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1. Introduction

*Helicobacter pylori* (*H. pylori*) is a gram negative, microaerophilic spiral bacterium with a clear role in the pathogenesis of gastric and duodenal ulcer, low grade B cell gastric lymphoma (MALT lymphoma) and gastric cancer. The bacterium was successfully cultivated by Warren and Marshall in 1982. Their discovery led to completely different therapeutic approach to patients with peptic ulcer disease and gastric MALT lymphoma. Despite initial skepticism concerning *H. pylori* role in gastric carcinogenesis, in 1991 two epidemiological studies [1,2] confirmed previously published reports suggesting higher incidence of gastric cancer in *H. pylori* infected individuals [3-5]. Furthermore, cohort studies in California, Hawaii and Great Britain confirmed increased risk for gastric cancer in *H. pylori* infected individuals. As a result of accumulated scientific evidence *Helicobacter pylori* was marked as human carcinogen by International Agency for Research on Cancer in 1994. Gastric cancer is the third most common cancer among males and fifth most common among females. The incidence of gastric cancer is declining in developed countries, but the global burden is rising due to cases occurring in developing countries. The five-year survival for advanced stage gastric cancer is below 20% even in developed countries [6].

Different outcomes of *Helicobacter pylori* infection are to be expected depending on distinct patterns of gastritis that were identified and described in detail. Namely, antrum-predominant gastritis leads to duodenal ulcer formation while chronic corpus predominant and multifocal atrophic gastritis lead to increased risk for gastric cancer formation [7-9].
The outcome of *Helicobacter pylori* infection depends on characteristics of the microorganism, characteristics of the host and environmental factors.

2. *Helicobacter pylori* virulence factors

*Helicobacter pylori* virulence depends on different factors enabling it to colonize gastric mucosa and induce tissue damage. Epidemiological studies revealed so far six distinct *H. pylori* strains (hpEurope, hpEastAsia, hpAsia2, hpAfrica 1, hpAfrica2 and hpNEAfrica) that are related to geographic regions and correlate well with the incidence of gastric cancer [10].

*H. pylori* is a highly heterogeneous bacterium [11,12]. Several *H. pylori* virulence factors are thought to contribute to gastric cancer development and this review will focus on cytotoxin-associated gene A and CagA protein (CagA), vacuolating cytotoxin (VacA) and outer inflammatory protein (OipA) (Figure 1) with a brief comment on possible role of duodenal ulcer-promoting gene (dupA).

![Figure 1. Mechanism of Helicobacter pylori induced host cell injury](image)
2.1. cagA

There are two types of clinical *H. pylori* isolate: CagA-producing (cagA positive) strains and CagA-nonproducing (cagA negative) strains. Results of studies obtained on animal model revealed that gastric cancer developed only in animals infected with cagA positive *H. pylori* strains or when CagA protein was artificially introduced into the host [13,14] and authors concluded that CagA protein was important factor in gastric carcinogenesis. Further investigations revealed that cagA gene polymorphisms could at least partly explain why only some cagA positive individuals develop gastric cancer. Namely, there are different number of repeat sequences located on 3’ region of the cagA gene in different *H. pylori* strains. It is now well known that sequences of these repeat regions differ significantly between strains isolated in East Asia and Western strains [15]. Each repeat region of the CagA protein contains Glu-Pro-Ile-Tyr-Ala (EPIYA) motifs. The first repeat region is termed EPIYA-A and EPIYA-B and second repeat region EPIYA-C or EPIYA-D segments [11]. Western type CagA has EPIYA ABC, ABCC or ABCCCC, while East Asian CagA has EPIYA ABD segments [11].

CagA protein is composed of a disordered C-terminal region that contains the EPIYA motifs and a structured N-terminal region with several conserved regions. When the bacterium contacts the host cell CagA is injected into the host cell via the cag pathogenicity island (cagPAI)-encoded type IV secretion system (T4SS). Upon injection, CagA is linked to the inner leaflet of the cell membrane probably via an electrostatic interaction with phosphatidylserine [16]. When injected into the cytoplasm via the T4SS, CagA can be phosphorylated by the host and alter host cell signaling in both phosphorylation-dependent and phosphorylation-independent manner. CagA is phosphorylated on EPIYA motifs [17]. Induction of heme oxygenase 1, which exhibits anti-inflammatory and antioxidant effects, reduced CagA phosphorylation during *H. pylori* infection of gastric epithelial cells *in vitro*. However, there is data suggesting that the bacterium developed a strategy to diminish heme oxygenase 1 gene expression in gastric epithelial cells [18].

Recently it was demonstrated that another protein component of T4SS termed CagL induces hypergastrinemia, which is a major risk factor for the development of gastric adenocarcinoma [17].

2.2. vacA

VacA induces vacuolation of the host cell, membrane-channel formation, cytochrome c release from mitochondria leading to apoptosis, induces autophagy and alters host immune response [17,20,21]. VacA also inhibits T-cell activation and proliferation [22]. All *H. pylori* strains have a functional vacA gene. There is a variation in the vacuolating activity among different *H. pylori* strains [23] related to differences in the vacA gene structure at the signal (s1 and s2) region, middle region (m1 and m2) and intermediate (i1 and i2) region. Combination of subtype of these three regions influences the levels of VacA activity and is related to risks for different gastrointestinal diseases.

The most cytotoxic are s1/m1 strains, followed by s1/m2 strains, whereas s2/m2 strains have no cytotoxic activity and s2/m1 strains are rare [24]. Individuals infected with s1 or m1 *H. pylori* strains have an increased risk of peptic ulcer or gastric cancer compared with individuals...
infected with s2 or m2 strains [11]. In East Asia most H. pylori strains are s1 type that led to conclusion (based on epidemiological data) that in East Asia presence of m1 region leads to increased risk for gastric cancer [11] and its typing is the best marker for gastric cancer risk out of all vacA regions.

Intermediate region of vacA is identified between s and m regions few years ago. All s1/m1 strains belong to the type i1, and all s2/m2 strains are type i2. Strains that contain s1/m2 can be type i1 or i2. Strains with i1 region are more pathogenic. In some populations typing of i-region is superior in predicting gastric cancer risk than typing of s region, while in other populations it has no predictive value [25].

The deletion (d) region—was identified between the i-region and the m region [26]. The d region is divided into d1 and d2. The study of Western strains demonstrated that d1 was a risk factor for gastric mucosal atrophy; however, almost all East Asian strains are classified as s1/i1/d1 [11].

2.3. oipA

Outer membrane proteins (OMPs) are coded by different genes in Helicobacter pylori genome. One of the OMPs that can function as adhesin is OipA, which was identified in 2000 [11]. OipA is a protein that induces proinflammatory response and its activity leads to increase in mucosal interleukin-8 (IL8) levels. OipA is involved in the attachment of H. pylori to gastric epithelial cells in vitro [27]. Results from animal studies demonstrated that OipA alone plays a role in the development of gastric cancer [13].

The production of CagA, VacA and OipA is linked and the majority of H. pylori strains produce either all of these proteins or none of them. Almost all East Asian strains of H. pylori are classified as CagA-producing, VacA-producing (vacA s1), and OipA-producing strains and are highly pathogenic. In addition, CagA, VacA and OipA are all thought to be involved in the development of both gastric cancer and duodenal ulcer [11].

2.4. dupA

Duodenal ulcer–promoting gene dupA was described in 2005. It is localized in the plasticity zone and it is involved in T4SS formation. Initial report suggested it to be the first disease-specific H. pylori virulence factor that induced duodenal ulcer formation and had a suppressive action on gastric cancer (DupA) [28]. After this report multiple studies failed to demonstrate correlation between dupA gene and specific gastroduodenal disease [10,29,30]. This gene is, however, highly polymorphic, and that could explain the conflicting data obtained in different studies [17].

2.5. Geographic differences in gastric cancer incidence related to Helicobacter pylori strains

Multilocus sequence typing (MLST) of the housekeeping genes revealed six H. pylori strains (hpEurope, hpEastAsia, hpAsia2, hpAfrica 1, hpAfrica2 and hpNEAfrica) that are related to geographic regions and correlate well with the incidence of gastric cancer.
Sequence differences among the examined housekeeping genes of the six major genotypes probably have no influence on the disease outcome but serve as a marker for other virulence factors related to the disease outcome (i.e. *cagA* and *vacA*) [10]. In 2003, Falush et al analyzed 370 *H. pylori* isolates and assigned the strains to four main clusters: hpEurope, hpEastAsia, hpAfrica1 and hpAfrica2 due to their obvious geographical associations [31]. HpEurope is common in Europe and countries colonized by Europeans and most isolates from East Asia belong to hpEastAsia. HpAfrica2 is very distinct and has only been isolated in South Africa. Four years later Linz et al expanded the analysis using 769 *H. pylori* isolates and assigned the isolates to six distinct groups, adding to previously described clusters (hpEurope, hpEastAsia, hpAfrica1 and hpAfrica2) two new clusters termed hpAsia2 and hpNEAfrica. Cluster hpAsia2 was isolated in South and Southeast Asia. Cluster hpNEAfrica is predominant among isolates from Northeast Africa [12].

Populations with high rates of gastric cancer correspond with regions presenting hpEastAsia strains. In contrast, incidence of gastric cancer is very low in Africa, where most strains are hpNEAfrica, hpAfrica1 or hpAfrica2, and in South Asia, where most strains are hpAsia2. The differences in *H. pylori* isolates in different populations are considered to be an explanation of the both African and Asian enigma. Namely, populations with high incidences of *Helicobacter pylori* infection in East Asian countries have high incidences of gastric cancer, as opposed to low incidence of gastric cancer in other highly infected populations in Africa (African enigma) and South Asia (Asian enigma) [10].

3. Host factors

Host characteristics and immune response to *Helicobacter pylori* infection also play a role in gastric carcinogenesis. Different gene polymorphisms that affect host immune response and extent of cell proliferation are described and linked to gastric carcinogenesis together with gene polymorphisms for growth factors and growth factor receptors. Majority of these polymorphisms are single nucleotide polymorphisms (SNP). In recent years, host genetic polymorphisms involved in inflammatory response, carcinogen metabolism, antioxidant protection, mucosal protection and cell proliferation regulation have been widely studied as potential biomarkers to predict gastric cancer risk.

4. Cytokine gene polymorphisms

Cytokines modulate inflammatory response to *Helicobacter pylori* infection and indirectly the risk for gastric cancer. Gene polymorphisms for *interleukin-1* beta and its receptor antagonist, tumor necrosis factor alpha, interleukin-8 and interleukin-10 were extensively studied and reported as relevant in determining gastric cancer risk.
4.1. Interleukin-1

Interleukin-1 beta (IL-1β) is a potent inhibitor of gastric acid secretion. Reduced gastric acid secretion on the other hand promotes development of *H. pylori* induced pangastritis, gastric mucosa atrophy and subsequently, in a subset of individuals, gastric cancer development. El-Omar et al described in 2000 polymorphisms in the pro-inflammatory IL-1B gene (encoding IL-1β) and IL-IRN (encoding IL-1β receptor antagonist) associated with elevated risk for hypochlorhydria and gastric cancer in the persons with *H. pylori* infection [32] (El-Omar). Presence of IL-1B-31C or -511T and IL-IRN*2/*2 polymorphisms is associated with a 2-3-fold increase in risk for intestinal and diffuse non-cardia gastric cancer among *H. pylori*-infected persons [33]. These genetic polymorphisms modulate gastric cancer risk by increasing expression of pro-inflammatory cytokine IL-1β. It was later established that these polymorphisms interact synergistically with bacterial virulence factors (*cagA* positive, *vacA*s1 and *vacA*m1). Gastric cancer risk is highest among those with both host and bacterial high-risk genotypes [34].

Results of meta-analysis provided by Persson et al. based on available data from epidemiological studies showed strongest association of IL1RN2 polymorphism with increased risk for gastric cancer [35] in non-Asian populations for both intestinal and diffuse cancers. The IL1RN22 genotype has been reported to cause high circulating IL-1 receptor antagonist and IL-1β levels resulting in a severe and prolonged inflammatory response. IL1B-511T carriers were found to have an increased risk of gastric cancer in non-Asian populations. Possible explanation is that stronger inflammatory reaction may increase the risk of cancer through damage to gastric cells and bacterial overgrowth and accumulation of toxic byproducts [32]. These findings are supported by results of some [36,37], but not all previously published meta-analysis [38]. Surprisingly, according to Persson et al IL-1B-31C polymorphism was associated with a reduced overall risk for gastric cancer in Asian populations [35].

4.2. Tumor necrosis factor alpha

Tumor necrosis factor alpha (TNF-α) is another proinflammatory cytokine produced in gastric mucosa in response to *H. pylori* infection. Polymorphisms in TNF-A gene, especially presence of the high producing A allele of TNF-A at position 308 (G→A) is considered to be associated with increased risk for non-cardia gastric cancer [6]. This finding is supported by majority [39-41], but not all of available meta-analysis [35].

4.3. Interleukin 10

Interleukin 10 is an anti-inflammatory cytokine that down-regulates IL-1B, TNF-A, and interferon-γ gene expression. Previously published studies suggested that individuals carrying the IL-10 ATA haplotype associated with low IL-10 production (-592, -819, -1082) have an increased risk for non-cardia gastric cancer [33]. Nevertheless meta-analysis failed to confirm these observations except for IL10-1082G where the effect of polymorphism depends on ethnicity of the host. It seems that this polymorphism increases gastric cancer risk in Asians, but has no significant effect in non-Asian populations [39].
4.4. Interleukin 8

IL-8 is a CXC family cytokine that is a potent chemoattractant for neutrophils and lymphocytes affecting proliferation, migration, and tumor angiogenesis. However, not all studies have replicated the positive associations between pro-inflammatory cytokines polymorphisms and gastric cancer risk [42]. A polymorphisms in IL-8 (-251 T→A) is associated with increased production of IL-8 in \textit{H. pylori} infected gastric mucosa and with precancerous gastric abnormalities in Caucasians and gastric cancer in Asian populations [43]. Nevertheless, studies in Asian populations [44,45] have failed to confirm relevance of this polymorphism and meta-analysis supported this finding [35], but this can be attributable to small sample size in these studies and limited number of the high quality studies available for the meta-analysis.

4.5. Presence of multiple high-risk cytokine gene polymorphisms

Possession of multiple high-risk host polymorphisms is associated with increased risk for gastric cancer. Presence of 3-4 of the polymorphisms (IL-B1-511*T, IL-IRN*2/*2, TNF-A-308*A and IL-10 ATA/ATA) confers a 27-fold increase in risk of non-cardia gastric cancer [33].

5. Polymorphisms in innate immune response genes

\textit{H. pylori} attaches to gastric epithelium via receptors. Thus, polymorphisms in the innate immune response genes, which interact with these receptors, could influence outcome of infection and potentially the risk of gastric cancer.

5.1. Toll like receptor 4 (TLR4)

TLR4 is a cell-surface signaling receptor involved in the recognition and host response to \textit{Helicobacter pylori}. Toll-like receptor 4 gene codes for a lipopolysaccharide (LPS) receptor molecule involved in innate immune recognition of microbe pathogen-associated molecular patterns. The TLR4+896A>G polymorphism linked with impaired reactivity to bacterial lipopolysaccharide may play a role in gastric carcinogenesis [44] and is associated with hypochlorhydria and upper gastrointestinal cancer. TLR4 896 polymorphisms results in changed conformation of the extra cellular domain of the TLR4 receptor and carriers are unable to adequately respond to LPS challenge. The defective signaling through TLR4 leads to an exaggerated inflammatory response with severe tissue destruction that causes gastric atrophy and severe hypochlorhydria. Two independent case-control studies have demonstrated that TLR4+896G carriers have eightfold increase in odds ratio for hypochlorhydria and gastric atrophy, and over two-fold increase for gastric cancer [44,45]

6. Cell proliferation-related gene polymorphisms

Meta-analysis by Gao et al identified 23 polymorphisms significantly related to gastric cancer in at least one published study suggesting the importance of polymorphisms in genes
implicated in cell proliferation in development of gastric cancer. The overall effect of these polymorphisms is probably modest but should not be neglected [46].

6.1. Cell cycle and apoptosis regulators

Cell cycle and apoptosis regulators are directly involved in the initiation of malignant proliferation of the cell. Polymorphisms of functional regulators of TP53, TP53BP2 (tumor protein P53 binding protein 2) and MDM2 (gene encoding Mouse double minute 2 homolog, an important negative regulator of the p53 tumor suppressor) were found to be related to the development of gastric cancer. TP3 gene encodes a multi-purpose protein (P53) that takes part in regulating the cell cycle, carrying out programmed cell death, initiating DNA repair, and regulating the transcription of a large number of genes that cells use for various biological purposes. Given its many essential functions, P53 is frequently found inactivated in tumor cells. Results of meta-analysis confirm that association of TP3 gene polymorphisms vary by population and type of gastric cancer. The TP3 Arg allele carriers of Asian origin have an increased risk for gastric cancer, while same polymorphism seems to have protective effects in Caucasians [43]. Possible explanation is difference in environmental factors that act together with either apoptotic or DNA repairing mechanisms. Data obtained for other two polymorphisms in cell-cycle related genes that were extensively studied are inconsistent, both for Lmyc (nuclear oncogen) EcoRI polymorphism and p21 (gene encoding P21/cyclin-dependent kinase inhibitor 1) polymorphism (Arg31Ser), probably due to relatively small sample sizes and underestimated importance of environmental factors and their interplay with host genetics.

PPAR-γ (peroxisome proliferator-activated receptors γ) is a member of the nuclear hormone receptor family that plays an important role in cell differentiation and regulation of metabolism. A potential interplay between PPAR-γ Pro12Ala polymorphism and H. pylori infection was observed in the development of gastric cancer [47,48].

6.2. Growth factors and growth factor receptors

Polymorphisms determining higher level of growth factors and related receptors, which are important for tissue repair, were associated with reduced risk of gastric cancer. Such associations were observed for gene encoding epidermal growth factor (EGF) EGF 5′ UTR 61G>A, [49], polymorphisms for transforming growth factor beta (TGFβ) i.e. TGFβ1 -509C>T, TGFBR2-875G>A [50] and gene encoding insulin-like growth factor-binding protein 3 IGFBP3 -202A>C and Gly32Ala [51,52].

7. Environmental factors

Environmental regulation of virulence factors could be an interesting concept explaining why not all infected individuals develop severe complications of disease despite infection with pathogenic Helicobacter pylori strains.

Recent study demonstrated that high salt diet could influence H. pylori protein expression [53].
7.1. Diet

In 2007 World Cancer Research Fund declared that high intake of vegetables and fruit probably decrease risk of gastric cancer, and that high intakes of salt and salty food probably increase risk of gastric cancer [54]. The proposed underlying mechanism for the inverse association of gastric cancer risk with vegetable and fruit/rich diet is related to the presence of antioxidants. Salt on the other hand acts directly on the stomach lining, destroying the mucosal barrier, causes gastritis and increased epithelial cell proliferation [55]. A synergistic interaction between diet and Helicobacter pylori infection with risk of gastric cancer has been proposed [56]. Recent study demonstrated that high salt diet could influence Helicobacter pylori protein expression leading to increased risk of gastric cancer [53]. Salt responsive increase in cagA expression attributable to increased CagA transcription was described that could lead to increased risk of gastric cancer.

7.2. Smoking

Tobacco smoking is the risk factor associated with the largest number of cancer cases worldwide and the causal link with stomach cancer is recognized [54]. A recent meta-analysis found significant positive associations of smoking with risk of both cardia and non-cardia gastric cancer among the majority of studies, overall increasing risk by 62% for male and 20% for female current smokers [57]. It is possible that tobacco smoke carcinogens affect gastric cancer risk directly through contact with the stomach mucosa or indirectly through the blood flow [54]. In a large population-based study in Europe (EPIC), 17.6% of gastric cancer cases were attributed to smoking [56]. The cancer risk in past smokers can remain up to 14 years after cessation of smoking [57,58]. The effect of smoking on gastric cancer is dose-dependent and additive in the presence of other risk factors [59,60]. However, passive smoking does not seem to increase the risk [61].

7.3. Non-steroidal anti-inflammatory drugs

Protective effect of regular use of non-steroidal anti-inflammatory drugs (NSAIDs) and particularly aspirin on risk of gastric cancer was repeatedly reported in observational studies and then results of meta-analysis [62] confirmed these finding. According to Algra et al regular NSAID users have up to 20% reduced risk of gastric cardia adenocarcinoma and up to 36% reduced risk of distal gastric adenocarcinoma [62]. NSAIDs suppress the production of cyclooxygenase enzymes. Data on clinical efficacy of NSAIDs in prevention of gastric cancer first suggested that aspirin reduces risk for both proximal and distal gastric cancer [63]. Recent results from population-based intervention trial by Wong et al revealed that celecoxib treatment or Helicobacter pylori eradication alone had beneficial effects on the regression of advanced gastric lesions. Nevertheless no favourable effects were seen for H. pylori eradication followed by celecoxib treatment [64]. Meta-analysis by Tian et al suggests significant protective effects of NSAIDs against gastric cancer [65].
7.4. Socioeconomic status

Lower socioeconomic status is associated with at least two-fold greater risk of gastric cancer irrespective of the country incidence of gastric cancer [54]. Possible explanation is related to increased likelihood of transmission and re-infection with *H. pylori* (large family, poor sanitation, less frequent use of antibiotics). Also, low socioeconomic status is related to a diet lower in fresh fruits and vegetables.

8. Importance of *Helicobacter pylori* eradication in prevention of gastric cancer development-current knowledge and evidence

*Helicobacter pylori* infection is mainly associated with distal forms and intestinal-type gastric carcinoma [7-9]. Namely, there are two distinct sites of gastric adenocarcinoma: proximal (cardia) and distal (non-cardia), with different epidemiological and clinical characteristics. Main risk factors for cardia gastric cancer are obesity, gastro-oesophageal reflux disease and Barrett’s oesophagus [66] and its incidence is increasing over last decades. On the contrary, main risk factors for development of distal gastric cancer are *H. pylori*, low socioeconomic status, smoking, salty and smoked food intake, low consumption of fruits and vegetables and a family history of gastric cancer. Clear decline in incidence of distal gastric cancer is observed in the last decades [67].

Histologically gastric cancer is, according to Lauren classification, divided into two subtypes: intestinal and diffuse type. The intestinal type is related to corpus-predominant gastritis with intestinal metaplasia and is closely related to long-lasting *H. pylori* infection, whereas the diffuse originates from superficial pangastritis without atrophy [7-9].

Intestinal type gastric adenocarcinoma results from a prolonged precancerous process. The link between gastric intestinal metaplasia and cancer was proposed by pathologists in Java and Sumatra in 1938. This idea was over time only occasionally revisited by scientists until in 1975 Correa et al proposed a model of gastric carcinogenesis. This model postulated that intestinal type of gastric cancer was a result of progressive changes in the gastric mucosa [68]. Authors updated their model in 1988 and 1992. In Correa cascade the following consecutive steps are now recognized: normal gastric mucosa, superficial (non-atrophic) gastritis, multifocal atrophic gastritis (MAG), complete (small intestine type) intestinal metaplasia followed by intestinal metaplasia of the incomplete (colonic) type, low-grade dysplasia, high-grade dysplasia and invasive adenocarcinoma. [7-9, 68]. Loss of normal glandular tissue is the first specific recognizable step in the precancerous cascade. However, the changes of the precancerous lesions over time remain an issue difficult to assess leading to the fact that the point of no return, although of critical importance for timely eradication, still remains unidentified.

Five randomized controlled trials (RCT) reported effects of *H. pylori* infection eradication on invasive gastric cancer or premalignant histological lesions of gastric mucosa. [69-75]

Study by Wong et al. was designed as RCT conducted in a high-risk gastric cancer region in China that evaluated gastric cancer incidence as a primary outcome [69]. Authors identified
healthy individuals with *H. pylori* infection and treated them either with eradication therapy or placebo. During a follow-up period of 7.5 years authors failed to demonstrate overall decrease in gastric cancer incidence. The study concluded that incidence of gastric cancer development in the general population was similar between subjects receiving *H. pylori* treatment and placebo. The study, however, suggested the possible protective role of *H. pylori* eradication in participants without precancerous lesions, including gastric atrophy, intestinal metaplasia, or dysplasia.

Other above cited trials were not designed to assess gastric cancer as a primary outcome [70, 71] or had low numbers of gastric cancer cases [69, 72] to provide an informative assessment of the effects of eradication on cancer occurrence. Nevertheless, recently an important study was published by Ma et al [74] with the long-term follow-up results of a randomized trial in which 2258 *H. pylori* seropositive adults from a general population in China were randomly assigned to three interventions (*H. pylori* eradication, garlic supplements, supplemental vitamins) or control groups. After 15 years there were 34 new gastric cancers in *H. pylori* eradication group and 52 in the corresponding control group (relative risk of 0.61, 95% confidence interval 0.38-0.96) that clearly demonstrated benefit of eradication therapy in gastric cancer prevention.

This year Lee et al [76] evaluated the benefit of mass eradication of *H. pylori* infection started in Taiwanese population with high incidence of *H. pylori* infection. Individuals who were aged 30 years and older were tested for the presence of *H. pylori* infection and those positive underwent endoscopic screening and subsequent eradication therapy. Authors demonstrated the success of eradication in 78.7%. Gastric atrophy incidence decreased in over 77% after successful eradication of *H. pylori* with no significant change in intestinal metaplasia. Gastric cancer incidence during the chemoprevention period was reduced by 25%. Based on these findings authors suggest that ultimate benefit in reducing gastric cancer incidence and its mortality should be validated by a further long-term follow-up.

Over a decade ago Uemura et al. [77] first reported that *H. pylori* eradication could reduce the subsequent development of metachronous gastric cancer after endoscopic resection of early gastric carcinoma. Namely, 132 *H. pylori* serology-positive patients who underwent endoscopic resection were assigned to the *H. pylori*-treatment group or the no treatment group according to the patients’ preference. Regular endoscopic follow-up for up to 4 years found no metachronous cancer in *H. pylori*-treated patients compared to 9% in the no treatment group.

Some years later, Fukase et al. [75] reported the results of a trial in which more than five hundred patients in Japan who had previously undergone endoscopic resection for treatment of early gastric cancer were randomized to either *H. pylori* eradication or usual care. They demonstrated statistically significant 65% reduction in the risk of metachronous gastric carcinoma compared with the control group.

The evidence from more recent trial reports [74-76], taken together with the epidemiological and experimental evidence for the carcinogenic activity of chronic *H. pylori* infection, provides support for a protective effect of *H. pylori* eradication in gastric cancer.
9. Conclusion

*H. pylori* infection contributes 5.5% to global cancer burden and is the single most important cause of infection-associated cancer globally [6]. It is therefore reasonable to consider eradication therapy as optimal preventive approach in infected individuals. Interrupting transmission of infection is primary prevention, secondary should be eradication therapy and tertiary prevention includes identification of the individuals with early gastric cancer. However data from large study in China that included 1630 healthy carriers of *H. pylori* infection that were given eradication therapy or placebo detected no difference of gastric cancer incidence between these two groups [69]. Benefit would probably be seen in patients where gastric cancer cascade did not cross the point of no return, which would explain results of recently published studies [74,76]. Namely, intestinal-type gastric cancer results from a multistep process of mucosal alterations leading from chronic inflammation (gastritis) to glandular atrophy followed by development of intestinal metaplasia and dysplasia resulting in invasive carcinoma. Clinical studies should aim to identify a ‘point of no return’, a situation when mucosal alterations are no longer reversible after *H. pylori* eradication and progression to gastric cancer became independent of presence of the bacterium.

Higher risk for gastric cancer might be modulated by an overall pro-inflammatory host genetic profile in the adaptive and innate immune systems genes (e.g. *IL-1B, TNFA, IL-10, IL-8*, and *TLR4*) [77]. This pro-inflammatory profile might drive the immune response to *H. pylori* infection to a severe chronic inflammatory phenotype, reduced gastric acid secretion, bacterial overgrowth, and oxidative stress to the gastric mucosa.

Therefore timely eradication of *H pylori* infection especially in individuals with proinflammatory genetic profile could prevent development and in long term possibly decrease incidence of gastric cancer.

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