Myofibroblasts in hollow visceral myopathy: the origin of gastrointestinal fibrosis?

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Abstract
A patient with hollow visceral myopathy is reported in whom light microscopical studies of the small and large intestine showed typical features of degeneration, thinning, and fibrous replacement of smooth muscle of the gastrointestinal tract. Electron microscopy showed a striking increase in collagen with minimal fibroblast proliferation. Smooth muscle fibres had a range of ultrastructural abnormalities including myofilament disarray, electron luency of the cytoplasm, and proliferation of the endoplasmic reticulum. Some fibres seemed to have typical ultrastructural characteristics of myofibroblasts, and others to be transition forms between typical smooth muscle cells and typical myofibroblasts. It seems likely that the fibrosis typical of this disorder has its origin in the transformation of smooth muscle fibres from a purely contractile to a myofibroblast collagen synthetic phenotype.

Case report
A 14 year old girl presented with constipation, intermittent abdominal pain, and vomiting. She underwent appendicectomy at age 17. She continued to experience abdominal pain and at age 18 was admitted with absolute constipation and intestinal obstruction. At laparotomy she was noted to have thickened bowel loops and a thickened dilated colon. No structural obstruction was noted. Investigation over the next two years showed intestinal hypomotility, with formal small bowel motility studies showing uncoordinated phase 3 propagation. She had difficulty with micturition, but no haematuria or dysuria. No other abnormalities were present on examination or investigation. At age 20 she became pregnant and the pregnancy was complicated by episodes of small and large bowel obstruction. A colostomy was performed in an attempt to relieve her large bowel symptoms, and subsequently closed. The labour was uneventful and the child, now aged 10, shows normal development.

During the next 10 years the patient suffered bouts of subacute obstruction and frequent faecal impaction with admissions for abdominal pain, absolute constipation, and vomiting, necessitating left extended hemicolectomy at the age of 22. At age 29 she underwent successful parathyroidectomy for primary hyperparathyroidism with nodular hyperplasia. No other endocrine abnormalities were present and calcium concentrations returned to normal after surgery. Her intestinal symptoms progressed and she vomited copious amounts of fluid every two to three hours. She had no appetite and her food intake declined to zero. A central venous line was inserted for home total parenteral nutrition. By the age of 30 she was still maintained on home total parenteral nutrition, and continued to vomit fluid. She also had gross abdominal distention and absolute constipation. She underwent further ileal resection to reduce her distention and tubal ligation for sterilisation.

Resection specimens of colon and ileum from the abdominal operations at ages 22 and 30 were studied histologically. Specimens were taken fresh and samples fixed in 2.5% glutaraldehyde for transmission electron microscopy and 10% formal saline for light microscopy. Samples from each specimen of bowel were taken at 5–10 cm intervals. All specimens of colon and ileum showed similar features. At light microscopy (haematoxylin and eosin, and elastic Van Gieson stains) there was thickening of the bowel wall by fibrous tissue with abnormalities of the muscularis propria, especially the external longitudinal layer (Fig 1). This showed loss of smooth muscle fibres, with remaining fibres showing...
hypertrophy, vacuolation, or degenerative features (Fig 2). Mucopolysaccharide and amyloid deposition was not detected with Alcian blue-periodic acid Schiff and Congo red stains. The submucosal and myenteric neural plexuses seemed normal in number and morphology with no intraneuronal inclusions or vacuolation present. Staining with neuron specific enolase and neurofilament antibodies (Sigma, UK) by indirect immunoperoxidase methods showed no abnormalities of distribution or morphology of neuronal cell bodies or axons.

Ultrastructural studies showed increased collagen in the muscularis propria with alterations in smooth muscle fibres including elongation, cytoplasmic vacuolation, proliferation of rough endoplasmic reticulum, myofilament disarray with increased lucency of myocyte cytoplasm, and prominent focal densities. Fibroblast infiltration was not prominent, with occasional fibroblast-like cells seen but not in abnormal numbers (Figs 3, 4). All nerve fibres and terminals present seemed normal in number and morphology. Similar ultrastructural features were present in the ileum and colon, with the most noticeable abnormalities present in the muscularis propria.

Light microscopical studies of smooth muscle of the fallopian tubes showed no histological abnormalities. Sections of the appendix and bladder biopsies were not available.

**Discussion**

Primary chronic intestinal pseudo-obstruction occurs without underlying systemic disease. This syndrome is characterised clinically by abnormalities of gastrointestinal and, in some cases, bladder motility. When thought to involve abnormalities of smooth muscle without abnormalities in myenteric neurons the syndrome is termed hollow visceral myopathy. This is considered to be the commonest cause of chronic primary intestinal pseudo-obstruction. Different patterns of clinical involvement with genetic transmission or sporadic occurrence are recognised. The histopathological findings are similar in all types. At light microscopy smooth muscle layers show fibrosis with vacuolar and other degenerative changes in smooth muscle fibres in the circular and longitudinal muscle layers. Ultrastructural abnormalities in smooth muscle fibres have been recognised, including myocyte damage with perinuclear vacuolation, disorientation and dissolution of myofilaments, electron lucency of the cytoplasm, and swelling of mitochondria. The origin of the fibrosis in this condition is unclear. Our findings suggest an explanation.

The typical clinical and histological features of hollow visceral myopathy, including pseudo-obstruction of the bowel with fibrosis, thinning, and vacuolation of the smooth muscle of the muscularis propria were present in our case. It is likely that our case represents the sporadic type of the disorder, in the absence of any family history. The familial form cannot be excluded, however, without full investigation of other family members, which is not possible here. At the ultrastructural level alterations of smooth muscle fibre included vacuolation, proliferation of the rough endoplasmic reticulum in the peri-nuclear region, disturbances in the distribution of myofilaments and electron-lucency of the myocyte cytoplasm. All of these changes are recognised in hollow visceral myopathy. We found, however, that the pronounced fibrosis was not accompanied by fibroblast infiltration. Also, the changes in smooth muscle cells consisted of a range of phenotypic alterations including forms showing features consistent with myofibroblast differentiation. These included elongation, notching or convolution of the nucleus, proliferation of rough endoplasmic reticulum, and electron lucent cytoplasm containing prominent 6–10 nm filaments with prominent focal densities. Pinocytic vesicles were present near the plasma membrane and basement membrane was present around these cells. Such myofibroblast type cells were focally surrounded by mature collagen fibres.
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The concept of a mesenchymal cell with features of both fibroblast and smooth muscle cells - termed a myofibroblast - was introduced in 1971 by Gabbiani et al. and Majno et al. Since that time there has been an increased awareness that myofibroblast type cells are part of the normal structure of many tissues, including the umbilical cord, lung, and colon. Myofibroblasts have been implicated in fibrous reactions in a wide range of pathological conditions, including fibrosing conditions of the large bowel. The origin of these cells has been the subject of much debate. It has been suggested that myofibroblast type cells may originate from fibroblasts, smooth muscle cells, pericytes, and, in thrombus, from circulating blood cells.

In our case the origin of the myofibroblasts is not clear but the presence of transitional forms of smooth muscle cells strongly suggests an origin from such cells rather than extension of pericyte cells or infiltration of circulating blood cells. Experimental models in which the oestrogen stimulated rat uterus was seen to develop smooth muscle cell alterations resembling the ultrastructural characteristics of fibroblasts emphasised the relation between the two cell types and the presence of intermediate forms, and these studies also documented the production of connective tissue proteins by such transformed smooth muscle cells. The changes seen in smooth muscle cells of the muscularis propria in our case may thus be adaptive, rather than of primary pathogenetic significance, as the inner-vation of the bowel seems morphologically normal and the response of the smooth muscle seems to follow a recognised pathophysiological response.

The precise stimulus to such alteration and the reversibility of the change, however, is not clear.

JEM is a Wellcome Trust Research Fellow and holds the Gillson Scholarship in Pathology of the Worshipful Society of Apothecaries. Our work is supported by the Motor Neuron Disease Association of Great Britain.