of the sphincter was recorded using a bipolar electrode consisting of two trimel-coated wires (diameter, 0-025 mm) with their ends bare and hooked and staggered so that they did not come into electrical contact.

After insertion of the probe, anorectal manometry was recorded under resting conditions for 15 minutes. After the resting period, subjects were asked to contract the anal sphincter maximally. This was repeated two more times with gaps of at least one minute between contractions. After a further gap of at least five minutes the rectal balloon was serially inflated with a rapid infusion of air (50 ml/s) at volumes of 10, 20, 40, 60, 100, 150, 200, 250, and 300 ml. If the subject experienced discomfort at any of these volumes then the higher volumes were not used. Inflations were maintained for one minute, the balloon was then deflated and inflated with the next volume after a gap of one minute. At each volume the subject was asked whether he or she felt the distension, and if so whether the sensation was that of gas, desire to defecate, or discomfort. The lowest balloon volumes required to induce initial perception, sensations of gas, desire to defecate and discomfort were noted as well as the lowest volume required to induce a sphincter relaxation.

After completing the series of distensions, subjects were requested to strain down as if they were attempting to defecate. This manoeuvre was repeated on three occasions.

Anorectal endosonography
Anal endosonography was performed by one radiologist (RJP) using a Bruel and Kjaer (Naerum, Denmark) ultrasound scanner equipped with a 7 MHz rotating probe, 6 cm long and 1·5 cm wide. The probe was inserted just inside the anal canal without any previous bowel preparation and cross-sectional ultrasound images were obtained at several sites in the anal canal. From these images, a value for the maximum thickness of the internal anal sphincter was obtained.

Medical treatment
The two symptomatic patients (PB and NC) were treated with a slow release formulation of the calcium antagonist, nifedipine (Adalat retard, 10 mg), which was administered at a
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Results
ANORECTAL FUNCTION TESTS

Anal pressure
The manometric data are presented in Table I. In each of the three patients there were prominent ultraslow waves with a periodicity of 0.5–1 min (Fig 2). Although maximum resting anal pressures (MRAP) were very high in each case, squeeze increments were within the normal range (Table I) and were similar whatever position on the ultraslow waves they occurred.

Rectal distension
Rectal sensitivity was blunted in two of our subjects. The rectal volumes that were required to induce a desire to defecate were abnormally increased in cases 1 and 2 but normal in case 3, who was asymptomatic (Table I). Rectal compliance was increased compared with normal data in all three cases, but it was lower in case 3 than in the two symptomatic subjects (Table I).

Straining
It was impossible to test the mechanics of defecation in these subjects because the sphincter was too tight to insert a simulated stool. Nevertheless, there was no electromyographic evidence for a paradoxical external anal sphincter contraction (anismus) when any of these subjects strained as if to defecate.

Anal and rectal endosonography
The maximum thickness of the internal anal sphincter was much higher than the normal range in each of the three patients (Table I).

Case 1 (PB)
After starting the drug, the anal pain frequency was decreased from 5–6 to 2–3 times a night and remained at this level for six months. In addition the severity of pain had improved to 65% of the original intensity within two weeks and this stabilised at around 45% after a further three months. These effects improved the patients’ sleep and general well being. Constipation, however, was not improved by this treatment. A repeat anal manometry while the patient was receiving treatment showed that the maximum resting anal pressure (MRAP) had decreased (before drug it was 150–250 cm H2O and afterwards 100–210 cm H2O), but was still abnormally high. Blood pressure stabilised at around 135/90 mm Hg during treatment with Adalat retard.

Case 2 (NC)
After two weeks of treatment, the frequency of anal pain was decreased from 7–8 to 3–4 times a night and the intensity was improved by 50%, causing a much better quality of sleep. There was, however, no obvious change in her bowel habit, which she regulated with fibre supplements, suppositories, and lactulose. A repeat anal manometry showed MRAP had decreased (before drug it was 150–200 cm H2O and afterwards 125–175 cm H2O) but was still abnormally high. Blood pressure had reduced from 160–110 mm Hg to a stable value of 130/90 mm Hg.

Discussion
This paper describes the clinical, physiological, and ultrasonographic features of a second family with proctalgia fugax and internal anal sphincter hypertrophy. The family tree is
similar to that shown in Kamm's report and is compatible with an autosomal dominant inheritance with incomplete penetration.

The clinical, ultrasonographic, and manometric findings in the subjects that form the basis of this paper were very similar to those in Kamm’s report. The subjects in both families had endosonographic evidence of a thickened anal sphincter and manometric recordings of an increased basal pressure with prominent ultraslow waves. Ultraslow wave oscillations are thought to be generated by activity in the smooth muscle of the internal anal sphincter, and occur in conditions such as haermorrhoids and anal fissures where the ‘resting’ anal pressures are raised. They also occur in normal subjects, but only when resting pressure is greater than 100 cm H2O. The resting pressure in the anal canal is normally increased by activity in the sympathetic nervous system. Thus, the coincidence of vascular hypertension and sphincter hypertension in two of our patients might suggest a common pathophysiology, but there was no evidence of autonomic dysfunction or any dysfunction in any other part of the gastrointestinal tract that might indicate autonomic neuropathy in our patients, or in Kamm’s patients.

Kamm et al reported that the muscle of the lower rectum as well as the internal anal sphincter was thickened in two of his three patients. Although we could not confirm this finding in the three patients we studied, another member of the same family had evidence of rectal muscle hypertrophy in the resected specimen.

Kamm concluded that the internal sphincter hypertrophy and hypotonia was caused by a primary myopathy on the basis of both histological and pharmacological data. Histology of the circular smooth muscle of the internal anal sphincter showed periodic acid Schiff positive polyglycan bodies in the smooth muscle fibres and increased endomyal fibrosis. Polyglycan bodies have never been described in smooth muscle, but similar features have been noted in a rare polycaccharide storage myopathy involving the striated muscles of the limb girdles. Pharmacological studies on strips of muscle from the internal anal sphinter were completely unresponsive to electrical field stimulation and a battery of pharmacological agonists and antagonists. These data contradict the findings of normal sphincter relaxation in response to rectal dilatation in vivo, but the authors emphasise that the tissue did not show any signs of deterioration or necrosis (Hoyle and Kamm, personal communications). In contrast to Kamm’s data, physiological data from two of our subjects showed no sphincter relaxation in response to rectal distension, although sphincter relaxation was present in the records from one of them (NC) five years previously. We cannot rule out that the lack of sphincter relaxation observed in our patients with up to 300 ml distension is secondary to a megarectum, but sphincter relaxation was absent even at volumes that produced a normal desire to defecate, and we have previously observed that rectal sensation is always accompanied by internal anal sphincter relaxation in elderly patients with a functional megarectum. The physiological findings are reminiscent of short segment Hirschsprung’s disease, which also presents with internal anal sphincter hypertrophy and hypotonia, and raise the possibility that this condition may be a neuropathic disorder of the sphincter, perhaps a dying back of the enteric nerve terminals that mediate sphincter relaxation. The major argument against this interpretation is the result of the pharmacological studies of Kamm and Heyle, and the identification of a wide range of neuronal effects on the sphincter. Perhaps our patients do not have the same condition as those described in Kamm’s paper.

The considerable improvements in pain after administration of a calcium antagonist were associated with a reduction in resting sphincter pressure and would be compatible with the notion that the episodes of pain were caused by exacerbations in the tension in the internal anal sphincter muscle. Kamm’s group were able to record the activity of the anal sphincter during a bout of pain, using real time endosonography, and observed fluctuations in the diameter of the sphincter, synchronous with pain. Such exacerbations could perhaps be induced by fluctuations in the activity of the sympathetic nervous system, which is known to increase the internal anal sphincter tone. Even during treatment, however, the resting pressure remaining abnormally high with prominent ultraslow waves, and constipation was not relieved. The results of treatment with slow acting calcium antagonists are quite different from the results of surgical excision of a strip of internal sphincter smooth muscle, which was not always successful in abolishing the pain, though constipation was relieved. Strip myectomy would not alter the tone generated in the internal anal sphincter smooth muscle although it would prevent the hypotonia from obstructing the anal canal.

1 Abrahams A. Proctalgia fugax. Lancet 1935; ii: 444.
2 Hurst AF. Constipation and allied intestinal disorders. London: Oxford University, 1909.
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