Index diagnoses of gastric intestinal metaplasia in the United States: patient characteristics, endoscopic findings, and clinical practice patterns at a large tertiary care center

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Abstract

Background: Gastric intestinal metaplasia (GIM) is a premalignant gastric mucosal change that is often incidentally detected during esophagogastroduodenoscopy (EGD). Despite the established higher risk of gastric cancer associated with GIM, the incidence, prevalence, and outcomes data for GIM are limited in the United States (US), and practice patterns are highly variable.

Objectives: Our primary objectives were to accurately identify incident histology-confirmed GIM cases and determine patient characteristics, endoscopy findings, Helicobacter pylori (HP) detection, and eradication treatment outcomes, as well as surveillance and follow-up recommendations.

Design: We conducted a retrospective cohort study using administrative data.

Methods: We first developed and validated a rule-based natural language processing tool to identify the patients with GIM on gastrointestinal pathology reports between 2011 and 2016. We then performed a manual chart review of all EGD procedures and associated pathology notes to confirm cases and obtain clinically relevant data.

Results: In all, 414 patients with an index diagnosis of GIM were confirmed (prevalence = 2.5% of patients undergoing any EGD). A majority (52.4%) of patients were non-Hispanic white. The most common indication for EGD was abdominal pain (46.9%). A majority (55%) did not receive specific follow-up recommendations or were asked to see their primary care provider. HP testing was documented in 86% of patients, and detected in 94 patients (prevalence = 26.4%). Treatment was documented in 94.7% of cases, and eradication confirmed in only 34.8% of these cases.

Conclusion: A large group of US patients with an index diagnosis of GIM was accurately identified. There was wide variability in clinical practice patterns including biopsy practice, HP treatment and eradication confirmation testing, and surveillance recommendations. This work demonstrates that there is a major unmet need for quality improvement efforts to standardize care for patients with GIM, a premalignant condition, and inform future prospective studies in a US population.

Keywords: endoscopy, gastric cancer, gastric intestinal metaplasia, Helicobacter pylori, screening and surveillance

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Introduction
Gastric cancer ranks as the fifth most common cause of cancer and fourth most common cause of cancer-related death worldwide, responsible for an estimated 1 million new cases and over 780,000 related deaths in 2019 alone, accounting for 8.2% of all cancer-related deaths. In the United States (US), gastric cancer is the second most common luminal GI malignancy after colorectal cancer, with an estimated 26,380 new cases and 11,090 related deaths estimated to occur in 2022. Importantly, although the US is considered a low-incidence country overall, certain racial and ethnic minority groups have a much higher incidence. It was recently demonstrated that immigrants from high- to low-incidence countries retained their elevated risk of gastric cancer and related mortality. Indeed, the incidence of gastric cancer is at least twofold to threefold higher than US-born non-Hispanic Whites, with the incidence increasing in certain demographic groups.

Gastric intestinal metaplasia (GIM) develops in response to chronic gastric inflammation, with the most common trigger *Helicobacter pylori* (HP) infection. GIM is considered a premalignant condition with an estimated 0.16% baseline-associated annual risk of gastric cancer, with certain modifying factors such as extent of GIM and persistent HP infection associated with an even higher risk of progression. Inflammation from HP infection is known to play a primary role in carcinogenesis, along with other factors including genetic (family history) and environmental (smoking, diet) factors. Because GIM can be reliably identified by gastroenterologists and pathologists, combined with the slow stepwise progression to dysplasia or cancer in a small minority of individuals, endoscopic surveillance of GIM offers a promising opportunity for early diagnosis of gastric cancer with markedly improved outcomes. Indeed, early gastric cancer prior to submucosal invasion is associated with greater than 95% 5-year survival compared to the dismal <30% 5-year survival associated with more advanced disease. In countries with a high burden of gastric cancer, identification and surveillance of patients with premalignant lesions (i.e. GIM) can be a cost-effective way of decreasing the morbidity and mortality associated with the disease.

In the US, due to low overall prevalence of GIM, surveillance has not been shown to be effective in preventing cancer, although some subgroups of the population may benefit. GIM is often detected incidentally from biopsies during esophagogastroduodenoscopy (EGD) performed for common symptoms. Incidence, prevalence, and outcomes data are limited in the US due to an inability to accurately identify patients with GIM in the electronic medical records (EMRs). In addition, endoscopy and pathology reports are often separate and data acquisition can be challenging. Thus, there have been very few studies evaluating large cohorts of patients with GIM in the US. Two small surveys of US physicians have highlighted the variability of clinical practice patterns after diagnosis of GIM, and identified significant knowledge gaps in regard to ethnic populations who may be at increased risk for gastric cancer. A recent large study which evaluated risk factors in US Veterans identified older age, male sex, nonwhite race/ethnicity, and current smoking status as non-endoscopic risk factors which were associated with GIM development.

Recently published professional society guidelines highlight the paucity of high-quality evidence guiding consistent recommendations surrounding GIM management, which is propagated further by difficulties in case ascertainment that would enable robust epidemiologic and outcomes studies. The recent American Gastroenterological Association (AGA) guideline on GIM recommends against routine endoscopic surveillance of GIM due to lack of high-quality evidence, but acknowledges that certain demographic groups may benefit. There was stronger evidence to support treatment of HP in patients with GIM. Thus, the primary objectives of our study included creation of a valid and accurate method to identify patients with GIM diagnosed by EGD, to describe characteristics as well as EGD indications and findings of patients incidentally diagnosed with GIM, and to describe the clinical practice habits after diagnosis, including biopsy patterns and follow-up recommendations. Secondary objectives included assessing the prevalence of GIM in a cohort of patients undergoing EGD, defining the proportion of patients with GIM tested and treated for HP, and delineating the associated pathologic characteristics of GIM, based on a large academic tertiary care setting.

Methods
The reporting of this study conforms to the STROBE statement.
Inclusion and exclusion criteria
We queried the institution’s data warehouse to identify all eligible patients. Eligible patients were those who underwent an EGD with current procedural terminology codes (43200–43259) at the University of Utah (a tertiary referral center) between September 2011 and November 2016 and had gastric biopsies obtained during the procedure. Only patients with new (index cases) diagnoses of GIM were included. Individuals aged <18 years, individuals who underwent EGD but did not have gastric biopsies, individuals with evidence of prior GIM, and individuals with evidence of low- or high-grade dysplasia or gastrointestinal malignancy at the time of GIM diagnosis were excluded. The first diagnosis of GIM was considered the index diagnosis.

Natural language processing
A rule-based Pycon Text natural language processing (NLP) tool was applied to the entire document corpus of endoscopy and pathology notes to identify patients with a diagnosis of GIM.23 NLP systems are a form of machine learning and are developed using a standard or ‘training’ set of terms or phrases that provide the ‘dictionary’ for the software to extract the information from text documents. In this case, phrases and terms from pathology results for intestinal metaplasia, including accounting for misspelling, were used to train the system. The software system was then validated iteratively to determine information extraction characteristics. The NLP system was developed using open-source code and is available to use and reproduce at this link: https://gitlab.chpc.utah.edu/endoqual/endoscopy-path-classify/-/tree/master

Data abstraction and case confirmation
The EMRs of all patients meeting initial inclusion criteria, inclusive of clinical notes, endoscopic reports, and pathology reports, were manually reviewed to confirm the diagnosis of GIM and abstract additional details. Using a standardized data collection form, we recorded the following details at the time of endoscopic evaluation: age, sex, race/ethnicity (non-Hispanic white, non-Hispanic black, Hispanic, Asian, etc.), smoking status (current, former, never), and family history of gastric cancer in a first-degree relative. Missing or unknown data were documented. The following endoscopic details were recorded: indication, gross findings, location of each biopsy site, including whether antrum/incisura and corpus biopsies were placed in separate containers. The following pathology details were recorded: locations of GIM (antrum, incisura, corpus/fundus, cardia), anatomic extent (corpus-extended GIM, defined as any GIM involving the corpus versus limited GIM, defined as GIM confined to the antrum/incisura8), and any additional gastric pathology (e.g. dysplasia, cancer). We also manually abstracted other HP testing and eradication therapy details at any time following patients’ diagnosis of GIM; details included test modality, result, eradication therapy (yes/no), and eradication outcomes (eradicated successfully versus persistent infection). Follow-up recommendations after a confirmed index diagnosis of GIM were documented. No biopsies were re-reviewed for the purposes of this study.

Analyses and descriptive statistics were performed using STATA 10.0.

Results
A total of 16,505 distinct patients underwent a total of 23,404 procedures during the study time frame. The NLP tool initially identified 516 patients with a likely diagnosis of GIM, 510 (98.8%) of whom were confirmed to have histologic evidence of GIM on manual review of associated pathology reports. The F-measure, a surrogate for specificity for information extraction, of the NLP tool was 92%. An additional 20 patients (3.9%) had dysplasia or evidence of malignancy at the time of diagnosis and were excluded. After additional exclusion criteria were applied (Figure 1), a total of 414 patients with an index diagnosis of GIM were included for analysis.

The overall prevalence of incidentally diagnosed GIM in all patients undergoing endoscopy during the study time period was 2.5%. Baseline characteristics of the patients diagnosed with GIM and no neoplasia (N=414) are included in Table 1, along with comparative demographics of those diagnosed with concomitant neoplasia (N=20). The mean age was 59.3 years (range, 18–91 years), and the majority of patients were female (60.1%). With respect to race/ethnicity, most were non-Hispanic white (52.4%), followed by Hispanic (26.1%), Asian (9.7%), and Native American/
Pacific Islander (5.6%), respectively. A majority were categorized as never smokers.

The most common indication for EGD was abdominal pain/dyspepsia (46.9%), followed by gastro-esophageal reflux disease (22.2%), dysphagia (16.9%), iron deficiency anemia (8.5%), and nausea/vomiting (7.7%) (Table 2). The most common endoscopic finding among patients with GIM was gastritis (69.8%), with a minority reporting ulcerations/erosions (13.5%) or normal gastric mucosa (13%). The most common specified location of gastric biopsies were antrum and corpus combined in the same container (47.6%), followed by antrum alone (23.2%), fundus alone (12.8%), and corpus alone (2.9%). Nearly 25% had an unspecified location for gastric biopsies.

Table 3 details the anatomic locations of confirmed GIM, as well as anatomic extent. An unspecified gastric location for GIM was observed in 43% of cases (e.g. ‘random gastric’ biopsies), followed by antrum alone (34.1%), antrum and body combined (10.9%), and body alone (8.7%).

Follow-up recommendations varied widely (Figure 2). A majority of patients (55%) did not receive specific follow-up recommendations or were simply asked to follow-up with their primary care provider as opposed to their GI provider, while 38% of patients were offered repeat EGD (23% being recommended within 1 year and the remaining 15% recommended an EGD at an interval greater than 1 year). A minority (7%) of patients were offered a gastroenterology clinic visit to discuss these findings and decide upon a future care plan. Of the 97 patients who were recommended to have a repeat EGD within 1 year, 51% had this exam completed at our institution within that time frame.

A majority of patients (86%) did have HP testing based on immunohistochemical testing of pathology specimens. Of these, HP was detected in 94 patients (Table 4), with a prevalence of 26.4% in patients with an index diagnosis of GIM. Treatment was documented in 89 of these patients (94.7%), but unfortunately, most patients did not have eradication confirmation testing. Eradication was confirmed based on repeat testing in 31 patients (34.8%). The most commonly used test to confirm eradication was repeat biopsy (16/31 patients, 51.6%), followed by stool antigen test (8/31 patients, 25.8%) and breath test (7/31 patients, 22.6%).

Figure 1. Flow diagram of cohort selection. GIM, gastric intestinal metaplasia; NLP, natural language processing.
In total, 20 patients were excluded because of findings of malignancy or dysplastic changes at the time of index diagnosis of GIM. The total malignancies (12) were as follows: adenocarcinoma (4), neuroendocrine tumor (4), mucosa-associated lymphoid tissue lymphoma (2), gastrointestinal stromal tumor (1), and metastatic hepatocellular carcinoma (1). The remaining eight patients with high-grade dysplasia were all recommended to have a repeat EGD within 1 year, and five of these patients had documented follow-up within our EMR.

Table 1. Patient characteristics.

|                          | GIM N = 414 | Dysplasia/malignancy N = 20 |
|--------------------------|-------------|----------------------------|
| **Demographics**         |             |                            |
| Age (mean, SD)           | 59.3 ± 14.3 | 64.2 ± 14.5                | \( p = 0.13 \) |
| BMI (mean, SD)           | 27.7 ± 6.9  | 27.2 ± 6.0                 | \( p = 0.384 \) |
| Sex                      |             |                            | \( p = 0.366 \) |
| Male N (%)               | 165 (39.9%) | 10 (50%)                   |
| Female N (%)             | 249 (60.1%) | 10 (50%)                   |
| **Race**                 |             |                            | \( p = 0.198 \) |
| White N (%)              | 217 (52.4%) | 11 (55%)                   |
| Asian N (%)              | 40 (9.7%)   | 2 (10%)                    |
| Native American/Pacific Islander N (%) | 23 (5.6%) | 1 (5%)                     |
| Black N (%)              | 14 (3.4%)   | 3 (15%)                    |
| Other N (%)              | 12 (2.9%)   | 0 (0%)                     |
| **Ethnicity**            |             |                            | \( p = 0.507 \) |
| Hispanic N (%)           | 108 (26.1%) | 3 (15%)                    |
| Non-Hispanic N (%)       | 306 (73.9%) | 17 (85%)                   |
| **Smoking status**       |             |                            | \( p < 0.001 \) |
| Never smoker N (%)       | 259 (62.6%) | 9 (45%)                    |
| Prior smoker N (%)       | 101 (24.4%) | 7 (35%)                    |
| Current smoker N (%)     | 50 (12.1%)  | 1 (5%)                     |
| Unknown N (%)            | 4 (1%)      | 3 (15%)                    |
| **Family history of gastric cancer** |             |                            | \( p = 0.051 \) |
| Yes N (%)                | 3 (0.7%)    | 1 (5%)                     |
| No N (%)                 | 411 (99.3%) | 19 (95%)                   |

GIM, gastric intestinal metaplasia.
Table 2. Endoscopy indications, findings, and biopsy practice patterns.

| Endoscopy indications                      | GIM (%) | Dysplasia/malignancy (%) |
|-------------------------------------------|---------|--------------------------|
| Abdominal pain/dyspepsia N (%)            | 194 (46.9) | 7 (35)                  |
| Reflux/GERD N (%)                         | 92 (22.2) | 3 (15)                  |
| Dysphagia N (%)                           | 70 (16.9) | 1 (5)                   |
| Iron deficiency anemia N (%)              | 35 (8.5)  | 4 (20)                  |
| Nausea/vomiting N (%)                     | 32 (7.7)  | 1 (5)                   |
| History of Helicobacter pylori N (%)      | 18 (4.4)  | 1 (5)                   |
| Melena/hematemesis N (%)                  | 14 (3.4)  | 4 (20)                  |
| Variceal screening or surveillance N (%)  | 18 (4.4)  | 0 (0)                   |
| Diarrhea N (%)                            | 17 (4.1)  | 0 (0)                   |
| Barrett’s esophagus N (%)                 | 14 (3.4)  | 0 (0)                   |
| Abnormal imaging N (%)                    | 12 (2.9)  | 1 (5)                   |
| History of peptic ulcer N (%)             | 12 (2.9)  | 0 (0)                   |
| Atrophic gastritis N (%)                  | 10 (2.4)  | 0 (0)                   |
| Eosinophilic esophagitis N (%)            | 2 (0.5)   | 0 (0)                   |
| Other N (%)                               | 42 (10.1) | 4 (20)                  |
| Endoscopy findings                        |         |                         |
| Gastritis/erythema N (%)                  | 289 (69.8) | 8 (40)                  |
| Ulcerations/erosions N (%)                | 56 (13.5) | 4 (20)                  |
| Normal N (%)                              | 54 (13)   | 0 (0)                   |
| Polyps N (%)                              | 40 (9.7)  | 7 (35)                  |
| Atrophic N (%)                            | 42 (10.1) | 3 (15)                  |
| Nodular/papular N (%)                     | 35 (8.5)  | 3 (15)                  |
| Mass N (%)                                | 0 (0)     | 3 (15)                  |
| Other N (%)                               | 5 (1.2)   | 1 (5)                   |
| Endoscopy biopsy sites                    |         |                         |
| Antrum and body N (%)                     | 197 (47.6) | 13 (65)                  |
| Antrum N (%)                              | 96 (23.2) | 2 (10)                  |
| Fundus N (%)                              | 53 (12.8) | 4 (20)                  |
| Body N (%)                                | 12 (2.9)  | 1 (5)                   |
| Cardia N (%)                              | 9 (2.2)   | 2 (10)                  |
| Unspecified N (%)                         | 103 (24.9) | 2 (10)                  |

GIM, gastric intestinal metaplasia.
Table 3. GIM location and histology characteristics.

| GIM location          | N=414 |
|-----------------------|-------|
| Antrum N (%)          | 141   (34.1%) |
| Antrum and body N (%) | 45    (10.9%) |
| Body N (%)            | 36    (8.7%) |
| Fundus N (%)          | 21    (5.1%) |
| Cardia N (%)          | 5     (1.2%) |
| Unspecified N (%)     | 178   (43%) |

| GIM characteristics   |       |
|-----------------------|-------|
| Focal N (%)           | 164   (39.6%) |
| Extensive N (%)       | 34    (8.2%) |
| Unspecified N (%)     | 216   (52.2%) |

GIM, gastric intestinal metaplasia.

Discussion
In this large retrospective observational cross-sectional study, we identified marked practice pattern variability and heterogeneity in the management of index diagnoses of GIM. Despite GIM being a well-established premalignant condition, with an estimated baseline 0.16% annual risk of gastric cancer, the vast majority of patients diagnosed with GIM were not recommended for any GI follow-up nor further risk stratification to better determine their need for ongoing surveillance. In addition, we successfully developed and validated an NLP tool among patients referred for routine EGD, and confirmed the diagnostic accuracy for incident GIM cases. If externally validated, this tool may have the potential value for creating larger cohorts of patients with GIM in an effort to better define the natural history of GIM and comparative outcomes based on practice patterns in the US. Finally, this study demonstrates the marked heterogeneity of biopsy techniques, pathology reporting, follow-up recommendations, and HP treatment strategies following diagnosis of GIM, which likely is the effect of previous lack of formal accepted guidance. GIM is a fairly common incidental finding during EGD but is associated with a significantly increased risk of gastric cancer, particularly in patients with additional risk factors including corpus-extended GIM, family history of gastric cancer, persistent HP, among others. Until recently, the lack of established US guidelines left clinicians without clear guidance when confronted with this clinical scenario.

Figure 2. Follow-up recommendations.
EGD, esophagogastroduodenoscopy; GI, gastrointestinal; PCP, primary care provider.
The prevalence of GIM diagnosis in this study was 2.5% of patients undergoing upper endoscopy during the study timeframe. While the prevalence of GIM in the US population is unknown, this prevalence is lower than that reported in previous US cohorts, which has ranged from 5% to about 19% in various studies.\textsuperscript{9,24,25} Reported prevalence in international cohorts has varied even more widely, from 3.4% to over 30%.\textsuperscript{9} These studies included patients undergoing EGDs for certain indications and had varying proportions of ethnic diversity included in their study populations compared to Utah which is ~85% non-Hispanic White. We also did not restrict our prevalence estimate to only those patients undergoing gastric biopsies, which presumably would bias toward a higher prevalence. Because this is a retrospective study, we could not control for the number or location of biopsies, which is relevant given that GIM is typically patchy and subject to sampling error.

As has been described in other studies, this study showed variability in recommendations from providers after diagnosis of GIM. A survey of gastroenterologists in the US in varying practice settings found confusion regarding the existence of guidelines, and variability in recommendations for surveillance.\textsuperscript{17,18} In a more recent retrospective study of patients with GIM, surveillance endoscopy was recommended in only about 17% of patients with intervals ranging from 1 to 4 years.\textsuperscript{26} A majority of patients (55%) in this study were not given clear guidance on when (or whether) to have an endoscopy repeated for surveillance. In addition, of ~1 in 4 patients who were recommended to have a surveillance endoscopy within a year, only about half completed this exam. This finding suggests that gastric cancer risk (such as country of origin, histology findings) is neither sufficiently characterized nor understood by providers who are making recommendations to patients. In addition, it is unknown whether patients are educated about their risk for developing gastric cancer. It is unclear what other factors endoscopists may have used in recommending surveillance endoscopies to some patients and not in others.

Endoscopy practice patterns, including biopsy sites and description of gross findings, along with histology practice patterns, including histologic subtyping of GIM and severity scoring, both vary widely. The risk of progression of GIM to gastric cancer has been demonstrated to be higher among patients with metaplastic changes in both the antrum and corpus.\textsuperscript{8,27} Despite this, the location of GIM was unspecified in over 40% of cases described. This can be explained by the fact that there has been shown to be a poor correlation between the endoscopic (visual) identification of GIM and a histologic diagnosis in the US particular.\textsuperscript{28} In addition, while histologic description of ‘limited’ or ‘corpus-extended’ GIM was used in 47% of cases, over half of the diagnoses were not specified; histologic subtype, complete versus incomplete, was not reported, even though this can be a marker of progression risk. While there are standardized biopsy approaches (updated Sydney protocol) and recommended quality performance measures for upper endoscopy, adherence to these is not evident in this study.\textsuperscript{29,30}

| Table 4. Details related to HP diagnoses and treatment outcomes. |
|---------------------------------------------------------------|
| HP tested | N = 414 |
| Yes N (%) | 356 (86%) |
| No N (%) | 58 (14%) |
| HP present | N = 356 |
| Yes N (%) | 94 (26.4%) |
| No N (%) | 320 (89.9%) |
| HP treated | N = 94 |
| Yes N (%) | 89 (94.7%) |
| No N (%) | 5 (5.1%) |
| HP eradication confirmation testing performed and eradication confirmed | N = 89 |
| Yes N (%) | 31 (34.8%) |
| No* N (%) | 58 (65.2%) |
| Test used for eradication confirmation | N = 31 |
| Biopsy N (%) | 16 (51.6%) |
| Stool antigen N (%) | 8 (25.8%) |
| Breath test N (%) | 7 (22.6%) |

*This number represents either those without confirmation testing, or with confirmation testing and not eradicated. HP, Helicobacter pylori.
HP testing was clearly described in pathology reports in 86% of cases, and treatment was documented in almost 95% of patients; however, eradication was confirmed in only 35%, owing to a very poor rate of performance for eradication confirmation testing. The European Society of Gastrointestinal Endoscopy strongly recommends to test for and eradicate HP once the diagnosis of GIM has been made. The recent AGA guidelines also strongly recommend HP treatment and confirmation of eradication based on high-quality evidence that treatment (regardless of GIM) is associated with reduced risk of gastric cancer. Whether or not histologic testing is sufficient in the setting of GIM to adequately assess for HP infection is unknown. In this study, there were no cases where a negative histologic examination for HP triggered further evaluation (i.e. via hydrogen breath test or stool antigen test).

The strengths of this study include the large cohort of patients diagnosed with GIM, which is among the largest in the US identified to date, and manual chart review to confirm the diagnosis and abstract patient-level data. The physicians involved in the study were all teaching faculty in a large university setting. An integrated EHR was available to obtain relevant patient and clinical data. The novel development of an NLP tool to meld endoscopy and pathology reports made identification of the study population achievable. One of the limitations of the study is its retrospective nature. As data were obtained from the EHR, and since the University of Utah serves a large catchment area, it is possible that not all follow-up data were captured. Although all endoscopies were performed using high-definition white light, the lack of standardization in EGD and biopsy practices (e.g. adherence to Sydney protocol, use of virtual chromoendoscopy) and reporting of pathology specimens is also a limitation. However, we will note that the practice patterns we identified are unlikely to be unique to our practice setting and may even be more pronounced in some centers.

Conclusion
In conclusion, this study identifies significant unmet needs and points of intervention in the management of GIM. Notably, it highlights the lack of standardized care and follow-up recommendations preceding recently published US-based guidelines. The study highlights the variability in clinical practice and future work should focus on understanding if the new US guidelines help providers and patients better understand GIM disease progression risk factors to allow for more standard follow-up recommendations. In addition, clinicians should be educated about the need for documentation of HP testing and confirmed eradication of infection, particularly given the rising rate of HP eradication failure. Further studies are needed to determine and solidify appropriate recommendations for the management and surveillance of GIM.

Declarations

Ethics approval and consent to participate
This retrospective cohort study was approved by the University of Utah Health Institutional Review Board.

Consent for publication
All listed authors consent to publication.

Author contributions
Sheeva K. Parbhu: Conceptualization; Data curation; Formal analysis; Methodology; Writing – original draft.
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Kathryn A. Peterson: Writing – review & editing.
Andrew J. Gawron: Conceptualization; Data curation; Investigation; Methodology; Resources; Software; Supervision; Writing – original draft; Writing – review & editing.

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Availability of data and materials
Data are not publicly available due to protected health information. The corresponding author can be contacted if additional materials and methods information is needed.

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Supplemental material
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