INTRODUCTION

Celiac disease (CD) is an immune-mediated enteropathy triggered by the presence of gluten in genetically susceptible individuals, with a prevalence in children as high as 1% (1–3). Currently, the only treatment for CD is a strict, lifelong gluten-free diet (GFD) (4). Diagnosis of CD in the United States is made through endoscopic proximal small-intestine biopsy to assess intraepithelial lymphocytosis, crypt hyperplasia, villous atrophy, and inflammation (1,5,6). While duodenal biopsy is the standard of care for CD diagnosis in North America, follow-up recommendations in pediatric patients with CD are limited (7). While current clinical guidelines do not recommend follow-up biopsies, there is no consensus on the value and role of repeat biopsy in the assessment of mucosal healing after beginning and maintaining a GFD for pediatric patients (5,8–10). Many pediatric practices rely solely on serologic findings such as tissue transglutaminase immunoglobulin A (tTG-IgA), and less commonly endomyosal antibody IgA (EMA-IgA) and deamidated gliadin immunoglobulin G, following initial CD diagnosis in the assessment of duodenal healing. However, the reliability of serologies to assess degree of duodenal damage is unclear (1,2,4,11,12). Therefore, the accuracy of using tTG-IgA as a marker of duodenal mucosal healing and its potential correlation with symptoms, adherence to a GFD, and other factors require further investigation (13).

Current standard of care in the management of uncomplicated CD is not to undergo multiple esophagastroduodenoscopies (EGDs).
Therefore, it was important to identify a subsection of pediatric CD patients who have had multiple EGDs to investigate the relationship of biopsies and antibody levels over time. In this study, patients with both CD and eosinophilic gastrointestinal disorders (EGID) including eosinophilic esophagitis (EoE) and eosinophilic gastritis (EG) were identified to explore this relationship, as it is standard for patients with EGID to undergo repeat EGDs for disease surveillance (14,15).

In this study, data from pediatric patients with CD and EGID who have had multiple EGDs and duodenal biopsies are presented to assess duodenal healing before and after starting a GFD and correlate it with tTG-IgA serology. In this way, the reliability of using serologies to monitor improvement of duodenal damage can be assessed in this patient cohort.

METHODS

A retrospective data analysis was conducted by obtaining patient data from the electronic medical record from the Children’s Hospital of Philadelphia (CHOP). According to the North American Society for Pediatric Gastroenterology, Hepatology, and Nutrition guidelines, a CD diagnosis is confirmed through characteristic pathology taken from the duodenal bulb and the distal duodenum (16). These microscopic features include infiltration of lymphocytes into the epithelium of the duodenum, progressive flattening of the villi, and deepening of the crypts (17). All patients included in this study met the North American Society for Pediatric Gastroenterology, Hepatology, and Nutrition criteria for CD and had positive levels of tTG-IgA at the time of diagnosis. All patients were started on a GFD after CD diagnosis. EGID consists of several EGID; the study reported IgA level. The final analyzed cohort was 39 patients (Fig. 1). Additionally, 10 were eliminated for being IgA deficient or having no EGD, 3 were excluded for not having repeat antibody levels, and 5 patients were eliminated due to insufficient data in the electronic medical record, 17 were excluded because they did not undergo a repeat diagnostic or International Classification of Diseases, Tenth Revision or International Classification of Diseases, Ninth Revision coded diagnoses of CD and EoE/EoG seen at CHOP between 2003 and 2018 were included. Of these 132 patients, 35 did not meet the diagnostic criteria of CD or EoE/EoG, as defined above. An additional 23 patients were eliminated due to insufficient data in the electronic medical record; 17 were excluded because they did not undergo a repeat EGD, 3 were excluded for not having repeat antibody levels, and 5 more were excluded for having EGD performed at outside hospitals. Additionally, 10 were eliminated for being IgA deficient or having no reported IgA level. The final analyzed cohort was 39 patients (Fig. 1). Eighty-two percent (32/39) of patients were diagnosed with CD and EGID at the same time.

Celiac Serologies

Laboratories reviewed and analyzed from CHOP and other commercial laboratories included tTG-IgA, total IgA, and EMA-IgA. The average length of time between collection of laboratories and EGD was 30 days (median = 0 days). It should be noted that celiac serologies were analyzed at several different laboratories, each with unique tTG-IgA normal scales.

Duodenal Biopsies

EGD pathology reports from diagnostic duodenal biopsies and repeat duodenal biopsies were reviewed and the presence of intraepithelial lymphocytes and degree of villous blunting was assessed. Villous blunting was classified as absent, mild, moderate, or severe based on pathologic report. The presence of increased intraepithelial lymphocytes was classified as either present or absent. Intraepithelial lymphocytes were considered to be increased if there were more than 40 lymphocytes per 100 enterocytes in the duodenum (17). Pathology reports did not consistently comment on crypt architecture/hyperplasia and this was therefore not analyzed.

Clinical Manifestations

Clinic notes from office visits before initial EGD diagnosis of CD and before start of the GFD, as well as at time of second EGD and after initiation of the GFD, were reviewed. Clinical manifestations including abdominal pain, nausea, vomiting, diarrhea, constipation, flatus, gastroesophageal reflux, rashes, joint pain, and malnutrition were recorded. Malnutrition was defined as a z score of ≤−1.0 based on weight/height-for-age, body-mass-index-for-age, weight-for-height-for-age, or mid-upper-arm-circumference. Patients were considered malnourished if they had mild, moderate, or severe malnutrition.

Statistical Analysis

The data were analyzed using the Fisher exact test and t test.

Institutional Review Board Approval

This study was approved by the Institutional Review Board at CHOP.

RESULTS

In total, 39 pediatric patients met the inclusion criteria of both CD and EGID and were IgA sufficient. All 39 had positive endoscopic findings indicative of CD and positive tTG-IgA at the time of diagnosis, and all 39 had repeat biopsies and serologies after beginning a GFD. As shown in Table 1, 92% (36/39) of patients were symptomatic at diagnosis, including 51% (20/39) with abdominal pain, and 38% (15/39) with malnutrition.

All study patients underwent at least 1 repeat EGD after CD diagnosis to assess EGID disease activity, as is standard of care for EGID. All patients had duodenal biopsies as well as esophageal biopsies during repeat EGD. At time of second EGD, 59% (23/39) reported ongoing symptoms and 33% (13/39) were asymptomatic. A symptom update was unavailable for 8% (3/39) of patients. Figure 2 depicts the patient-reported symptoms at time of first and second EGD. The average length of time between the first and second EGDs for the cohort was 324 days (the median was 175 days; the range was 31–1897 days). Fifty-six percent (22/39) of patients had a repeat endoscopy within 6 months. All symptoms improved for all patients between the first 2 EGDs. Abdominal pain, constipation, and malnutrition were the most frequently reported ailments at time of CD diagnosis. At time of second EGD, abdominal pain, nausea, and malnutrition were most commonly reported. Eighty-seven percent (34/39) were on a proton pump inhibitor at the time of second EGD, and 97% (38/39) were treated with a proton pump inhibitor at some point during the study timeframe. Fifty-nine percent (23/39) were treated with elimination diets for eosinophilic disease. Five percent (2/39) were treated with swallowed steroids, and 1 patient was treated with an inhaled steroid. Sixty-two percent (23/37) of patients still had evidence of active EoE at the time of second EGD (≥15 eosinophils per high powered field). Both patients with EG (2/2) still had evidence of eosinophilia in the stomach at the time of the second EGD.

If available, data from up to 5 EGDs was collected for each patient. All 39 patients had at least 2 EGDs, 72% (28/39) had 3 EGDs, 33% (13/39) had 4 EGDs, and 21% (8/39) had 5 EGDs. Multiple EGDs were performed due to persistently active EGID. At the tie of second EGD, 36% (14/39) had mild to severe villous blunting; at the third EGD, 7% (2/28) had mild to severe villous blunting; and
Forty-four percent of patients (17/39) had a normal second duodenal biopsy after CD diagnosis and starting a GFD (Table 2). Repeat biopsies were performed a mean of 324 days or 10.5 months after initial EGD. Fifteen of the 17 patients with normal repeat duodenal biopsies had tTG-IgA levels taken at the time of the second EGD, and of these 15 patients, 60% (9/15) had an abnormal tTG-IgA level. The relationship between tTG-IgA and duodenal pathology was completely independent in these 15 patients ($P = 1.000$). Of the nine patients with abnormal tTG-IgA and normal biopsies, 78% (7/9) had down-trending levels of tTG-IgA, 11% (1/9) had a higher tTG-IgA, and 11% (1/9) were not able to be compared due to subsequent bloodwork being drawn at different laboratories with unique reference ranges.

Forty-three percent (9/21) of patients with a positive tTG-IgA at the time of the second biopsy had a normal duodenal biopsy. Of these 9 patients with elevations of tTG-IgA, 44% (4/9) had active EoE, 22% (2/9) had gastritis, and the remaining 33% (3/9) had no evidence of active EoE, gastritis, or CD.

Thirty-six percent (14/39) of patients had negative tTG-IgA values at the time of their second biopsy. Of these, 57% (8/14) had evidence of active histologic duodenal disease at the time of second biopsy despite negative tTG-IgA levels.

**TABLE 1.** Patient characteristics at diagnosis of CD (n = 39)

| Characteristic               | % (n) |
|------------------------------|-------|
| Gender                       |       |
| Female                       | 31 (12)|
| Male                         | 69 (27)|
| Mean age at diagnosis, y ± SD| 7 ± 3  |
| Type 1 diabetes              | 13 (5 )|
| Asthma                       | 15 (6 )|
| Symptoms at diagnosis        |       |
| Abdominal pain               | 51 (20)|
| Nausea and/or vomiting       | 13 (5 )|
| Diarrhea                     | 18 (7 )|
| Constipation                 | 31 (12)|
| Flatus                       | 5 (2 ) |
| Gastroesophageal reflux      | 23 (9 )|
| Rash                         | 13 (5 )|
| Joint pain                   | 8 (3 ) |
| Malnutrition                 | 38 (15)|

CD = celiac disease; SD = standard deviation.
In a subanalysis, patients whose repeat endoscopies were completed at least 1 year after CD diagnosis were evaluated. Sixty-seven percent (26/39) had a repeat biopsy taken at least 1 year after diagnosis and initiation of GFD, and 46% (18/39) had a repeat biopsy taken at least 2 years after diagnosis and initiation of GFD. Of these 18 patients, 94% (17/18) had normal duodenal biopsies after 2 years, and 83% (15/18) had normal tTG-IgA values after 2 years. Of note, the 1 patient with an abnormal biopsy had a normal tTG-IgA value, and the 3 patients with abnormal laboratory values had normal biopsies.

The reference values for tTG-IgA are not standardized. Rather, they vary between laboratories. In this patient cohort, 59% (23/39) had their initial bloodwork, and at least 1 repeat set of bloodwork, at CHOP’s laboratory which quantified “normal” range as 0–20 U/mL, and “abnormal” as above 20 U/mL. A subanalysis of these 23 patients was conducted to compare tTG-IgA levels over time after initiating a GFD. The average tTG-IgA value at diagnosis in this sub-group was 145.6 U/mL. Fifty-seven percent (13/23) had repeat laboratories drawn within 6 months of CD diagnosis (mean = 118 days). The average repeat tTG-IgA level within 6 months of starting a GFD was 42.3 U/mL. Sixty-five percent (15/23) of patients in this subgroup had repeat laboratories taken between 6 and 12 months from diagnosis (mean = 279 days). The average tTG-IgA level taken within 6–12 months of initiating a GFD was 26.8 U/mL. Forty-three

FIGURE 2. Frequency of symptoms at the time of initial CD diagnosis and second EGD after initiating a gluten-free diet. CD = celiac disease; EGD = esophagogastroduodenoscopy.

FIGURE 3. Relative proportions of patients with villous blunting on subsequent EGDs. EGD = esophagogastroduodenoscopy.
### TABLE 2. Symptoms, serology, and histopathology results at first and second EGD

|                      | N (%) | \(P\) value |
|----------------------|-------|-------------|
| **First EGD**        |       |             |
| Symptomatic at diagnosis | 36/39 (92) |       |
| Positive tTG-IgA     | 39/39 (100) |       |
| Positive EMA-IgA     | 30/30 (100) |       |
| Mild to severe duodenal villous blunting | 39/39 (100) |       |
| **Second EGD**       |       |             |
| Ongoing symptoms     | 23/39 (59) |       |
| On PPI               | 34/39 (87) |       |
| On an elimination diet | 39/39 (100) |       |
| Mild to severe duodenal villous blunting | 14/39 (36) |       |
| Active eosinophilic disease | 23/39 (59) |       |
| Normal repeat duodenal biopsy | 17/39 (44) |       |
| Positive tTG-IgA     | 9/15* (60) | 1.000†    |
| Negative tTG-IgA     | 6/15* (40) |       |
| Active CD on repeat duodenal biopsy | 22/39 (56) |       |
| Positive tTG-IgA     | 14/22 (64) |       |
| Negative tTG-IgA     | 8/22 (36) | 1.000†    |
| Both increased intraepithelial lymphocytes and villous atrophy | 8/39 (21) |       |
| Positive tTG-IgA     | 6/8 (75) | 0.324      |
| Negative tTG-IgA     | 2/8 (25) |       |
| Negative tTG-IgA     | 14/39 (36) |       |
| Active CD on repeat duodenal biopsy | 8/14 (57) | 1.000    |
| Positive tTG-IgA     | 21/39 (54) |       |
| Normal repeat duodenal biopsy | 9/21 (43) | 1.000     |
| Active EoE           | 4/9 (44) |       |
| Gastritis            | 2/9 (22) |       |
| No histologic pathology in esophagus, stomach or duodenum | 3/9 (33) |       |
| Negative EMA-IgA     | 17/32 (53) |       |
| Abnormal repeat duodenal biopsy | 2/17 (12) | 0.106    |
| Positive EMA-IgA     | 15/32 (47) |       |
| Normal repeat duodenal biopsy | 9/15 (60) | 0.106     |

CD = celiac disease; EGD = esophagogastroduodenoscopy; EMA = endomysial antibody; EoE = eosinophilic esophagitis; PPI = proton pump inhibitor; IgA = immunoglobulin A; tTG = tissue transglutaminase.

*15 of 17 had tTG-IgA levels recorded.
†No correlation between tTG-IgA level and duodenal pathology.

percent (10/23) had repeat laboratories taken 12 months or more after diagnosis (mean = 826 days). The average tTG-IgA value of these patients was 22.8 U/mL.

At the time of the first EGD, 30 of 39 patients had an EMA-IgA collected. All 30 patients had a positive EMA-IgA. At the time of the second EGD, 32 of 39 total patients had an EMA-IgA collected. Forty-seven percent (15/32) had a positive EMA-IgA. Sixty percent (9/15) had a positive EMA-IgA and normal duodenal pathology on the second EGD. Twelve percent (2/17) had a negative EMA value at the time of the second EGD and also had an abnormal duodenal biopsy \( (P = 0.106) \).

Twenty-three percent (9/39) of patients in the study never had a normal biopsy indicating CD remission. Of these 9 patients, 5 became symptom-free on a GFD, and therefore did not undergo additional endoscopies. Three patients were lost to follow-up. The final patient had a follow-up EGD but the physician did not take duodenal biopsies. Rather, the physician noted that the duodenum was visually normal, compared with previously abnormal visual findings. All patients that had 4 or 5 EGDs achieved histologic remission. The average length of time from initial abnormal biopsy and CD diagnosis to duodenal healing in those that did achieve duodenal healing was a mean of 565 days and a median of 297 days (the mean is skewed due to outliers with prolonged healing times). Eight percent (3/39) of patients had a normal duodenal biopsy, but later had an abnormal biopsy during repeat EGD for persistent EGD.

### DISCUSSION

CD is a life-long immune-mediated enteropathy with the potential to lead to many chronic health complications without proper management. While the current standard of care in pediatric gastroenterology is to monitor serologic markers following diagnosis to predict gastrointestinal healing, there is conflicting evidence as to whether serology appropriately reflects the degree of duodenal damage, and therefore whether a child’s CD is being adequately controlled. Current guidelines from the American Gastroenterological Society recognize the potential unreliability of tTG-IgA, and recommend repeat biopsy in patients with persistent or relapsing symptoms even with negative serology (20).

Long-term consequences of uncontrolled CD, including nutritional deficiencies, increased risk of lymphoma, low bone density, dermatologic rashes, and ongoing gastrointestinal symptoms, indicate that systematic follow-up and monitoring are important in the management of pediatric CD (4,11,21,22). Data collected in the United States regarding mucosal healing are highly variable and suggest that serology levels may not be reliable markers of pathology (1,4,10). Since the standard of care for patients diagnosed with CD alone does not currently include repeat EGD, evaluating pathology data for patients diagnosed with both CD and EGID provides biopsy data on CD patients before and after initiation of a GFD.

This pilot study demonstrated that in patients with pediatric CD and EGID, abnormal tTG-IgA and EMA-IgA antibody levels do not correlate with abnormal duodenal pathology after initiation of a GFD. If a duodenal biopsy was found to be normal on the second EGD, only 40% of the patients had a normal tTG-IgA and 60% had a normal EMA-IgA. Some pediatric patients still demonstrated characteristic changes on duodenal biopsy at the time of repeat EGD, such as increased intraepithelial lymphocytes and villous blunting, and yet had normal tTG-IgA and EMA-IgA values. Thus, for pediatric patients with CD and EGID, abnormal celiac serologies after initiating a GFD may not correlate with duodenal mucosal injury and may not be a reliable predictor of duodenal healing. Therefore, healthcare providers should be cautious of relying solely on celiac serologies, particularly tTG-IgA, as markers to assess duodenal mucosal healing in children with CD. Further consideration should be made regarding repeat duodenal biopsies to confirm mucosal healing.

The vast majority of patients who had biopsies taken at least 2 years after their initial diagnosis of CD had tTG-IgA levels and duodenal structure that normalized. The normalization of previously damaged duodenal mucosa in this study reinforces the principle that small bowel inflammation is reversible for children with CD who maintain a strict lifelong GFD (23,24). Furthermore, these findings reinforce that clinicians should temper expectations of mucosal healing and serologic normalization in children with CD before 2 years on a GFD.

Questions arise as to what could be driving elevated tTG-IgA in patients who no longer have evidence of active CD on duodenal biopsies. Of the patients with elevated tTG-IgA and normal duodenal
biopsies, 66% (6/9) had inflammation elsewhere in the upper gastrointestinal tract, including 4 patients with active EoE and 2 patients with gastritis. This raises the question as to whether EoE or other inflammatory conditions of the upper gastrointestinal tract could also elevate tTG-IgA in CD patients, even after their CD has gone into remission. Several studies have investigated and hypothesized potential causes of elevated tTG-IgA in pediatric patients without CD, including increased polyclonal IgA production in children with juvenile idiopathic arthritis, an acute immunologic phenomenon in response to infectious disease, and in response to fibrogenesis and apoptosis in patients with end-stage heart failure (25–28). Additionally, recent studies have reported elevated serum tTG-IgA in non-CD patients with EoE (29). The findings of this study support the aforementioned literature. Further investigation into the relationship of TTG-IgA and eosinophilic disorders of the gastrointestinal tract is warranted.

tTG-IgA remains an important laboratory test to screen for CD. High levels of TTG-IgA in a patient who has not yet been diagnosed with CD should warrant further workup with an EGD. CD serologic surveillance with TTG-IgA, with or without EMA-IgA, may not be sufficient to monitor CD status in children with CD and EGID, and further consideration should be made regarding repeat duodenal biopsies to assess mucosal healing.

Limitations of this study include its retrospective nature, relatively small cohort size, and potential confounding variable of concomitant EGID diagnosis. Additionally, TTG-IgA levels were analyzed at several different laboratories, each with unique normal range scales. The study participants were primarily male, possibly due to EoE being more common in males, which is a possible confounding factor (30). Given the data from this retrospective study, additional prospective research is needed to further validate these findings.

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