Interaction of *Helicobacter Pylori* with Other Microbiota Species in the Development of Gastric Cancer

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Received date: March 06, 2017; Accepted date: March 27, 2017; Published date: April 03, 2017

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Abstract

Gastric adenocarcinoma is the second-leading cause of cancer-related death in the world. *Helicobacter pylori* is a Gram-negative bacterial species that selectively colonizes gastric epithelium. The resulting chronic infection is the most significant risk factor for gastric adenocarcinoma identified so far. In the host, *H. pylori* interacts with microorganisms, which is likely involved in the development of gastric cancer. To identify *H. pylori*-associated bacteria and explore the contributions of these inter-bacterial events in the process of gastric cancer, we examined the interactions of *H. pylori* with other microbes in the microbiota community that are implicated in the development and progression of gastric cancer. A better understanding of the complex interactions between *H. pylori* and altered microbiota is critical to develop personalized therapies and approaches.

Keywords: *Helicobacter pylori*; Gastric cancer; Microbiota

Introduction

The human gut microbiota comprises approximately 100 trillion microbial entities consisting of numerous bacteria, archaea and viruses [1]. The gastrointestinal microbiota is involved in numerous biological functions, such as energy harvest, regulation of the gut mucosal immune system, detoxification of xenobiotics and carcinogens, and protection against pathogens [2].

In this complex microbial ecosystem, *Helicobacter pylori* is a Gram-negative spiral bacterium that may colonize the gastric mucosa. Indeed, the bacterium is found in more than half of the world population. *H. pylori* is a prominent member of the human gastric microbiota. The microbe resides on the mucus layer of the gastric mucosa and is implicated in chronic gastritis, gastric and duodenal ulcers and even gastric cancer [3]. Here we aim to provide a comprehensive literature review on the roles of *H. pylori* and its interaction with other gastric microbes in gastric cancer, which may provide potential biomarkers for risk assessment of gastric cancer and new therapeutic targets.

Dysbiosis of Gastric Microbiota in *H. pylori* Associated Gastric Cancer

Gastric cancer is a multifaceted disease associated with different etiologies, genetic changes and phenotypes. The adhesion capacity, enzymes and metabolic activity in the gastrointestinal tract could contribute largely to variation of different strains. A number of gastrointestinal conditions have been reported to be correlated with alterations in the gut microbiota, which result from changes in the life styles and dietary habits. Approximately 1000 species from relatively few phyla have been reported to be present in gut microbiota population [4]. The species are members of the phyla Firmicutes and Bacteroidetes, with very minimal number being representatives of the Proteobacteria, Fusobacteria, Cyanobacteria, Verrucomicrobia and Actinobacteria, among other species [5].

Epidemiologic study showed that gastric cancer was the fourth most prevalent cancer and the second leading cause of cancer-resulted death [6]. *H. pylori* infection is considered the single most important risk factor for gastric cancer [7]. Andersson et al. showed that the stomach displayed a diverse microbiota composition when *H. pylori* is absent or insignificant whereas samples positive for *H. pylori* were predominated by this bacterium [8]. It was higher in *H. pylori*-infected patients compared with those uninfected individuals (P=0.005) [9]. The unweighted principal coordinate analysis showed that the communities of microbiota in gastric cancer were more diversified. Five genera of bacteria with potential carcinogenic activities were enriched in patients with gastric cancer, including Lactobacillus, Escherichia-Shigella, Nitrospirae, Burkholderia fungorum, and unculturable Lachnospiraceae species. Of particular interest, Nitrospirae was present in all patients with gastric cancer but absent in patients with chronic gastritis [9].

Eun et al. examined the microbial composition of gastric mucosa from patients with gastric cancer in different stages of disease and revealed that the relative abundance of
Helicobacteraceae family was significantly lower in gastric cancer patients than those with chronic gastritis and intestinal metaplasia and that the opposite pattern was true for the relative abundance of Streptococcaceae family [10]. Compared with the intestinal metaplasia group, the relative abundances of Epsilonproteo bacteria in the gastric cancer group were significantly decreased. On the contrary, the homogeneity and diversity of gastric microbiota in the gastric cancer group were increased. In addition, two studies showed an overall lower prevalence of *H. pylori* in gastric cancer patients than in that in the control group [11,12]. Examination on the gastric microbiota of 63 antral mucosal and 18 corpus mucosal samples via pyrosequencing of the 16S rRNA gene revealed that the abundance of nitrate-reducing bacteria other than *H. pylori* was two times higher in the cancer groups than in the control groups [13] As such, *H. pylori* is a crucial microbe affecting the clinical manifestations and prognosis of gastric cancer [14].

**Alterations of Gastric Microbiota after *H. pylori* Infection**

Recent studies have revealed significant differences between the microbiota of individuals with and without *H. pylori* infection. For example, interaction between *H. pylori* and the normal gut microbiota was found during early stages of life using germ-free and specific pathogen free mice models [15]. In *H. pylori*-negative individuals, the most abundant phyla are typically Firmicutes, Bacteroidetes, and Actinobacteria [16]. The common phytopotypes found in *H. pylori*-negative individuals include Streptococcus, Prevotella, and Gemella [8]. Consistent with these results, the composition of the microbiota between *H. pylori*-positive subjects and *H. pylori*-negative controls were different in clostridia and the total number of anaerobes [17]. *H. pylori* accounts for >90% of all sequence reads and greatly reduces the overall diversity of the gastric microbiota in *H. pylori*-colonized individuals. Proteobacteria, Firmicutes, and Actinobacteria were the most abundant phyla in *H. pylori*-colonized human stomachs [18]. *H. pylori* infection increased the relative abundance of Proteobacteria, Spirochaetes, and Acidobacteria while decreasing the that of Actinobacteria, Bacteroidetes and Firmicutes (Table 1) [19].

<table>
<thead>
<tr>
<th>Group</th>
<th>Microbiota</th>
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<tr>
<td><em>H. pylori</em>-negative</td>
<td>Firmicutes, Bacteroidetes, and clínobacteria</td>
</tr>
<tr>
<td><em>H. pylori</em>-uninfected</td>
<td>Streptococcus, Prevotella, Gemella</td>
</tr>
<tr>
<td><em>H. pylori</em>-colonized</td>
<td>Proteobacteria, Firmicutes, and Actinobacteria</td>
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Atrophy is one of the key steps in the histologic progression culminating in intestinal-type gastric cancer [19]. In the context of atrophic gastritis, non-molecular analyses have shown that in the absence of *H. pylori*, urease-producing members of the gastric microbiota, such as *Proteus mirabilis*, *Klebsiella pneumonia*, *Staphylococcus aureus*, *Staphylococcus capitis*, and *Micrococcus* species can bloom to levels that cause false positive results in *H. pylori* urea breath tests [20]. Specifically, the microbiota of patients with gastric cancer was dominated by species of *Streptococcus*, *Lactobacillus*, *Veillonella*, and *Prevotella*. Likewise, another study reported that *Lactobacilli* was increased in *H. pylori*-infected patients [21]. In addition, *S. mitis* was found to be significantly enriched in the stomach of atrophic gastritis and gastric cancer patients. Using a co-culture method, *S. mitis* produced and released one or more diffusible factors that directly or indirectly induce coccoid conversion of *H. pylori* cells [22]. Compared to peptic ulcer samples, coccoïd *H. pylori* cells were abundant in gastrectomy specimens from adenocarcinoma [23]. The involvement of coccoïd *H. pylori* in carcinogenesis might due to its stronger effect on proliferation and weaker effect on apoptosis of gastric epithelial cells [24]. Another bacterium commonly present in the stomach, *L. fermentum* could promote survival of *S. mitis* via secreting diffusible factors [25].

From these observations, it is tempting to hypothesize that the interactions between *H. pylori* and other bacteria might play important roles in *H. pylori*-associated diseases, especially gastric cancer.

**Mechanisms of *H. pylori* Interaction with other Microbiota Involved in the Progress of Gastric Cancer**

One critical event of gastric cancer is the rise of pH which may increase abundance of bacteria in the gastric environment. It has been acknowledged that either acute or chronic *H. pylori* infection can induce perturbations of acid secretion in the gastric environment [26]. Changes of gastric pH due to the decrease in acid secretion can encourage colonization of non-*H. pylori* bacterial species from oral mucosa, the upper-respiratory tract, or the intestines that normally cannot persist in the healthy stomach [27]. Persistent infection of the gastric mucosa by *H. pylori* initiates an inflammatory cascade that progress into atrophic gastritis and produces insufficient acid that increases the risk of developing gastric cancer [11,28].

*H. Pylori* uses different mechanisms to inhibit acid secretion. *H. pylori* can indirectly inhibit parietal cell function as a result of changes in cytokines as well as hormone levels, such as interleukin 1β (IL-1β), and tumor necrosis factor-α [29]. CagA-positive *H. pylori* strains have been shown to affect the secretion of several hormones, including 5-HT, ghrelin, dopamine, and gastrin, and altered levels of these hormones might be the cause of the psychological disorders of functional dyspepsia patients [30]. Our study has indicated that *H. pylori* eradication was associated with long-term disturbance in three hormones (active amylin, PP and total PYY) both pre- and post-prandially [31]. Indeed, as the gastric acidity reduced in late stages of chronic *H. pylori* infection, other bacterial species appeared to outgrow *H. pylori* [32]. Thus, the beneficial effects of antibiotic eradication regimens could be explained in part by the effects of interbacterial competition. The gastrointestinal tract is a complex and dynamic network that includes interactions between intestinal mucosal cells, the immune system, food particles, and the commensal microbiota. An intricate and mutualistic symbiosis modulates relationship between the host...
and the gut microbiota. The interaction of *H. pylori* and the gut microbiota needs to be further studied.

The risk of developing gastric cancer is associated with *H. pylori* strains, variations in host responses governed by genetic diversity, and/or specific interactions between host, microbial, and environmental determinants [20]. It has been shown that the carcinogenic effects of *H. pylori* infection have been mainly linked to the cag pathogenicity island (cag PAI) and the vacuolating cytotoxin gene A (vacA) [33]. The cytotoxin-associated gene A (cagA) is the most investigated gene of the cag PAI. CagA protein, encoded by cagA, could be delivered into mammalian cells and affect cytoskeletal and tissue structure, as well as cell proliferation. Infection with cagA-positive *H. pylori* strains is associated with high risk of peptic ulcers and gastric carcinoma. *H. pylori* strains that harbor the cag PAI (Cag+ strains) are associated with a significantly increased risk of gastric cancer compared to cag−strains. Non-phosphorylated CagA also exerts effects within host cells that affect oncogenesis. Following translocation, phospho-CagA by Src and Abl kinases subsequently interacts with and activates several host cell proteins, result in morphological alterations such as cell scattering and elongation.

CagA can activate the transcription factor NF-1 and stimulate expression of IL-8, leading to neutrophil infiltration into the gastric mucosa [34]. In addition, CagA also induces DNA damage in vitro and in rodent models of infection which have been validated in human subjects colonized with *H. pylori* cag+ strains. Cag+ strains contract with host cells and thereby activate multiple signaling pathways that may increase the risk for malignant transformation during prolonged colonization. *H. pylori* CagA induce TWIST1 expression and EMT in gastric cancer cells by regulating PDCD4 [35].

VacA is another *H. pylori* component linked to the development of gastric cancer. The toxin is secreted by *H. pylori* as an 88 kDa monomer (p88) consisting of two domains (p33 and p55) [36]. Infection of epithelial target cells by *H. pylori* has been demonstrated to be associated with both increased and reduced levels of apoptosis [37]. VacA causes multiple alterations in human cells, including autophagy, vacuolation, altered plasma and mitochondrial membrane permeability and apoptotic cell death. All *H. pylori* strains possess vacA and produced marked variation in the 5' region of VacA sequences, which encodes the signal. VacA and CagA can mutually regulate each other to affect host cell responses [38]. CagA antagonizes VacA-induced apoptosis, which activates the survival pathway mediated by the MAPK ERK [39]. CagA also activates EGFR signaling inhibited by VacA [40]. Recently, it was shown that the opposing effects of CagA and VacA may be cell-lineage specific.

A growing body of evidence has suggested that inflammation is a critical component of tumor progression in stomach. For example, many cancers arise from sites of infection, chronic irritation and inflammation [41]. *H. pylori* induced chronic inflammation is believed to be one of the mechanisms in the development of gastric cancer. It is revealed that colonization with both *H. pylori* and complex intestinal flora resulted in a greater inflammatory response, manifested by higher proinflammatory gene expression levels compared with those treated with *H. pylori* alone or with restricted Altered Schaedler’s Flora [42]. Various studies have demonstrated that the interaction between *H. pylori* and the microbial community in the stomach may stimulate inflammatory signals, which contribute to the development and progression of cancer. Overgrowth of some bacterial species may enhance inflammatory responses accelerating atrophy, metaplasia, and cancer [43]. The alterations of gastric environment, hormones levels, immunity and inflammatory response might attribute to *H. pylori*-associated gastric cancer.

Production of IL-1, which may inhibit gastric acid secretion, is increased in the gastric mucosa of infected individuals in comparison with uninfected controls [20]. Polymorphisms within the IL-1 gene cluster that increase IL-1 production are associated with significant increases in risk for hypochlorhydria, gastric atrophy, and distal gastric adenocarcinoma compared with low-expression IL-1 genotypes [44]. Virulent strains of *H. pylori* in a genetically susceptible person further increase the risk for gastric cancer. High-expression IL-1 alleles infected with *H. pylori* cagA+ or vacA-type strains increases in risk for gastric cancer compared to uninfected persons at 25-fold or 87-fold [20].

Diet may also be a factor influencing gastric cancer development. Iron depletion accelerates the development of gastric dysplasia and cancer by promoting assembly and activity of the *H. pylori* cagPAI [45]. *H. pylori*-induced expression of TLR-2 and TLR-5 can qualitatively shift cag PAI-ependent to cag PAI-independent pro-inflammatory signaling pathways with possible impact on the outcome of *H. pylori*-associated diseases [29]. *H. pylori* predominantly activates the NLRP3 inflammasome through a process that requires TLR2-dependent licensing. *H. pylori* can activate the inflammasome and caspase-1 in antigen-presenting and other cells, in result to the processing and release of caspase-1-dependent cytokines [46]. It is reported that activation of the TLR2/NLRP3/caspase-1 axis is required for acid adaptation (Figure 1).

**Figure 1** Network of *H. pylori* interact with other factors.

**Conclusions**

Gastric cancer is still a major challenge worldwide; because detection is frequently made only at an advanced stage, mortality remains high. Gastric cancer prevention programmes via *H. pylori* eradication have been shown to benefit high-risk patients.
The best results from gastric cancer prevention strategies are obtained when \textit{H. pylori} eradication is performed before advanced atrophic gastritis with changes becomes established and thus implementation of \textit{H. pylori} screening and treatment in early adulthood is required. \textit{H. pylori} infection could decrease acid secretion by the production of ammonia and bicarbonate from urea. Additionally, these two products could serve as substrates for other bacteria and the gastric microbiota changed [35].

In contrast to studies comparing the composition of the gastric microbiota between \textit{H. pylori}-infected and -uninfected subjects, few studies have examined differences in microbial composition and outcomes from diseases such as gastric cancer [48]. The composition of the gastric microbiota in well-characterized human populations with and without gastric cancer, are needed further clearly studied. Simultaneously, the alteration of gastric microbiota during \textit{H. pylori} infection might in turn exert a dramatic effect on \textit{H. pylori} itself and thus promote carcinogenesis. This will help to find more personalized therapies and potential approaches to treat gastric cancer.

It is reported that \textit{H. pylori} eradication from the residual gastric mucosa after the endoscopic treatment of early gastric cancer may result in an inhibitory effect on the occurrence of metachronous gastric cancer [49]. Triple therapy (proton-pump inhibitors+amoxicillin+clarithromycin) is the first-line regimen for the eradication of \textit{H. pylori}. Quadruple therapy (proton-pump inhibitors+bismuth subcitrate potassium+metronidazole +tetracycline) remains the best second-choice treatment. Conversely, several meta-analyses have shown that the progression of atrophic gastritis and intestinal metaplasia to gastric cancer can indeed occur following \textit{H. pylori} eradication, suggesting that other factors contribute to the progression of pre-neoplastic lesions. Thus, the actual role of \textit{H. pylori} eradication in the prevention of gastric cancer is still a matter of wide debate.

In conclusion, it can be seen that various stages of the development of gastric cancer involve the progression from \textit{H. pylori} superficial gastritis to atrophic gastritis and then the further progression to development of dysplasia and cancer. Numerous co-factors are involved in promoting or inhibiting this progression. However, \textit{H. pylori} infection does appear to be an obligatory co-factor in most cases of non-cardia gastric cancer. Consequently, prevention of \textit{H. pylori} infection or its eradication at an early stage should markedly reduce the incidence of this common and usually fatal cancer.

References


