Proton pump inhibitor-responsive oesophageal eosinophilia: an entity challenging current diagnostic criteria for eosinophilic oesophagitis

Javier Molina-Infante,1 Albert J Bredenoord,2 Edaire Cheng,3 Evan S Dellon,4 Glenn T Furuta,5 Sandeep K Gupta,6 Ikuo Hirano,7 David A Katzka,8 Fouad J Moawad,9 Marc E Rothenberg,10 Alain Schoepfer,11 Stuart J Spechler,12 Ting Wen,10 Alex Straumann,13 Alfredo J Lucendo,14 From the PPI-REE Task Force of the European Society of Eosinophilic Oesophagitis (EUREOS)

ABSTRACT

Consensus diagnostic recommendations to distinguish GORD from eosinophilic oesophagitis (EoE) by response to a trial of proton pump inhibitors (PPIs) unexpectedly uncovered an entity called ‘PPI-responsive oesophageal eosinophilia’ (PPI-REE). PPI-REE refers to patients with clinical and histological features of EoE that remit with PPI treatment. Recent and evolving evidence, mostly from adults, shows that patients with PPI-REE and patients with EoE at baseline are clinically, endoscopically and histologically indistinguishable and have a significant overlap in terms of features of Th2 immune-mediated inflammation and gene expression. Furthermore, PPI therapy restores oesophageal mucosal integrity, reduces Th2 inflammation and reverses the abnormal gene expression signature in patients with PPI-REE, similar to the effects of topical steroids in patients with EoE. Additionally, recent series have reported that patients with EoE responsive to diet/topical steroids may also achieve remission on PPI therapy. This mounting evidence supports the concept that PPI-REE represents a continuum of the same immunological mechanisms that underlie EoE. Accordingly, it seems counterintuitive to differentiate PPI-REE from EoE based on a differential response to PPI therapy when their phenotypic, molecular, mechanistic and therapeutic features cannot be reliably distinguished. For patients with symptoms and histological features of EoE, it is reasonable to consider PPI therapy not as a diagnostic test, but as a therapeutic agent. Due to its safety profile, ease of administration and high response rates (up to 50%), PPI can be considered a first-line treatment before diet and topical steroids. The reasons why some patients with EoE respond to PPI, while others do not, remain to be elucidated.

HISTORICAL BACKGROUND AND DEFINITIONS

Eosinophilic oesophagitis (EoE) and GORD are the most prevalent chronic oesophageal inflammatory conditions in children and adults in the Western world.1 Whereas the first is an allergen-driven disease,2 the latter develops as a consequence of pathological exposure of the oesophageal mucosa to acid-predominant gastric contents.3 Distinguishing both disorders is important because of their different aetopathogenesis; natural history and monitoring.2 However, a rigid distinction between EoE and GORD is difficult due to overlapping clinical and histological features, not to mention their frequent coexistence and potential partially shared pathogenic pathways.3 The presence of heartburn and marked oesophageal eosinophilia, for instance, might be fairly common in both entities.1,4 In paediatric patients, this differentiation is even more complex due to a wider spectrum of clinical manifestations, difficulties in expressing symptoms and subtle or absent endoscopic abnormalities.5

In order to solve this diagnostic conundrum, the first consensus recommendations for diagnosis and management of EoE were published in 2007.6 These guidelines advocated a diagnosis of EoE in patients with symptomatic oesophageal eosinophilia (>15 eosinophils per high power field (eos/HPF)) showing either lack of response to proton pump inhibitor (PPI) therapy or a normal acid exposure on oesophageal pH monitoring. Accordingly, a diagnosis of GORD was recommended for those patients who were either responders to PPI therapy or had objective evidence of pathological oesophageal acid exposure. This distinction was based on the assumption that only GORD, as an acid-related disorder, could respond to the acid-suppressive effect of PPIs. As such, these guidelines equated GORD with symptomatic and histological response to PPI therapy. Far from fulfilling the expectation of distinguishing GORD from EoE, the recommended PPI trial unexpectedly uncovered a third intriguing category of patients apparently sharing features of EoE and GORD.4

Updated consensus recommendations in 20115 included changes to these findings: (1) the description of a novel phenotype, PPI-responsive oesophageal eosinophilia (PPI-REE), referring to patients with features of EoE who achieve clinical and histological remission on PPI therapy (2) response to PPI therapy in patients with PPI-REE was not necessarily considered a manifestation of GORD and (3) the retraction of recommending oesophageal pH monitoring as a diagnostic criterion, due to its low accuracy to predict response to PPI.4 Nonetheless, support for a PPI trial was maintained as a diagnostic criterion, since PPI-REE and EoE were still considered separate clinical entities as they showed a different response to the PPI trial.2

At this stage, it is crucial to ascertain the accurate location of PPI-REE within the spectrum between EoE and GORD, the therapeutic mechanisms
leading to responsiveness to PPI therapy in patients with suspected EoE and whether the response to a PPI trial has any validity as a means of excluding EoE.

DIFERENCES AND SIMILARITIES BETWEEN GORD, PPI-REE AND EOE
The need to distinguish among GORD, EoE and PPI-REE in clinical practice, pharmaceutical trials and research studies has led to careful investigations to distinguish these entities. The results of these studies are summarised in table 1.

Symptoms
In adults, the clinical presentations of GORD and EoE are typically distinct.3 Patients with GORD present with heartburn, regurgitation and bitter/sour taste of gastric content. Dysphagia as a dominant symptom is rare in GORD, unless a peptic stricture is present. GORD symptoms are exacerbated after consumption of large meals, rapid eating, acidic foods, alcohol, obesity, tobacco and body position changes. In contrast, adult patients with EoE present predominantly with intermittent dysphagia during consumption of solid foods commonly associated with food impactions. While heartburn and chest pain may be present in EoE, they are characteristically not the dominant complaints reported by adult patients and if present, usually accompany dysphagia. Available studies have identified that demographics, atopic history and clinical manifestations do not reliably discriminate EoE from PPI-REE.47

Endoscopic features
Most patients with GORD have a normal appearance of the oesophageal mucosa on endoscopy, whereas erosive oesophagitis or Barrett’s oesophagus is identified in the minority.6 Endoscopically, nearly all adult patients with EoE demonstrate one or more characteristic features of loss of vascular markings, rings, white exudates, longitudinal furrows, narrow calibre oesophagus and strictures, whereas some children may have a visually normal mucosa.11 12 Reflecting the natural history of oesophageal remodelling, rings and strictures are common in adults but rare findings in children with EoE.3 Typical EoE endoscopic signs are useful in distinguishing GORD from EoE, but not PPI-REE from EoE.4 7 8 10 11

Histological findings
Histological characteristics of GORD include basal cell hyperplasia, papillary elongation, dilated intracellular spaces and a paucity of intraepithelial inflammatory cells.13 Eosinophils may be present in GORD but typically are in low numbers (<10 eos/HPF), although we lack prospective studies defining numbers and extent and numbers of eosinophils observed in GORD. Histological features of EoE include all of the above GORD features with the addition of a marked, eosinophil-predominant, cellular infiltration of the mucosa. Superficial squamous epithelial distribution, eosinophil degranulation, eosinophil microabscesses and lamina propria fibrosis are also commonly identified in EoE, but not in GORD. Mast cells have been recognised in the mucosa of both patients with GORD and patients with EoE.14 15 Multiple studies have noted that these histological features are found in both EoE and PPI-REE. These include evidence of superficial distribution of epithelial eosinophils, eosinophil degranulation and microabscess formation.4 7 8 11

Table 1 Updated similarities and differences between GORD, PPI-REE and EoE

<table>
<thead>
<tr>
<th></th>
<th>GORD</th>
<th>PPI-REE</th>
<th>EoE</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Age</strong></td>
<td>Adults &gt; children</td>
<td>Children and young adults</td>
<td>Children and young adults</td>
</tr>
<tr>
<td><strong>Gender</strong></td>
<td>Male = Female</td>
<td>Male predominance</td>
<td>Male predominance</td>
</tr>
<tr>
<td><strong>Dominant symptom</strong></td>
<td>Heartburn, regurgitation</td>
<td>Dysphagia</td>
<td>Dysphagia</td>
</tr>
<tr>
<td><strong>Food impaction</strong></td>
<td>Uncommon</td>
<td>Common</td>
<td>Common</td>
</tr>
<tr>
<td><strong>Endoscopic findings</strong></td>
<td>Normal endoscopy (70–80%)</td>
<td>Normal endoscopy (&lt;10%)</td>
<td>Normal endoscopy (&lt;10%)</td>
</tr>
<tr>
<td></td>
<td>Erosions, ulcers, strictures</td>
<td>Oedema, rings, exudates, furrows, strictures, crêpe-paper oesophagus, narrow calibre oesophagus</td>
<td>Oedema, rings, exudates. furrows, strictures, crêpe-paper oesophagus, narrow calibre oesophagus</td>
</tr>
<tr>
<td></td>
<td>Barrett’s oesophagus</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>oesophageal adenocarcinoma</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Histology and inflammatory</strong></td>
<td>Usually &lt;5–10 eos/HPF</td>
<td>&gt;15 eos/HPF</td>
<td>&gt;15 eos/HPF</td>
</tr>
<tr>
<td><strong>cells</strong></td>
<td>Neutrophils, lymphocytes, low-grade eosinophilia</td>
<td>Eosinophils and mast cells</td>
<td>Eosinophils and mast cells</td>
</tr>
<tr>
<td><strong>Oesophageal acid exposure on</strong></td>
<td>Mostly positive</td>
<td>Positive and negative</td>
<td>Negative and positive</td>
</tr>
<tr>
<td><strong>pH monitoring</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Primary treatment</strong></td>
<td>Inhibitors of gastric acid secretion, including PPIs, surgical fundoplication</td>
<td>PPI therapy, unclear whether other inhibitors of gastric acid secretion are effective</td>
<td>Topical steroids</td>
</tr>
<tr>
<td><strong>Aetiology</strong></td>
<td>Reflux of gastric contents</td>
<td>Unclear</td>
<td>Food/airborne allergens</td>
</tr>
<tr>
<td><strong>Type of immune response/involved chemo/cytokines</strong></td>
<td>Th1</td>
<td>Th2</td>
<td>Th2</td>
</tr>
<tr>
<td></td>
<td>IL-8, MCP-1, RANTES</td>
<td>Eotaxin-3, IL-5, IL-13</td>
<td>Eotaxin-3, IL-5, IL-13</td>
</tr>
<tr>
<td><strong>EoE transcriptome panel</strong></td>
<td>Not expressed</td>
<td>Similar expression to EoE</td>
<td>Similar expression to PPI-REE</td>
</tr>
<tr>
<td><strong>Specific molecular effect of</strong></td>
<td></td>
<td>PPIs downregulate Th2 inflammation and normalise EoE gene expression</td>
<td>Topical steroids downregulate Th2 inflammation and normalise EoE gene expression</td>
</tr>
<tr>
<td><strong>therapy</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

EoE, eosinophilic oesophagitis; IL, interleukin; MCP-1, monocyte chemotactic protein-1; PPI, proton pump inhibitor; REE, responsive oesophageal eosinophilia.
basophil infiltration\(^{10}\) and the expression of major basic protein and tryptase.\(^{15}\) Interestingly, a lower rate of response to PPI therapy has been reported in patients with more severe histological findings, including either \(\geq 15\) eos/HPF at three levels of biopsies\(^{16}\) or increasing degrees of oesophageal eosinophilia.\(^{9}\)

**Molecular and genetic features**

GORD promotes a proinflammatory response characterised by innate immunity with overexpression of cytokines, such as interleukin (IL)-8 (CXCL8), CCL2 (monocyte chemoattractant protein-1) and CCL5 (Regulated on Activation, Normal T Expressed and Secreted (RANTES)).\(^{17}\) These cytokines and chemokines promote active recruitment of neutrophils and lymphocytes and sometimes a mild eosinophilic infiltration, normally \(< 5–10\) eos/HPF. Unlike GORD, EoE is a chronic immunoinflammatory disorder characterised by an aberrant Th2 inflammatory response involving IL-5 and IL-13 and local production of CCL26 (eotaxin-3), a chemokine that specifically attracts eosinophils to the oesophageal mucosa. When activated, the eosinophils cause local tissue damage and recruit and/or activate other effector cells, such as mast cells, which have a role in oesophageal fibrotic remodelling.\(^{18}\) By using whole-genome transcript expression profiling of oesophageal tissue, a molecular EoE diagnostic panel has been recently identified.\(^{19}\) This panel is made of 94 EoE genes and accurately distinguishes patients with EoE from GORD or control subjects.\(^{19}\)

Over the past years, an increasing number of papers have tried to further characterise PPI-REE. Baseline markers of eosinophilic inflammation in oesophageal tissue (eg, eosinophil-derived major basic protein and CCL26) have been shown to be increased in PPI-REE similar to EoE. In addition, the expression of mast cell signature genes (eg, tryptase),\(^{15}\) as well as the expression of genes involved in type 2 (Th2)-associated allergic inflammation (including CCL26, IL-5, IL-13, thymic stromal lymphopoietin (TSLP) and periostin (POSTN))\(^{9} 10 20 21\) have demonstrated largely overlapping patterns between EoE and PPI-REE, although PPI-REE typically has more modest overexpression levels. One of the key findings in the past year is that PPI-REE, unlike GORD, has a transcriptome that nearly completely overlaps with the EoE transcriptome, including the hallmark EoE genes for eosinophil chemotaxis (CCL26), barrier molecules (desmoglein DSG1), tissue remodelling (POSTN) and mast cells (CPA3).\(^{22}\) Overall, these findings suggest PPI-REE and EoE are alike and both associated with allergic inflammation (4). In addition, recent clinical studies have shown that PPI monotherapy in patients with PPI-REE can almost completely reverse the Th2 signature of PPI-REE (CCL26, IL-5, IL-13, POSTN)\(^{9, 20 21}\) and concurrently induce a normalisation of the mast cell genes (CPA3, TPSAB2), Th2 inflammation indicators (TNFAIP6, ALOX15), epithelial barrier genes (DSG1, CDH26, FLG), tissue fibrosis markers (eg, KRT13) and IL-13/IL-4-induced genes (POSTN, MUC4).\(^{22 22 22 23}\) Since these effects are similar to those of topical steroids in patients with EoE,\(^{9}\) these striking data pose the possibility that EoE and PPI-REE represent a common disorder.

Recent genome-wide association studies in EoE have identified two replicated susceptibility loci at 2p23 and 5q22, regions that encode the epithelial gene products CAPN14 and TSLP.\(^{24–26}\) The presence of susceptibility loci was shown to not depend upon response to PPI, reinforcing the idea that oesophageal eosinophilia, independent of PPI stratification, likely shares genetic aetiology.

**THE EFFICACY OF PPI THERAPY IN PATIENTS WITH SUSPECTED EOE**

The evidence for PPIs inducing either clinical or histological disease remission in patients with suspected EoE was initially reported by one case series and three retrospective studies published between 2005 and 2009.\(^{16, 27–29}\) The case series reported clinicohistological response in all three patients,\(^{27}\) whereas a 50–86% clinical response and a 40% histological response were reported in the retrospective cohorts.\(^{16, 28–29}\) In the first large prospective study in adults with clinical, endoscopic and histological features of EoE, an 8-week course of PPI therapy led to complete response in 50% (5/10) of cases.\(^{9}\) Of note, response to PPIs was observed not only in 80% (23/29) of patients with endoscopic evidence of GORD or normal pH monitoring, but also in 33% of those with a normal pH study.\(^{4}\)

Two randomised controlled trials comparing PPIs with topical steroids in patients with an EoE phenotype reported a similar efficacy (33%) for PPI therapy.\(^{30, 31}\) The latter trial demonstrated a response to PPIs in 100% (4/4) and 18% (3/17) of patients with a pathological pH study and normal pH study, respectively.\(^{31}\) In three recent prospective studies, 35–47% of adult and paediatric patients achieved histological remission (defined by < 5 eos/HPF) on PPI therapy.\(^{32 33 34}\) Of note, response to PPI therapy increased up to 50%,\(^{30, 57}\) and 68%\(^{34}\) when histological remission was redefined as < 15 eos/HPF.

A recent systematic review with meta-analysis, including 33 studies with 619 patients with suspected EoE, revealed that PPIs achieved histological remission (defined by < 15 eos/HPF) in 51% (95% CI 42.2% to 58.7%) and symptomatic improvement in 61% (95% CI 48.3% to 72.2%) of cases.\(^{34}\) No significant differences were noted in patients’ age, study design and types of PPIs assessed. However, a trend towards increased efficacy was observed when PPIs were administered twice daily compared with once daily, and among patients with increased oesophageal acid exposure on pH monitoring. Noteworthy, a significant publication bias in favour of studies reporting histological responses to PPI therapy was observed in this meta-analysis.

The sustained efficacy of PPIs in children has been evaluated in two retrospective small series and a recent prospective study, with most patients (11/14, 78.6%) remaining in clinicopathological remission at 1-year follow up while on maintenance PPI therapy.\(^{35–37}\) As for adults, the first long-term follow-up multicentre study including 75 patients with PPI-REE demonstrated that the majority of patients (55/75, 73%) maintained histological remission 1 year after tapering dosage to the lowest effective clinical dose.\(^{35}\) Among relapsers, most regained histological remission with dose escalation, suggesting some patients with PPI-REE continue to require high-dose maintenance PPI. Allergic rhinoconjunctivitis and a CYP2C19 rapid metabolizer genotype predicted long-term relapse, hinting at the influence of pharmacogenomic and environmental factors on the long-term efficacy of PPI therapy.

**POTENTIAL MODE OF ACTION OF PPIS IN EOE**

It is widely appreciated that PPIs block gastric acid secretion, and this antisecretory effect is assumed to underlie their great efficacy in treating GORD. However, PPIs do not prevent the reflux of non-acidic material, and up to 20% of patients with GORD have symptoms that are refractory to PPIs. It is less well known that PPIs have anti-inflammatory actions (independent of antisecretory effects) that also might contribute to healing...
PPIs have antioxidant properties, inhibit immune cell functions, decrease adhesion molecule expression by endothelial cells and reduce inflammatory cytokine expression by epithelial cells. PPIs also have anti-inflammatory effects that might be especially pertinent to the allergen-driven eosinophilia of EoE.

In EoE, eosinophils accumulate in the oesophagus when allergens induce production of Th2 cytokines like IL-4 and IL-13, which stimulate oesophageal secretion of CCL26 (eotaxin-3). Omeprazole, in concentrations achieved in blood with conventional dosing, inhibits Th2 cytokine-stimulated eotaxin-3 secretion in isolated oesophageal epithelial cells by blocking binding of the transcription factor STAT6 to the eotaxin-3 promoter. Lansoprazole exhibits similar actions, suggesting that this inhibition of Th2 cytokine-stimulated eotaxin-3 secretion is a PPI drug class effect. In one study of children with oesophageal eosinophilia, PPI treatment significantly decreased eotaxin-3 protein expression by epithelial cells in the proximal but not distal oesophagus. In three recent studies primarily in adult patients with PPI-REE, PPIs reduced oesophageal expression of eotaxin-3, IL-5 and mast cell density, suggesting that PPIs downregulate Th2-mediated events. Moreover, gene transcriptome analyses of oesophageal biopsies from adult and paediatric patients with PPI-REE have shown a pronounced and specific effect of PPIs on reducing expression of genes related to allergic inflammation. Impaired oesophageal mucosal barrier function, likely mediated by reduced expression of desmoglein-1, is a common feature of EoE, and PPIs have been shown to restore mucosal barrier function and improve desmoglein-1 expression in patients with PPI-REE.

All of the therapeutic effects of PPIs on oesophageal inflammation, gene expression and mucosal integrity in patients with PPI-REE are similar to the responses seen with topical steroid therapy in patients with EoE. Collectively, these data support a trial of PPIs for virtually any patient with oesophageal eosinophilia, regardless of the underlying mechanism. If eosinophilia is caused solely by GORD and is not antigen driven, then PPI antiserective effects can improve eosinophilia by limiting acid reflux. If oesophageal eosinophilia is solely antigen driven, anti-inflammatory PPI effects might improve eosinophilia by attenuating Th2-associated responses. If GORD causes or exacerbates an antigen-driven oesophageal eosinophilia, both the antiserective and anti-inflammatory effects of PPIs might combine to ameliorate the condition. Finally, hypersensitivity to acid in the oesophagus has been reported in patients with EoE. During perfusion of the oesophagus with acid, patients with EoE felt the burning sensation evoked by the acid earlier than those with concomitant reflux or healthy volunteers. This phenomenon might explain why PPI-mediated acid suppression may improve symptoms in some patients with EoE, despite the absence of histological remission on PPI therapy.

PPI-REE: IS IT GORD OR IS IT EOE?

The above-mentioned data all point in the same direction suggesting that PPI-REE and EoE are indistinguishable except that PPIs have a more robust effect on patients with PPI-REE than patients with EoE. Subjects with EoE and PPI-REE have similar symptoms, demographics, endoscopic findings, histology and response to other treatments besides PPIs. Most striking, perhaps, is that the transcriptomes of EoE and PPI-REE largely overlap. Furthermore, recent data reveal that patients with EoE responsive to diet and topical steroid therapy were eventually found to respond to PPI therapy as well, providing further data that an allergic inflammatory cause is important in PPI-REE.

All of these data provide no rational basis to make a distinction between patients with symptomatic oesophageal eosinophilia based on a different response to PPI therapy. At the present time, phenotypic, molecular, mechanistic and therapeutic features cannot reliably distinguish EoE from PPI-REE. As such, the requirement of a distinct name among indistinguishable patients for the subgroup responding to PPIs is questionable. We therefore propose not to include the responsiveness to a given drug as a diagnostic criterion and, consequently, avoiding the term PPI-REE for those subjects who have an EoE phenotype with both histological and clinical responses to PPI therapy. Given all of the above-mentioned arguments, we suggest viewing the PPI trial not as a diagnostic tool for EoE, but rather as a potential therapy in all patients with clinical, endoscopic and histological features suitable for EoE.

REAPPRAISAL OF THE PPI TRIAL AS A DIAGNOSTIC TOOL AND POSITION OF PPIS IN THE TREATMENT OF EOE

Currently, either swallowed topical steroids or dietary elimination are considered an appropriate first-line therapeutic options after the diagnosis of EoE is established. But these modalities have limitations and neither is universally effective. Therefore, it is important to consider where PPIs might fit in the treatment algorithm for EoE. Respecting their favourable safety profile, the simplicity of administration of the compounds and high response rates, PPIs could be considered as first-line therapy for patients with EoE. The use of PPIs would therefore, instead of deciphering which patients do not have EoE, will likely identify a substantial proportion of patients with EoE who achieve remission on PPI therapy and will not need topical steroid or dietary therapy. As with topical steroid use, it is important to note that this represents off-label use of these medications.

PROPOSAL FOR UPDATED DIAGNOSTIC CRITERIA FOR EOE

EOE represents a chronic, immune/antigen-mediated oesophageal disease characterised clinically by symptoms related to oesophageal dysfunction and histologically by eosinophilic predominant inflammation. Eosinophilic inflammation is restricted to the oesophagus and other causes of local and systemic oesophageal eosinophilia should be excluded (box 1). After a diagnosis of EoE, clinical and histological features of EoE may respond in the majority of patients to treatment with PPIs, topical steroids or elimination diets.

UNSOLVED ISSUES

Can we positively state that PPI-REE is EoE?

No, we cannot. EoE is formally defined as an immune/antigen-mediated disease, but we currently lack evidence on the ultimate aetiology of PPI-REE. Solid evidence corroborates it is a Th2-mediated disease with significant molecular overlap with EoE, but we do not know whether this immune response is triggered by reflux-mediated epithelial injury, food/airborne allergens or the combination of both factors.

In addition, a diagnosis of EoE in patients with no clinical or endoscopic features of EoE might be questionable, given the fact we know patients with GORD might also have Th1-mediated oesophageal eosinophilia. However, this subset of patients is likely to represent a minority of adult patients. A recent study performed a thorough subanalysis of 75 patients with PPI-REE on long-term follow-up and 86% of patients had typical clinical and endoscopic features of EoE,
Recent advances in clinical practice

Box 1 Proposal for updated diagnostic criteria for eosinophilic oesophagitis (EoE)

1. Symptoms of oesophageal dysfunction (dysphagia/food impaction in adults; abdominal pain, nausea, reflux-like symptoms, feeding difficulties, growth failure, dysphagia in children)
2. Baseline oesophageal eosinophil-predominant inflammation (characteristically consisting of a peak value of ≥15 eos/HPF) limited to the oesophagus
   - Baseline endoscopy should be preferably performed off proton pump inhibitor (PPI) therapy to better understand the patient profile in case of further response to PPI therapy
   - Other local and systemic causes of oesophageal eosinophilia should be ruled out: eosinophilic gastroenteritis, Crohn’s disease, hyper eosinophilic syndrome, parasites, drug hypersensitivity, achalasia, vasculitis, pemphigoid, connective tissue disorders and graft-versus-host disease
   - Biopsies from the antrum and/or duodenum should be obtained in all children and in adults with GI symptoms or endoscopic abnormalities
   - A diagnosis of EoE in patients based solely on histology, without clinical and endoscopic features compatible with EoE, might be questionable
   - Routine oesophageal pH monitoring is not recommended in the diagnostic work-up of EoE
   - A majority of patients with EoE will achieve symptom response and histological remission (<15 eos/HPF) on PPI, topical steroid or dietary intervention

with only one single patient showing a pure GORD phenotype. The bulk of evidence on PPI-REE comes from adult patients, so we need further prospective studies corroborating these findings in children as well. Based on the high population prevalence of GORD, it is inevitable that many patients with EoE will have coexisting GORD. In such cases or atypical clinical presentations, comprehensive consideration of the clinical criteria listed in Table 1, endoscopic features, ambulatory pH monitoring and responsiveness to PPI therapy may have clinical utility in patient management.

Molecular biomarkers distinguishing EoE and PPI-REE would be helpful to distinguish between both entities. KCNJ2 has been recently identified as the only gene with significant differential expression between PPI-REE and EoE, showing a 72% sensitivity/specificity to predict PPI-REE at baseline. KCNJ2 encodes a potassium channel which is abundant in GI mucosa and localises with the proton pump. Therefore, the authors proposed a potential interaction between this potassium channel and proton pump in the upper GI epithelium to explain PPI-REE. A genome-wide approach currently underway may reveal alternative mechanisms that might differentiate the two entities.

Considerations for paediatric patients

A distinction between EoE and GORD may be especially complex in children, where EoE symptoms tend to overlap more substantially with GORD (feeding difficulties, regurgitation, heart burn) and endoscopic findings are not so prototypical as in adults. Concerns about endoscopic procedures in children often lead to treatment with PPIs before any diagnostic procedures are completed. A symptomatic response to PPIs will lead to most paediatricians considering a diagnosis of GORD, but a diagnosis of PPI-REE might be missed since biopsies were not obtained. Furthermore, a significant dissociation between oesophageal symptoms and inflammation has been reported in EoE, so a clinical response to PPI therapy does not necessarily rule out EoE. Unfortunately, EoE is a clinicohistological entity requiring objective confirmation of histological abnormalities for diagnosis and for remission after therapeutic interventions.

Performing an additional baseline endoscopy off PPI therapy raises concerns for practitioners, parents and patients, but it is critical to remember that normal endoscopic and histological oesophageal features on PPI therapy in children with suspected EoE could create a lack of diagnostic clarity as well as short-term and long-term therapeutic uncertainties. For instance, children with GORD, PPI-REE, functional dyspepsia or recurrent abdominal pain might have similar symptoms (regurgitation, vomiting, abdominal pain), experience a therapeutic-related or a placebo-related response to PPIs and exhibit normal endoscopic and histological features on PPI therapy. Questions of the duration, dose and frequency of PPI treatment will remain unanswered. Overall, reconciling concerns about endoscopic procedures and anaesthesia with the current need of endoscopy for diagnosis and monitoring EoE will continue to be challenging in paediatric patients.

Mechanisms underlying response to PPI therapy

The precise mechanism(s) by which PPIs accomplish their effects on oesophageal eosinophilia in EoE remains unclear. Anti-inflammatory effects of PPIs have been only proven in experimental studies. While omeprazole in vitro is present in the culture media for up to 48 h, the short half-life for PPI drugs (1–2 h active) makes it unclear if a sustained anti-inflammatory effect is maintained in vivo. PPI therapy have recently shown their ability to downregulate Th2 allergic oesophageal inflammation, but it is not certain whether this is a direct (primary anti-inflammatory effect) or indirect (primary acid inhibition leads to secondary inflammation healing) effect.

On the other hand, the role of GORD in PPI-REE is unclear. PPIs can reverse dilation of epithelial intercellular spaces and restore mucosal integrity in patients with GORD and PPI-REE suggests reflux may be the initial trigger in some patients with PPI-REE. This hypothesis might be supported by a greater likelihood of GORD in patients with PPI-REE. However, the demonstration of pathological oesophageal acid exposure in patients with PPI-REE does not prove a causal role for GORD, whereas lack of response to PPIs does not necessarily rule out GORD as a primary trigger for EoE. It will be important to eventually determine if patients with PPI-REE would also respond to other classes of anti-acid drugs such as histamine receptor 2 (H2R) antagonists, as it would be informative of the acid-suppressive effects as a primary driver of the PPI-REE designation. It is important to acknowledge that no complete response of another allergic disease with PPI therapy has been documented so far today.

How do we define response to PPI therapy?

The effect of PPIs in patients with suspected EoE is not an all or none effect, but a gradient varying between no response, some response and near-complete or complete response. It should be emphasised that, currently, a diagnosis of EoE which
and duration recommendations.

Hence, recommendations is poor and thus appropriate. However, evidence supporting the recommendation may play a smaller or larger role in these patients.

Molina-Infante J, 20114 Different cytokines in EoE and GORD when compared with inflammatory response. There are also data that PPIs inhibit cytokine secretion.56 While there does not seem to be a relation between the medication dose and response rate in prospective studies, it is clear that any of the PPI agents can be effective when used at a ‘high daily dose’ (table 2). The first meta-analysis on this issue has recently suggested a non-statistically significant advantage of a twice daily administration, with no differences between drugs or doses.34 Future prospective dose-ranging studies of PPIs in patients with oesophageal eosinophilia would be helpful in providing more definitive dose and duration recommendations.

Natural history and long-term prognosis of responders to PPI therapy

The similarities between PPI-REE and EoE also raise the question of whether oesophageal fibrotic remodeling is present if left unmanaged, or whether PPI therapy can lead to reversal of oesophageal fibrosis in PPI-REE.38 Further studies should address this issue.

Combination therapy: PPIs plus steroid/diet therapy

Another area of speculated use is in combined therapy with steroids, thus potentially and synergistically enhancing an anti-inflammatory response.39

Author affiliations

1Department of Gastroenterology, Hospital San Pedro de Alcantara, Caceres, Spain
2Department of Gastroenterology and Hepatology, Academic Medical Centre, Amsterdam, The Netherlands
3Department of Pediatrics and Internal Medicine, Children’s Health Children’s Medical Center, and the University of Texas Southwestern Medical Center, Dallas, Texas, USA
4Center for Esophageal Diseases and Swallowing, University of North Carolina School of Medicine, Chapel Hill, North Carolina, USA
5Section of Pediatric Gastroenterology, Hepatology and Nutrition, Digestive Health Institute, Children’s Hospital Colorado, Aurora, USA
6Section of Pediatric Gastroenterology, Hepatology and Nutrition, Indiana University School of Medicine, Riley Hospital for Children, Indianapolis, Indiana, USA
7Division of Gastroenterology and Hepatology, Department of Medicine, Northwestern University Feinberg School of Medicine, Chicago, Illinois, USA
8Division of Gastroenterology and Hepatology, Mayo Clinic, Rochester, Minnesota, USA
9Gastroenterology Service, Department of Medicine, Walter Reed National Military Medical Center, Bethesda, Maryland, USA
10Division of Allergy and Immunology, Cincinnati Children’s Hospital Medical Center, Cincinnati, Ohio, USA
11Division of Gastroenterology and Hepatology, Centre Hospitalier Universitaire Vaudois, Lausanne, Switzerland
12Department of Internal Medicine, VA North Texas Health Care System, and the University of Texas Southwestern Medical Center, Dallas, Texas, USA
13Swiss EoE Research Network, Olten, Switzerland
14Department of Gastroenterology, Hospital General de Tomelloso, Tomelloso, Spain

Contributors All authors have equally contributed to drafting of the manuscript and critical revision of the manuscript for important intellectual content. Competing interests AIB has received research funding by AstraZeneca, MMS, Endosim, Nutricia and Shire and received fees for development of educational material and speaker fees from MMS, AstraZeneca, Nutricia, Takeda and Astellas. EC and SJS: The Office of Medical Research, Departments of Veterans Affairs (SJS), the National Institutes of Health (R01-DK63621 to SJS, R01-CA134571 to SJS and K08-DK099383 to EC) and NASPGHAN Foundation/AstraZeneca Award (EC). ESD is consultant for Apatalis, Novartis, Regeneron, Roche. Research funding from Meritage; Miraca; Receptos; Regeneron. Funding—NIH R01 DK101856. SKG is a consultant for Receptos and QOL. IH is a consultant for Shire, Regeneron, Roche and Receptors. He receives grant support from NIH U54 AI117804 (part of the Rare Disease Clinical Research Network (RDCRN), an initiative of the Office of Rare Disease Research (ORDR), NCATS and is funded through collaboration between NCATS, NIAID and NIDDK). JM-I is a consultant for Casen-Recordati. MER is a consultant for Novartis, Genetech, Receptos, Immune Pharmaceuticals, Celsus Therapeutics and NKt Therapeutics and has an equity interest in the latter three. He is an inventor of EoE-related patents owned by Cincinnati Children’s Hospital Medical Center. He has a royalty interest in reslizumab, a drug being developed by Teva Pharmaceuticals. MER’s component to this review was supported in part by NIH R37 AI045898, U19 AI070235, R01 A057803, USA AI177804, the Campaign Urging Research for Eosinophilic Disease (CURED), the Buckeye Foundation and...
Sunshine Charitable Foundation and its supporters, Denise A and David G Bunning. The US4 AI117804 is part of the Rare Disease Clinical Research Network (RDCRN), an initiative of the Office of Rare Disease Research (ORDR), NCATS and is funded through collaboration between NCATS, NIAID and NIDDK, which have collectively resulted in the Consortium of Eosinophilic Gastrointestinal Disease Researchers (CEGIDR). ASC and SST: Swiss National Science Foundation (32003B_160115/1). ASl has consultant contracts with Actelion, Falk, Novartis, Receptos, Regeneron and Roche-Genentech.

Provenance and peer review. Not commissioned; externally peer reviewed.

REFERENCES


