Observational Study

Structure of the myenteric plexus in normal and diseased human ileum analyzed by X-ray virtual histology slices

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Abstract

BACKGROUND
The enteric nervous system (ENS) is situated along the entire gastrointestinal tract and is divided into myenteric and submucosal plexuses in the small and large intestines. The ENS consists of neurons, glial cells, and nerves assembled into ganglia, surrounded by telocytes, interstitial cells of Cajal, and connective tissue. Owing to the complex spatial organization of several interconnections with nerve fascicles, the ENS is difficult to examine in conventional histological sections of 3-5 μm.

AIM
To examine human ileum full-thickness biopsies using X-ray phase-contrast nanotomography without prior staining to visualize the ENS.

METHODS
Six patients were diagnosed with gastrointestinal dysmotility and neuropathy based on routine clinical and histopathological examinations. As controls, full-thickness biopsies were collected from healthy resection ileal regions after...
Hemicolecction for right colon malignancy. From the paraffin blocks, 4-µm thick sections were prepared and stained with hematoxylin and eosin for localization of the myenteric ganglia under a light microscope. A 1-mm punch biopsy (up to 1 cm in length) centered on the myenteric plexus was taken and placed into a Kapton® tube for mounting in the subsequent investigation. X-ray phase-contrast tomography was performed using two custom-designed laboratory setups with micrometer resolution for overview scanning. Subsequently, selected regions of interest were scanned at a synchrotron-based end-station, and high-resolution slices were reported. In total, more than 6000 virtual slices were analyzed from nine samples.

RESULTS
In the overview scans, the general architecture and quality of the samples were studied, and the myenteric plexus was localized. High-resolution scans revealed details, including the ganglia, interganglional nerve fascicles, and surrounding tissue. The ganglia were irregular in shape and contained neurons and glial cells. Spindle-shaped cells with very thin cellular projections could be observed on the surface of the ganglia, which appeared to build a network. In the patients, there were no alterations in the general architecture of the myenteric ganglia. Nevertheless, several pathological changes were observed, including vacuolar degeneration, autophagic activity, the appearance of sequestosomes, chromatolysis, and apoptosis. Furthermore, possible expulsion of pyknotic neurons and defects in the covering cellular network could be observed in serial slices. These changes partly corresponded to previous light microscopy findings.

CONCLUSION
The analysis of serial virtual slices could provide new information that cannot be obtained by classical light microscopy. The advantages, disadvantages, and future possibilities of this method are also discussed.

Key Words: Enteric nervous system; Immunohistochemistry; Neuropathy; Synchrotron; Virtual histology; X-ray phase-contrast nanotomography

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Core Tip: Full-thickness biopsies of 1 mm diameter and up to 1 cm length from the human ileum were scanned using two laboratory-based µ-computed tomography setups to study the architecture of the enteric nervous system (ENS) and further scanned by a synchrotron-based end-station for histopathological studies, without any staining. Several pathological neuronal changes, such as vacuolar degeneration, autophagic activity, appearance of sequestosomes, chromatolysis, and apoptosis, were identified in diseased patients. Phenomena that were undetectable by light microscopy were observed. The relationships among various tissue components could be followed in all directions. Thus, this method provides a unique analysis of the ENS.

Citation: Veress B, Peruzzi N, Eckermann M, Frohn J, Salditt T, Bech M, Ohlsson B. Structure of the myenteric plexus in normal and diseased human ileum analyzed by X-ray virtual histology slices. World J Gastroenterol 2022; 28(29): 3994-4006

URL: https://www.wjgnet.com/1007-9327/full/v28/i29/3994.htm
DOI: https://dx.doi.org/10.3748/wjg.v28.i29.3994

INTRODUCTION
The neurons of the enteric nervous system (ENS) assemble into ganglia, forming the submucosal and myenteric plexuses[1]. The ganglia, which also contain glial cells, are connected by nerve fascicles and are surrounded by telocytes and interstitial cells of Cajal (ICC), as well as by other cells and fibers of connective tissue. Various diseases of the ENS may lead to severe bowel motility dysfunction in the form of chronic intestinal pseudo-obstruction (CIPO) and enteric dysmotility (ED)[2,3]. Enteric neuropathy is seldom studied because only a few centers are available for the evaluation of gastrointestinal motility, access to the myenteric plexus requires full-thickness bowel biopsies, and pathologists subspecializing in neuromuscular diseases of the ENS are scarce. Subsequently, the neuropathy diagnosis may be ignored or delayed for several years, which prevents proper healthcare of the patients[4].
We recently presented a methodology to use X-ray phase-contrast nanotomography to study the volume and structure of the ENS[5]. Neurons and surrounding structures are visible using this method without any prior staining, allowing the possibility to study large sample volumes[5,6]. The aim of the present qualitative analysis of virtual slices was to describe in detail the structure of human ileal myenteric ganglia in the normal human ileum as well as in patients with CIPO or ED.

**MATERIALS AND METHODS**

**Patients and samples**

Six patients with severe gastrointestinal pain and dysmotility were examined using esophageal manometry, gastric emptying scintigraphy, antroduodenojugal manometry, and colonic transit time, after exclusion of organic disease. Patients were diagnosed with CIPO when they fulfilled the following three criteria: a medical history compatible with pseudo-obstruction, documented events or chronic signs mimicking mechanical obstruction (bowel dilatation and/or air/fluid levels), and absence of mechanical obstruction or other organic causes for these symptoms and findings[3]. The criteria for ED were documented abnormal contractile activity, but no history of episodes or current signs mimicking mechanical obstruction and the absence of any medication that could lead to the observed motor abnormalities[2].

A previously described laparoscopy-assisted technique for ileal full-thickness biopsies and the preparation of biopsies was used[4,7]. Briefly, 1.0 cm × 1.5 cm large full-thickness biopsies were cut from the ileum and embedded in paraffin. Transversal and horizontal 3-5-mm thick serial sections were stained according to the protocol for CIPO analysis[8]. Histopathological findings were evaluated by a pathologist (B.V.) and classified as previously described[9]. The criteria for neuropathies were based on criteria suggested by the Gastro 2009 International Working Group for Gastrointestinal Neuromuscular Pathology[10]. The histopathological diagnoses and various stains, such as CD117 for ICC, CD34 for telocytes, S100 for Schwann cells, periodic acid-Schiff (PAS) with or without diastase (PAS-D) for carbohydrates or their compounds (e.g., glycogen and mucin), and p62 for sequestosome, were obtained from the medical records (Table 1). For all patients, the disease initially presented when they were adults without any known heredity. Genetic analysis was not performed. There was no correlation between the varying histopathological findings described in Table 1 and the clinical presentation or other functional findings of patients (data not shown).

Three men aged 45, 54, and 82 years were used as controls, all of whom had undergone hemicolectomy due to malignancy in the right colon. A full-thickness biopsy specimen was collected and stained from the healthy resection region of the ileum. The sample size was adopted based on the beam time available in the laboratory.

**Sample preparation**

From the paraffin blocks of these nine individuals, 4-µm thick sections were prepared and stained with hematoxylin and eosin (H&E) for the localization of the myenteric ganglia under a light microscope. A biopsy punch of 1 mm diameter and up to 1 cm in length centered on the myenteric plexus was taken from the paraffin blocks of patients and controls and placed into a Kapton® tube (Paramount, Indiana, United States) for mounting in the subsequent tomographic investigation[11].

**X-ray phase-contrast tomography**

X-ray phase-contrast tomography was performed on two custom-designed laboratory setups with micrometer resolution, located at the Institute for X-Ray Physics, University of Göttingen, Göttingen, Germany[11] and a Göttingen Instrument for Nano-Imaging with X-rays (GINIX), installed at the P10/PETRAIII beamline (Hamburg, Germany) with sub-micron resolution[12]. All setups used propagation-based, phase-contrast methods, in which the sample is placed between the source and detector, and the phase information is obtained by free-space propagation and self-interference of the coherent X-ray beam without the need for additional optical elements. Phase retrieval is necessary to correctly extract phase information before tomographic reconstruction[13-15].

The two laboratory setups were installed at a liquid metal jet source (Excillum, Kista, Sweden) or a microfocus rotating anode (mm007, Rigaku, Austin, TX, United States). Both setups used high-resolution scintillator-based detectors (Xsight, Rigaku) and employed broad bandpass radiation peaking at the characteristic lines of Ga Ka (9.25 keV) and Cu (8.05 keV) for the liquid jet and rotating anode sources, respectively. Phase retrieval was performed using the Bronnikov-aided correction algorithm[11, 15,16]. As a result, these home-built setups allowed us to image the whole 1-mm wide and several-mm long sample with an isotropic effective voxel size of approximately 1 µm (approximately 15 h scan time per sample), enabling identification of the neural tissue structure (further information about the experimental parameters is provided in Table 2). The two systems could run in parallel and were used to optimize time consumption.
Table 1 Basic characteristics of subjects

| Age, sex | Disease (years) | Clinical diagnosis | Histopathological findings | Histopathological diagnosis | X-ray |
|----------|-----------------|-------------------|----------------------------|-----------------------------|-------|
| 45, M    | NA              | Control           | Normal neurons             | Healthy                     | Healthy |
| 54, M    | NA              | Control           | Normal neurons             | Healthy                     | Healthy |
| 82, M    | NA              | Control           | Normal neurons             | Healthy                     | Healthy |
| 27, W    | 4               | ED/type 1 diabetes| Vacuolated, apoptotic neurons, chromatolysis | Lymphocytic ganglioneuritis | Vacuolated, shrunken neurons, chromatolysis, autophagia |
| 27, W    | 6               | Idiopathic ED     | Vacuolated neurons, chromatolysis | Degenerative neuropathy     | Pre-apoptotic and dead nucleus |
| 32, W    | 2               | Drug-induced CIPO | Vacuolated, shrunken, chromatolysis, hyperplasia ICC | Visceral degenerative neuropathy with axon vaculization, hyperplasia, and vuculization of ICCs, and hypertrophy of the longitudinal and internal circular muscle layers | Only fascicle, no ganglia for evaluation |
| 43, W    | 3               | Idiopathic ED     | Amphophilic shrunken neurons, Vacuolated neurons, chromatolysis | Lymphocytic ganglioneuritis with neurondegeneration | Amphophilic degeneration, vacuolated neurons |
| 52, W    | 4               | Idiopathic ED     | Vacuolated, shrunken neurons, hypoplasia of ICC | Lymphocytic ganglioneuritis | Vacuolated, shrunken neurons, chromatolysis |
| 56, W    | 50              | ED/Ehlers-Danlos  | Vacuolated neurons, chromatolysis, hyperplasia ICC | Lymphocytic ganglioneuritis with vacuolar neurodegeneration and hyperplasia of the ICC | Severe atrophy with reduced volume |

Data obtained from the medical records; disease duration assessed in years. CIPO: Chronic intestinal pseudo-obstruction; ED: Enteric dysmotility; ICC: Interstitial cells of Cajal; M: Man; W: Woman; NA: Not available.

Table 2 Experimental and reconstruction parameters

| Scan configuration                      | Effective pixel size (µm) | Energy (keV) | Number of projections | Phase retrieval scheme | Phase retrieval parameter | Ring removal algorithm |
|-----------------------------------------|---------------------------|--------------|-----------------------|------------------------|--------------------------|------------------------|
| Laboratory setup (liquid metal jet)     | 0.920                     | 9.25 (Kα)    | 1 × 1000 (50 s) or 1 × 700 (40 s) | BAC                    | α = 0.008, β = 0.160     | Wavelet               |
| Laboratory setup (rotating anode)       | 1.072                     | 8.05 (Kα)    | 1 × 1000 (50 s) or 1 × 700 (40 s) | BAC                    | α = 0.07, β = 0.16      | Wavelet               |
| GINIX waveguide (all samples except for ED/Ehlers-Danlos patient) | 0.169                     | 8.00         | 4 × 1500 (1 s) or 3 × 1500 (1 s) | Non-linear Tikhonov    | δ/β = 50, lim1 = 8 × 3, lim2 = 0.5 | Additive              |
| GINIX waveguide (ED/Ehlers-Danlos patient) | 0.176                     | 7.50         | 4 × 1000 (2 s)        | Non-linear Tikhonov    | δ/β = 50, lim1 = 1 × 4, lim2 = 0.1 | Additive              |

1Due to time constraints, some samples were scanned with a lower number of projections and shorter exposure times (in the overview scans) or with less distances (in the high-resolution scans of the drug-induced chronic intestinal pseudo-obstruction patient and of the 40 years old control).
2After voxel binning of 2 × 2 × 2.
3No support, no restriction on phase shift.
GINIX: Gottingen Instrument for Nano-Imaging with X-rays; ED: Enteric dysmotility.

Selected regions of interest (ROIs) of 320 µm × 320 µm × 320 µm were scanned with an isotropic effective voxel size of 169 nm at the synchrotron-based end-station (approximately 2.5-3.0 h scan time), using inline holography based on a coherent divergent beam [8 keV, Si (111) monochromator] exiting from an X-ray waveguide[11,12]. Holographic phase retrieval based on contrast-transfer-function (CTF) approach, ring removal, and tomographic reconstructions were performed using in-house reconstruction pipelines[16]. Further details can be found in Table 2.

Complete information regarding the experimental setups and reconstruction pipelines, which were performed by physicists unaware of the diagnosis, is provided in a previous publication[5]. The analysis of the slices reported in this paper was performed on the data acquired at the synchrotron end-station, on the high-resolution scans.
**Image analysis**
All the obtained volumetric datasets were digitally sectioned along any arbitrary slicing plane, enabling virtual histology of the samples. In the histological evaluation, the term “spindle-shaped cells” was used to describe the cells around the ganglia, which represented either telocytes, ICCs, or fibroblasts/cytes because no differentiation could be made without immunohistochemistry. In cases where identification can be made from double immunohistochemical staining of the same regions[17], the cells are called telocytes or ICC. In total, more than 6000 virtual slices were analyzed from the nine samples.

**RESULTS**
In the overview scans performed with the laboratory setup, the general architecture of the bowel samples was examined with respect to the quality of the sample and the presence and localization of the myenteric plexus. High-resolution scans revealed details of the myenteric plexus, including the ganglia, interganglionic nerve fascicles, and surrounding tissue.

**Normal structure of the ileal myenteric plexus**
**Ganglion:** The shape of the ganglia was uneven toward both the muscle layers and interganglionic nerve fascicles. The smaller or larger irregularities contained one or more neurons and glial cells (Figures 1, 2A, 4A-D, and 3).

In the surface view of the ganglia, a parallel arrangement of telocytes with long thin “primary” telopodes originating directly from the cellular bodies could be observed, from which shorter “secondary” telopodes radiated sidewise, building up a network (Figures 1A-C, 3B, 4A and 8). Occasionally, at the optimal transverse view of the ganglion, a double layer of thin cytoplasmic projections was present (Figure 2B), corresponding to the immunohistochemical demonstration of telocytes and ICC[17].

Both larger and smaller neurons had mostly rounded nuclei and nucleoli, where larger neurons had finely granulated cytoplasm with a paler network between the granules, whereas the smaller neurons had slightly denser and more homogeneous cytoplasm (Figures 2B and 3B-D). Axons were occasionally observed (Figure 2A). Small glial cells were present around and between the neurons (Figures 2A, 3C and D). No degenerating neurons were found in either of the two younger individuals. In the 82-year-old control, however, one neuron with a shrunken, dark body, pyknotic small almost black nucleus and larger autophagic vacuoles could be observed in addition to numerous normal neurons (not shown).

**Nerve fascicles and spindle-shaped cell layer**
The nerve fascicles had normal architecture. Parallel thin telopodes separated the individual axons (Figures 2C and 3A-C). Some spindle-shaped cells had very thin, dark nuclei, whereas the nuclei of others were larger and cigar-like.

**Pathological findings of the ileal myenteric plexus**
Four patients had lymphocytic ganglioneuritis or lymphocytic neuritis, and two patients had only visceral degenerative neuropathy. No alterations were observed in the general architecture of the myenteric ganglia. Nevertheless, several pathological changes could be observed, partly corresponding to the light microscopic findings, as shown in the insets of the figures. Cellular changes occurred in all six patients with dysmotility. Suspected lymphocytes were found within the ganglion only in one patient. The lack of lymphocytic attack on the ganglion in other patients can be explained by the focal nature of inflammation.

**Pathological neuronal changes**
The cytoplasm of several neurons contained vacuoles. Many of the vacuoles contained small dark granules (Figure 5A, C and D) (autophagic vacuoles/Lipofuscin; 5D-inset: PAS-D). In one patient, several neurons exhibited pale cytoplasm (Figure 4A-D) (chromatolysis), and a large homogeneous irregular dark area was present (Figure 4F) (4F-inset: Sequestosome p62). Apoptotic shrunken neurons had compact, dark cytoplasm with small, irregular, pyknotic nuclei (Figures 4C-E, 5B and 6) (4E-inset H&E; 5B-inset: Apoptosis H&E). These neurons were observed in three patients at the ganglion border without telopodes in a few virtual slices (Figure 4B-D), possibly representing the ejection of a dead neuron (Figures 4F-D and 6). Very dense dark inclusions with similar substructures were found in two patients, which are foreign structures for the normal ganglion; one of these bodies is shown in serial virtual slices (Figure 4A-C) (4A-C: Hyaline bodies; 4A and B-insets: PAS-D’). At 6.8 μm in Figure 4A, an inner structure with a central pale area surrounded by an irregular dark ring can be seen within the body (Figure 4B). When viewing 4.0 μm deeper, there is a semi-dense “half-moon” at the edge of the hyaline body (Figure 4C).
Figure 1 Ganglion from healthy human ileum. A: Part of the myenteric ganglion in two portions. Quite regular, almost parallel, thin cytoplasmic projections of telocytes (telopodes) with short, thin “spines” on the surface are observed (between arrows). The scale bar (50 μm) was applied to all subfigures; B: A small portion of the ganglion containing two neurons. On the surface of the ganglion (between the two lines), the cellular nuclei of the two telocytes were observed as dark spots (empty arrowheads). Telopodes radiate from their body. Arrows indicate the telopodes around the ganglion. A telopode can be followed toward the surface (arrowhead); C: 16.9 μm deeper from Figure B, two portions appeared with thin telopodes on the surface; D: 10.1 μm deeper from Figure C, SMCs are observed between the two portions. B-D are representative virtual slices from a series covering 27.0 μm thickness of the ganglion. Ggl: Ganglion; SMC: Smooth muscle cells.

Figure 2 Ganglion and fascicle from healthy human ileum. A: Within a portion of the ganglion, there is a neuron with an axon (thick arrow). Arrowheads indicate glial cells. Thin arrows indicate a single layer of telopodes at the border of the ganglion. The scale bar (50 µm) was applied to all subfigures; B: Part of a ganglion with several large (thick arrows) and medium-large (short thin arrow) neurons. The nuclei contain dark nucleoli. The cytoplasm is slightly granulated. Above the straight line, a small area of the neuropil was observed. Long thin arrows indicate the double-cell layer at the border; C: Normal nerve fascicle and thicker nerve with telopodes separating axons (arrows). Note the single layer of telopodes around the fascicle and nerve (arrowheads). Inset: Normal nerve fascicles stained with double immunohistochemistry. Telocytes run parallel to Schwann cells (light microscopy; S100: Schwann cells red; CD34: Telocytes brown).

Pathological change of the nerve fascicles and periganglional spindle-shaped cell layer
In one patient who was diagnosed with lymphocytic ganglioneuritis, a segment of the periganglional spindle-shaped cell layer showed vacuolization and/or was absent within a 19.9 μm distance along the length axis of the bowel sample (Figure 5). The largest window in this absence was 16.1 μm in the virtual slices (Figure 5B). At a 7.8 μm deeper level (Figure 5C), a double layer of thin cellular projections appeared at the border of the defect, whereas at an additionally 4.7 μm deeper, one intact layer of cellular projections was again present between the ganglion and the surrounding tissue (Figure 5D).

Video-film
In the video film from one patient, more than one dozen neurons and glial cells could be observed (Supplementary material). Almost all the neurons showed various types of degeneration. Furthermore, one pre-apoptotic neuron in the lower part of the ganglion was probably in contact with the tip of the incomplete septum, and hence, extended into the surrounding tissue (Supplementary material).
DISCUSSION

The main qualitative finding in the present study was that X-ray phase-contrast nanotomography could analyze the different cellular components of the myenteric plexus in more than 6000 serial virtual slices, which could reveal minor changes not seen under the light microscope. We could image large volumes with finer sampling resolution with shorter distances between sections along the sample direction, that is, below the standard thickness of a histological scan.

A study of the normal ganglia revealed that their shape is not usually seen in the transverse sections under a light microscope with a well-defined border. In serial virtual slices, the ganglia have smaller or larger irregularities, with one or a few neurons and glial cells protruding into the muscle layers or the intermyenteric connective tissue. Within the ganglia, neurons of various sizes and glial cells could be clearly differentiated. The absence of thin cytoplasmic projections in the neuropil corroborates our previous finding in the ileal ganglia, namely the lack of telopodes in the ileum in contrast to the colon [17]. The presence of one degenerating neuron in the 82-year-old control may be the result of aging starting at around 40-50 years of age [18].

The optimal transverse sections showed two clear layers of cellular projections surrounding the ganglion, in accordance with our previous immunohistochemical findings [17]. The inner layer consisted of cytoplasmic projections of telocytes called telopodes, which were situated almost parallel to each other, in agreement with immunohistochemical observations [17]. The outer layer is composed of an ICC [17].

Pathological changes from the patients revealed different types of neuronal degeneration (vacuolization, autophagic vacuoles/Lipofuscin, chromatolysis, and apoptosis) and the presence of a hyaline body within the ganglion, as observed under a light microscope by the golden standard of various staining techniques [8]. Furthermore, p62+ sequestosomes were also observed in accordance with the findings of Alafuzoff et al. [19].

Three new phenomena were discovered with the present phase-contrast nanotomography, which cannot be observed under a light microscope because of the thickness of the paraffin sections. First, the presence of a semi-dense “half-moon” attached to the hyaline body could hypothetically be one of the end stages of cellular death. Second, the phenomenon of a probable expulsion of dead neurons could be followed in the virtual serial slices because of the occurrence of a short “opening” of the telopode-layer between the surrounding connective tissue and shrunken apoptotic neurons. Third, the total absence of the single telopode layer within the 16.1 µm “window” could possibly have negatively affected the stability of the ganglion.

The virtual histology of the myenteric ganglion using X-ray nanotomography has advantages, disadvantages, and future possibilities. The main advantage is that the method can exactly measure the volumes of chosen tissue components following segmentation, as have been previously published [5]. The second advantage is the possibility to analyze several hundred serial virtual slices at subcellular, isotropic resolution, and through this, discover “nano-changes” which cannot be seen using the light microscope. Because the neuron size is 15-40 µm, the ENS is difficult to examine in conventional histological sections of 3-5 µm. Slow videos make it possible to follow the relationships between the various tissue components. The third advantage is the flexibility in sample preparation; samples can be examined with or without paraffin embedding (e.g., in liquid or epoxy resin) or staining. Thus, the...
Figure 4 Series of virtual slices covering 17.2 μm thickness of the ganglion from one patient. A: Four portions of the ganglion. There is a dark body at the center of one of these areas (arrowheads). An arrow indicates a neuron with a very pale homogeneous cytoplasm below the nucleus. Between the straight lines, a network of telopodes is observed on the ganglion surface. Incomplete septa between portions. Scale bar (50 μm) is applied to subfigures A-D. Inset: Hyaline body in the ganglion [1st level of serial sections; light microscopy; periodic acid-Schiff with diastase (PAS-D) staining]; B: 6.8 μm from Figure A., three portions of the ganglia are completely separated, and the surface area is smaller (between straight lines). The central dark body shows an irregular dense outer layer and central light area (arrowhead). A new neuron with pale cytoplasm is seen in the middle portion (short arrow). Intact telopodes form the border between the ganglion and the periganglional connective tissue rim (thin arrows). An almost triangle-shaped moderately dense area is seen below the telopode telopode corresponding to an apoptotic neuron (long arrow; see also C and D). Inset: The central region of the hyaline body is negative, whereas the outer layer is positive with PAS-D staining (2nd level of serial sections, light microscopy; PAS-D); C: 4.0 μm from Figure B. The hyaline body is without a central pale area (arrowhead), with a semi-dense irregular half-moon to the left. The two neurons had a pale cytoplasm (short arrows). A dark pyknotic nucleus appeared in the semi-dense cytoplasm at the border showing an apoptotic neuron corresponding to the "dark triangle" in Figure B (long arrow); D: 6.4 μm deeper from Figure C. Two of the portions are united. Nuclei are present in neurons with pale cytoplasm (short arrows). Intact telopodes are present between the degenerated apoptotic cells (long arrow) and the periganglional space (thin arrow); E: Digital magnification of a shrunken neuron with a darker cytoplasm, small vacuoles, and enlarged nucleolus. Scale bar: 25 μm. Inset: Pre-apoptotic neurons with pyknotic nucleus and vacuole-containing amphiphilic cytoplasm (light microscopy; H&E staining); F: There is a homogeneous circumscribed "inclusion" in the neuron, suggesting a sequestosome, as seen in the inset (arrow). Scale bar: 25 μm. Inset: Large p62+ aggregates in neurons (arrow; light microscopy; p62 immunohistochemistry). Scales bars of 50 or 25 μm included. Ggl: Ganglion; n: Neuron; PAS-D: Periodic acid-schiff with diastase.

samples require less handling and fewer preparation steps[20].

The current established method to diagnose enteric neuropathy uses immunohistochemistry with a wide range of staining according to a standardized protocol[8,21]. The strength of this method lies in the exact identification of different cell types and tissue structures. The weakness is that there is a wide range of different molecules and structures that can vary in abundance among different patients with the same diagnosis. We do not know whether these changes are causal or primary, or whether they are secondary or compensatory. Due to the large variation between patients, no uniform pattern has been determined for the diagnosis of enteric neuropathy, and thus, some of the information did not add substantial value to the examination. The current description of neurons, glial cells, and spindle-shaped cells appears to be sufficient for diagnosis.
Figure 5 Series of virtual slices covering 19.9 μm thickness of the ganglion from one patient. A: Two portions of the ganglion with smooth muscle cells between them. Long arrows indicate the continuous layer of telopodes with tiny vesicles bordering the ganglion. A string of glial cell nuclei was present within the left portion. The thick arrow shows the vacuole above the nucleus of the neuron. Four small dark granules were present in the other neurons. Inset: Several vacuoles fill the cytoplasm of the two degenerating neurons (light microscopy and H&E staining); B: 7.4 μm deeper from Figure A. Between the thin arrows there is no continuous layer of telopodes, instead some vesicles are seen. The left portion has disappeared. The remaining portion is part of a pre-apoptotic neuron with a dark cytoplasm and pyknotic nucleus (arrowhead). Above this neuron is the nuclei of normal glial cells. The thick arrow indicates a vacuole in the neuron. Inset: Apoptotic neurons with strongly amphophilic cytoplasm and rest of the pyknotic nuclei (light microscopy; H&E staining); C: 7.8 μm deeper from Figure B. The defect of the telopodes (between the thin arrows) was shorter. Double layer of telopodes below the defect. In the upper part of the ganglion, there is a large neuron with a few small dark, dense dots, whereas the pre-apoptotic neuron from the middle of the ganglion is no longer present; D: 4.7 μm deeper from Figure C. There is a continuous layer of telopodes (between arrows) with a “remnant” of the double layer, as shown in Figure C. The dark granules in the neurons were larger and more numerous. Arrowhead shows a neuron with both dark granules and vacuoles. Inset: The cytoplasm of the neurons was filled with diastase-resistant PAS+ lipofuscin granules (light microscopy; PAS-D staining). The scale bar (50 µm) applies to all the subfigures. Ggl: Ganglion; H&E: Hematoxylin & eosin; n: Neuron; PAS: Periodic acid-Schiff; PAS-D: Periodic acid-Schiff with diastase; SMC: Smooth muscle cells.

Fluorescence confocal microscopy has been performed with serial sectioning to mimic three-dimensional (3D) imaging; however, the maximal thickness of viewing is still limited, and the space between sections is larger[22-25]. Recently, another method using immunohistochemistry and confocal microscopy was developed for the quantitative analysis of full-thickness bowel biopsies[26]. These different efforts to improve the analysis of ENS prove the importance of clinicians obtaining more information about the function and dysfunction of the gastrointestinal tract.

Among the disadvantages were the limited access to synchrotron beamtime and the cost of the entire process in terms of money and manpower. Furthermore, because the ganglia of the ENS are very sparse, especially in patients with dysmotility, and the myenteric plexus can fold and curve beneath what is visible on the surface of the block, it is very difficult to correctly punch the tissue of interest based on a histological section. Hence, multi-scale X-ray phase-contrast tomography, which involves scanning a tissue block with a mm-sized field-of-view at µm-resolution and then imaging specific features of interest in greater detail using local nanotomography on the full block (instead of extracting biopsy punches as in this work), would be preferable and has been proven to be feasible at different synchrotron beamlines, such as TOMCAT[27,28] and P10/GINIX[29-31].

In the future, the combination of immunohistochemistry with X-ray phase-contrast nanotomography, such as immuno-electron microscopy[32], or with joint X-ray phase-contrast tomography and focused ion beam scanning electron microscopy (FIB-SEM) could provide unique possibilities to study targeted cells and their internal and external relationships[33]. In addition, this technique can determine the exact volume of the target tissue components[5].

CONCLUSION

Virtual sectioning by X-ray phase-contrast nanotomography is a method that provides a unique analysis of the myenteric ganglion, revealing minor pathological changes that cannot be discovered using light microscopy or other two-dimensional scanning methods to study the relationship between various tissue components.
Figure 6 Series of virtual slices covering 6.4 μm thickness of the ganglion from one patient. A: Neuron with shrunken pyknotic nucleus and dark cytoplasm containing vacuoles at the edge of the ganglion (arrowhead). A part of the cytoplasm of the pre-apoptotic neuron is present between the telopodes (arrows); B: 2.0 μm deeper from Figure A. The lower telopode was discontinuous (arrows); C: 0.7 μm deeper from Figure B. The two telopodes are intact, and almost all of the cytoplasm between them has disappeared; D: 0.7 μm deeper from Figure C. Pre-apoptotic neurons seem to be in contact with the lower telopode; E: 3.0 μm deeper from Figure D. The pre-apoptotic neuron (arrowhead) lies within the ganglion, without contact with the telopodes. Arrows point to two telopodes on the figures. The scale bar (25 μm) applies to all the subfigures.

ARTICLE HIGHLIGHTS

Research background
The enteric nervous system (ENS) is difficult to study because of its deep localization in the intestinal wall. The limited sections visualized by light microscopy cannot show the size of the neurons or the interconnection between different parts of the ENS or different cell types.

Research motivation
Since the ENS is difficult to examine, patients with gastrointestinal dysmotility are not properly examined, diagnosed, and treated. They can experience symptoms for several years before a proper diagnosis is established.

Research objectives
The main objective of this study was to examine whether X-ray phase-contrast tomography could improve the visualization of the ENS. We realized that the ENS could be visualized without any prior staining, and the information provided was superior to conventional light microscopy in some respects. Thus, this method should be further evaluated in future studies.

Research methods
Full-thickness biopsies from the ileum of patients with dysmotility and controls were examined using X-ray phase-contrast tomography. For comparison, the same samples were examined using immunohistochemistry. This is the first time that bowel biopsies have been examined using X-ray tomography, and the histopathology has been described in detail.

Research results
This study showed that X-ray phase-contrast tomography can be used to study the ENS in detail to describe normal and pathological cells and tissue structures. The interconnections between the cells and
structures can be visualized. Some new findings could not have been observed using conventional light microscopy.

**Research conclusions**
Virtual sectioning by X-ray phase-contrast nanotomography provides a unique analysis of the myenteric ganglion, revealing minor pathological changes that cannot be discovered by light microscopy, to follow the relationship between various tissue components.

**Research perspectives**
This method needs to be further studied in larger cohorts, with scanning of whole paraffin blocks combined with different staining methods to try to improve the identification of different cell types and tissue structures. More healthy controls should be examined to obtain reference values for health and disease.

**ACKNOWLEDGEMENTS**
The authors thank Maria Teresa Moreira Jara and Maria Nilsson for their technical assistance and the pathologists for control sample collection. The valuable discussion with Anca Dragomir, MD, PhD, Uppsala University, regarding possible artifacts, is greatly appreciated.

**FOOTNOTES**
**Author contributions:** Veress B performed the histochemical and pathological analyses of the X-ray data and wrote the initial draft together with Ohlsson B; Peruzzi N, Eckermann M, and Frohn J scanned the samples; Peruzzi N analyzed the technical data; all authors planned and designed the study, contributed to intellectual analysis of the results, and approved the final version of the manuscript.

**Supported by** the Development Foundation of Region Skåne, No. REGSKANE-818781 and No. 2018-Projekt0024; and the Foundation Skåne University Hospital, No. 2020-000028.

**Institutional review board statement:** The study was reviewed and approved by the Regional Ethics Review Board at Lund University (2009/209, Date of approval 28/04/2009; 2012/527, Date of approval 16/10/2012 and 2018/132, Date of approval 02/08/2018) and the Swedish Biobank Act.

**Informed consent statement:** All study participants provided informed written consent prior to study enrollment.

**Conflict-of-interest statement:** There are no conflicts of interest to report.

**Data sharing statement:** No additional data is available.

**STROBE statement:** The authors have read the STROBE Statement—checklist of items, and the manuscript was prepared and revised according to the STROBE Statement—checklist of items.

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**S-Editor:** Chen YL
**L-Editor:** A
**P-Editor:** Chen YL
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