Validation of New Cine Magnetic Resonance Imaging Parameters For The Differential Diagnosis of Chronic Idiopathic Intestinal Pseudo-Obstruction

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Abstract

Chronic idiopathic pseudo-obstruction (CIPO) is a severe and refractory intestinal motility disorder whose diagnosis currently relies on subjective imaging assessments. Cine magnetic resonance imaging (MRI) may potentially improve the quantitative analysis of gastrointestinal motility; however, suitable CIPO detection parameters should be determined.

Cine MRI was performed in seven patients with CIPO and 11 healthy controls. The logarithm of the Mahalanobis distance ($x_1$) and luminal distance variation per time ($x_2$) were used as the original parameters to determine CIPO diagnostic thresholds. Furthermore, the correlation between cine MRI findings and CIPO severity was investigated.

Threshold values of $\alpha=1.10$ and $\beta=0.15$ for $x_1$ and $x_2$, respectively, produced a CIPO diagnosis sensitivity of 1.00 (7/7) and specificity of 0.82 (9/11). The resulting error was 0.11 (2/18). The two parameters were correlated (Pearson’s correlation coefficient: -0.52). The intestinal tracts of patients with severe CIPO requiring home parenteral nutrition belonged to the category defined by $x_1 \geq 1.10$ and $x_2 \leq 0.15$.

Cine MRI is effective for the quantitative evaluation of small intestinal motility and CIPO diagnosis when using the abovementioned parameters and can be useful for treatment decision-making. However, these parameters have a wide distribution in healthy volunteers; this may complicate the detection of other disorders.

Introduction

Chronic intestinal pseudo-obstruction (CIPO) is a rare refractory disorder of the intestine wherein intestinal motility is severely impaired. It causes patients to experience abdominal pain, distention, vomiting, and weight loss despite the absence of organic disorders in the intestinal lumen. Recurrent paralytic ileus is a complication that causes social discomfort and necessitates repeated hospitalization for intestinal tract decompression or (at least) bowel rest. Furthermore, intestinal strangulation requires surgical resection of the necrotic segment, as that is the only strategy that can save the patient's life. However, patients who require resection of the intestine risk developing adhesive ileus or short bowel syndrome when extensive resection of the small intestine is performed. If their symptoms become more severe owing to intestinal motility failure, or malabsorption occurs owing to short bowel syndrome, home parenteral nutrition (HPN) becomes necessary to supplement nourishment.

Histological examination of the impaired intestine is a common approach to investigate CIPO of unknown etiology. Based on previously reported patients, CIPO is considered a collection of several pathologies that include neuropathy, abnormalities of intestinal cells of Cajal, and myopathies. However, in most cases, histological investigation does not yield a diagnosis, suggesting that CIPO has molecularly heterogeneous underlying causes. Furthermore, researchers are able to obtain specimens only from patients with advanced disease given that obtaining samples for histology will not influence
treatment decision-making, and such procedures may worsen the underlying dysmotility (as mentioned above).

Histological investigation can only be performed for critical patients and the subsequent clinical course does not necessarily improve (and can even worsens in some cases); therefore, radiological imaging is preferable for diagnosing CIPO. In typical cases, non-obstructive ileus findings are observed on radiography and computed tomography. However, these modalities carry a risk of radiation exposure; moreover, radiography is not sufficiently sensitive or specific to detect ileus, particularly in its early stages when symptoms are relatively light and/or the duration since disease onset has been relatively short. Therefore, other medical methodologies for the quantitative analysis of intestinal motility are recommended for CIPO diagnosis. Several modalities have been reported to be suitable for this purpose, although none are considered a gold standard owing to varying advantages and disadvantages. For example, intestinal manometry can analyze the peristaltic wave at a certain point; however, the limited length of the catheter makes it impossible to analyze the distal parts of the intestine. Small bowel transit time on capsule endoscopy has intra-and inter-individual discrepancies, and capsule endoscopy carries the risk of retention and is contraindicated in patients with CIPO. On the other hand, recent advances in magnetic resonance imaging (MRI) has allowed for the development of cine MRI, a noninvasive radiation-free tool to assess organ function and diagnose certain diseases (particularly cardiovascular conditions). As such, cine MRI can be applicable to gastrointestinal disorders including CIPO; however, parameters for cine MRI readings have not been validated to date, and remain entirely subjective. Moreover, the inter- and intra-evaluator discrepancy remains a hinderance for objective diagnosis.

Therefore, the aim of this study was to establish new, validated parameters for cine MRI that would assist in the diagnosis of CIPO.

Results

Seven patients with CIPO including three men (42.9%) and four women (57.1%) whose age was 52 (38.5–54.5) years underwent cine MRI; their characteristics are shown in Table 1. The disease duration was 40.1 (36.6–118.8) months, and all the patients were underweight (body mass index = 17.3 [16.2–18.2] kg/m²) and had abdominal distention. Cine MRI examination was also performed on 11 healthy volunteers comprising six men (54.5%) and five women (45.5%) with an age of 35 (29.0–41.5) years.
Table 1
Clinical course and cine magnetic resonance imaging findings in patients with chronic intestinal obstruction

| Patient No. | Age (y) | Sex | BMI (kg/m²) | CIPO duration (months) | Symptoms | Cine MRI↑ | Treatment                      |
|-------------|---------|-----|-------------|------------------------|----------|----------|-----------------------------|
| 1           | 37      | M   | 18.2        | 180.3                  | D, V     | 0.24     | 1.54 Medication              |
| 2           | 52      | M   | 19.4        | 35.7                   | D, V, P  | 0.10     | 2.68 Medication, Surgical bypass, HPN |
| 3           | 61      | M   | 17.1        | 6.5                    | D, V     | 0.16     | 1.12 Medication              |
| 4           | 55      | F   | 18.2        | 81.0                   | D, V     | 0.04     | 3.02 Medication, surgical bypass, HPN |
| 5           | 35      | F   | 11.1        | 37.6                   | D, P, C  | 0.10     | 3.24 Surgical bypass        |
| 6           | 54      | F   | 15          | 40.1                   | D, P     | 0.12     | 2.72 Surgical bypass        |
| 7           | 40      | F   | 17.3        | 156.6                  | D, C     | 0.04     | 4.15 Medication              |

Abbreviations: BMI, body mass index; CIPO, chronic intestinal obstruction; HPN, home parenteral nutrition; MRI, magnetic resonance imaging. ↑ D: abdominal distention, V: vomiting, P: Abdominal pain, C: constipation. ↑ Jt: the variation of luminal diameters, Dt: Mahalanobis distance.

Threshold value for separating patients with CIPO from healthy volunteers

The logarithm of the Mahalanobis distance as well as the distance variation per time were calculated for both patients with CIPO and healthy volunteers, and the data are shown graphically in Fig. 1.

As the values of α and β are critical for classifying individuals, we determined the optimal α and β pairing (i.e., that which minimized error) from among candidate pairs (Fig. 2). To determine the most robust pair that optimally takes into account the variability of samples, we used the leave-one-out method\(^\text{14}\), which generates artificial variability among the subjects with the use of resampling\(^\text{15}\). There were 18 samples available; according to the leave-one-out method, one sample was excluded, and the remaining 17 were used as training samples. Note that the test sample was not used in optimization. Next, an α and β pair was selected from candidate pairs. Thereafter, we determined whether each intestinal tract of a training sample had severe impairment of motility using the selected α and β pair. If \(x_1 \geq \alpha\) and \(x_2 \leq \beta\), the intestinal tract was classified as impaired one, and the number of misclassified intestinal tracts was counted from among intestinal tracts of 17 training samples to estimate the error. This was conducted for each candidate pair. Ultimately, we selected the pair that produced the lowest error among the intestinal tracts of 17 training samples. The above operation was repeated until each of the 18 samples had been excluded once. Finally, we determined the optimal pair that was most frequently selected from among the 18 attempts: \(\alpha = 1.10\) and \(\beta = 0.15\), which were considered to be robust for the variability of samples. The parameters with these values produced a sensitivity of 1.00 (7/7) and specificity of 0.82.
(9/11) (Fig. 3a); the error was 0.11 (2/18). On the other hand, a specificity of 1.00 (11/11) and sensitivity of 0.86 (6/7) were obtained when values of $\alpha = 2.00$ and $\beta = 0.15$ were used (Fig. 3b).

**Severity of CIPO and cine MRI findings**

The severity of CIPO were determined based on the treatment required: mild CIPO cases were controlled by medication only; moderate cases required surgical intervention; and severe cases required permanent HPN (Table 1). As a result, the intestinal tracts of severe cases (patients 2 and 4) belonged to the category defined by $x_1 \geq 1.10$ and $x_2 \leq 0.15$ in the two-dimensional feature space. Furthermore, in patients with CIPO, Pearson's analysis of the logarithm of the Mahalanobis distance and the distance variation per time revealed a mild inverse correlation of -0.52 (Fig. 4). When analyzing all subjects including patients and volunteers ($n = 18$) as well as volunteers alone ($n = 11$), no correlation was found (the Pearson's correlation coefficients were -0.15 and 0.01, respectively).

**Discussion**

We successfully established cine MRI parameters that could differentiate patients with CIPO from healthy volunteers. Our analysis also revealed that all patients with CIPO had impaired small intestinal motility and that CIPO severity correlated with the luminal diameter.

We used the original parameters 'logarithm of the Mahalanobis distance' and 'distance variation per time' in this study because other previously reported parameters, such as intestinal 'amplitude' and 'contraction cycle' were difficult to determine during our previous efforts. This was presumably because of the various types of small intestinal contraction that occur concurrently, including peristalsis, segmentation, and phase III migrating motor complex. This makes it difficult to distinguish the type of each contraction definitively; moreover, the 'amplitude' and 'contraction cycle' are not considered single or regular. Although the feature vectors of healthy volunteers had widely distributed values in this study, the threshold value of 1.10 for the parameter $x_1$ was found to be satisfactory for CIPO diagnosis. Furthermore, the luminal diameters were selected for analysis from all parts of the small intestine because we could not always measure these diameters in specific areas (i.e., the proximal jejunum, middle intestine, and distal ileum) given that the analyses were performed on sliced views, wherein certain parts of the small intestine were often in a blind spot and thus invisible. Another drawback was that intestines sliced horizontally or perpendicularly were not always captured given the small intestine's tortuousness in the three-dimensional abdominal cavity.

A key finding in this study was that all patients with CIPO had impaired small intestinal motility. In some cases, however, lower colonic propulsion may have been impaired, which further inhibited small bowel propagation. Furthermore, the severity of CIPO was reflected in the cine MRI findings. Therefore, our calculated parameters may potentially be useful for clinical decision-making. In contrast, the parameters calculated for healthy volunteers had a wide range, which was a notable observation. Further studies of cine MRI are required to reliably differentiate CIPO from other disorders, such as irritable bowel syndrome,
although determining the definitive parameters are likely to be challenging. At a minimum, longer cine MRI observation is necessary, and the extent of small intestinal contraction (peristalsis, segmentation, and migrating motor complex), should be analyzed.

Based on our results, using cine MRI together with computer-assisted or artificial intelligence-driven diagnosis of gastrointestinal motility may have potential to accurately identify patients with CIPO, although several drawbacks should be addressed (such as the limitations of sliced-based analysis). Our study focused on small intestinal motility, and colonic motility should be further explored in additional investigations. The small intestine usually functions to move and clear its luminal contents before more food is ingested. In contrast, the colon is idle up to 90% of the time. As such, cine MRI may be unsuitable for the evaluation of colonic movement given its glacial pace.

Our study had several limitations that should be acknowledged. First, only a small number of patients with CIPO were analyzed because of the rarity of this disorder. Second, we classified the severity of CIPO based on the required treatment; however, severity is determined using a complicated scale in the real-world clinical settings. For example, patient 7 had severe symptoms that were controlled with many medications and well-balanced meals.

In conclusion, our new cine MRI parameters have the potential to assist in the differential diagnosis of patients suspected of having CIPO, including the severity thereof. The wide ranges of distributions for feature vectors of healthy volunteers may reflect the complicated function of the small intestine, which should be elucidated using additional cine MRI studies in the future.

**Methods**

**Cine MRI protocol**

After 8 hours of fasting, the participants underwent MRI in the supine position after orally ingesting 1,000 mL of water to fill the small intestine. Patients who were unable to ingest the entire amount because of severe abdominal symptoms were encouraged to drink as much as possible to prevent worsening of symptoms.

Imaging was performed using a 1.5-T MRI unit (SIGNA™ Creator, GE Healthcare, US). Before performing cine MRI, coronal images of the entire abdomen were obtained to identify an appropriate imaging plane. At least three optimal images were selected to cover the maximum length of the small bowel loops during each examination. A steady-state free precession sequence (FIESTA sequence: repetition time = 3.6 msec, echo time = 1.6 msec, flip angle = 70°, slice thickness = 10 mm, matrix = 196×230, field of view = 400×400 mm, number of excitations = 1.0, bandwidth = 100 kHz, and fat suppression = special [inversion time = 200 msec]) was used for imaging, which allowed for continuous scanning without intervals for each image. Sequential scanning was performed every 0.571 seconds with 35 images.
All procedures involving human participants were performed in accordance with the principles of the 1964 Declaration of Helsinki. Niigata university hospital review board approval was obtained (approval no: 2018-0403) and written informed consent was obtained from all patients.

**Data analysis**

Cine MRI imaging analysis was performed using the Centricity DICOM Viewer software v. 2.2 (GE Medical Systems). An unsharp masking filter was used to improve the quality of all cine MRI scans degraded by noise (Fig. 5). At the corresponding location of the small intestinal tract, a line perpendicular to the long axis was drawn; this task was repeated for all sequential cine MRIs (Fig. 6). The luminal diameters of each intestinal tract were measured on 35 cine MRI scans; using these diameters, the criterion $J_t$, which evaluates the motility of an intestinal tract, was defined by

$$J_t = \frac{L_t}{L_{t-1} + L_t}$$

where $L_t$ is the luminal diameter at time $t$ and $L_{t-1}$ is that at time $t-1$. The $J$ values ranged from 0 to 1. As an intestinal tract expands, the $J$ value falls in the range of $0.5 \leq J \leq 1.0$. Conversely, as the intestinal tract contracts, the value falls in the range of $0 \leq J \leq 0.5$. Moreover, $L_t = L_{t-1}$ leads to a $J$ value of 0.5. Therefore, this value indicates that the intestinal tract has no motility (Fig. 7). Thus, by using the relative index calculated from sequentially adjacent images, we made the most of the motion characteristic of the cine MRI.

Next, we defined two 34-dimensional vectors; one was a motility vector whose components were the values of $J$, while the other was a non-motility vector whose components were all 0.5. The magnitude of the intestinal tract motility was determined by the distance between the motility and non-motility vectors; as motility increases, the distance lengthens. Moreover, the mean of 35 luminal diameters was estimated. We measured the Mahalanobis distance $D(x)$ to obtain the invariance to the scaling of the coordinate axes. Given that luminal diameters of patients with CIPO have diagnostic significance only when they are compared with those of healthy controls, the Mahalanobis distance was thought to be the optimal method since it normalized the luminal diameter of patients with CIPO using that of healthy volunteers. Furthermore, the other advantage of using the Mahalanobis distance is that it is not affected by the scan resolutions and can be applied to scans from different types of MRI equipment, which is different from the Euclidean distance.

The Mahalanobis distance $D(x)$ between the mean $x$ of 35 luminal diameters of the intestinal tract of a patient and the volunteer distribution was defined as follows:

$$D(x) = \sqrt{x^T S^{-1} x}$$
\[ D(x) = \frac{(x - \mu)^2}{\sigma^2} \]

where \( \mu \) and \( \sigma^2 \) are the sample mean and sample variance of the volunteer distribution, respectively. The sample mean and sample variance were estimated using the means for each of intestinal tracts of healthy volunteers. Then, we calculated the logarithm of the Mahalanobis distance \( D(x) \); thus, any intestinal tract was represented as a two-dimensional vector whose components \( x_1 \) and \( x_2 \) were the logarithm of the Mahalanobis distance and distance variation per time, respectively. In pattern recognition, the parameters \( x_1 \) and \( x_2 \) were called features. Each intestinal tract was represented as a feature vector in a two-dimensional feature space.

Once the intestinal tract of the patient was selected, it was deemed to have impaired motility if \( x_1 \geq \alpha \) and \( x_2 \leq \beta \), where \( \alpha \) and \( \beta \) are the threshold values (mentioned below). An intestinal tract that had no impairment in motility satisfied the condition \( x_1 < \alpha \) or \( x_2 > \beta \). The patient was deemed to have CIPO if any parts of the intestinal tract with impaired motility was detected.

Statistics

Continuous variables, such as patient age and body mass index are presented as medians and interquartile ranges, while categorical variables such as sex are expressed as numbers and percentages. All statistical analyses were performed using EZR (Saitama Medical Center, Jichi Medical University, Saitama, Japan), which is a graphical user interface for R (The R Foundation for Statistical Computing, Vienna, Austria). More precisely, it is a modified version of R commander designed to add statistical functions frequently used in biostatistics\(^\text{19}\).

Declarations

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Author contributions:

H.S. wrote the main manuscript text and performed volunteers’ recruitment. H.O. and Y.H. analyzed data. K.T., Y.K., Y.K., K.T., and J.Y. recruited cases. S.T. supervised the study. All authors reviewed the manuscript.

Additional Information:

Competing interests

The authors declare no competing interests.
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Data availability

All data used in the study are already provided in the tables, figures, and online supplementary materials.

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**Figures**
**Figure 1**

Scatter diagram of logarithm of the Mahalanobis distance (horizontal axis) and distance variation per time (vertical axis) in patients with chronic intestinal pseudo-obstruction and healthy volunteers.
Figure 2

The scheme of optimization of threshold values.
Figure 3

Scatter diagram for determining sensitivity and specificity in two-dimensional feature space. (a) $\alpha=1.10$ and $\beta=0.15$ can be used to obtain the best sensitivity for chronic intestinal pseudo-obstruction (CIPO) diagnosis. (b) $\alpha=2.00$ and $\beta=0.15$ can be used to obtain the maximum specificity for CIPO diagnosis.
Figure 4

In patients with chronic intestinal obstruction (CIPO), the logarithm of the Mahalanobis distance and distance variation per time were inversely correlated.
Figure 5

The edge of the small intestine can be enhanced automatically using software. Shown are (a) before enhancement and (b) afterward.

Figure 6

The luminal diameters of the small intestine (yellow line) are calculated using the software. This task was repeated for all sequential cine MRIs. The typical findings are shown as a (1st of the series), b (17th of the series), and c (35th of the series).
Figure 7

Time variation of parameter J in a healthy volunteer (a) and a case of chronic intestinal pseudo-obstruction (b).

Supplementary Files

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