Early-Life Malnutrition Causes Gastrointestinal Dysmotility That Is Sexually Dimorphic

Running title: Malnutrition-Induced GI Dysmotility

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ABSTRACT

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**Background:** Slow gastrointestinal (GI) transit occurs in moderate-to-severe malnutrition. Mechanisms underlying malnutrition-associated dysmotility remain unknown, partially due to lack of animal models. This study sought to characterize GI dysmotility in mouse models of malnutrition.

**Methods:** Neonatal mice were malnourished by timed maternal separation. Alternatively, low-protein, low-fat diet was administered to dams, with malnourished neonates tested at two weeks or weaned to the same chow and tested as young adults. We determined total GI transit time by carmine red gavage, colonic motility by rectal bead latency, and both gastric emptying and small bowel motility with fluorescein isothiocyanate-conjugated dextran. We assessed histology with light microscopy, *ex vivo* contractility and permeability with force-transduction and Ussing chamber studies, and gut microbiota composition by 16S rDNA sequencing.

**Key Results:** Both models of neonatal malnutrition and young adult malnourished males but not females exhibited moderate growth faltering, stunting, and grossly abnormal stomachs. Progression of fluorescent dye was impaired in both neonatal models of malnutrition, whereas gastric emptying was delayed only in maternally separated pups and malnourished young adult females. Malnourished young adult males but not females had atrophic gastrointestinal mucosa, exaggerated contractile responses, and increased gut barrier permeability. These sex-specific abnormalities were associated with altered gut microbial communities.

**Conclusions & Inferences:** Multiple models of early-life malnutrition exhibit delayed upper GI transit. Malnutrition affects young adult males more profoundly than females. These models will facilitate future studies to identify mechanisms underlying malnutrition-induced pathophysiology and sex-specific regulatory effects.

**KEYWORDS:** Malnutrition, gastroparesis, gastrointestinal motility, permeability, microbiome, sex differences

1. INTRODUCTION

Malnutrition accounts for 1.7 million child deaths per year.\(^1\) Over 50 million children under age five years (8%) show signs of wasting and 151 million (22%) are stunted.\(^2\) Malnutrition creates a...
“vicious cycle”\textsuperscript{3,4} of gastrointestinal (GI) pathophysiology that further erode health. Among the least understood comorbidities is GI dysmotility, which is observed in children with severe acute malnutrition,\textsuperscript{5-7} small-for-gestational-age neonates,\textsuperscript{8,9} and young adults with anorexia nervosa.\textsuperscript{10-14} Slow GI motility might impair weight gain, given that previously non-malnourished patients with gastroparesis\textsuperscript{15,16} and chronic intestinal pseudo-obstruction\textsuperscript{17,18} lose weight. Mechanisms underlying these associations are poorly understood, in part due to lack of animal models.

This study aimed to characterize GI dysmotility in early-life malnutrition. Two neonatal models – timed maternal separation (TmSep)\textsuperscript{19} and introduction to dams of a low-protein, low-fat diet (LPLFD) mimicking the macronutrient composition of diets typical in rural Northeast Brazil\textsuperscript{20} – as well as a young adult LPLFD model\textsuperscript{21} were employed to ensure that dysmotility was not specific to the means by which malnutrition was modeled. We assessed total GI transit time with carmine red gavage, colonic motility by rectal bead latency, and gastric emptying and small bowel transit by quantifying the distribution of fluorescein isothiocyanate (FITC)-conjugated dextran after gavage. Although many central, peripheral, and other endogenous and exogenous factors affect GI motility,\textsuperscript{22-25} we focused our initial investigation of possible mechanisms underlying malnutrition-induced dysmotility by determining whether muscle tone and contractility, intestinal barrier permeability, and fecal microbial communities are affected in our mouse models. We assessed tone and contractility with \textit{ex vivo} force-transduction experiments, permeability with Ussing chamber studies, and gut microbiota alterations with 16S rDNA sequencing. Each malnutrition protocol affected the gross appearance of the stomach, decreased the length of the small intestine, impaired gastric emptying or small bowel transit, and altered the gut microbiota. Findings in malnourished young adult mice, including abnormal contractile responses to cholinergic stimulation and increased gut barrier permeability, were sex-specific. These results suggest novel interactions between nutritional status and sex that influence GI motility.

\section*{2. MATERIALS AND METHODS}

\subsection*{2.1 Modeling malnutrition in mice}

The Institutional Animal Care and Use Committee at Baylor College of Medicine approved all aspects of this study. C57BL/6J mice were obtained from the Center for Comparative Medicine at Baylor College of Medicine and maintained on control chow containing 20\% kcal protein, 15\%
kcal fat, and 65% kcal carbohydrate (#D09051102, Research Diets, New Brunswick, NJ). Malnutrition was induced as previously described\textsuperscript{19,26,27} by TmSep of the same half of litters for 4 hours on day of life (DOL) 5, 8 hours on DOL 6, and 12 hours DOL 7-14, with overnight nursing \textit{ad libitum}. In accordance with our previously published studies, separated pups from multiple litters were combined within one large nest to obviate the need for external warming prior to random redistribution to dams for overnight feeding. Alternatively, LPLFD chow containing 7% kcal protein, 5% kcal fat, and 88% kcal carbohydrate (#D09081701B, Research Diets), specially formulated to be isocaloric to the control chow, was introduced to dams on DOL 8. Previous studies have modeled malnutrition by introducing this LPLFD to dams on DOL 3\textsuperscript{21} or DOL 10.\textsuperscript{28} We introduced LPLFD on DOL 8 so that TmSep pups and LPLFD pups were equally underweight and stunted when they underwent motility testing at 2 weeks. Alternatively, on DOL 21, mice were weaned to the same diets as their dams (either control diet or LPLFD) and were maintained on these diets for another 6 weeks (Fig. 1A).

Daily chow consumption by cage was calculated by determining the weight of chow consumed over a 24 hr period divided by the summed daily body weight for mice in that cage. Testing was performed for neonatal mice at 2 weeks of life, one day after ending TmSep. Eight-week old young adult mice were tested after an overnight fast of 15 hours. Each experiment included mice (with \( n \) ranging from 4 per group for Ussing chamber studies to 10 per group for microbiota studies) originating from two or more randomized litters. Each experiment was repeated twice on different occasions using mice derived from distinct mating groups. Female estrous cycles were not tracked. Although our data did not reveal litter-specific effects, previously we have reported subtle changes in motility that depend on the time of day;\textsuperscript{29} thus, all motility testing was performed between the hours of 09:00-13:00. Nose-to-rump body length was measured in the prone position, and intestinal length measurements were made by gently stretching intact, freshly harvested tissue.

\subsection*{2.2 Assessing GI motility in vivo}

Upper GI tract motility was assessed as previously reported\textsuperscript{29} by determining the distribution of 100 \( \mu \)L of 10 mg mL\(^{-1} \) FITC conjugated to 70 kDa dextran (Sigma, St. Louis, MO) 30 minutes after gastric gavage, with no food or water access post-gavage. Following euthanization for 2 minutes in \( \text{CO}_2 \), 14 GI luminal samples (stomach, 10 small intestinal segments of equal length, cecum, and proximal and distal colon) were collected and flushed with 3 mL phosphate-buffered...
saline (PBS). After centrifugation, fluorescence from each flushing was quantified (excitation 485 nm, emission 525 nm), and mean geometric center (MGC) and % gastric emptying were computed.\textsuperscript{29} The assay for neonatal mice was modified as previously reported. Briefly, nursing was permitted up until gavage with 50 µL FITC-dextran, after which pups were isolated from dams to prevent further nursing.\textsuperscript{29} Thirty minutes later, following cervical dislocation, GI segments were harvested immediately from pups. In young adults, total GI transit time was recorded as the number of minutes between gastric gavage of 100 µL carmine red (6% weight volume\textsuperscript{-1} in 0.5% methylcellulose; Sigma) and first appearance of red dye in the stool.\textsuperscript{30} Rectal bead latency\textsuperscript{31} was recorded as the number of minutes needed to expel a 2 mm-diameter glass bead (Sigma) inserted to a depth of 2.5 cm under light isoflurane anesthetization.

2.3 Evaluating GI histology

For histology, a separate set of mice that had not undergone motility testing were used. All of these animals, including TmSep pups, had fed \textit{ad libitum} overnight and until sample harvest. Tissues were fixed in formalin, embedded in paraffin, cut, and stained with hematoxylin and eosin. Images were captured on an Eclipse 90i Ni-E microscope (Nikon, Tokyo). Measurements were made by an observer blinded to the treatment group using NIS-Elements software (Nikon) for villus height (small bowel) or mucosa thickness (stomach and colon), crypt depth, and muscularis thickness using ImageJ. For each histologic parameter, 5-20 measurements per GI region were averaged for each animal.

2.4 Quantifying intestinal contractility

Intestinal contractile activity was measured in the longitudinal axis as previously described\textsuperscript{32,33} with some minor modifications. Briefly, 1 cm segments of duodenum, jejunum, proximal colon, and distal colon were harvested and immediately mounted in organ baths filled with 25 mL Krebs solution (in mM: 120.9 NaCl, 5.9 KCl, 2.5 CaCl\textsubscript{2}, 14.4 NaHCO\textsubscript{3}, 1.2 NaH\textsubscript{2}PO\textsubscript{4}, 1.2 MgCl\textsubscript{2}, 11.5 glucose) and infused continuously with 95% O\textsubscript{2}/5% CO\textsubscript{2} gas. Isometric force was monitored by an external force displacement transducer connected to a PowerLab recorder (ADInstruments, Colorado Springs, CO). Tension of 0.4 g or 0.5 g was applied to small bowel or colonic segments, respectively. After 30 min of equilibration, baseline contractile activity was recorded for 5 min. Responses to increasing doses of cholinergic stimulation were determined, first by adding 1 µM carbachol (Sigma) to each organ bath with activity recorded for 5 min, then
by increasing the dose to 10 µM with activity recorded for another 5 min. Subsequently, the length of each intestinal segment was measured, and tissue was removed, dried, and weighed. Contractile activity was calculated as the integral from minimum, and the amplitude and frequency of contractions were determined by normalizing readings to tissue cross-sectional areas using the measured weight and length and an assumed tissue density of 1.05 g cm\(^{-3}\) using LabChart software (ADInstruments).

### 2.5 Measuring intestinal permeability

Intestinal permeability was quantified in a P2300 Ussing chamber system (Physiologic Instruments, San Diego, CA). Briefly, full-thickness sections of duodenum, proximal colon, and distal colon segments were harvested and immediately flushed with KBR buffer (in mM: 115 NaCl, 1.2 CaCl\(_2\), 25 NaHCO\(_3\), 2.4 K\(_2\)HPO\(_4\), 0.4 KH\(_2\)PO\(_4\), 1.2 MgCl\(_2\), 10 glucose). Each segment was opened along its mesenteric border, rinsed, and mounted on a slide with an exposure area of .30 cm\(^2\). The mucosal and serosal chambers each were filled with 5 mL KBR buffer maintained at 37 °C and infused with 95% O\(_2\)/5% CO\(_2\) gas. Permeability was assessed by flux of 4 kDa FITC-dextran (Sigma) and transepithelial resistance (TER). After an equilibration period of 10 min, FITC-dextran was added to the mucosal chamber to a final concentration of 2.2 mg mL\(^{-1}\) and sampled from the serosal side after 0, 30, 60, 90, and 120 min. The fluorescent intensity of each sample was measured by a Cytation5 Imaging Reader (BioTek Instruments, Winooski, VT). TER was calculated from the spontaneous potential difference and short-circuit current via Ohm's law. Aggregate data for FITC-dextran flux and TER were normalized to the average measurements over all time points.

### 2.6 Evaluating gut microbial communities

Stool was obtained from individual mice that had not undergone motility testing, and 16S rDNA sequencing was performed in the Baylor College of Medicine Alkek Center for Metagenomics and Microbiome Research as described in detail previously.\(^{27}\) Briefly, DNA was extracted using MoBIO PowerSoil DNA Isolation Kit (Qiagen, Germantown, MD) according to the manufacturer’s protocol. The 16S V4 region was amplified by polymerase chain reaction and sequenced on the MiSeq platform (Illumina, San Diego, CA) using the 2 x 250-bp protocol. A custom pipeline, using the QIIME software package,\(^{34}\) was used for analysis. Reads were trimmed, quality-filtered, clustered into operational taxonomic units (OTUs) with a 97% similarity
cutoff using the UPARSE algorithm,\(^35\) and aligned using the SILVA database.\(^36\) The Fisher index was used to compare taxonomic diversity between samples.

### 2.7 Statistical analysis

All data were confirmed to be normally distributed by the Shapiro-Wilk test, and are presented as mean ± standard deviation (SD). Data from control, TmSep, and LPLFD pups were analyzed by one-way ANOVA and, when the global test was significant \((P < .05)\), by Tukey’s multiple comparisons test. Only adjusted \(P\) values are presented for the three groups of mouse pups. Alternatively, male or female LPLFD young adults were compared to their respective controls by two-tailed \(t\) test. Associations between gastric emptying and MGC were identified using two-tailed Pearson correlation tests. All data are representative of at least two identical experiments performed on different sets of animals on different dates.

### 3. RESULTS

#### 3.1 Malnutrition causes sex-specific changes in weight and length

Impaired weight gain was observed immediately after initiating TmSep (DOL 5) or LPLFD chow (DOL 8, Fig. 1B). At 2 weeks of life, both TmSep pups and LPLFD pups weighed < 70% of control pups (adjusted \(P < .001\)). However, young adult males and females were affected differently by malnutrition (Fig. 1C); LPLFD males weighed just 67% of control males \((P < \ldots .0001\)), whereas LPLFD females weighed 89% of control females \((P < .01\)). Growth impairment was not due to decreased food intake; compared to controls, malnourished young adult males and females consistently consumed a greater amount of chow relative to body weight (Fig. 1D). Similarly, nose-to-rump length, small bowel length, and total GI tract length were decreased in TmSep pups, LPLFD pups, and LPLFD males (Fig. 1E, adjusted \(P < .01\) for each comparison) but not LPLFD females. Colon length was not altered by malnutrition in any model, and consistent with our previous findings,\(^29\) sex-specific differences among neonatal mice were not identified.

#### 3.2 Malnutrition causes GI dysmotility

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Total GI transit time and colonic motility could not be assessed reproducibly in neonates, which rely on maternal stimulation to defecate. In young adults, neither whole GI transit (Fig. 2A) nor rectal bead latency (Fig. 2B) times differed significantly between malnourished and control mice, although a trend toward prolonged bead expulsion was observed in LPLFD vs control females ($P = .09$). To more precisely assess upper GI tract motility, we determined the distribution of FITC-dextran 30 minutes after gavage. Congruent with our previous findings,$^{29}$ our modified neonatal mouse protocol yielded markedly reduced variance in comparison to the standard adult FITC-dextran protocol (Fig. 3A-B and Supporting Fig. S1). Relative to control pups, MGC was decreased in TmSep pups (-1.8 segments, adjusted $P < .001$) and in LPLFD pups (-.9 segment, adjusted $P < .05$; Fig. 3C). No significant changes in MGC were observed in LPLFD young adults. Mean gastric emptying was decreased from 97% to 87% in TmSep pups (adjusted $P < .05$) and from 88% to 75% in LPLFD young adult females ($P < .001$, Fig. 3D). Correlational analysis revealed an association between gastric emptying and MGC only in TmSep pups ($P < .05$), suggesting that decreased MGC in LPLFD models cannot be attributed exclusively to delayed gastric emptying. These data indicate that multiple models of early-life malnutrition are associated with impaired upper GI motility in a sexually dimorphic manner.

3.3 Malnutrition alters gross stomach appearance and GI mucosal structure

To gain initial insights into mechanisms by which malnutrition-associated dysmotility occurs, we examined the gross and microscopic appearance of the malnourished GI tract. Stomachs from TmSep pups were grossly dilated, whereas stomachs from LPLFD pups and LPLFD young adult males were shrunken and nodular (Fig. 4A). Stomachs from LPLFD young adult females appeared less abnormal. Examination by light microscopy (Fig. 4B-C) revealed thinning of the gastric mucosa in two of the malnutrition models. Mucosa was atrophic in stomachs of both TmSep pups (57.2 vs 108.5 µm, adjusted $P < .01$) and LPLFD males (166.5 vs 220.1 µm, $P < .05$). Although gastric muscularis thickness was not significantly altered in any model, we observed a trend toward smooth muscle atrophy in TmSep pups (17.6 vs 27.6 µm, adjusted $P = .07$). Likewise, a broad histologic survey of the duodenum, jejunum, ileum, and proximal and distal colon (Supporting Fig. S2) revealed that malnutrition spared muscularis thickness while impacting mucosal histology. Ileal villi were blunted in LPLFD pups (79% vs control pups, adjusted $P < .05$) and LPLFD males (85% vs control males, $P < .01$) but not LPLFD females (Fig. 5A). Similarly, crypt depth was diminished in jejunum, ileum, and proximal colon in multiple
malnutrition models (Fig. 5B). A preliminary examination using whole mount immunostaining did not reveal obvious changes in the density or relative proportions of glia or neurons expressing Hu, nNOS, or calbindin in the myenteric plexus of the duodenum or colon (Supporting Figs. S3-S4). A more detailed quantitative analysis of the malnourished enteric nervous system, which is outside the scope of the present study, needs to be conducted. Taken together, these findings indicate that malnutrition grossly alters gastric and intestinal mucosal morphology, but smooth muscle layer thickness and enteric nervous system structures were not affected to the same extent.

3.4 Malnutrition Increases spontaneous and cholinergic-mediated contractile activity

To determine whether additional functional changes are present in the malnourished intestine, we measured *ex vivo* contractile activity at baseline and in response to increasing concentrations of cholinergic stimulation (Fig. 6A). Unexpectedly, duodenal segments from LPLFD young adult males demonstrated increased contractility compared to controls. Spontaneous contractile activity was increased 1.8-fold (*P* < .01), and contractile responses to 1 µM and 10 µM carbachol were each increased 1.7-fold (*P* < .05) in duodenum from LPLFD males but not females compared to controls (Fig. 6B). In contrast, malnutrition did not induce changes in contraction amplitude (Fig. 6C) or frequency (Fig. 6D). This result was specific to the duodenum; increased contractility was not observed in the malnourished male jejunum, proximal colon, or distal colon (Supporting Fig. S5-S7). Rather, in the distal colon, malnourished females exhibited decreased contractile responses to cholinergic stimulation (Supporting Fig. S7). These results indicate that cholinergic contractile responses are intact in malnutrition, with sex-specific differences including exaggerated contractile activity in the duodenum of males and decreased contractile activity in the distal colon of females.

3.5 Intestinal permeability is increased in malnourished males

Malnutrition impairs gut barrier function in humans and animal models, and our histological survey revealed decreased thickness of the intestinal mucosa. Thus, we assessed barrier function by quantifying the translocation of FITC-dextran across full-thickness sections of duodenum, proximal colon, and distal colon mounted in Ussing chambers. Consistent with previous studies that fed LPLFD to weanling mice, we observed increased permeability in intestine harvested from LPLFD males. Translocation was increased across the duodenum 1.9-
fold ($P < .05$; Fig. 7A-B) and across the proximal colon 2.1-fold ($P < .05$; Figs 7A, 7C). Increased permeability occurred without a significant decrease in TER (Supporting Fig. S8). Although translocation appeared to increase across the distal colon in LPLFD males (Fig. 7D), this trend was not significant. Strikingly, gut permeability was not increased in LPLFD females. Taken together, these results demonstrate that malnutrition increases gut barrier permeability in the proximal intestine in a sex-specific manner.

3.6 Gut microbial communities are altered in malnutrition

Given that altered gut microbiota are associated with changes in GI motility and permeability,$^{23}$ we sought to determine whether fecal microbial community changes are present in our malnourished models with 16S rDNA sequencing. Consistent with our previous findings,$^{26,27}$ stool from TmSep pups contained fewer unique taxa (adjusted $P < .05$, Fig. 8A) and decreased microbial community diversity (adjusted $P < .05$, Fig. 8B) compared to stool from control pups. Conversely, stool from LPLFD males and females had a greater number of unique taxa and increased gut microbial community diversity relative to their respective controls ($P < .01$ for each comparison), which could reflect the increased carbohydrate content of the LPLFD chow. At the phylum level, a greater relative abundance of Bacteroidetes was observed in LPLFD males ($P < .01$, Fig. 8C) and a greater abundance of Firmicutes was observed in LPLFD females ($P < .05$, Fig. 8D). Strikingly, both neonatal malnutrition models exhibited increased abundance of Proteobacteria (Fig. 8E), a phylum which contains numerous proinflammatory taxa. Compared to control pups, the relative abundance of Proteobacteria was increased 5.6-fold in TmSep pups (adjusted $P < .0001$) and 4.1-fold in LPLFD pups (adjusted $P < .01$). This increase in Proteobacteria was driven by expansion of a single species, *Muribacter muris* (Fig. 8F), a rodent commensal bacterium in the family Pasteurellaceae.$^{38}$ These results indicate that malnutrition-induced changes in GI motility, morphology, contractility, and permeability are associated with altered gut microbial communities and are dependent on host sex.

4. DISCUSSION

The goals of this study were to recapitulate malnutrition-induced GI dysmotility in mouse models and to gain initial insights into potential underlying mechanisms by identifying associated histologic and physiologic changes. Our results demonstrate that multiple mouse models of
early-life malnutrition exhibit GI dysmotility, namely delayed gastric emptying and small bowel progression of gavaged FITC-dextran. Our data also demonstrate that malnutrition alters mucosal structure (to a greater extent than smooth muscle or enteric nervous system structure), increases the baseline tone and contractility of intestinal segments, increases barrier permeability, and alters the gut microbiota. Most strikingly, our data show that the effects of malnutrition on intestinal physiology in young adults are dependent on the sex of the animal.

Among the key findings of these studies is that malnutrition slows GI motility, mimicking features observed in small-for-gestational-age neonates,\textsuperscript{8,9} children with severe acute malnutrition,\textsuperscript{5-7} and young adults with anorexia nervosa.\textsuperscript{10-14} Although the influence of high-fat diet and obesity on GI motility is well documented,\textsuperscript{39-42} this study is the first to our knowledge to recapitulate undernutrition-associated dysmotility in the laboratory. We employed multiple models to ensure that our results are independent of the means by which malnutrition is induced. TmSep pups receive the same diet as control pups, with the amount of breast milk limited to what is consumed during nocturnal nursing. Given the wide variation in timing and duration of existing maternal separation protocols,\textsuperscript{43} the motility impairment observed in our TmSep pups might not be recapitulated in similar models. Recent studies of maternal separation in rodents reveal subtle alterations in both maternal caregiving\textsuperscript{44} and offspring stress response.\textsuperscript{45} Although we did not observe any obvious behavioral changes, we did not study maternal and offspring behaviors systematically. Thus, because we cannot rule out separation-induced stress as contributing to the dysmotility, we also modeled malnutrition through dietary manipulation without separation. In young adult mice, testing was performed following a prolonged overnight fast to minimize the possibility that luminal differences in macronutrient concentrations between LPLFD and control diet drives the dysmotility. We performed our investigations in mice because the two neonatal mouse models of malnutrition are well characterized and are equally underweight and stunted when testing is performed, and because of the greater number of genetic tools that will facilitate further mechanistic investigations in mice as opposed to rats.

The two techniques most often used to assess GI motility, carmine red gavage and rectal bead latency, did not reveal significant differences between malnourished and healthy young adult mice. Given the previously reported role of rodent sex steroid hormones on gastric emptying and GI transit,\textsuperscript{46} we might have increased our ability to detect subtle motility changes among young adult female mice had we controlled for estrous cycle fluctuations. Nonetheless, first appearance of red in a fecal pellet and expulsion time of a bead from the distal colon are crude
and incomplete evaluations of whole GI transit and large bowel motility, respectively. We used an adaptation of the FITC-dextran technique recently validated in neonatal mice\textsuperscript{29} to facilitate the detection of subtle changes in upper GI motility. These models highlight a novel role of host nutritional status in the regulation of GI motility and will serve as important tools to further explore mechanisms underlying these effects.

Smooth muscle atrophy was not observed in any of our malnourished models, except for the trend toward thin gastric muscularis in TmSep pups. Similarly, preliminary examination of stained myenteric plexus did not reveal any obvious neurochemical or architectural differences between our malnourished models and their controls. Given that sufficient smooth muscle mass and an intact enteric nervous system are required for normal propulsive transit, these results are consistent with our unexpected finding of increased contractile activity at baseline and following cholinergic stimulation in duodenal segments from LPLFD males. It is possible that increased cholinergic responses compensate for other prokinetic signals that are diminished. For example, we previously reported that plasma serotonin and luminal bile acids are depleted in TmSep pups.\textsuperscript{19} We cannot exclude the possibility that strong but poorly-coordinated contractions contribute to malnutrition-induced dysmotility. Because the smooth muscle and enteric ganglia appear to be intact, one might hypothesize that refeeding the malnourished mice would normalize the dysmotility as soon as the underlying metabolic, hormonal, or other signaling abnormalities are corrected. If true, this finding would provide hope that treating malnutrition-induced dysmotility might not require extensive smooth muscle mass accumulation or restructuring of the enteric nervous system. However, our broad initial characterization might have missed subtle changes in enteric nervous system structure; more detailed analyses of the various neurochemical subtypes of the enteric nervous system and their activity are required in future studies to adequately address the involvement of the enteric nervous system in malnutrition-associated dysmotility.

Our finding that malnutrition increases gut barrier permeability confirms a previous study that reported increased translocation of FITC across jejunal sections from malnourished mice using a similar model of malnutrition.\textsuperscript{28} Our results extend these findings to duodenal and proximal colon (but not distal colon) segments, and to young adult males but not females. It remains unclear whether increased gut permeability influences motility in our LPLFD models, although increased gut permeability is associated with increased symptom severity in a recent study of adults with irritable bowel syndrome.\textsuperscript{47}

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Unsurprisingly, we found gut microbial community composition to be altered in each mouse model of malnutrition, confirming our previous results with TmSep pups\textsuperscript{26,27} and extending these data to our LPLFD pup and young adult malnourished models. Gut microbiota changes are associated with increased barrier permeability, altered GI transit times, and numerous other pathophysiologies that underlie functional GI disorders\textsuperscript{23}. Whether microbiota alterations promote or simply result from the altered intestinal niche in malnutrition, and whether our observed sex-specific gut microbiota alterations drive any aspect of malnutrition-associated GI pathophysiology, remains unknown. We are further investigating these models with mechanistic studies that seek to define a potentially causative role of gut microbiota changes in malnutrition-induced GI pathophysiology.

It has been hypothesized that dysmotility in malnutrition may reduce food intake by promoting abdominal pain and decreasing hunger. We did not observe this phenomenon in our young adult models of malnutrition. Consistent with previous reports using similarly aged C57BL/6J mice\textsuperscript{48}, we observed relatively constant chow consumption over time, with the exception of a small number of outlying days on which all groups simultaneously had decreased chow intake in the absence of an identifiable stressor. Surprisingly, LPLFD young adults consumed increased feed per gram body weight compared to mice receiving isocaloric control chow, suggesting that other metabolic abnormalities contribute to growth faltering in these models. We cannot exclude the possibility that malnourished pups consume less milk than control pups, given the impaired gastric emptying in TmSep pups and the shrunken stomachs in LPLFD pups which might not be able to accommodate a full feed. Additional techniques must be developed to monitor rates of milk intake in malnourished neonatal mice. Alternatively, slow GI motility might act as a host adaptation to malnutrition, given that slowing the rate of transit of ingested food could increase its contact time with the intestinal epithelium, thereby increasing opportunities for absorption of scarce nutrients. Furthermore, preservation of female body mass to maintain healthy endocrine and reproductive axes\textsuperscript{49}, thereby maintaining species fitness during periods of famine, offers a possible evolutionary explanation for our observation of delayed gastric emptying in malnourished young adult females but not males. This intriguing hypothesis requires additional investigation.

The most striking and unexpected finding from these studies is the sexual dimorphism that results from malnutrition. Our data show that sex differences are present with respect to
anthropometric indices, gastric emptying, and gross and histologic stomach appearance. The exaggerated contractile activity and increased barrier permeability observed in LPLFD males is completely absent in LPLFD females. These sex differences are prominent even without controlling experiments for various stages of the estrous cycle. It is unclear why LPLFD females have delayed gastric emptying, yet do not exhibit the same alterations in contractile activity and barrier permeability found in LPLFD males. Females have a higher prevalence of irritable bowel syndrome and certain motility disorders including gastroparesis compared to males, although mechanisms underlying these sex differences are poorly understood. In mice, GI transit times of female and male mice differ not only at baseline, but also in their responses to high-fat chow and to the 5-HT3 receptor antagonist, alosetron. We recently showed that motility indices are identical in healthy male and female neonatal mice, that sex differences emerge soon after puberty, and that motility is more sensitive to numerous variables including the time of day at which motility testing is performed in females compared to males. Future studies will use these models to help determine the mechanistic basis of these sex differences, the specific role host nutritional status plays in GI dysmotility including any structural or neurochemical or functional changes in the malnourished enteric nervous system, and the potential contributions of sex and nutritional status to pathophysiology in functional GI disorders.

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DISCLOSURES

No competing interests declared.
AUTHOR CONTRIBUTION

KGS and JHS designed the research study, performed experiments, analyzed the data, conducted the statistical analysis, and wrote the paper. PND, TH, PTE, and JPPF performed experiments and wrote the paper. MEC designed the study, provided technical and scientific support for the study, and wrote the paper. GAP designed the study, obtained funding, performed experiments, analyzed the data, conducted the statistical analysis, and wrote the paper. All authors approved the final version of this manuscript.

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FIGURE LEGENDS

**FIGURE 1** Multiple mouse models of early-life malnutrition induce characteristic whole-body and GI tract changes. A, Dietary manipulation begins in the neonatal period by feeding dams LPLFD (blue) or control diet (gray) with or without TmSep (orange); pups are tested at 2 wk of life in the fed state or are weaned to their dams’ diet and tested as 8 wk old young adults. B, TmSep pups and LPLFD pups are equally underweight when tested at 2 wk. C, At the time of testing all malnourished models are underweight. D, Decreased chow consumption does not occur in malnourished young adult mice. E, All malnourished models except LPLFD young adult females exhibit decreased length of body, small bowel, and total GI tract. Individual data points are shown along with mean ± SD; **P < .01; n = 7 mice per group; TmSep = timed maternal separation; LPLFD = low-protein low-fat diet.

**FIGURE 2** Malnourished young adults do not exhibit significant changes in whole gut transit time or distal colon motility. A, Elapsed time between carmine red gavage and appearance of the first red fecal pellet. B, Elapsed time between rectal insertion of a glass bead under light...
isoflurane anesthesia and expulsion of the bead onto the cage floor. Individual data points are shown along with mean ± SD; n = 5-9 mice per group; LPLFD = low-protein low-fat diet.

**FIGURE 3** Multiple malnutrition models demonstrate impaired progression of FITC-dextran 30 minutes after gavage. A, Modified neonatal gavage technique reveals minimal variance. B, The small intestinal segment containing peak mean intensity is more distal in control pups compared to TmSep pups and LPLFD pups, but no differences are observed among young adults. C, Progression of the MGC of FITC-dextran is impaired in both malnourished pup models. D, Gastric emptying is delayed in TmSep pups and LPLFD young adult females. Individual data points are shown along with mean ± SD; **P < .01; *P < .05; n = 5 pups and 7 young adult mice per group; FITC = fluorescein isothiocyanate; LPLFD = low-protein low-fat diet; MGC = mean geometric center; TmSep = timed maternal separation; segment 0 = stomach, 1 = proximal duodenum, 10 = terminal ileum. No dye was observed in the cecum or colon.

**FIGURE 4** Multiple malnutrition models affect gross and histologic stomach structure. A, Representative stomachs photographed immediately after harvest reveal grossly abnormal appearance in each malnutrition model; scale bar = 1 cm. B, Representative hematoxylin- and eosin-stained gastric sections reveal well-delineated mucosa and muscularis; scale bar = 100 µm. C, Malnutrition thins the gastric mucosa in TmSep pups and LPLFD males relative to respective controls. Individual data points are shown along with mean ± SD; **P < .01; *P < .05; n = 7-8 mice per group; LPLFD = low-protein low-fat diet; TmSep = timed maternal separation.

**FIGURE 5** Malnutrition alters the intestinal mucosa. A, Malnutrition blunts ileal villus height in LPLFD pups and LPLFD males relative to respective controls. B, Multiple malnutrition models exhibit decreased crypt depth in jejunum, ileum, and proximal colon. Individual data points are shown along with mean ± SD; **P < .01; *P < .05; n = 7-8 mice per group; LPLFD = low-protein low-fat diet; TmSep = timed maternal separation.

**FIGURE 6** Malnutrition increases ex vivo contractility of duodenal segments. A, Representative force-transduction tracings showing equilibration at baseline and after exposure of first 1 µM then 10 µM carbachol. B, Contractile activity, calculated as the integral from minimum, is

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increased in duodenum from LPLFD males at baseline and in response to 1 µM and 10 µM carbachol. C, Average contraction amplitude of duodenal segments does not differ between experimental groups. D, Frequencies of spontaneous or cholinergic-stimulated contractions of duodenal segments are not different between experimental groups. Individual data points are shown along with mean ± SD; **P < .01; *P < .05; n = 7-8 mice per group; LPLFD = low-protein low-fat diet.

**FIGURE 7** Malnutrition increases intestinal permeability to 4-kDa FITC-dextran in Ussing chambers. A, Average translocation across all time points reveals increased permeability in duodenum and proximal colon harvested from LPLFD males. B-D, Real-time measurements of translocation throughout the two-hour experiment for duodenum, proximal colon, and distal colon sections. Individual data points are shown along with mean ± SD; *P < .05; n = 4 mice per group; FITC = fluorescein isothiocyanate; LPLFD = low-protein low-fat diet.

**FIGURE 8** Gut microbial community composition is altered in multiple models of malnutrition. A, Microbial community richness, defined as the number of unique operational taxonomic units, is decreased in TmSep pups but increased in LPLFD males and females relative to controls. B, Fisher’s diversity index reveals a similar pattern. C, Relative abundance of the phylum Bacteroidetes is increased in LPLFD males. D, Relative abundance of the phylum Firmicutes is increased in LPLFD females. E, Relative abundance of the phylum Proteobacteria is increased in both neonatal models of malnutrition. F, Relative abundance of the species *Muribacter muris* drives the observed changes of Proteobacteria. Individual data points are shown along with mean ± SD; ****P < .0001; **P < .01; *P < .05; n = 7-10 mice per group; LPLFD = low-protein low-fat diet; OTU = operational taxonomic unit; TmSep = timed maternal separation.
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