Microbiome and Gastroesophageal Disease: Pathogenesis and Implications for Therapy

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Abstract

There is growing evidence that gastroesophageal disease is influenced by the esophageal microbiome, and that commensal bacteria of the oropharynx, stomach, and colon are thought to have a role in modulating pathogenesis. These emerging hypotheses are based on observed changes in the composition of the esophageal flora, notably, repeated observations: 1. There is an abundance of gram-positive bacteria in the healthy esophagus. are more gram positive prevalent 2. The esophageal bacterial population becomes increasingly gram negative with disease progression. Associated with this shift to a more gram negative prevalence is an increase in the potential for the presence of antigenic lipopolysaccharide (LPS). The immunoreactivity of LPS endotoxin thought to promote susceptibility to inflammation and disease.

The pathogenesis of the more common diseases of the esophagus e.g. gastroesophageal reflux disease (GERD), esophageal dysmotility (achalasia), eosinophilic esophagitis (EoE), Barrett’s esophagus (BE), and esophageal cancer, are well-established. Emerging data suggest however, that these are all characterized by an immune-mediated inflammatory cascade, propagated by a dysbiotic state. Thereby, the ability of the healthy “normative state” to protect against foreign bacteria is compromised. This dysbiosis thereby can create adverse inflammatory or immunoregulatory responses with progression of disease.

In the normal healthy state, the esophageal microbiome is constituted in-part, by a multitude of gram positive bacteria, many of which produce antibacterial peptides called bacteriocins. Bacteriocins are selective and used to maintain population integrity by killing off foreign bacteria. When the “normative biome” is interrupted (e.g. antibiotics, medcations, diet, environmental factors), the constitutional changes may allow a more hospitable imbalance favoring the proliferation of opportunistic pathogens. Therefore it seems rational that defining, perhaps that defining, perhaps cultivating, a protective bacterial community that could help prevent or mitigate inflammatory diseases of the esophagus. Furthermore, in conjunction with evidence demonstrating that some bacteriocins are cytotoxic or antiproliferative toward cancer cell lines, further exploration might provide a rich source of effective peptide-based drug targets.

Therapeutic options targeting the microbiome, including prebiotics, probiotics, antibiotics and bacteriocins, have been studied, albeit the attributable effects on the esophagus for the most part, have been unrecognized by clinicians. This review focuses on the current knowledge of the involvement of the microbiome in esophageal diseases (most notably GERD/Barrett’s esophagus/esophageal cancer) and identifies emerging new concepts for treatment.

Introduction

Gastroesophageal disease is a major source of health and economic cost worldwide, and, in 2015, annual expenditures for esophageal disorders were estimated at $18.1B annually in the United States [1]. Gastroesophageal reflux disease (GERD) is the most common esophageal pathology, with 31% of the United States population reporting heartburn or reflux symptoms within the past week [1,2]. Barrett’s esophagus (BE), a premalignant complication of GERD, has an estimated prevalence of 5.6% within the United States [3]. Esophageal cancer also poses a significant burden of disease, with > 18,000 new diagnoses and > 16,000 deaths estimated in 2020 [4]. Changes in understanding of the pathogenesis of the esophageal disease have contributed to the development of new possible therapeutic options.

There has been a recent meteoric rise in the literature demonstrating the significance of the gut microbiome and dysbiosis (defined as microbial imbalance or maladaptation), in the pathogenesis of gastrointestinal (GI) disease [5]. This article aims to review the current literature for microbiome-
related pathogenesis of gastroesophageal disease and to
discuss disease-mitigation strategies and areas for future
research.

“Normal” gastroesophageal microflora

The esophageal microbiome is shaped by the oral cavity,
oropharynx, and stomach due to migration of oral bacteria to
the esophagus and reflux of gastric microbiota. Recognizably,
this varies considerably from person to person, even in the
apparently healthy population. There have been multiple
attempts to classify the healthy esophageal flora into different
cluster types. One study evaluated healthy and GERD patients
demonstrated two microbiome types, referred to as type I
and type II [6]. Type I microbiome associated with healthy
subjects, consisted primarily of gram-positive microbes,
dominated by those within the Streptococcus genus.
Conversely, type II microbiome had a greater presence of
gram-negative anaerobes/microaerophiles and primarily
correlated with esophagitis (odds ratio, 15.4) and Barrett’s
esophagus (odds ratio, 16.5) [6].

Conceivably, the dysbiosis antedates the inflammation.
In this hypothesis, there is a notable activation of Toll-like
receptors (TLRs) by gram-negative bacterial products
compounds produced by gram-negative bacterial, and a
subsequent propagation of the inflammatory cascade [7]
(Figure 1). Another study further classified esophageal
biomes into three community types or clusters: one
type dominated by *Streptococcus* spp., one by *Prevotella*
spp., and another with an intermediate predominance of
streptococcus, prevotella, haemophilus and rothia spp
[8]. These clusters were associated with a variation in
metabolic function. The streptococcus cluster associated
with pentose phosphate metabolism, the prevotella cluster
was associated with lipopolysaccharide (LPS) production,
and, the intermediate cluster was associated with glycolysis
and short chain fatty acid (SCFA) production [8]. For all three
clusters, progression to reflux-related esophageal disease
was associated with increase in relative abundance of gram-
negative flora. This supports the TLR/inflammatory cascade
hypothesis which is substantiated by the observation that
composition of the esophageal microbiome is closely linked
to the gastric and oral flora. The prevalent genera including
streptococcus, prevotella, haemophilus, fusobacterium, and
neisseria, appear similar in both the stomach and oral cavity,
suggesting that the oronasal compartment is the primary
source of bacteria in the aerodigestive tract [9].

Dysbiosis in disease states

Gastroesophageal reflux disease: It is well recognized
that GERD is an inflammatory disease state affecting
the lower esophagus related to inappropriate transient
relaxation or chronic hypotensisity, of the lower esophageal
sphincter. Retrograde reflux of gastric acid with or without
bile causes symptoms and inflammatory changes associated
with GERD [10]. The most frequent treatment medications
are intended to buffer (antacids) or reduce gastric acid
secretion are intended using proton-pump inhibitors
(PPIs) or histamine-2 receptor antagonists [11]. Untreated
GERD may progress and manifest with complications of
erosive esophagitis, esophageal stricture, BE, or esophageal

Inflammatory pathogenesis of GERD: Histologically, the
mucosal inflammation observed in GERD is classically thought
to be a consequence of direct chemical injury from gastric acid
reflux. The specific factors contributing to epithelial insult
were gastric acid and duodenal bile salts. Recent literature,
has demonstrated that a concomitant immunogenic pathway
exists. It remains unclear however, whether erosive injury or
cytokine-mediated inflammation (via response to antigenic

![Figure 1: Factors influencing pathogenesis in GERD, Barrett’s esophagus, and esophageal adenocarcinoma. Abbreviations: GERD: Gastroesophageal Reflux Disease; GI: Gastrointestinal; LES: Lower Esophageal Sphincter; LPS: Lipopolysaccharide; TLR: Toll-Like Receptor.](https://www.heighpubs.org/hcg)
bacterial LPS) occurs first. Biopsies of tissue from the distal esophagus in GERD patients frequently show submucosal (cytokine-induced) inflammation, but with intact epithelial cells. These findings are inconsistent with initial surface epithelial chemical injury (erosion). Therefore, a non-errosive mechanism is more likely, and would be the result of immunoreactivity to a foreign biomaterial. In this case, the literature implicates a pathway involving LPS, a well-recognized cell wall constituent of gram-negative bacteria. This LPS is critical to maintain the bacterial cell integrity and viability, but also is a potent inducer of inflammatory responses and can directly modulate the immune system and susceptibility to disease [12]. The interaction of bacterial LPS and TLR-4 directs the innate immune system via NF-κB and cytokine cascade initiating the inflammatory response [12].

There are however, also many protective factors against an inflammatory response, including the stratified squamous epithelial barrier, paracellular adhesion and intracellular buffering. Bypassing or overwhelming these protective mechanisms leads to cellular injury and an inflammatory cascade [10]. In vitro and in-vivo exposure of lower esophageal keratinocytes to acidified-bile salts such as those from duodenogastric reflux promotes local cytokine production and migration of lymphocytic cells, primarily T-cells [13]. Progression of exposure leads to inflammation of the mucosa but preservation of the epithelial cell layer, implying that the main insult to the mucosa is deep rather than superficial. This suggests that the pathogenesis of reflux esophagitis is driven more by the *innate immune system* than by chemical insult. A study examining biopsies from patients with GERD before treatment with PPIs confirmed histologic damage (proliferation of T-cells, hyperplasia of basal cells, and papillary elongation) without damage to surface epithelial cells [14].

Various inflammatory mediators produced by the mucosa also contribute to lower esophageal sphincter (LES) relaxation. Production of interleukin (IL)-8, among other induced factors (including transient receptor potential channel vanilloid subfamily member-1, substance P, calcitonin gene related peptide, and platelet activating factor) stimulates migration of immune response cells. These factors also induce further IL activation and subsequent NADPH oxidase production of hydrogen peroxide. This peroxide effect on local smooth muscle leads to LES relaxation [15]. It is also hypothesized that nitric oxide, another downstream result of cytokine-induced inflammation, is responsible for the relaxation of the LES and decreases gastric emptying [16]. The production of IL-8 is also inhibited by PPIs through a mechanism involving nuclear factor kappa-light-chain-enhancer of activated B cells (NF-κB) and activator protein-1 [17]. This may be a contributing factor to the therapeutic effect of PPIs in GERD.

**Role of the microbiome in GERD:** Recognizably, LPS is the most abundant and important cell wall constituent of gram-negative bacteria. It is vital for bacterial cell integrity, viability, and defense against environmental stress [18]. The TLR-4 protein site in humans is the best characterized of several sensing receptors that mediate LPS-induced signal transduction. Following disruption of the epithelial barrier, increased LPS-TLR-4 binding activates production of IL-18, which induces a cascading inflammatory response. This activation of TLR-4 is pivotal for both infectious and noninfectious (e.g. allergic or autoimmune) related inflammation and is a major mechanism for pathogenesis of inflammatory disease states by gram-negative flora [19]. Further TLR-based signaling promotes transcription of pro-inflammatory chemokines, including IL-1, IL-6, IL-8, and tumor necrosis factor-alpha (TNF-α) and mediators such as nitric oxide synthase [20].

Used as a standard for inflammation in models of septic shock, LPS (aka endotoxin) is an intrinsic cell membrane component in gram-negative bacteria. It is a potent pathogen-associated molecular pattern (PAMP) which alerts the host immune system that something non-self is present [21]. This occurs first by interaction with TLR-4, which recognizes the PAMP, and signals cells to produce a series proinflammatory cytokines, in particular, TNF-α and interferon (IFN)-γ [12]. The former induces nitric oxide synthase [22] and the latter affects the sensitivity to LPS [23]. The subsequent release of NO triggers both vasodilation and potentiates response by activated macrophages [24]. In smaller amounts, LPS triggers a localized inflammatory response. When the concentration of LPS is high, as it would be in septic shock, hypotension, thrombosis, and mortality can occur. Additional mechanisms for the recognition of LPS including TLR-2, TRPV1 and 2, and others are being investigated [25].

Chemokines also lead to production of nitric oxide and cyclooxygenase (COX)-2, which promote lower esophageal sphincter relaxation and delayed gastric emptying, respectively [26]. This is exemplified in literature discussing *Campylobacter* spp., specifically *Campylobacter concisus*, which is observed frequently in patients with esophageal pathology [27]. *C. concisus*, a common oral flora, is not often found in the healthy esophagus. However, colonization by the organism was observed at the site of histologic changes due to GERD and BE as well as inflammatory bowel disease [28]. Additionally, there is a strong correlation between *C. concisus* and an increased level of the proinflammatory cytokine IL-18 [27, 29].

The focal effects of bacterial colonization suggest a local bacterial adherence, facilitated by a biofilm formation. Biofilm develops via a structurally organized community of flora that stimulate local microbial secretion of an environmentally protective coating (frequently e.g. mucopolysaccharide, extracellular polysaccharide, dextran and adhesion molecules) [30]. Biofilms have been observed in association with GI disease, most notably in oral and colonic pathology [31, 32]. Biofilm-associated proliferation may present a framework for understanding esophageal pathology although further research regarding the

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**References:**

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2. [https://www.heighpubs.org/hcg](https://www.heighpubs.org/hcg)
Therapeutic regimens, especially PPIs, have been demonstrated to alter both the gastroesophageal as well as colonic microbiomes. The use of PPIs in GERD patients has been demonstrated to affect diversity of gastric, esophageal, and fecal flora, mainly by allowing more proximal organisms to populate more distal areas [34-36]. It is unclear if this is a protective or injurious effect although this could be a contributing factor for increased infection risk and may play a role in the association with fecal microbiome-related disease such as clostridium difficile infection [37]. Reduced diversity, (e.g. chemotherapy, radiation, antibiotics) promotes the proliferation of *Clostridium difficile*. The spores (typical to all *Clostridium spp.*) are able to survive when other bacteria (that would have otherwise inhibited *Clostridium difficile* proliferation) have been eradicated.

It has been theorized that metabolic activity of the colonic biome may further contribute to GERD progression [38]. Colonic breakdown of fermentable oligo-, di-, monosaccharides and polyols (FODMAPs) produces SCFAs, some of which contribute to LES relaxation [39,40]. A common FODMAP, fructooligosaccharides have been demonstrated to increase the frequency of transient lower esophageal sphincter relaxations (TLESRs) and reflux episodes [39]. The mechanism for this may be through stimulation of Peptide YY (PYY) production or GLP-1 production [39,41]. Some SCFAs as well as PYY act to inhibit gastric motility as well as to relax lower esophageal sphincter tone. The result is retention of gastric contents as well as susceptibility to reflux, which has implications for dietary intake and colonic microbiome as well as motility alteration within the scope of GERD [40]. These SCFAs, however, do not all have the same effect (at least on the motility of the guinea pig colon). Butyrate increases frequency of propulsive colonic contraction, whereas propionate and acetate decrease this frequency [42].

**Barrett’s esophagus**

Barrett’s esophagus is an intestinal metaplasia of the distal esophageal epithelium characterized by transition from normal stratified squamous epithelial composition to columnar mucosa [43,44]. Development is thought to be in response to chronic inflammation of the mucosa secondary to gastroesophageal reflux. The associated risks for esophageal adenocarcinoma (EAC) are well recognized [6,26,45].

**Inflammatory pathogenesis of BE:** As seen in the pathogenesis of GERD, expression of proinflammatory cytokines such as IL-1B, IL-6, and IL-8 is linked to the transition to metaplasia seen in BE. A mouse model of BE demonstrated that IL-1B and IL-6 are overexpressed at the squamocolumnar junction of the esophagus and promote inflammation of gastric cardiac stem cells [46]. The inflammation provides a further pathway for progression to dysplasia.

Another mouse model of Barrett’s esophagus has revealed that a high fat diet (HFD) is associated with esophageal dysplasia through alteration of the microbiome [47]. The HFD increases production of IL-8/C-X-C Motif Chemokine Receptor 1 chemokines, known to be upregulated in a pro-inflammatory state, and these stimulate the migration of immature granulocytic cells in the esophagus, promoting local inflammatory responses [47].

With the increased abundance of gram-negative species in the distal esophagus, the related increase in LPS may trigger a pro-inflammatory response through an increased release of chemokines/cytokines. The host’s response to the increased level of LPS (intrinsic to gram-negative organisms) results in NF-κB activation of the epithelial cells [48]. The NF-κB molecular pathway serves as an initial response step to noxious stimuli (chemical, bacterial, and viral) and assumes a role of upregulating inflammation, innate immune responses, adaptive responses, apoptosis inhibition, cell proliferation and differentiation. IL-1B and IL-8 are increased as a result of NF-κB activation and the secreted cytokines create a positive feedback loop eliciting a more robust innate immune response in BE [26].

**Role of the microbiome in BE:** As seen in patients with GERD, patients with BE have a distinct microbiome composition [49]. Type II microbiomes, as described earlier, are associated with progression of GERD to BE [50]. These type II microbiomes demonstrate a decrease in Streptococcus spp. and an increase in gram-negative anaerobes/microaerophiles including Veillonella, Prevotella, Haemophilus, Neisseria, Rothia, Granulicatella, Campylobacter, Porphyromonas, Fusobacterium, and Actinomyces spp [6]. This transition from gram positive to gram negative relative abundance is thought to be associated with the pathogenesis of esophagitis as well as the metaplastic progression to BE [50,51]. Expression of LPS from gram-negative bacteria, and subsequent activation of the TLR-4-NF-κB pathway is associated with expression of IL-8 and COX-2. The levels of both are directly correlated with transition from metaplasia to dysplasia [52]. It is also possible that gastric acid could contribute to conversion from type I to type II microbiome by killing acid-sensitive bacteria in the esophagus [53]. Gram-negative organisms are more acid-resistant, and in many cases highly tolerant of bile, which is toxic for gram-positive organisms [54,55].

Molecular products secreted by these flora or components of the bacterial wall such as LPS interact with TLRs and continue the inflammatory cascade seen in reflux esophagitis, preventing resolution of mucosal changes. During the ongoing inflammatory process, changes in the local flora predispose the local squamous epithelial
tissue towards metaplasia to columnar epithelium. Wild-type murine models given a HFD demonstrated increase in goblet cell prevalence and relative neutrophil presence compared to germ-free models [47]. This suggests that local microbe-epithelium interactions, presumed to be through microbe product-TLR binding, is a possible mechanism for the metaplastic process. Furthermore, there may be a role of the colonic microbiome in this process as an increase of the Firmicutes:Bacteroidetes ratio within the colon is seen in BE [47]. Many firmicutes metabolize FODMAPs and dietary fiber into SCFAs. As previously described, SCFA production is associated with decreased LES tone as well as decreased gastric motility through stimulation of PYY mediated smooth muscle effects [40].

There is considerable overlap between periodontal and esophageal flora, with a similar firmicutes/bacteroidetes ratio [56]. It is presumed that this similarity in composition is mainly from movement of microbes in a distal direction. A recent study found a distinct oral microbiome that is associated with presence of BE [49]. This case-control study analyzed the oral microbiome using a three-taxon model, (lautropia, streptococcus, and unspeciﬁed genus in the order of bacteroidales) which distinguished the microflora of patients with BE from healthy individuals. Albeit a small number of patients, this model of clustering predicted patients with BE with a sensitivity of 97% and speciﬁcity of 88% [49]. The model suggests decreased abundance of Lautropia and increased abundance of streptococcus and enterobacteriaceae in BE patients. The increased presence of Lautropia and Enterobacteriaceae, both gram negative, correlates with the LPS/TLR-4 hypothesis, but increased abundance of Streptococcus spp. suggests that pure gram-positive/negative ratio may provide an incomplete explanation. More broadly, increased firmicutes suggests that SCFA production may also play a role in BE pathogenesis. Larger investigations are needed to further evaluate the role of oral dysbiosis and the diagnostic implications for BE because environmental factors can influence the oral microbiome [57].

Esophageal cancer

Esophageal cancer is growing in incidence in many areas of the world [58,59]. Squamous cell carcinoma and adenocarcinoma comprise the two major histological subtypes [60]. Globally, esophageal squamous cell carcinoma (ESCC) is the most common of the two, making up approximately 88% of esophageal cancers [60,61]. Risk factors for both malignancies overlap, although etiologies and incidences vary [62].

Inflammatory pathogenesis

Adenocarcinoma: The etiology of EAC is associated with longstanding inflammation or mucosal injury, such as that seen in reflux esophagitis. The pathophysiology contributing to this disease state is complex involving an interplay between environmental factors, genetic susceptibilities, and host dynamics. Approximately 80% of cases can be attributed to GERD, cigarette smoking, obesity, and low fruit and vegetable consumption [63]. GERD is one of the strongest risk factors for EAC, correlating strongly with duration and frequency of symptoms. Other risk factors include BE, motor disorders of the esophagus, other malignancies, medications, and environmental exposures [64]. BE is the risk factor that has the strongest association with EAC with estimates that BE progresses to high-grade dysplasia at a rate of 0.5% to 0.9% per year [65]. However, EAC, like the complications of GERD discussed above, can arise without preceding symptoms. As discussed previously, the pathogenesis of GERD follows an inflammatory-mediated cascade rather than through direct surface mucosal chemical injury. Conversely, GERD is not a risk factor for ESCC.

Recent epidemiological studies have observed that the incidence of EAC to be on the rise in the Western world, with a 6-fold increase in the United States alone [66]. The recent rise in incidence in the Western world suggests an environmental etiology at play. Studies have explored factors such as diet, smoking, obesity, H. pylori infection, and antibiotics [53].

Squamous cell carcinoma: Esophageal squamous cell cancer is a complex disease, with many predisposing factors, involving both genetic as well as environmental components. Incidence of the disease is influenced by environmental exposure, but there is regional variation to the nature of exposure. The highest areas of incidence include East Asia, Southeastern Africa, and Southeastern South America [61]. Smoking and tobacco use are the most common risk factors in wealthier populations, while hot-liquid consumption, dietary carcinogens, and poor dentition contribute enough to incidence that poverty itself is a risk factor for pathogenesis [61,67,68]. There is a growing body of literature evaluating genetic susceptibility that may increase carcinogenesis following an environmental trigger [61].

Heavy alcohol consumption is a strong risk factor for ESCC. Acetaldehyde, a direct metabolite of ethanol oxidation, inhibits DNA repair through a variety of mechanisms. Ethanol may also directly induce production of reactive oxidation species and promotes aberrant epigenetic modification, in particular DNA methylation [69]. Abnormal methylation of genes associated with carcinogenesis inhibits expression of tumor-suppressor genes, promotes oncogene transcription, and is a the major proposed mechanism for the direct carcinogenic effect of ethanol [70].

Pathogenesis of ESCC is associated with overexpression of inflammatory mediators. Persistent production of NF-κB and activation of TLR-4 have both been demonstrated to be present in early stage ESCC whereas production/activation decreases with progression to advanced stages [71]. TLR-4, in particular, activates an innate inflammatory response with subsequent activation of an acute to chronic inflammatory cascade [72]. This suggests that the presence of external factors affect the local mucosa-microbe interaction. This leads
to a localized inflammatory reaction. The persistence of this inflammation, coupled with a genetic predisposition, triggers hyperproliferation of squamous tissue and progression to carcinoma.

**Role of the microbiome in esophageal cancer**

There is increasing literature evaluating the relationship between the microbiome and esophageal cancer [73]. The microbiome is altered in precursors to esophageal carcinoma, such as the abnormal type II microbiomes enriched with gram-negative bacteria that are mainly associated with GERD and BE (Table 1). This alteration of the microbiome is potentially involved in carcinogenesis. The microbiota of cancerous esophageal tissue has been characterized to be profoundly affected by the oral microbiome, and periodontopathic species derived from the oral cavity. Oral microbial composition has been associated with the risk of EAC and ESCC [73]. In fact, many of the studies implicate oral bacteria in the etiology of oral, esophageal, gastric and other GI cancers [74]. Due to bacteria migration, the oral and gastric microbiota shape the esophageal microbiome and therefore may contribute to esophageal carcinogenesis [73]. However, there are variations in microbiota even between ESCC and EAC.

**Adenocarcinoma:** The microbiome of EAC has been characterized as similar to the type II microbiome, but individual periodontopathic species derived from the oral microbiome are found to be associated with disease: treponema denticola, streptococcus mitis, and streptococcus anginosus [75]. The latter two are gram positive and this suggests a different pathway involving migration or SCFA metabolism, as described in the BE pathogenesis rather than the previously discussed LPS/dysbiosis pathway. This suggests that periodontitis and inadequate oral hygiene may be associated with increased esophageal cancer risk [76]. In particular, fragments of s. anginosus have been isolated in head and neck carcinomas as well as in early dysplastic changes of esophageal and gastric cancer [77]. This implies that s. anginosus is associated with numerous malignancies of the upper digestive tract. The exact mechanism underlying this process has not been delineated. However, induction of inflammatory cytokines by infection with s. mitis and s. anginosus has been demonstrated [75]. Other periodontal species have been associated with esophageal cancer [73]. For instance, it has been found that the periodontal pathogens tannerella forsythia, veillonella, selenomonas, and treponema spp. are associated with higher risk of EAC [78,79].

Decreased streptococcus prevalence is associated with a lower risk of EAC implying that specific flora are inversely related to malignant transformation [73]. Further evidence of this inverse correlation is provided by lower EAC risk is associated with H. pylori infection [73]. In one study, the microbiome in both normal subjects and EAC was characterized to be more alike than to BE comparisons, with an increased relative abundance of bifidobacteria, bacteroides, fusobacteria, veillonella, staphylococcus and lactobacilli and decreased relative abundance of campylobacter when compared with BE samples [27]. Other protective factors such as bacterial biosynthesis of carotenoids by oral neisseria spp. were also associated with protection against EAC [73].

This suggests that commensal colonization by specific flora may be protective from the metaplastic process by inhibiting proliferation of pro-inflammatory flora as well as through synthesis of vitamins or perhaps by eliciting a generally protective immune response.

**Increased EAC risk has also been associated with depletion of certain bacteria. For instance, depletion of the commensal genus neisseria and the species streptococcus pneumoniae are associated with higher EAC risk [73]. This is corroborated by other studies wherein microbial diversity has also been shown to be decreased [79,80]. It has been postulated that once carcinogenesis has begun, streptococci leave the local environment to invade surrounding tissue [81]. Conceivably, etiology from GERD to BE to EAC is a progression of microorganisms. Each stage opens the door to a new microbiota behavior, for example, migration into tissue and cells. Thus, a “snapshot” assessment and comparison at any one time, may not be all encompassing and truly reflective of the entire progressive changes.

**Toll-like receptors** are a class of proteins that play a key regulatory role within the innate immune system. A potential mechanism by which the microbiome participates in carcinogenesis, is via TLRs [79]. TLRs 1-3, 6, 7, and 9 are significantly upregulated in EAC [81]. Both TLR-4 and

| Table 1: Comparison of parameters of inflammatory pathogenesis and microbiome changes seen in GERD, Barrett’s esophagus, and esophageal cancer. |
|-------------------------------|-----------------------------|
| **Inflammatory Pathogenesis** | **Role of the Microbiome** |
| GERD | Initially esophageal bacterial dysbiosis activates a localized submucosal cytokine inflammatory response leading to subsequent direct chemical damage by acid bile. | Associated with transition from to Type I to Type II microbiome (more gram-negative anaerobes/microaerophiles). |
| Barrett’s Esophagus | Following presence of GERD and continued local expression of interleukins such as IL-1β and IL-6. | Interaction of LPS with TLR-4 activates NF-κB and subsequent cytokine cascade. |
| Esophageal Cancer | EAC: GERD and Barrett’s esophagus increase risk due to ongoing mucosal inflammation. | LPS from gram-negative bacteria activates the TLR-4-NF-κB pathway and COX-2 expression. |
| | ESCC: Environmental factors such as tobacco and alcohol use increase local inflammation and risk of malignant transformation. | Both IL-8 and COX-2 are associated with progression of metaplasia to dysplasia. |
| | EAC: Associated with specific bacteria, carcinogenic role of cytokines via TLR pathways, COX-2 isoform increase. | EAC: Decreased microbial diversity, carcinogenic role of cytokines, environmental factor alterations (e.g. alcohol, tobacco) alter biome and microbe-immune crosstalk. |

**Abbreviations:** COX: Cyclooxygenase; EAC: Esophageal Adenocarcinoma; ESCC: Esophageal Squamous Cell Carcinoma; GERD: Gastroesophageal Reflux Disease; IL: Interleukin; LPS: Lipopolysaccharide; NF-κB: Nuclear Factor Kappa-Light-Chain-Enhancer of Activated B Cells; TLR: Toll-Like Receptor
TLR-5 have also been suggested as potential mediators of the progression from reflux disorders to EAC. This suggests an association between the TLR signaling pathway and the altered microbiome. In tissue biopsies from the esophagus, TLR-4 (whose natural ligand is LPS) expression is significantly increased in EAC and BE when compared to normal esophagus [53]. Further, activation of the TLR-4-NF-κB pathway is evident in reflux disorders and may contribute to malignant transformation [26]. Therefore, with abnormal type II microbiomes, there is an increased relative abundance of gram-negative bacteria, over resultant stimulation of TLR-4. This may initiate a larger may trigger a larger and more carcinogenic inflammatory cascade. Further, expression of the COX-2 isofrom, an LPS-TLR-4-NF-κB pathway downstream gene, is elevated in esophageal carcinomas. It has been found that there is an increase of COX-2 that occurs along the progression from low-grade dysplasia to high-grade dysplasia in the EAC pathway [82]. This implies that the activation of the LPS-TLR-4-NF-κB pathway may contribute to malignant transformation. The theory has been experimentally tested in a murine model wherein the presence of E. coli induced activation of TLRs implicated in EAC [81].

Squamous cell carcinoma: As with esophageal adenocarcinoma, ESCC has been shown to be associated with periodontal pathogens and poor oral hygiene [61,72]. Specifically, the abundance of the periodontal pathogen Porphyromonas gingivalis trended with higher risk of ESCC [83]. In one study, poor oral health was reported as a risk factor for esophageal squamous dysplasia [84]. To this point oral SCC, also associated with poor oral hygiene, has been linked to changes in the oral microbiome ( firmaicutes, streptococcus, actinobacteria, and rothia), which were substantially decreased in relation to normal tissue [85]. Oral SCC has been shown to be accompanied with other squamous cell carcinomas of the digestive tract [86]. It has been suggested that a region of epithelial cells can be affected by carcinogenic alterations [87].

Subjects with ESCC have also been shown to exhibit decreased microbial diversity [72]. Interestingly, this decrease in microbial diversity has been replicated in other anatomical sites of the GI system such as the stomach with gastritis [74], and the colon with colorectal cancer [88]. Gastric microbiota changes have also been associated with ESCC, and Clostridiales and Erysipelotrichales orders have been particularly implicated [89].

In addition to direct carcinogenic effects, environmental factors such as alcohol and tobacco may alter the local microbiome and contribute to carcinogenesis. Consumption of alcohol in patients with oral microflora abundant with oxidizing flora leads to production of acetaldehyde, and subsequent DNA repair inhibition, leading to increased susceptibility to oral squamous cell carcinoma [90,91]. Tobacco use is associated with increased abundance of Streptococcus spp. and yeast capable of metabolizing alcohol to acetaldehyde, as well as inhibiting acetaldehyde breakdown, suggesting that persistence of salivary aldehyde can contribute to esophageal carcinogenesis [92]. Alcohol use is associated with an increased firmicutes: bacteroidetes ratio in murine fecal samples [93]. This increased ratio alters local nutrient metabolism and increases serum LPS levels, and may suggest further means of potentiating pathogenesis.

Alcohol may also influence both local and systemic response to microbe-immune crosstalk. Locally, ethanol inhibits epithelial cell expression of tight junction-associated proteins, zonula occludens-1 and claudin-1, increasing barrier permeability and susceptibility to the LPS-mediated inflammatory response [94]. Systemically, both heavy acute and chronic ethanol consumption may also decrease clearance of LPS from the bloodstream, potentiating a systemic pro-inflammatory effect [95].

**Esophageal dysmotility**

**Inflammatory pathogenesis:** Mucosal inflammation leading to altered neuromotor function and GI dysmotility has been described in esophagitis and ulcerative colitis [96]. In patients with GERD, an increase in cytokines and chemokines such as IL-1B, IL-6, IL-8, IL-10, interferon-γ, monocyte chemoattractant protein-1, and Regulated-upon-Activation, Normal-T-Cell-Expressed-and-Presumably-Secreted (RANTES) are observed [97]. IL-6, a cytokine released as a result of mucosal damage, affects the circular smooth muscle cells in the lower third of the esophagus and ultimately disturbs muscle contractility. Although the exact mechanism of cytokine effects on afferent neural pathways are unknown, it is proposed that increased cytokine production occurs as a consequent response to gastric refluxate exposure or the dysbiosis/LPS evident with GERD. These cytokines, including IL-6, are able to alter normal esophageal contractility [98].

**Role of the microbiome in esophageal dysmotility:** While there are no studies directly evaluating the role of the local microbiome in the pathogenesis of esophageal motility disorders, there are some investigations that have characterized the microbiome in achalasia, Chagas disease and connective tissue connective tissue diseases such as systemic sclerosis [99]. Additionally, a large volume of literature characterizes the effects of local flora on smooth muscle elsewhere along the GI tract, particularly the colon.

Although Chagas disease is a well-recognized consequence of a tropical parasitic infection caused by Trypanosoma cruzi, samples of flora grown from patients with chagas related megaesophagus has demonstrated a predominance of nitrite- and nitrate-reducing bacteria [100]. Further investigation of patients with achalasia and megaesophagus demonstrate an overgrowth of streptococcus spp., many of which may act as nitrite-/nitrate-fermenters [101]. As previously described, type I esophageal microbiomes are typically seen as ‘normal’ flora and consist primarily of Streptococcus spp. It is possible that the decrease in interaction between esophageal and gastric flora in various megaesophagus states leads to
overgrowth of these type I flora, although this has not clearly been characterized. Furthermore, increase in nitrite-/nitrate-fermentation, coupled with increased esophageal retention of food products may contribute to development of squamous cell carcinoma and may explain the increased risk that is seen in these patients [64, 100].

As described previously in relation to GERD and BE, an increase in the fecal firmicutes/bacteroidetes ratio corresponds with an increase in firmicutes that produce SCFAs. These SCFAs and downstream PYY decrease the contraction of gastric smooth muscle [40]. This SCFA-mediated effect following changes in the colonic firmicutes:bacteroidetes ratio has also been demonstrated in cases of decreased colonic motility leading to constipation [102]. While their effect in the terminal ileum appears to be mainly stimulation of peristalsis following ileocolic reflux, effects elsewhere in the GI tract manifest as a decrease in motor tone [103]. More investigation is needed to isolate the effect of SCFA production on esophageal peristalsis.

Connective tissue diseases are associated with gut dysbiosis [104]. Most notably, systemic sclerosis, which can lead to fibrosis of the muscular layers of the GI tract and subsequent dysmotility, most frequently manifests in the esophagus [99]. Multiple studies investigating compositional changes in colonic flora have demonstrated decrease in commensal flora such as bacteroides, clostridium, and faecalibacterium, and an increase in potentially invasive flora, such as fusobacterium and desulfovibrio [105-108]. While the mechanism for dysbiotic contribution to fibrosis still requires more investigation, it is believed that epithelial barrier dysfunction leading to microbial inflammatory cascade activation may play a role [108].

**Eosinophilic esophagitis**

Eosinophilic esophagitis (EoE) is a chronic immune mediated disorder that is characterized endoscopically by fixed esophageal rings, esophageal narrowing, and mucosal friability and diagnosed by biopsy findings of eosinophilic infiltration of the esophageal mucosa [109]. This is frequently associated with atopic disease, most frequently in male patients. Initial treatment once, confirmed not PPI responsive, is with topical steroids and dietary modification to remove potential allergens [109].

**Inflammatory pathogenesis:** EoE is a multifactorial disease, with several genetic components, frequently atopic condition associations, as well as environmental factors that are presumed to affect pathogenesis. The role of these various factors in development of EoE is still being characterized, and there are several explanations for its inflammatory origins. One theory is that repeat allergen exposure in susceptible individuals may contribute to eosinophil-driven inflammation [110]. This is highlighted in some data that demonstrates childhood PPI use with EoE as theoretically decreasing allergen digestion and prolonging exposure [111]. The progression of EoE is believed to follow genetic or environmental disruption of the epithelial barrier exposing the underlying mucosal tissue to local allergens and bacterial products. These products stimulate secretion of IL-1, IL-8, and migration of T-helper 2 (Th2) cells, which produce IL-5 and stimulate recruitment and activation of eosinophils, as well as IL-13, which stimulates downregulation of desmoglein-1, a cell adhesion molecule integral to the maintenance of the epithelial barrier [112,113]. Local activation of eosinophils leads to toxic degranulation that further stimulates local inflammation and transforming-growth factor (TGF) expression, which mediates structural protein deposition and leads to the characteristic endoscopic and histologic fibrotic findings [112,114].

**Role of the microbiome in EoE:** Esophageal mucosal eosinophilia and its associated diseases are characterized by a change in the local microbiome [115]. It has been reported that compared to healthy controls without EoE, the esophageal microbiome of patients with active EoE contained a significantly greater abundance of *Haemophilus spp* [115]. Further, this change was reversed by following standard treatment, in-essence, restoring the bacterial composition to one more typical of GERD and healthy subjects. Additionally, the bacterial load, but not the diversity, was increased in subjects with EoE and GERD relative to healthy controls [115]. However in another study, patients with active EoE had a significantly greater abundance of *Neisseria* and *Corynebacterium spp.* when compared with controls (restored EoE patients, now apparently healthy) [116], *haemophilus* and *neisseria* are both genera within the proteobacteria phylum and are associated with activation of the inflammatory cascade [117]. Composition of more proximal flora may also contribute to pathogenesis of EoE. A study of the salivary microbiome in pediatric patients demonstrated significant differences in bacterial composition in EoE patients compared to non-EoE controls [118]. Specifically, there was an association with *haemophilus spp.* with active disease, similar to the changes seen in esophageal samples. This suggests a close interaction between the flora of both locations, and that characterization of salivary dysbiosis may be a surrogate marker for EoE disease activity [115,118].

**Role of bacteriocins**

Bacteriocins are small peptide molecules that are expressed by bacteria in response to stress. Their primary role is presumed to be inhibition of competitive local flora as well as newly introduced foreign competitors for the same niche. This may explain why most studies of probiotic consumption observe transient effects and lack of long-term colonization. They have been traditionally used with the food industry internationally for preservation [119]. They may also play a larger role in regulation of the microbiota within the human GI tract. These bacterial products are ribosome derived peptides typically classified by the nature
of bactericidal activity (target and potency), amino acid sequence and peptide structure (including post-translational modifications), bacteriocin operon, molecular weight, and charge, and are separated into four classes [120-122].

Bacteriocins are potent antibacterial agents against very specific strains of bacteria. Exceptions do occur, however, and broad-spectrum antibacterial peptides are known, and use is made of them as preservatives for food (e.g. nisins and pediocins). Several bacteriocins have been noted to be exquisitely toxic to human pathogens, both gram positive, (e.g. MRSA) and gram negative, (e.g. listeria monocytogenes) [123,124]. These inhibitory effects tend to add resilience to the established composition of the local flora.

Another potential role of bacteriocins is in their antineoplastic effect. Research studying the effect of nisin Z on head, neck, and squamous cell carcinoma (HNSCC) has found that bacteriocins may trigger apoptosis in vitro, and have been demonstrated to reduce the size of tumor xenografts [125,126]. Studies have also observed that such bacteriocins appeared benign to normal healthy cells at the concentrations that were effective against the targeted cancer cells. However, greater study of the cytotoxic effects of bacteriocins is needed. Their role in inhibition of carcinogenesis more generally is also still developing [119].

**Implications for therapy**

**Prebiotics:** There is increasing evidence that dietary intake has a profound effect on microbiome balance and activity. This effect plays a key role in reducing dysbiosis related induction of inflammatory signaling cascades [127-129]. As the pathogenic role of esophageal dysbiosis microbiota becomes more clear, targeted therapeutic intervention using specific prebiotics appears promising- specifically when focused on enhancing the gram-positive/gram-negative ratio. It may be possible, by selectively promoting the activity of certain organisms that produce metabolites, proteins or peptides, that a local microbiome may be curtailed to a more favorable state. Isoomaltooligosaccharides have been demonstrated to increase the number of gram-positive flora, especially bifidobacterium and lactobacillus spp. Prebiotics, including Maltosyl-isomaltooligosaccharide (MIMO), are aimed at improving gram-positive/gram-negative ratio in the esophagus. This type of intervention has demonstrated some potential in reducing or eliminating GERD symptoms in case series [130].

**Probiotics:** The use of probiotics to modify the gut microbiome has been studied in a variety of GI disease states, and there are several investigations into usage to decrease GERD symptoms. Probiotic formulations that include strains mainly within the genera lactobacillus and bifidobacterium have been studied and have demonstrated reduction in symptoms when used as monotherapy [131-134]. However, many studies are limited by quality due to limitations in experimental design such as producing an adequate placebo for control [135]. A randomized-controlled study utilizing bacillus subtilis and enterococcus faecium with PPI demonstrated decrease in diarrhea symptoms and small intestinal bacterial overgrowth but did not reduce GERD symptoms or healing rate compared to PPI alone [136]. Additionally, recent evidence demonstrated that although orally administered probiotics can remain viable, there may be a marked resistance by the host to mucosal colonization [137].

**Antibiotics:** Antibiotics are also a possible therapeutic option for modifying the microbiome, given their efficacy in the treatment of GI infectious diseases. While routine use of antibiotics for the treatment of esophageal disease has not been investigated, it has been used with success elsewhere in the GI tract for dysbiosis related disease states. In patients with small intestinal bacterial overgrowth (SIBO), a condition characterized by proliferation of commensal flora within the small bowel, antibiotics with poor oral bioavailability such as rifaximin have been used with efficacy, though data on the use of systemic antibiotics is limited [138]. Given the implications of emerging antibiotic resistant pathogens and risk for c. difficile colitis, it seems unlikely that this approach will be viable for esophageal diseases [139,140].

**Bacteriocins:** Another possible avenue for intervention involves application of a bacteriocins, whether directly or indirectly, via selectively providing prebiotics to specific organisms. As previously mentioned, bacteriocins have a potential use through two mechanisms: (1) direct antibiotic effect as well as (2) cytotoxic effect towards neoplastic cells.

While this is an active area of research, and inhibitory isolates that target methicillin-resistant staphylococcus aureus [123] and vancomycin- resistant enterococcus [141]. have been isolated, more research is needed regarding products that target specific organisms as well as subsequent literature to evaluate for efficacy and safety [141].

Bacteria have been found within tumors exhibiting cooperative activity with cancer cells—they can metabolically deactivate chemotherapeutics (gemcitabine) and induce chemotherapeutic resistance in pancreatic cancer [142]. A direct therapeutic bacteriocin that could disable these organisms with minimal cytotoxicity to healthy cells could have great potential as adjuvant therapy.

Bacteriocins have two potential weaknesses as potential therapeutic agents:

1. **Cytotoxic effects on non-target tissues.**
2. **Targeted pathogenic species to become bacteriocin resistant.**

Fortunately, one promising solution involves improvements in targeted drug delivery via encapsulation or attachment of bacteriocins to macromolecule-based, metal, or polymer-based nanoparticles [143]. This has the added potential for modulating or increasing intended effect.
through attachment of adjuvants to the same nanodelivery particles.

**Conclusion**

Dysbiosis of the local flora within the esophagus is associated with progression to various gastrointestinal disease states including GERD, BE, EAC, ESCC, esophageal dysmotility and EoE. Dysbiosis may be caused by multiple factors including diet, lifestyle, antibiotic use, and/or food-borne pathogens and remains to be further studied. At present, prebiotics and perhaps direct bacteriocin therapies, have the most promising new potential for treatment of common diseases of the esophagus.

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