PrimeView Helicobacter pylori infection

Helicobacter pylori infection causes several gastroduodenal disorders, most commonly chronic gastritis, which can progress to peptic ulcer, gastric cancer or, rarely, gastric mucosa-associated lymphoid tissue lymphoma. Unique properties of the bacterium enable colonization of the gastric epithelium.

Epidemiology

H. pylori is usually transmitted in childhood and infection mostly persists without treatment. Around 50% of the world population was infected by H. pylori a decade ago but prevalence has since decreased by ~10%, possibly owing to socioeconomic development, improved hygienic conditions and antibiotic eradication therapies. Despite an overall decline in peptic ulcer disease (PUD), H. pylori infection remains the predominant cause of this disorder with infection detected in up to 90% of patients. Importantly, ~90 % of gastric cancer diagnoses can also be attributed to *H. pylori* infection. In individuals with the infection, lifetime risk of the malignancy is up to 5%. Worldwide, the incidence and mortality of gastric cancer vary widely and are highest in regions of Asia and Eastern Europe.

• *H. pylori* is the most prevalent carcinogenic pathogen, as gastric cancer accounts for ~37% of all cancers caused by a chronic infection.

Diagnosis

Most acute H. pylori infections remain undiagnosed owing to predominantly absent or mild symptoms. In children, endoscopic examination is only advised when PUD is suspected or complications occur. The recommended diagnostic tests in individuals aged <50 years (range 45-55 years) and in those with dyspepsia without alarm symptoms, including anaemia, weight loss or family history of gastric cancer, are the non-invasive ¹³C-urea breath test and stool antigen test. Endoscopic diagnosis is advised in those aged >50 years or when alarm symptoms are present to exclude mucosal changes.

• Some data suggest that *H. pylori* eradication can reduce gastric cancer incidence and mortality on the population level but further research is required to determine the utility of this prevention strategy.

Pathophysiology

H. pylori is exclusively adapted to colonize the gastric mucus layer. Infection-associated may progress to pathological the Correa cascade, which

Normal epithelia

Mucosal colonization by H. pylori depends on unique bacterial properties, including enhanced motility, acid neutralization, adhesion, epithelial damage via vacuolization and induction of inflammation enhanced by cytotoxin-associated antigen pathogenicity island (CagPAI). Structural variations, including of flagellins and lipopolysaccharides, may support immune evasion.



submucosal influx of diverse immune cells, leading to chronic gastritis, the severity of which depends on bacterial virulence, host genetics and environmental factors, and that often remains asymptomatic for decades.

Atrophic gastritis

The Correa cascade describes the progress from chronic gastritis initiated by H. pylori infection to gastric cancer. Gastritis-induced increased levels of reactive oxygen species and nitric oxide metabolites, and reduced antioxidant levels may be involved in the development of subsequent pathologies.

and/or metaplasia Dysplasia



Gastric cancer



Management

Adults with clinical symptoms or complications of *H. pylori* infection as well as those without symptoms but at risk of complications require therapy. Eradication regimens comprise a strong acid suppressant and antibiotics. Owing to increasing antibiotic resistance of *H. pylori*, antibiotic susceptibility testing or, if not available, local resistance patterns define the choice of first-line therapy. A combination of a proton pump inhibitor with clarithromycin and amoxicillin or, alternatively, metronidazole in case of resistance to one of the former agents is predominantly used. Bismuth-containing regimens, comprising a proton pump inhibitor, bismuth, a tetracycline and a nitroimidazole antibiotic, are useful empiric first-line therapies and are effective after failure of previous treatments.

• Treatment success needs to be tested and confirmed from 4 weeks after therapy to inform further therapy needs and regional utility of employed regimens.

Outlook

Population-level test-and-treat strategies might reduce *H. pylori* infection-related complications but financial costs and risk of widespread antibiotic resistance highlight the need for tailored approaches in those with a high risk of gastric cancer. Those with a high risk of gastric cancer include first-degree family members of patients and populations in regions with high gastric cancer incidence. To overcome increasing antibiotic resistance new agents are needed, such as those interfering with the bacterium's acid neutralization, motility or adhesion mechanisms. In addition, the interactions of H. pylori with the gastrointestinal microbiome require further study to better understand their roles in the development of infection-related pathologies and the possibly beneficial potential of probiotic treatments.

Check for updates

Helicobacter pylori infection

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Abstract

Helicobacter pylori infection causes chronic gastritis, which can progress to severe gastroduodenal pathologies, including peptic ulcer, gastric cancer and gastric mucosa-associated lymphoid tissue lymphoma. H. pylori is usually transmitted in childhood and persists for life if untreated. The infection affects around half of the population in the world but prevalence varies according to location and sanitation standards. H. pylori has unique properties to colonize gastric epithelium in an acidic environment. The pathophysiology of *H. pylori* infection is dependent on complex bacterial virulence mechanisms and their interaction with the host immune system and environmental factors, resulting in distinct gastritis phenotypes that determine possible progression to different gastroduodenal pathologies. The causative role of H. pylori infection in gastric cancer development presents the opportunity for preventive screen-and-treat strategies. Invasive, endoscopy-based and non-invasive methods, including breath, stool and serological tests, are used in the diagnosis of *H. pylori* infection. Their use depends on the specific individual patient history and local availability. H. pylori treatment consists of a strong acid suppressant in various combinations with antibiotics and/or bismuth. The dramatic increase in resistance to key antibiotics used in H. pylori eradication demands antibiotic susceptibility testing, surveillance of resistance and antibiotic stewardship.

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Introduction

Helicobacter pylori is the most frequent cause of chronic gastritis and variably leads to severe gastroduodenal pathologies in some patients, including gastric and duodenal peptic ulcer disease (PUD), gastric cancer, and gastric mucosa-associated lymphoid tissue (MALT) lymphoma¹⁻³. The diverse pathologies attributed to *H. pylori* infection are caused by complex interactions of bacterial virulence, host genetics and environmental factors^{4,5}, which result in different phenotypes of chronic gastritis (Table 1). These phenotypes are defined as antral-predominant, corpus-predominant gastritis or pangastritis according to the highest gastritis severity within gastric anatomical compartments.

The milestone discovery of *H. pylori* invalidated the dogmatic assumption of the acidic stomach as a sterile organ. This finding required a fundamental revision of gastric pathophysiology and gastroduodenal pathologies. Although spiral microorganisms in the stomach had been reported⁶, it was not until 1982 that Warren and Marshall identified a bacterial infection as the cause of chronic gastritis and succeeded in isolating the responsible microorganism⁷ (Fig. 1). The proof of concept that H. pylori infection causes gastritis was obtained by voluntary self-experiments with ingestion of a bacterial broth and cure of gastritis following H. pylori eradication (that is, fulfilment of the Koch's postulates)^{8,9}. The Koch's postulates requires proof of causality for a pathogen to induce disease and cure of disease when the causal agent is removed - this finding was eventually confirmed in clinical trials¹⁰. The bacterium originally referred to as *Campylobacter pylori* (C. pyloridis) became reclassified as H. pylori in 1989 (ref. 11). Peptic ulcer, considered an acid-driven disease in the traditional pathophysiological concept, became an infection-driven disease¹²⁻¹⁴. The standard therapy with long-term acid suppression became short-term H. pylori eradication therapy¹⁴. For the discovery that eventually led to the permanent cure of peptic ulcers by H. pylori eradication, Marshall and Warren were awarded the Nobel prize in Physiology or Medicine in 2005 (ref. 15). To this day, continuous scientific progress and new clinical developments have led to frequent modifications and updates to the clinical management of H. pvlori¹⁰.

H. pylori infects nearly half of the population in the world, with strong differences between geographical areas but with consistent trends towards a decreasing incidence¹⁶. Around 80% of individuals with *H. pylori* infection remain asymptomatic, but gastritis develops in all individuals with the infection, with unpredictable and potentially severe individual outcomes as well as high morbidity and mortality^{17,18}.

This Primer provides an update on current epidemiological trends of *H. pylori* infection, key aspects of its pathogenicity and its role in gastroduodenal pathologies. An important focus is also on gastric cancer prevention by *H. pylori* eradication. The diagnostic and therapeutic management of *H. pylori* infection is discussed according to current international guidelines. The dramatic increase in antibiotic resistance requires special measures, including the incorporation of new molecular methods for antibiotic susceptibility testing, the adaptation of individual treatment regimens and the implementation of antibiotic stewardship.

Epidemiology H. pylori infection

Once individuals acquire *H. pylori* infection, the pathogen usually persists throughout their lifetime². However, spontaneous clearance was reported in 9 of 58 (15.5%) children during the 20 years of follow-up of a retrospective cohort study from 2002 (ref. 19). Clearance of *H. pylori* does often occur in patients with advanced atrophic gastritis²⁰. The global prevalence of *H. pylori* infection in adults has declined from 50–55% to 43% during 2014–2020 (refs. 16,17), mostly attributed to improvement of socioeconomic status, living standards and hygiene conditions^{16,21-23}. The increased use of antibiotics, including eradication therapies, in individuals with the infection might be a further contributor.

Prevalence varies substantially with age, ethnicity, associated diseases, geographic regions, socioeconomic status and hygiene conditions^{16,21}. For young age groups, the 2002 study showed that most newly acquired H. pylori infections occurred before the age of 10 years¹⁹. The overall crude incidence rate was 1.4% per year, ranging from 2.1% at 4-5 years, 1.5% at age 7-9 years, to 0.3% at 21-23 years of age¹⁹. During 2014-2020, the prevalence of infection in children and adults was higher in low-income and middle-income countries, including in Africa, the Eastern Mediterranean, Russia, and Middle America and South America, than in high-income countries but was reduced in Western Pacific regions¹⁷ (Fig. 2). The prevalence of infection is higher in adults than in children. It is also higher in rural developing areas than in urban developed regions². Prevalence of *H. pylori* infection in children has been decreasing owing to improvements in socioeconomic status and hygiene conditions; however, the global prevalence in children remained as high as 34% during 2014-2020 (refs. 17,24). The higher prevalence in older individuals compared with children is explained by most (90%) of H. pylori infections being acquired in childhood and persisting throughout life rather than by a higher risk of infection at older age.

Some studies suggest increased susceptibilities to *H. pylori* infection in certain populations based on genetics and ethnicity; however, food sharing and housing habits may also have a role²²⁻²⁴. For example, in the Sumatra islands of Indonesia, the prevalence of *H. pylori* infection is very low in the Malay and Java populations, but is high in Batak

Phenotype	Frequency	Localization	Effect on secretory function	Possible outcomes ^a		
Mild gastritis phenotype	Most patients	No specific gastric compartment predominantly affected	Normal acid secretion	Asymptomatic in most patients, no significant clinical outcome		
Duodenal ulcer phenotype	10–15% of patients	Antral-predominant gastritis	High gastrin and acid secretion and impaired inhibitory control of acid secretion	Dyspeptic symptoms, duodenal ulcer		
Gastric cancer phenotype	~1% of patients	Corpus-predominant gastritis	Low or absent acid secretion; variable gastrin secretion	Severe atrophic gastritis and intestinal metaplasia, gastric cancer		
"Sastric ulcer and mucosa-associated lymphoid tissue lymphoma are not associated with a distinct dastritis phenotype. Gastric ulcer is associated more frequently with predominant						

Table 1 | Disease phenotypes of H. pylori infection

^aGastric ulcer and mucosa-associated lymphoid tissue lymphoma are not associated with a distinct gastritis phenotype. Gastric ulcer is associated more frequently with predominant corpus-type gastritis and low acid secretion. H. pylori, Helicobacter pylori.

populations, indicating that genetic factors may contribute to differential host susceptibility²⁵. Gene and genome-wide association studies have identified that polymorphisms in IL-1B, Toll-like receptor 1 (TLR1) locus and the FCGR2A locus are associated with *H. pylori* seroprevalence^{26,27}. However, a 2022 study has cast doubt on a role of the TLR1/6/10 locus in *H. pylori* seroprevalence²⁸, and further studies are needed²⁹.

Faecal–oral and oral–oral routes are considered the most likely routes of transmission^{30,31}. Contaminated water may be a source of infection in developing countries³². *H. pylori* can be cultivated from the vomitus, stool and saliva of individuals with infection, indicating the potential transmissibility via these routes³³. However, future studies about transmission pathways and their relative importance are urgently needed.

Person-to-person transmission within families, especially from mothers and siblings with the infection, is common in developing countries³⁴. Genotyping studies have shown that strain concordance was detected in 10 of 18 (56%) mother-offspring and in 0 of 17 fatheroffspring relations³⁵. Concordant strains in siblings were detected in 29 of 36 (81%) families³⁵. Nevertheless, transmission within couples or spouses remains controversial³⁵⁻³⁷. In two studies, the ribopatterns of H. pylori strains were similar in 8 of 18 (44%) and 5 of 23 (22%) couples with H. pylori infection^{35,36}. However, another study showed that, although restriction fragment length polymorphism patterns were similar in 5 of 13 couples, further restriction fragment length polymorphism using restriction endonucleases revealed distinct patterns in these 5 couples, indicating that transmission between spouses is infrequent³⁷. Due to the extremely high genetic diversity of *H. pylori*, even short nucleotide sequences can be highly informative about transmission pathways and the direction of transmission between two individuals. Seven-gene multilocus sequence typing³⁸ and, more recently, whole-genome sequence analysis³⁹ have enabled the reconstruction of the spread of H. pylori in families and have great potential to answer open questions in H. pylori epidemiology.

The annual reinfection or recrudescence rate after successful eradication is low (<2%) in adults in developed countries but is higher (5–10%) in adults in developing countries and in children^{17,40}. Some randomized trials showed that a strategy of family-based *H. pylori* screening and treatment can reduce the recurrence rate more than a single-patient approach⁴¹. Further well-designed, large-scale randomized trials are warranted to validate whether family-based screening and eradication may reduce the transmission of *H. pylori* within families.

H. pylori infection-related diseases

H. pylori infection is an important causal factor of gastric cancer, duodenal ulcer and gastric ulcer⁴².

Peptic ulcer disease. Lifetime prevalence of PUD in individuals with *H. pylori* infection is estimated at around $10\%^{14,43,44}$. After 10 years, >11% of individuals with the infection develop PUD compared with 1% of individuals without the infection⁴⁵. In a prospective study, the lifetime risk of developing duodenal ulcer and gastric ulcer was respectively increased by 18.4-fold and 2.9-fold in individuals with infection with *cagA*-positive *H. pylori* strains⁴⁶.

Since the 2000s, the global prevalence of PUD is declining⁴⁷⁻⁵⁰ in parallel with a decreasing prevalence of *H. pylori* infection¹⁶ for various reasons^{47,51-54}. The epidemiological trend indicates an increasing role of NSAIDs, including acetylsalicylic acid, which independently increase the risk of gastroduodenal ulcer and ulcer bleeding^{44,53}.

Despite changing trends in PUD worldwide, *H. pylori* remains the most prevalent cause of PUD. A study from Denmark showed an odds ratio of 4.3 (95% CI 2.2–8.3) for the association between *H. pylori* infection and PUD⁵⁷. In a meta-analysis including endoscopic surveys and national screening programmes in unselected population samples from Europe and China, the pooled prevalence of PUD was 6.8% and PUD was associated with *H. pylori* infection in 91% of cases⁵⁸. Around 3,000,000 diagnoses of PUD per year are estimated to relate to *H. pylori* infections and -90% of patients with duodenal ulcers and 70–90% of patients with gastric ulcers have *H. pylori* infection with variation according to geographical areas^{52,53,58–60}.

Gastric cancer. Around 90% of gastric cancer cases can be attributed to H. pylori infection⁶¹. In 2018, 812,000 gastric cancers, including non-Hodgkin lymphoma of gastric location, were recorded, accounting for ~37% of all cancers driven by a chronic infection, which makes H. pylori the most frequent carcinogenic pathogen⁶². Gastric cancer incidence and mortality differ significantly between regions, with the highest rates in Asia and Eastern Europe. The lifetime risk of gastric cancer is 1-5% in individuals with H. pylori infection, depending on ethnicity and environmental factors^{2,17,63}. Some populations are at an increased risk of gastric cancer following H. pylori infection, probably due to genetic factors, housing situation and dietary habits, for example, increased consumption of salted or pickled foods in East Asian populations. In addition, substantially higher gastric cancer incidence is found in indigenous populations worldwide⁶⁴ and in ethnic groups in the USA, including Asian Americans⁶⁵. Socioeconomic, dietary and lifestyle factors, such as smoking and extent of salt intake, are contributing factors to gastric cancer development, but they are all subordinate to the presence of *H. pylori* infection^{66,67}.

Extra-gastric diseases. Unexplained iron-deficiency anaemia, vitamin B_{12} deficiency and some cases of idiopathic thrombocytopenic purpura can be related to *H. pylori* infection^{68,69}. Antigen mimicry-induced autoimmunity related to *H. pylori* has been suggested in idiopathic thrombocytopenic purpura^{70,71}. Furthermore, other associations of *H. pylori* infection with diseases localized outside the stomach have been reported, including cardiovascular diseases, ischaemic heart disease, metabolic syndrome, diabetes mellitus, hepatobiliary diseases, non-alcohol fatty liver disease and neurodegenerative diseases, which have been attributed to persistent and low-grade systemic inflammation⁷²⁻⁷⁴. Most of these associations are based on limited and inconsistent data and remain inconclusive, and only a few, mostly observational, studies have documented a significant decrease in some of these manifestations when *H. pylori* is eradicated⁷³.

In children, particularly in the USA and Europe, an inverse association between *H. pylori* infection and asthma and allergy has been reported⁷⁵⁻⁷⁷, although this link has not been unequivocally confirmed⁷⁸. The often reported inverse association between *H. pylori* infection and the risk of gastro-oesophageal reflux disease (GERD), Barrett oesophagus and oesophageal adenocarcinoma remains highly controversial^{79,80}, and evidence for positive and negative associations exists⁷⁹⁻⁸⁵. Explanations for the discrepancies might lie in differing study protocols and *H. pylori* testing methodologies as well as in heterogeneity in the selection of patient and control populations. At present, the controversial findings and debates about a potential benefit of *H. pylori* for specific clinical scenarios have no confirmation nor impact on the management of the infection⁸⁶.

	— 1982: ·	Discovery of Helicobacter pylori by Marshall and Warren
	— 1989: «	H. pylori renamed from Campylobacter pylori
	— 1990: .	Other <i>Helicobacter</i> species discovered in human pathologies First trial confirming that <i>H. pylori</i> eradication cures duodenal ulcer
	— 1991:	 First epidemiological evidence of <i>H. pylori</i> as a risk factor for gastric cancer New gastritis classification system (Sydney system) published
	— 1992:	 Start of experimental H. pylori vaccine research Gastric mucosa-associated lymphoid tissue lymphoma curable by H. pylori eradication
	— 1994: •	 WHO declares H. pylori a class I carcinogen US National Institutes of Health recommend H. pylori eradication in peptic ulcer diseases
	— 1996: •	 Updated gastritis classification system (Sydney–Houston system) Maastricht I Consensus Report introduces the first international guidelines for management (including triple therapy for 7 days)
	— 1999: ·	PPI-triple therapy for H. pylori eradication with high efficacy confirmed in randomized controlled trials becomes global standard for peptic ulcer disease
	2000:	 Maastricht II Consensus Report recommends a test-and-treat approach in adult patients <45 years of age presenting in primary care with persistent dyspepsia and no alarm symptoms (treatment efficacy should always be confirmed by ¹³C-urea breath test or endoscopy)
	— 2000 c	 Antibiotic resistance of <i>H. pylori</i> against clarithromycin and levofloxacin increases Quadruple therapies become alternatives to PPI-triple therapy
	— 2001: 	• First prospective clinical study to prove <i>H. pylori</i> as key factor in gastric cancer development
	— 2004: 	• First randomized controlled trial of <i>H. pylori</i> eradication for gastric cancer prevention
	— 2005:	 Nobel prize for Marshall and Warren Maastricht III/Florence Consensus Report recommends selected extragastric diseases as indications for H. pylori eradication
	2008:	OLGA and, since 2010, OLGIM systems to predict gastric cancer risk in histological staging of gastritis
	— 2010: •	Bismuth quadruple therapy becomes first-line option in regions with high clarithromycin resistance PPI-triple therapy duration extended to 14 days Maastricht IV/Florence Consensus Report presents a series of innovations in management; screen-and-treat for consideration in areas/communities with high gastric cancer incidence First randomized controlled trials of <i>H. pylori</i> vaccines for prevention of infection start
	— 2012: •	Management of precancerous conditions and lesions in the stomach (MAPS) guidelines for surveillance of atrophic gastritis and early gastric cancer detection
	— 2013 o	nwards: • Main trials of gastric cancer prevention with <i>H. pylori</i> screen-and-treat in general populations
	— 2015: • •	Kyoto Gastritis Consensus defines <i>H. pylori</i> -associated gastritis as infectious disease Maastricht V/Florence Consensus Report recommends eradication therapy in individuals with <i>H. pylori</i> infection, even if asymptomatic, to prevent infection-related complications
	— 2016: •	Potassium-competitive acid blockers become more effective alternatives to PPIs in dual and triple therapy (first available in Japan)
		Taipei consensus on screen-and-treat for gastric cancer prevention recommends eradication therapy to be offered to all individuals infected with <i>H. pylori</i> and mass screening and eradication of <i>H. pylori</i> to be considered in populations at increased risk of gastric cancer MAPS guidelines update (MAPSII)
•	<u> </u>	Maastricht VI/Florence Consensus Report sets the focus on antibiotic susceptibility-based treatment, strategies in gastric cancer prevention and new insights into the relationship between <i>H. pylori</i> and gut microbiota

Fig. 1 | **Key developments in** *H. pylori* **clinical research and management.** *Helicobacter pylori* was discovered and reported at a conference in 1982 but the finding was not further disseminated before the first publication in 1983 (ref. 7). The timeline highlights key developments in clinical research of *H. pylori*

Mechanisms/pathophysiology

H. pylori microbiology

H. pylori is Gram-negative, microaerophilic curved or S-shaped bacteria that are highly motile due to a unipolar bundle of sheathed flagella. The cell envelope has a characteristic Gram-negative structure, but many other components have unique features adapted to the habitat of *H. pylori* in the human stomach². In comparison with many other pathogenic bacteria, H. pylori has a small ~1.6-Mbp genome consisting of a single circular chromosome that encodes ~1,600 proteins^{87,88}. The H. pylori core genome consists of ~1,100 genes present in all H. pylori strains, whereas the remaining accessory part of the genome comprises genes variably found in strain subsets⁸⁹, for example, a large number of diverse restriction-modification systems (genetic elements that provide protection against foreign DNA), providing variable DNA methylation⁹⁰. Extensive variation of genome content and gene sequences between strains, and even within the bacteria present in the stomach of one individual^{39,91,92}, is a prominent characteristic of *H. pylori* and results from the unusual combination of very high mutation and recombination rates⁹³. H. pylori has a high mutation frequency due to lack of a classical mismatch repair pathway in combination with the pro-mutagenic properties of its DNA polymerase I94,95. H. pylori is naturally competent and can take up DNA by means of the unique ComB DNA uptake system with similarities to a type IV secretion system (T4SS)^{96,97}.

DNA sequence diversity can rapidly spread through *H. pylori* populations due to recombination between strains^{98,99}. After import, DNA can be integrated into the chromosome based on homology, and such chromosomal imports have a unique bimodal length distribution, enabling *H. pylori* to adapt its genome to new environments in an extremely efficient way¹⁰⁰.

H. pylori strains show a characteristic population structure that reflects their coevolution with their human hosts and has led to conclusions about the history of its association with humans¹⁰¹⁻¹⁰³. H. pylori was acquired by modern humans in Africa at least ~100,000 years ago, possibly by a host jump from an unknown animal source. The most ancestral phylogeographic population of H. pylori is hpAfrica2, mostly found in Southern Africa. Further important, widespread and more recently evolved populations include hpAfrica1, hpNEAfrica, hpEurope, hpEastAsia, hpAsia2 and hpSahul^{104,105}. A major step in the evolution of H. pylori from the ancestral hpAfrica2 population to the populations that have spread over the globe was the acquisition of the cag pathogenicity island (cagPAI) by ancestral H. pylori from an unknown source. cagPAI encodes components of the Cag T4SS^{106,107}, which is a protein complex that spans the bacterial cell envelope and can directly deliver diverse effector molecules into host cells following adherence. Hence, whether strains possess an active Cag T4SS has substantial effects on their interaction with hosts. cagPAI-positive strains elicit far more inflammation than cagPAI-negative strains.

Bacterial factors involved in colonization and pathogenesis

H. pylori is highly adapted to the colonization of a unique ecological niche in the deep gastric mucus layer. Several mechanisms, including motility, urease production, adhesion and others, are important in *H. pylori* colonization (Box 1).

infection and its therapeutic management since 1982 (refs. 1,10–12,14,15,17,21,70, 184,224,241,311,336,377–383). OLGA, Operative Link on Gastritis Assessment; OLGIM, Operative Link on Gastritis/Intestinal Metaplasia Assessment; PPI, proton pump inhibitor.

Motility. Flagella-driven motility is essential for the entry of *H. pylori* into the mucus layer and for maintaining a swimming reservoir in the mucus¹⁰⁸ (Fig. 3). *H. pylori* has a unipolar bundle of rotating sheathed flagella, with filaments composed of two flagellin proteins¹⁰⁹ that evade activating the innate immune system via TLR5 due to specific adaptation of their amino acid sequences^{110,111}. The direction of movement is controlled by chemotaxis and energy taxis, enabling bacteria orientation through pH and bicarbonate (and possibly other) gradients in the gastric mucus¹¹². Motility can be inhibited in vitro by small molecule compounds that reduce *H. pylori* colonization density, which may be a future treatment approach¹¹³.

Urease. *H. pylori* produces abundant amounts of urease, aided by a unique system of accessory proteins that procure the required nickel, which is essential for urease holoenzyme activity, protects the bacterium from nickel toxicity and regulates urease activity by controlling urea influx into bacterial cells^{114,115}. Urease is essential for colonization, most likely because the enzyme, by cleaving urea into ammonia and carbon dioxide, enables the bacteria to survive brief periods of exposure to very low pH values, which *H. pylori* may encounter in the gastric lumen during transmission¹¹⁶. Through urease activity, urea provides an always available nitrogen source for the organism.

Adhesion. H. pylori can adhere to gastric epithelial cells by attaching surface molecules that are anchored on its outer membrane (adhesins) to host cell receptors. Adherence enables H. pylori to achieve high colonization despite epithelial cell shedding, mucus layer turnover and the physical force involved in gastric emptying, all of which act to reduce colonization¹¹⁷. The best-studied adhesins are encoded by members of the large hop superfamily of outer membrane protein-encoding genes. BabA mediates binding to Lewis b blood group antigens that are expressed on gastric epithelial cells¹¹⁸. The related SabA adhesin binds to host sialyl-Lewis x antigens, which are mainly expressed on epithelial cell surfaces under inflammatory conditions¹¹⁹. HopQ binds to multiple carcinoembryonic antigen-related cell adhesion molecules and seems to be important for Cag T4SS functionality^{120,121}. AlpA and AlpB mediate binding to the extracellular matrix glycoprotein laminin¹²². Expression of adhesins varies widely between strains; the contribution of individual adhesins to bacteria-cell adherence and to pathogenesis continues to be studied.

*cag*PAI and its translocated effectors. The -37-kb *cag*PAI¹⁰⁶ comprises -26 genes that encode the elements of T4SS¹⁰⁷. After *cag*PAI-carrying *H. pylori* attaches to a host cell, T4SS can translocate bacterial effector molecules into the host cell cytoplasm¹²³, including the CagA protein, which is also encoded by the *cag*PAI. In addition, several other molecules can be translocated via T4SS, including heptose-containing lipopolysaccharide core precursors^{124,125}, peptidoglycan fragments¹²⁶ and bacterial DNA¹²⁷. These molecules can interact with intracellular target molecules and profoundly affect intracellular signalling and cell function (Fig. 3).

After translocation, CagA undergoes tyrosine phosphorylation by cellular kinases¹²⁸. The phosphorylated form can interact with



Fig. 2 | **Prevalence of** *H. pylori* infection in adults and children. a,b, Global map of *Helicobacter pylori* infection prevalence in adults during 1970–2016 (part **a**) and in children and adolescents (<20 years) during 2000–2021 (part **b**). In adults, the prevalence was highest in Africa, Eastern Mediterranean regions, Russia, Middle America and South America. In children, the prevalence was

lower than that in adults in Russia, Western Pacific regions and European regions. However, the prevalence of *H. pylori* infection was similarly high in children and adults in Africa, Eastern Mediterranean regions, and Middle America and South America^{16,24}.

multiple target molecules in the host cell, including SHP2 (ref. 129), PAR1 (refs. 130,131) and ASPP2 (ref. 132), contributing to increased cell motility, reduced cellular tight junctions, genome instability, nucleotide damage and activation of the Wnt signalling pathway that is relevant in local neoplasia formation¹³³. Translocation of heptosecontaining lipopolysaccharide core intermediates may be important in inducing pro-inflammatory responses by both epithelial and immune cells through the ALPK1–TIFA signalling pathway and may also induce mutagenic and oncogenic processes^{134–136}. In addition, intracellular heptose signalling in macrophages may hamper antigen-presenting properties and subsequent T cell responses¹³⁶.

Vacuolating cytotoxin. Many *H. pylori* strains secrete vacuolating cytotoxin A (VacA), which is an oligomeric autotransporter protein toxin that can form anion-selective membrane channels¹³⁷. The effects of VacA on cells include induction of large intracellular vacuoles derived from late endosomes, induction of apoptotic cell death (following mitochondrial membrane perturbation) or necrosis, induction of autophagy, and inhibition of T cell and B cell proliferation and effects on other immune cells¹³⁸⁻¹⁴⁰. Together, these effects downregulate immune responses to *H. pylori* infection and promote host tolerance to the organism. Expression of VacA is not essential for colonization, and its contribution to illness remains controversial.

Immune responses to H. pylori

Innate immune evasion. The flagellins and lipopolysaccharides of *H. pylori* have evolved substantially differently from those of other Gram-negative bacteria and are largely not recognized by the human pattern recognition receptors TLR5 and TLR4, which signal danger to the host^{110,111}. These and other structural variations may contribute to immune evasion by *H. pylori* and its success as a persistent colonizer.

Innate immune activation. Contact between H. pylori and gastric epithelial and myeloid cells induces signalling through multiple innate pathways, leading to changes in cellular homeostasis and the release of cytokines and chemokines that trigger local and systemic inflammatory responses¹⁴¹⁻¹⁴³. As canonical TLR4-dependent and TLR5-dependent signalling is evaded, most inflammatory signalling depends on the activity of an intact *cag*PAI¹⁴⁴. The bacterial components transported into epithelial cells through T4SS engage multiple intracellular receptors. Many of the affected pathways converge on the activation of nuclear factor (NF)-κB, which leads to increased expression and release of IL-8 and other chemokines and cytokines¹⁴⁵⁻¹⁴⁷. IL-8 is a powerful attractant of neutrophils, which enter the gastric mucosa and are the defining element of the active component of chronic-active gastritis, the histological hallmark of H. pylori presence in the stomach^{148,149}. Monocytes, macrophages and dendritic cells are also attracted to the H. pylori-colonized mucosa. Activation of phagocytic monocytes and macrophages seems to strongly depend on the delivery of heptosecontaining lipopolysaccharide core intermediates via T4SS and the resulting signalling to the ALPK1-TIFA axis¹³⁵. Dendritic cells can be reprogrammed by contact with the bacteria, for example, to produce IL-18, which drives the conversion of T cells to regulatory T (T_{reg}) cells, suppressing immune activation¹⁵⁰.

Adaptive immune response. H. pylori invariably elicits a combined adaptive humoral and cellular immune response that is generally incapable of eradicating the bacteria. Colonization leads to formation of antibodies to many H. pylori antigens that have little effect on bacterial numbers¹⁵¹. In agreement with this apparent lack of a role of antibodies in protection against H. pylori, mice lacking antibody production can be successfully immunized against H. pylori¹⁵². H. pylori also induces the recruitment of T cells to the human gastric mucosa, including T helper 1 (T_H 1), T_H 17 and T_{reg} cells. Experimental vaccination in mouse models suggests that both $T_{\mu}1$ cells and $T_{\mu}17$ cells can be important in mediating protection against H. pylori infection¹⁵³. Furthermore, in mouse models, a protective effect of very early (neonatal) H. pylori infection against asthma was mediated by Treg cells accumulating in the lungs150,154,155, consistent with the hypothesis that H. pylori may downregulate systemic allergic responses through its recruitment of immunosuppressive T_{reg} cells to the gastric mucosa and, potentially, other body sites such as the lung.

From chronic H. pylori colonization to illness

H. pylori colonization of the gastric mucosa induces a pro-inflammatory response of gastric epithelial cells, which recruits diverse immune cells to the submucosa¹⁵⁶. The resulting condition is chronic–active gastritis, which is predominantly asymptomatic for decades of colonization in most patients. The severity of inflammation varies widely between individuals, depending on bacterial, host and environmental factors¹⁵⁷ (Box 1).

The single most important determinant of the pro-inflammatory activity of an *H. pylori* strain is its possession of a functional *cag*PAI¹⁵⁸. Expression of additional host-interaction factors, such as a portfolio of adhesins that fits the variable host receptor makeup and promotes strong binding to epithelial cells and, therefore, promotes crosstalk between the bacterium and the host cell, contributes to the response that a strain elicits in an individual host. Tolerogenic signalling contributes to the unusual accumulation and proliferation of gastric mucosaassociated lymphoid tissue. The decades-long inflammation in the gastric mucosa is thought to be an important driving force leading to gastric atrophy and, ultimately, gastric cancer as outlined by the Correa cascade¹⁵⁹ (Fig. 4). The Correa cascade describes a multistage, multifactorial process starting with superficial gastritis, progressing to atrophic gastritis, intestinal metaplasia and dysplasia, and culminating in gastric adenocarcinoma. A key emerging concept is that chronic inflammation, gastric atrophy and consequent achlorhydria lead to an aberrant and dysbiotic gastric microbiome that drives the process towards gastric neoplasia¹⁶⁰⁻¹⁶². Accumulating evidence suggests that, following H. pylori eradication, newly emerging components of the gastric microbiota might be involved in the oncogenic transformation of gastric epithelial cells^{160,163}. In other individuals, peptic ulcer disease or the rare H. pylori-associated MALT lymphoma can develop^{4,5,14,164}. The reasons why most individuals remain apparently asymptomatic

Box 1

Bacterial, environmental and host factors contributing to *H. pylori*-induced gastric cancer pathogenesis

Bacterial virulence factors

- Cag type IV secretion system^{106,123,167}
- vacA allelic genotypes linked to disease, for example, s1/m1/i1 alleles^{139,165}
- Adhesins, for example, BabA, SabA and HopQ¹¹⁸⁻¹²⁰

Environmental factors

- Smoking
- Dietary factors (low iron, high salt, and/or low fresh fruit and vegetable intake)^{167,383}

Host genetic factors

• Single-nucleotide polymorphisms in cytokine and growth factor genes encoding proteins that have been implicated in pathogenesis (IL-1 β , IL-2, IL-6, IL-8, IL-10, IL-13, IL-17A/B, IFN γ , TNF, TGF β) and their receptors (IL-RN, TGFR), innate immune receptors shown to be activated by *Helicobacter pylori* (TLR2, TLR4, CD14, NOD1, NOD2), enzymes involved in signal transduction cascades (PLCE1, PKLR, PRKAA1), glycoproteins (MUC1, PSCA) and DNA repair enzymes (ERCC2, XRCC1, XRCC3)^{29,387}

Gastric inflammatory phenotypes and associated gastric functions^{165–167}

- Corpus-predominant gastritis
- Atrophic gastritis (Operative Link on Gastritis Assessment (OLGA) III–IV)
- Hypochlorhydria
- High gastrin levels
- Low pepsinogen I levels and ratio of pepsinogen I to pepsinogen II

Gastric dysbiosis of microbes other than H. pylori^{160,364,384}

throughout their lifetime, whereas others proceed to clinical sequelae of varying severity, remain to be fully elucidated. Clinically useful, early bacterial predictive markers that could inform the decision to prescribe eradication therapy have not been identified.

It is now well established that the clinical outcome of *H. pylori* infection depends largely on the distribution and severity of *H. pylori*-induced gastritis¹⁶⁵ (Table 1). Thus, peptic ulcers are more likely in individuals with an antral-predominant pattern of gastritis characterized by high acid secretion and relative sparing of gastric corpus with its high parietal cell mass. Parietal cells secrete gastric acid and patients with peptic ulcers have a higher parietal cell mass than healthy individuals without ulcers. By contrast, gastric cancer develops in the context of corpus-predominant gastritis, gastric atrophy

and a profound loss of acid secretory capacity that precedes cancer by decades¹⁶⁶. The chronically inflamed and achlorhydric environment is further exacerbated by an aberrant pro-inflammatory and genotoxic gastric microbiota that drives the neoplastic process even after loss of *H. pylori* infection^{160,161,167}. Indeed, experimental work suggests that transplantation of the gastric microbiota from humans with intestinal metaplasia or gastric cancer into germ-free mice leads to the development of precancerous gastric changes¹⁶⁸.

Diagnosis, screening and prevention Diagnosis

Presentation. In daily routine, acute infection with *H. pylori* remains mostly undiagnosed at any age. Naturally occurring acute infection in childhood is usually not captured and is supposed to frequently present with abdominal complaints with potentially diverse aetiologies¹⁶⁹. In adults, the clinical presentation of acute infection can entail hypochlorhydria, epigastric pain and mild-to-moderate dyspeptic symptoms as described in case reports and from challenge studies in volunteers with *H. pylori* for vaccine development^{170–172}. By contrast, most children with *H. pylori* infection remain asymptomatic and complications are infrequent¹⁷³.

Once established, *H. pylori* infection is a persisting and not selflimiting condition in adults with the potential of severe complications in some individuals. PUD, gastric cancer and MALT lymphoma¹⁷⁴, in decreasing order of incidence, are the most important complications in adults⁸⁶.

Diagnostic tests. An accurate diagnosis of *H. pylori* infection is required before commencing treatment^{42,175}. Diagnostic methods for *H. pylori* detection include invasive and non-invasive test procedures^{70,176-181} (Fig. 5) (Tables 2 and 3).

Invasive tests require biopsy samples obtained during gastroduodenoscopy and include the rapid urease test (RUT), histological assessment, bacterial culture and direct detection of *H. pylori* genetic material using PCR, quantitative PCR or fluorescence in situ hybridization. Non-invasive methods include the ¹³C-urea breath test (UBT), serological detection for anti-*H. pylori* antibodies, the stool antigen test (SAT) and direct detection of *H. pylori* genetic material in stool via PCR¹⁷⁹.

RUT is a low-cost test with a specificity of 95–100%. False positive results are rare and can be explained by the presence of other urease-positive organisms such as *Proteus mirabilis*¹⁷⁹. Current use of a proton pump inhibitor (PPI) may lead to false negative results in RUT as well as in all other diagnostic tests except for serological assessment¹⁷⁵. Thus, PPI therapy should be interrupted 14 days before testing⁴².

Histological assessment on formalin-embedded samples is made according to the updated Sydney system, which provides information on *H. pylori* presence via direct visualization and on the extent of active and chronic inflammation and atrophy^{148,149,182}. The histochemical method for assessment of *H. pylori* gastritis relies on haematoxylin and eosin and Giemsa stains for detection of *H. pylori*¹⁴⁸. Gastritis severity is defined by the degree and extension of atrophy and/or intestinal metaplasia. Severe gastric atrophy is associated with an increased risk of gastric cancer and risk is best determined by changes according to the gastritis severity staging systems Operative Link on Gastritis Assessment (OLGA) and Operative Link on Gastritis/Intestinal Metaplasia Assessment (OLGIM)^{183,184}.

Culture of *H. pylori* is 100% specific but has a relatively low sensitivity (<80%) strongly dependent on transport media and logistics and laboratory proficiency owing to the required laboratory expertise

with special culture media. Limitations include costs and time constraints but microbial culture also enables phenotypical antimicrobial susceptibility testing (AST)¹⁸⁰.

Molecular testing from formalin-embedded biopsy or RUT samples with PCR methods, of which quantitative PCR is most appropriate, and fluorescence in situ hybridization is highly accurate in the detection of *H. pylort*¹⁸⁵ and can also be combined with molecular resistance testing¹⁸⁶.

Serological assessment of serum IgG levels is used as a screening test in specific clinical scenarios but it cannot distinguish active and previous infections because of the prolonged persistence of *H. pylori* antibodies. A positive serological test should be confirmed with a test that indicates active infection¹⁸⁷. Serological testing is the only method not influenced by current PPI intake. Next-generation blood tests to be used for screening in the consulting room became available in 2022 (ref. 187).

UBT uses stable isotope-labelled ¹³C-urea ingested with citric acid, which is then hydrolysed by bacterial urease and releases carbon dioxide and ammonia. UBT has high sensitivity and specificity (95–100%)¹⁸¹. SAT has a similar diagnostic accuracy as UBT. SAT is an immunological method based on monoclonal antibodies with which *H. pylori* antigens can be detected in stool samples¹⁸⁰. UBT, SAT and histological assessment are the commonly used tests in clinical practice for diagnosis of *H. pylori* as they enable the detection of active infection. However, the availability of these tests depends on the status of regional health-care services, which can directly affect treatment decisions^{175,179}.

The increase of antibiotic resistance to *H. pylori* worldwide^{185,189} demands AST in the individual patient to enable effective therapy choices following failed eradication treatment and to monitor antimicrobial resistance at the regional community level⁴². AST can be performed using phenotypic and genotypic approaches. Culture-based phenotypic testing requires fresh biopsy samples, but PCR-based genotypic testing can be done on fresh, formalin-embedded or RUT samples as well as on stool samples^{186,190–192}. Clarithromycin resistance should be excluded before its empirical use in regions with known clarithromycin resistance rates of >15% or unknown resistance against frequently used antibiotics. Clarithromycin resistance conferred by mutations





gene A (CagA) antigen. The host immune response is characterized by initial mucosal invasion with polymorphonuclear cells followed by activation of the innate and adaptive immune system with complex T helper 1 (T_H1), T_H17 and regulatory T (T_{reg}) cell interactions. Le^b, Lewis b blood group antigen; sLe^x, sialyl-Lewis x antigen.



Fig. 4 | Pathogenesis of gastric adenocarcinoma triggered by H. pylori. The Correa cascade describes the dynamic progress of gastric carcinogenesis along the stepwise evolution of chronic gastritis initiated by Helicobacter pylori infection. H. pylori causes chronic gastritis that is associated with the generation of reactive oxygen species and nitric oxide metabolites and a reduction in antioxidant vitamin Clevels. The risk of gastric cancer is highest in individuals who have infection by more virulent H. pylori strains, have pro-inflammatory host genetic factors, poor diet (high salt, smoked foods), low iron levels, unhealthy lifestyle and/or smoking habit. In these individuals, sustained chronic inflammation leads to damage and loss of acid-producing parietal cells, which leads to hypochlorhydria and finally achlorhydria. The loss of acidity facilitates colonization by harmful pro-inflammatory gastric microbiota, which in turn may produce more genotoxic pro-inflammatory metabolites and carcinogens that act directly on malignant epithelial cell transformation in the stomach³⁸⁴⁻³⁸⁶

in the gene encoding 23S rRNA are predominantly related to A2143G, A2142G and A2142C¹⁹³. Levofloxacin resistance is conferred by point mutations in the gyrase gene *gyrA*^{194,195}. The accuracy of the molecular detection methods for predicting antibiotic resistance varies between antibiotics, favouring clarithromycin and quinolone resistance detection^{194,196}. Formalin-embedded biopsy samples enable genotypical resistance testing at a later time point after endoscopy^{197–200}.

Infections with *Helicobacter* species other than *H. pylori* are rare and those most relevant refer to *Helicobacter heilmannii*, *Helicobacter felis* and *Helicobacter suis*^{201–203}. Standard *H. pylori* diagnostic tests (UBT, SAT, serological assessment and immunohistochemistry) have low sensitivity for the detection of these species²⁰¹. The clinical relevance of these often incidentally detected rare infections is low because of the low rates of complications.

Testing for eradication success after 4–6 weeks of antibiotic treatment is primarily – with some specific exceptions – performed with noninvasive diagnostic tests UBT and SAT (see Management section)⁴². PPI use has to be stopped 14 days before testing to exclude recrudescence of reduced bacterial density under acid suppressive therapy.

Indications for diagnostic testing. Based on the rarity of complications in childhood, a diagnostic endoscopic examination and treatment is recommended only in those with suspected peptic ulcer disease¹⁶⁹. In general, H. pylori detection in children is only recommended when complications arise^{169,204}. In the group of patients with dyspepsia without alarm symptoms, such as anaemia, loss of weight or family history of gastric cancer, and age <45 years (45-55 years according to age-related gastric cancer incidence variation among world regions), non-invasive testing with UBT or SAT is the strategy of choice^{70,205,206}. In patients aged >45 years or in the presence of alarm symptoms, endoscopy-based diagnosis is recommended to exclude mucosal changes^{207,208}. H. pyloriassociated dyspepsia is an independent entity that resembles but is distinct from functional dyspepsia^{1,208}. A test-and-treat strategy is the most cost-effective approach in patients with H. pylori infection and dyspepsia if H. pylori prevalence in the population is >5%. This strategy is superior to alternative therapies including PPIs^{70,209,210}, and the therapeutic gain of *H. pylori* eradication for symptom relief compared with other therapeutic options is substantial. A randomized, doubleblind, placebo-controlled trial for primary prevention of peptic ulcer bleeding in older patients who were prescribed aspirin in primary care lends support to an *H. pylori* test-and-treat strategy in patients starting aspirin treatment. Gastrointestinal bleeding episodes within a 2-year period were reduced by 65% in the *H. pylori* eradication group²¹¹.

Screening and prevention

H. pylori eradication as a strategy for preventing gastric cancer. Gastric cancer incidence and mortality at the population level are reduced by *H. pylori* eradication but more epidemiological data are required. Meta-analyses of randomized controlled trials and observational studies have concluded that moderate evidence suggests that H. pylori eradication therapy reduces the incidence of gastric cancer in healthy individuals^{212,213}, with an overall risk reduction of 46%²¹². In individuals with H. pylori infection and a family history of gastric cancer in firstdegree relatives, H. pylori eradication treatment reduces the risk of gastric cancer, with an overall risk reduction of 55%²¹⁴. A meta-analysis of randomized and observational cohorts that included five studies that considered baseline histological findings suggests that H. pylori eradication seems to be a primary preventive strategy in individuals with non-atrophic gastritis or multifocal atrophic gastritis without intestinal metaplasia, but not in those with intestinal metaplasia or dysplasia²¹⁵. In another meta-analysis, H. pylori eradication was associated with improvement in the severity of atrophic gastritis with and without intestinal metaplasia compared with placebo²¹⁶. Notably, eradicating H. pylori in patients treated for early-stage gastric cancer reduces rates of metachronous gastric cancer by ~50% (range 20-70%) according to two meta-analyses of randomized trials^{217,218}. In a pivotal trial, the reduction in metachronous gastric cancer incidence following endoscopic removal of early gastric cancer in patients receiving H. pylori eradication compared to placebo²¹⁹ suggests that eradication therapy may even work in the condition of severe atrophic gastritis²²⁰.

Diffuse and intestinal types of gastric cancer²²¹ are two major histological entities that differ in epidemiology, pathogenesis and clinical

course²²². However, randomized and observational studies have been unable to separately calculate the risk effects for these histological types. Further data on the benefits or adverse effects of *H. pylori* eradication will come from ongoing trials in China²²³, UK (HPSS study)²²⁴, Korea (HELPER Study)²²⁴ and Latvia (GISTAR study)²²⁵.

Targeted test-and-treat strategies for *H. pylori* infection. *H. pylori* test-and-treat strategies aim to decrease morbidity and mortality related to gastroduodenal disease (Box 2) according to the 2022 Maastricht VI/Florence guidelines⁸⁶. This strategy is appropriate for individuals with non-investigated dyspepsia. Testing for *H. pylori* infection should also be performed in persons who use NSAIDs and have a history of peptic ulcer. In addition, evidence is accumulating that supports the eradication of *H. pylori* in individuals with non-ulcer dyspepsia²²⁶, idiopathic thrombocytopenic purpura²²⁷, and iron and vitamin B₁₂ deficiency anaemia^{228,229}. Consensus exists for eradicating *H. pylori* in all cases of MALT lymphoma, regardless of disease stage and prognostic factors^{230,231}. Cure of *H. pylori* infection results in complete histological remission in most patients with localized MALT lymphoma²³².

Serological assessment for gastric cancer screening. A large body of research, particularly from East Asian populations at high risk, suggests that measurement of circulating pepsinogen levels is the most useful non-invasive test to define the status of the gastric mucosa (that is, whether it is atrophic)^{233,234}. Experts from the Kyoto Global Consensus agreed that pepsinogen levels in conjunction with anti-*H. pylori* antibody levels are useful for identifying individuals at increased risk for gastric cancer¹. Although there are still possibilities for optimization, the ABC (gastritis A, B, C and D) screening method based on this combined measurement is useful for the detection of an increased risk for both intestinal and diffuse types of gastric cancer²³⁵. The specific

groups are defined as follows, where Hp indicates *H. pylori* infection and PG indicates pepsinogen: A [Hp⁻PG⁻], individuals without infection; B [Hp⁺PG⁻], without chronic atrophic gastritis (CAG); C [Hp⁺PG⁺], with CAG; and D [Hp⁻PG⁺], with severe CAG; the latter two groups carry the highest risk for gastric cancer^{236,237}.

Population endoscopic screening for gastric cancer. Around 75% of all new gastric cancer cases are diagnosed in East Asian populations²³⁸. Consequently, Japan and South Korea have established successful national screening programmes in individuals aged \geq 40 years using either upper gastrointestinal series or upper endoscopy, depending on participant preference or comorbidities. Endoscopy has been the primary method for gastric cancer screening in Japan since 2017, and a study published in 2022 reported the benefits of this approach in the reduction of gastric cancer mortality²³⁹. In South Korea, the use of upper endoscopy has increased as this method is more accurate than upper gastrointestinal series for gastric cancer screening²⁴⁰.

Endoscopic surveillance of individuals at high risk. Although *H. pylori* eradication can reverse multifocal gastric atrophy and, to some extent, intestinal metaplasia, some patients with these histological lesions might benefit from surveillance at regular intervals. According to the management of precancerous conditions and lesions in the stomach (MAPS II)²⁴¹ European guidelines and the Maastricht VI/Florence consensus⁸⁶, individuals with advanced stages of atrophic gastritis (severe atrophic changes with and without intestinal metaplasia in both antrum and corpus, OLGA/OLGIM stages III and IV) should be followed-up with a high-quality endoscopy every 3 years. Based on growing evidence, endoscopic surveillance should also be considered in individuals with intestinal metaplasia at a single location but with a family history of gastric cancer, in those with incomplete-type intestinal



Fig. 5 | *H. pylori* **diagnostic procedures.** Diagnostic procedures are selected according to clinical scenarios. Non-invasive testing with the ¹³C-urea breath test and stool antigen test enables diagnosis of a current infection. Serological *Helicobacter pylori* antibody detection does not enable differentiation between current and previous *H. pylori* infection, necessitating confirmation by ¹³C-urea breath test or stool antigen test. All invasive tests are based on biopsy samples



from gastroscopy. These enable histological assessment for gastritis grading and staging, direct *H. pylori* detection via PCR, microbial culture, rapid urease test, and molecular examinations. Antibiotic susceptibility testing (AST) can be performed from stool or biopsy samples using microbial culture, next-generation sequencing (NGS) or real-time PCR (RT-PCR) techniques. FISH, fluorescence in situ hybridization; qPCR, quantitative PCR.

Table 2 | Indications for H. pylori testing

Indications for testing	Recommendation ^a		Refs.
	Strong	Weak	
Active or history of peptic ulcer disease	х		42,86
Low-grade gastric mucosa-associated lymphoid tissue lymphoma	x		86,337,338
History of endoscopic resection of early gastric cancer	x		86,219
Non-investigated dyspepsia in patients <50 years of age with no alarm symptoms	x		42,86
Investigated non-ulcer dyspepsia (functional dyspepsia)	x		42,86
First-degree relatives of patients with gastric cancer	x		42,86,214
First-generation immigrant from an area with high prevalence of <i>Helicobacter pylori</i> infection	x		42,86
Unexplained iron-deficiency anaemia when other causes have been excluded	x		42,86
Immune thrombocytopenia in adults	х		42,86
Long-term proton pump inhibitor use	x		42,86
Long-term acetylsalicylic acid and long-term NSAIDs, in consideration of individual additional risks		x	42,86,211

^aLevels of recommendations are adapted and based on current international guidelines and consensus reports and selected randomized controlled trials^{42,86}.

metaplasia and in those with persistent *H. pylori* gastritis. These recommendations are primarily intended for regions with low-to-moderate gastric cancer burden, where population-based screening is not practical or economically feasible but where subgroups at risk can be identified. Although the American Gastroenterological Association does not recommend routine use of endoscopic surveillance in patients with intestinal metaplasia²⁴², a common denominator between American Gastroenterological Association and European MAPS II guidelines is that they are based on low-quality evidence, highlighting the need for well-designed, large and long-term trials.

Management

General aspects

H. pylori gastritis is an infectious disease and all adult individuals with the infection require therapy for cure if clinical symptoms and complications are present or for prevention if at risk for complications even if asymptomatic^{1,42,243}. *H. pylori* test-and-treat strategies are selected according to diverse clinical scenarios^{17,42,206} (Box 2). In the paediatric population, *H. pylori* infection rarely leads to complications and requires specific management addressed in the joint ESPGHAN/ NASPGHAN Guidelines that were updated in 2016 (ref. 169). All treatment discussions in this section relate to the disease in adults.

Treatment regimens for *H. pylori* eradication are based on the combination of a strong acid suppressant and antibiotics. First-line therapy is selected according to locoregional or individual *H. pylori* antibiotic resistance patterns^{244,245}. Treatment failures induce resistance to several of the antibiotics used in first-line regimens and render further therapies more complex and costly^{42,206,246,247}. Second-line therapy needs to consider the first-line regimen and antibiotic resistance status (Fig. 6). Confirmation of treatment success not earlier than 4 weeks after end of therapy is mandatory to guide further management and provides important information on the effectiveness of treatment regimens in defined regions⁴².

PPI triple therapy

First-line setting. The introduction of PPI-based triple therapies (PPI-TT) marked a turning point in the treatment of H. pylori infection owing to their superior efficacy compared with previous approaches. The three components of PPI-TT include a PPI, clarithromycin and amoxicillin or, alternatively, metronidazole as a substitute for either amoxicillin or clarithromycin. Seven-day PPI-TT obtained initial eradication rates of >90%^{248,249} and, between 1997 and 2005, became the most widely recommended first-line therapy globally^{42,206,247,250}. Treatment duration has since been recommended to be extended to 14 days owing to a substantially higher efficacy compared to the 7-day duration 42,244,247. Antibiotics used in first-line PPI-TT are clarithromycin, amoxicillin and metronidazole or, more restrictive, levofloxacin and, in selected cases, furazolidone. Treatment failures with PPI-TT occur with increasing frequency and are primarily related to antibiotic resistance, insufficient acid suppression and inadequate adherence to medications^{10,251-253}. Acid suppression with PPI (omeprazole, esomeprazole, lansoprazole, pantoprazole or rabeprazole in double standard dose) is essential and aims to raise intragastric pH to 6 or higher, which optimizes the stability, bioavailability and efficacy of antibiotics^{254,255}. A modestly higher acid-inhibiting effect is shown for second-generation PPIs (esomeprazole, rabeprazole)²⁵⁶. Increased intragastric pH (optimum pH >6) enables bacterial replication, which increases the susceptibility of H. pylori to antibiotics. This is particularly important for amoxicillin, which is highly acid sensitive^{254,255}. Less effective acid suppressants, such as histamine 2 receptor antagonists, are no longer considered in H. pylori eradication regimens^{246,257}. PPI efficacy is further increased by doubling the PPI standard dose and should always be considered if first-line therapy fails^{258–261}.

Rapid metabolization of PPIs leads to reduced efficacy^{253,262,263}. Rapid and ultrarapid metabolization of PPIs varies considerably among ethnic groups and occurs more frequently in white and African American populations, whereas slow metabolization is more frequent in Asian, including lapanese and Chinese, populations^{264–267}. The efficacy of PPI metabolism depends on various genetic mutations related to CYP2C19 polymorphism and, to a minor extent, on CYP3A4 and gastric H⁺,K⁺-ATPase genotypes^{255,268}. Apart from the use of PPI double standard dose in rapid metabolizers, better control of acidity has been reported by increasing PPI dosing frequency up to four times or by switching to a PPI less influenced by CYP2C19 genotypes²⁶⁸⁻²⁷¹. No guideline is yet available recommending CYP2C19 genotyping to guide PPI prescription in clinical practice. The pharmacokinetic and pharmacodynamic properties of the second-generation PPIs esomeprazole and, in particular, rabeprazole are less influenced by variant CYP2C19 genotypes^{268,270,272,273}.

To overcome inadequate adherence, careful patient instruction is appropriate on how to take medications and how to proceed in case of adverse events^{274,275}. A history of penicillin allergy, availability of susceptibility testing, local prevalence of antibiotic resistance and history of prior eradication therapies should be considered when deciding on the initial therapy (Fig. 6).

Antibiotic resistance. Antibiotic resistance is the most important factor in PPI-TT failure²⁴⁴. Clarithromycin resistance and metronidazole resistance are the most relevant resistances for PPI-TT failure^{42,247,276}.

Clarithromycin resistance has increased from 3% to 11% around the turn of the century and is now up to 15–30% worldwide^{188,189,248,277-279}. In 2,852 treatment-naive patients from a European registry on *H. pylori* management (Hp-EuReg), resistance to clarithromycin, metronidazole and levofloxacin were 25%, 30% and 20%, respectively^{261,278}. Resistances to tetracycline and amoxicillin were <1% in the same study. A WHO global priority list qualifies clarithromycin-resistant *H. pylori* infection as a high threat among community-acquired infections²⁸⁰, and international guidelines recommend abandoning clarithromycin-based regimens if regional resistance exceeds 15%²⁴⁴.

Among several modifications developed to overcome clarithromycin resistance, including sequential therapy (PPI-dual followed by PPI-TT) and hybrid therapy (PPI plus three antibiotics), only concomitant therapy (PPI plus three antibiotics simultaneously administered)^{244,281-283} was found to be superior to clarithromycin-based PPI-TT^{281,282}. Concomitant therapy as an empirical first-line option should cautiously be considered in regions of clarithromycin resistance >15% and only used if individual AST or bismuth-based quadruple therapy (BiQT) are not locally available^{42,247,250}. Levofloxacin as a component of PPI-TT is effective in first-line and second-line regimens in regions with low levofloxacin resistance²⁸⁴⁻²⁸⁶. However, levofloxacin resistance is now up to 20% in Europe and 18% in the Asia-Pacific region^{188,278,287}. Although levofloxacin is not recommended as a first-line option, the high resistance restricts its use even in second-line regimens^{42,244,247}. AST before using levofloxacin in empirical second-line regimens is advised^{189,244,278,287}. Other quinolones, such as ciprofloxacin and moxifloxacin, which have reduced efficacy and/or less consistent results, are not an alternative to levofloxacin^{286,288}. Sitafloxacin-based triple and dual regimens that have been successfully tested in Japan²⁸⁹ are not used as an alternative to levofloxacin in western countries^{42,244,247}.

Metronidazole resistance is >25% in most areas of the world^{189,278} but has a minor effect on eradication efficacy when used in triple or quadruple regimens because of inconsistency between in vitro AST results and clinical efficacy and the synergism with co-administered drugs, in particular bismuth^{224,290,291}. Resistance to amoxicillin and tetracycline is low (<2%) and these antibiotics remain a key component in standard PPI-TT and in BiQT, respectively, without the need for routine AST^{244,291}. Rifabutin resistance is <1% and the *H. pylori* eradication rate of rifabutin-containing regimens is 73% according to a meta-analysis from 2020 (ref. 292). A rifabutin delayed-release preparation, combined with amoxicillin and omeprazole, obtained an eradication rate of 89%²⁹³ and FDA approval for use as a first-line therapy was granted in 2019 (ref. 294). Outside of the USA, rifabutin-containing regimens are recommended as rescue therapy only owing to the need of this drug for other critical infections and the risk of myelotoxicity in rare cases^{42,247}. Furazolidone resistance is <5% and the drug is effective in triple and quadruple combinations; its use is limited to a few countries in Asia and South America^{195,295} and it may serve as rescue therapy in individual cases²⁹⁶.

For these antibiotic classes, mechanisms of resistance are related to drug-specific target gene mutations (macrolides and quinolones)

Test	Sensitivity	Specificity	Clinical use	Comments	Refs.			
Invasive methods								
Rapid urease test	84-95%	95–100%	Important for initial diagnosis; testing two biopsy samples improves sensitivity; provides rapid results	PPIs need to be stopped 14 days before testing; current or recent antibiotic therapy needs to be excluded	86,179			
Microbial culture	76–90%	100%	Important for phenotypic susceptibility testing	Absolute specificity but costly; PPIs need to be stopped 14 days before testing; current or recent antibiotic therapy needs to be excluded	86,179			
Histological assessment	60-93%	>95%	Gold standard for diagnosis and assessment of mucosal changes	Based on updated Sydney system	86,373			
Molecular testing (PCR methods and FISH)	80-95%	100%	Useful in initial diagnosis and follow-up; provides rapid results	High sensitivity and specificity; useful in gastrointestinal bleeding, virulence typing and detection of antibiotic resistance	86,179, 373–375			
Non-invasive methods								
UBT	95–100%	95–100%	Gold standard for non-invasive diagnosis; higher sensitivity and specificity than stool antigen test and serological assessment; for initial diagnosis and follow-up	PPIs need to be stopped 14 days before testing; current or recent antibiotic therapy needs to be excluded	86,179,373			
Stool antigen test	>95%	>95%	Useful for initial diagnosis and follow-up; slightly lower sensitivity than UBT	Rapid, simple and inexpensive	86,373			
Serological antibody detection	74.4%	59%	Useful for initial diagnosis in specific cases	Cheap, simple and rapid; highly variable results; ideal for epidemiological purposes; no need to stop PPI and useful in patients with gastrointestinal bleeding; cannot distinguish between active and previous infection	86,179,376			

Table 3 | Diagnostic methods for H. pylori detection

FISH, fluorescence in situ hybridization; H. pylori, Helicobacter pylori; PPI, proton pump inhibitor; UBT, ¹³C-urea breath test.

Box 2

Test-and-treat or endoscopy-based diagnosis in clinical management of *H. pylori* infection

Test-and-treat

This strategy refers to non-invasive testing of patients with dyspeptic symptoms and without alarm symptoms, such as vomiting, weight loss or anaemia, at age 50 years (range 45–55 years because of increased individual risk of gastric cancer). The non-invasive ¹³C-urea breath test or stool antigen test are highly accurate in diagnosing current *Helicobacter pylori* infection³⁷⁶ and surrogate markers for the histological detection of *H. pylori* gastritis. Serious upper gastrointestinal lesions in patients with dyspepsia in this age group are very rare; thus, non-invasive testing as an initial management step is appropriate in areas of low or intermediate gastric cancer risk^{388,389}. Test-and-treat is superior to other management options, including empirical proton pump inhibitor therapy in patients with dyspepsia, and is more cost-effective than empirical therapy and endoscopy-based management^{390,391}.

Endoscopy-based diagnosis

This approach is required to exclude gastric preneoplastic conditions or malignant disease in patients with dyspeptic or other symptoms referred to the upper abdomen at age >50 years or at any age in the presence of alarm symptoms. A patient with

or to detoxication (nitroimidazoles)²⁹⁶. *H. pylori* colonies may carry single-drug, multidrug or hetero resistance^{296,297}. Isolates from antrum and corpus are reported to differ by up to 15% in AST, which may account for treatment failure if biopsy samples for AST are only taken from a single site in the stomach^{298,299}.

Bismuth-based quadruple therapy

Bismuth has multiple beneficial properties in peptic ulcer healing that include a stimulating effect on prostaglandin synthesis, inactivation of pepsin, and bile acid binding but, most relevant in *H. pylori* eradication, is its bactericidal effect^{300,301}. Bismuth subcitrate upregulates the expression of genes involved in *H. pylori* growth and metabolism and impedes proton entry, thereby preventing lowering of the bacterial cytoplasmic pH. These mechanisms are suggested to render antibiotics more effective³⁰². Bismuth-based quadruple therapy (BiQT; PPI, bismuth, tetracycline and a nitroimidazole antibiotic), available either as individual components or as PPI plus a capsule containing all antibacterial components, has an eradication efficacy of 90%^{261,291,303,304}.

BiQT is recommended as an empirical first-line therapy, does not require AST, is not affected by clarithromycin resistance and overcomes metronidazole resistance owing to synergism with bismuth, as documented by its consistently high therapeutic efficacy^{244,291,300}. Bismuth added to clarithromycin-containing regimens increases eradication efficacy also in the presence of clarithromycin resistance but, in these combinations, offers no advantage over standard BiQT^{300,305,306}. BiQT does not contain antibiotics that are essential for cure of other symptoms related to ulcerogenic drug (NSAIDs) use should also be considered for endoscopy^{70,392}. Endoscopy-based investigations are the most reassuring and should be considered in patients with anxiety³⁹³.

Test-and-treat for gastric cancer prevention

This strategy targets asymptomatic individuals at increased risk of gastric cancer owing to a first-degree relative with this malignancy. The non-invasive ¹³C-urea breath test or stool antigen test are appropriate for younger adults. Endoscopy-based investigations should be considered in individuals >45 years of age or earlier according to the age at which gastric cancer was diagnosed in the index patient³⁹⁴.

Population-based test-and-treat

This approach is recommended in regions with a high gastric cancer incidence. For this purpose, serological assessment combining the detection of anti-*H. pylori* antibodies with measurement of pepsinogen levels provides useful information on the aetiology and atrophy stage of chronic gastritis and helps direct further management of the disease^{17,395,396}.

infections. BiQT is an effective rescue option with a success rate of >90% following previous treatment failures 303,307,308 .

Regimens with potassium-competitive acid blockers

Potassium-competitive acid blockers (P-CABs), a new class of acid inhibitors, have a more potent and durable effect on acid suppression than PPIs^{309,310}. Vonoprazan-based triple therapy (V-TT) with clarithro-mycin and amoxicillin in first-line achieved an eradication rate of 92.6% versus 75.9% with PPI-TT, and 98% in second-line in Japan³¹¹, which was also confirmed in western countries³¹². In network meta-analyses, V-TT ranked best among all current first-line empirical therapies, which was also confirmed after the inclusion of a trial conducted in western countries^{313,314}. Vonoprazan dual therapy, consisting of vonoprazan plus amoxicillin, provides an eradication rate of *H. pylori* similar to that of V-TT²⁵². Increasing resistance and absence of new antibiotics set major expectations on P-CAB-based regimens, which are being investigated in several trials³⁰⁹ (Supplementary Table 1).

H. pylori eradication and rescue therapies

Management of refractory *H. pylori* needs to consider individual or local antibiotic resistance, facilities for AST, logistics, and drug availability^{244,246} (Fig. 6). Following PPI-TT failure, BiQT or a regimen with antibiotics selected following AST is recommended^{42,244,247}. Empirical therapy, with careful consideration of previously taken medications, is a valid alternative to genotypic resistance-guided therapy of refractory *H. pylori* infection³¹⁵. BiQT is currently the best empirical approach as it is not influenced by antibiotic resistance³¹⁶. If BiQT fails

in first-line, levofloxacin-based triple therapy is recommended. A metaanalysis including 25 trials with levofloxacin-based triple therapy in second-line treatment reports a cumulative eradication rate of 74.5% (95% CI 70.9–77.8)³¹⁷. The PPI-amoxicillin, high-dose, dual therapy is another option, with a \geq 81% eradication rate achieved as second-line treatment and with an efficacy comparable to other recommended therapies^{318,319}.

P-CABs in triple therapies and in dual combination with amoxicillin already effectively used in Asian countries will become an important option once generally available and properly adapted to regional demands in first-line and second-line eradication regimens^{311,313}. Rifabutin-based triple therapy is well documented as effective and should be kept as rescue therapy²⁹². Surveillance programmes at the regional level, the introduction of antibiotic stewardship, regulations in the use of antimicrobials and increased public awareness are advised to control the increasing resistance of *H. pylori*^{244,320,321}.

Adverse events

Overall, eradication regimens have a favourable safety profile, with usually mild and very few severe adverse events. Mostly mild-to-moderate adverse effects occur in 30–70% of patients and include

taste disturbances, nausea, headache, diarrhoea and non-specific gastrointestinal symptoms with some variations according to type of eradication therapy^{282,291,322-324}.

Diarrhoea varies in prevalence from >1% to 15% according to definitions applied, the population treated and type of therapy³²⁵. The non-recording of adverse effects as primary criteria in clinical trials accounts for the high variations. Darkening of the tongue and faeces is characteristic of bismuth salts³²⁶. Antibiotics affect gut microbiota and lead to mostly transient dysbiosis, bacterial resistance and overgrowth of opportunistic pathogens; however, rarely of *Clostridioides difficile*^{40,325,327}.

Probiotics added to *H. pylori* therapies have a small and inconsistent effect on eradication rates but reduce adverse effects⁴², which has been shown in meta-analyses for individual probiotics as well as for mixtures^{322,328,329}. In a new randomized controlled study, *Saccharomyces boulardii* combined with a mixture of probiotic bacteria modestly increased the eradication efficacy and reduced adverse effects³³⁰, whereas *S. boulardii* alone had no effect on eradication but remained effective in reducing adverse effects such as severe diarrhoea³³¹. Defined probiotic mixtures have been shown to antagonize the harmful effects of antibiotics on the gut microbiota and their metabolic functions³²⁵.



^aIndividual AST should be performed; levofloxacin-based regimen can be used if *H. pylori* is susceptible or community resistance is <15%; otherwise, use rescue therapy.

Fig. 6 | **Suggested** *H. pylori* **therapy algorithm.** *Helicobacter pylori* therapy algorithm with the indication of regimens that consist of triple or quadruple combinations to be used in first-line and subsequently in case of failure. Proton pump inhibitors (PPIs) or, where available, potassium-competitive acid blockers are essential components for acid suppression to render antibiotics more effective. PPI can be substituted by potassium-competitive acid blockers where available. Antibiotics are selected according to individual antibiotic susceptibility testing (AST) or according to regional antibiotic susceptibility based on surveillance as well as according to local availability. Clarithromycin-based PPI

triple therapy (PPI-TT) is a first-line therapy if local clarithromycin resistance prevalence is <15%. If clarithromycin resistance exceeds 15% or is unknown, the recommended first-line regimen is BiQT (PPI, bismuth, tetracycline and a nitroimidazole antibiotic). Levofloxacin-based regimens are recommended as second-line treatments if a first-line regimen with BiQT fails. Levoflaxin-based regimens include amoxicillin and PPI. If levofloxacin resistance in regional surveillance exceeds 15%, it is advisable to directly select third-line or fourth-line regimens as rescue therapy. The fourth-line regimen (rescue therapy) consists of PPI, rifabutin and amoxicillin (or clarithromycin in case of penicillin allergy).

Effects on peptic ulcer and MALT lymphoma

Successful H. pylori eradication achieves ulcer healing rates of >90% and continued acid inhibition with PPI is not required for uncomplicated duodenal ulcer⁴². Gastric ulcer requires prolonged acid inhibition for healing and endoscopic follow-up is needed to ensure complete ulcer healing and to exclude underlying gastric malignancy³³². Management of bleeding peptic ulcers, both duodenal and gastric ulcers, requires immediate care by controlling and/or restoring cardiocirculatory and respiratory function and by performing emergency diagnostic endoscopic examination and endoscopic interventions according to standardized protocols^{333,334}. PPI treatment is continued until complete healing is endoscopically documented⁴⁴. *H. pylori* eradication should be initiated after the active bleeding phase is under control and oral nutrition can be resumed^{70,334}. Patients with *H. pylori* infection exposed to ulcerogenic medications, in particular NSAIDs, are at an increased risk of complications^{56,335} and benefit from *H. pylori* testing and treatment^{42,70}. Patients at high risk for rebleeding after *H. pylori* eradication, for example, those with continued NSAID use, require PPI maintenance therapy³³⁶.

H. pylori eradication is the standard-of-care initial therapy for MALT lymphomas in all stages and obtains 70–80% long-term remission in stage I disease^{337,338}. Eradication therapy in patients negative for *H. pylori* after exclusion of the infection with routine diagnostics obtains cure in 30% of patients and should, therefore, always be considered as a first management step³³⁹.

Quality of life

Despite the vast number of H. pylori treatment studies, surprisingly few investigations have measured quality of life (QoL) outcomes. Several different questionnaires have been used to ascertain QoL metrics across the spectrum of diseases associated with H. pylori, and results have shown that eradication of H. pylori can either improve or worsen QoL, which may depend on the type of treatment used³⁴⁰. In a study from Japan, participants were included to survey improvement of GERDrelated OoL measures following H. pylori treatment using a Japanese version of the QoL in reflux and dyspepsia score (QOLRAD-J) and Carlsson-Dent guestionnaires³⁴¹. GERD-related QoL scores improved following treatment and these were magnified among individuals with severe reflux symptoms. In another study, an 8-item Short-Form Health Survey and a modified Frequency Scale for GERD symptoms were used following H. pylori eradication³⁴²; QoL improved irrespective of treatment outcome. Finally, in a study from Thailand, patients with functional dyspepsia indicated that H. pylori infection, anxiety or depression were common, occurring in 23.3%, 23% and 7.3% of patients, respectively³⁴³. These findings suggest that eradication of *H. pylori* might not only improve functional dyspepsia but also potentially prevent the development of gastric cancer in some patients with functional dyspepsia by eliminating the chronic inflammatory process in the stomach.

In the UK, a randomized controlled study of QoL was conducted in 39,929 patients with dyspepsia following *H. pylori* therapy using a validated dyspepsia questionnaire and the psychological well-being index (PGWB) and reported no effect on QoL following therapy³⁴⁴. A further study in a smaller number of patients with functional dyspepsia similarly noted no improvement in QoL after eradication³⁴⁵. Other studies from Europe have reached different conclusions. In a study from Hungary, using the Functional Digestive Disorder Quality of Life system adapted from France to determine QoL in patients with functional dyspepsia, improvement of QoL was dependent on *H. pylori* therapy³⁴⁶. In a study from Croatia, the Gastrointestinal Symptom Rating Scale questionnaire was employed and improvement in QoL of patients with dyspepsia was found as early as 1 month into the year-long study³⁴⁷.

A group from Africa used the Short-Form Leeds Dyspepsia Questionnaire and the Short-Form Nepean Dyspepsia Index in health-care workers with dyspepsia and found reduced QoL in those with high dyspepsia prevalence³⁴⁸. In a study from Rwanda, dyspepsia was assessed within the general population using the Short-Form Nepean Dyspepsia Index questionnaire and noted improved QoL, which was dependent on *H. pylori* treatment³⁴⁹.

Finally, in a study examining potential detrimental consequences of eradication therapy, it was reported that patients with a duodenal ulcer in whom *H. pylori* eradication was successful were more likely to develop oesophagitis in the first year after treatment than those without *H. pylori* eradication³⁵⁰; however, in the subsequent 2 years, there was no difference between the groups. Collectively, these disparate results probably reflect differences in *H. pylori* treatment regimens, the QoL scoring systems used, varying genetic backgrounds of the host populations and differences in infecting *H. pylori* strains³⁵⁰.

Outlook

A vision for the future is to provide a healthy stomach free from H. pylori to all individuals. The expectation that H. pylori incidence will decrease to the point that the bacterium will disappear spontaneously within a foreseeable time frame is unlikely to occur³⁵¹. A populationwide test-and-treat strategy should therefore remain a consideration. This strategy could confer a health benefit with the prevention of H. pylori-related complications in a considerable number of individuals. However, logistic limitations, substantial health costs and risks related to the massive use of antibiotics with the fear of aggravating antibiotic resistance would be disadvantages. Thus, the identification of individuals and population subsets with a higher-than-average risk of gastric cancer should be, for now, the primary target in prevention strategies. This is the case for first-degree family members of patients with gastric cancer and populations in world regions with high gastric cancer incidence. This approach is supported by favourable cost-effectiveness and advised by expert consensus reports and guidelines^{17,86}. A new concept for comprehensive intrafamilial H. pylori management has been proposed for regions of high *H. pylori* prevalence³⁵². It advises actively proceeding with test-and-treat strategies in family members living in the same household as the index patient diagnosed with H. pylori based on the rationale of predominant intrafamilial spreading of the infection, mainly in childhood.

Antibiotic resistance, with its dramatic increase, demands new antimicrobial drugs that can specifically target *H. pylori* and avoid cross-resistance effects and induction of antibiotic resistance in *H. pylori* and other bacteria (Box 3). Colloidal bismuth subcitrate remains a candidate, with a direct bactericidal effect by inducing cellular swelling, vacuolization, structural degradation and cell wall eruption of *H. pylori*³⁵³. For now, bismuth has the limitation of cure rates of <20% if used as monotherapy³⁰¹.

Urease is an essential factor for the survival of *H. pylori* in the acidic gastric environment, producing ammonia and carbonic acid to neutralize the acidic surroundings³⁵⁴. However, urease is a complex target composed of two subunits with 12 active sites³⁵⁴. The deline-ated structure of the proton-gated urea channel of urease³⁵⁵ and the development of a drug that can block the rapid influx of urea into the bacterium may offer a solution to inhibit the activity of urease.

Inhibition of motility by blocking the flagellar function of *H. pylori* might be worth considering as a therapeutic target. *H. pylori* flagella

Box 3

Potential therapeutic targets for non-antibiotic drugs against *H. pylori* infection

Urease

Block the proton-gated urea channel, inhibit the activity of urease and block the production of urease.

Flagella

Inhibit motility, impair structure and production of flagella.

Adhesion factors

Reduce the adhesion of Helicobacter pylori to gastric mucosa.

Drug delivery into gastric mucus

Increase the delivery of antibiotics or new drugs into the firmly adherent mucus.

enable rapid movement through the viscous mucus, and disruption of a gene encoding cardiolipin synthase of *H. pylori* strain G27, shown in vitro, may abolish the biosynthesis of flagellum³⁵⁶. Furthermore, interference with the outer membrane proteins of *H. pylori* that confer adhesion to host glycans, mucins or gastric mucosa, such as BabA and SabA³⁵⁷, may impair *H. pylori* survival in the gastric mucosa. An anti-adhesion nanomedicine composed of *H. pylori*-mimicking outer membrane nanoparticles could compete with *H. pylori* and reduce its adhesion to gastric epithelial cells³⁵⁸.

Another approach is to increase the penetration of drugs into the gastric mucus layer, which includes the loosely adherent and firmly adherent mucus layers³⁵⁹. Conventional mucoadhesive particles usually attach to the loosely adherent layer only and are easily moved downstream to the lumen upon peristalsis³⁵⁹. Some potential mucuspenetrating polymeric nanoparticles can penetrate into the firmly adherent mucus layer³⁵⁹. Thus, delivery of selective antibiotics or other agents against *H. pylori* may be rendered more effective through the development of polymeric nanoparticles. H. pylori is a microaerobic organism and susceptible to increased oxygen levels, thus, the delivery of nanoparticles that include oxygen into the gastric mucus layer could render the bacterium vulnerable. In addition, multiple aspects of the interaction of H. pylori with the gastrointestinal microbiome are addressed by current research and will influence future research³⁶⁰⁻³⁶⁴. The exclusive property of *H. pylori* to colonize and infect the gastric mucosa affects the biodiversity of other gastric bacteria and their role in either enhancing or mitigating H. pylori-induced gastric inflammation; this needs to be further explored. H. pylori dominates the mucosa-associated community in the stomach but is less influential on the composition of bacterial communities in gastric juice³⁶⁵. In individuals with H. pylori infection, and even following eradication, the gastric microbial composition is dependent on the extent of gastric mucosal damage previously induced by H. pylori and gastric acidity^{366,367}. In patients with atrophic gastritis, bacterial clusters

comprising *Peptostreptococcus*, *Streptococcus*, *Parvimonas*, *Prevotella*, *Rothia* and *Granulicatella* with carcinogenetic potential become predominant, whereas the probiotic *Faecalibacterium prausnitzii* is depleted³⁶⁷. The development of targeted probiotics in the direction of antagonizing carcinogenic clusters in patients with atrophic gastritis following *H. pylori* eradication will serve an important purpose. It will be a further point of interest to understand the role of bacteria carried in the saliva and transiting the stomach and whether, in the absence of *H. pylori*, some of them become candidates for resilience and explain the entity of *H. pylori*-negative gastritis. The contributing role of bacteria in the development of severe gastritis into gastric cancer is shown in animal experiments and humans, and more insights are expected in this area^{161,368,369}. Vaccine development should be pursued and lessons from previous failures will help in new approaches¹⁷².

Field and challenge studies should both be conducted after identification of effective vaccine candidates. A vaccine for preventive and therapeutic purposes might consider multiple epitopes to direct the immune response towards essential *H. pylori* functions such as epithelial cell adherence, proliferation and survival in the specific gastric niche^{230,370}. Preliminary evidence shows that *H. pylori* reduces the efficacy of treatment with immune-checkpoint inhibitors in malignant diseases^{371,372}. If these findings are confirmed in further investigations, a test-and-treat strategy for *H. pylori* could become a requirement before starting immune therapies in oncological diseases. Immune mechanisms involved in this phenomenon deserve further exploration.

Published online: 20 April 2023

References

- Sugano, K. et al. Kyoto global consensus report on *Helicobacter pylori* gastritis. Gut **64**, 1353–1367 (2015).
- Suerbaum, S. & Michetti, P. Helicobacter pylori infection. N. Engl. J. Med. 347, 1175–1186 (2002).
- Peek, R. M. Jr & Blaser, M. J. Helicobacter pylori and gastrointestinal tract adenocarcinomas. Nat. Rev. Cancer 2, 28–37 (2002).
- Amieva, M. R. & El-Omar, E. M. Host-bacterial interactions in *Helicobacter pylori* infection. Gastroenterology 134, 306–323 (2008).
- van Amsterdam, K., van Vliet, A. H., Kusters, J. G. & van der Ende, A. Of microbe and man: determinants of *Helicobacter pylori*-related diseases. *FEMS Microbiol. Rev.* 30, 131–156 (2006).
- Kidd, M. & Modlin, I. M. A century of *Helicobacter pylori*: paradigms lost-paradigms regained. *Digestion* 59, 1–15 (1998).
- Warren, J. R. & Marshall, B. Unidentified curved bacilli on gastric epithelium in active chronic gastritis. *Lancet* 1, 1273–1275 (1983).
 Discovery of *H. pylori* that starts worldwide research leading to the cure of peptic
- ulcer disease and identification of the main risk factor for gastric cancer. 8. Marshall, B. J., Armstrong, J. A., McGechie, D. B. & Glancy, R. J. Attempt to fulfil Koch's
- postulates for pyloric Campylobacter. Med. J. Aust. **142**, 436–439 (1985).
 Morris, A. & Nicholson, G. Ingestion of Campylobacter pyloridis causes gastritis and raised fasting gastric pH. Am. J. Gastroenterol. **82**, 192–199 (1987).
- Malfertheiner, P., Link, A. & Selgrad, M. Helicobacter pylori: perspectives and time trends. Nat. Rev. Gastroenterol. Hepatol. 11, 628–638 (2014).
- Goodwin, C. S. & Armstrong, J. A. Microbiological aspects of *Helicobacter pylori* (Campylobacter pylori). Eur. J. Clin. Microbiol. Infect. Dis. 9, 1–13 (1990).
- Rauws, E. A. & Tytgat, G. N. Cure of duodenal ulcer associated with eradication of Helicobacter pylori. Lancet 335, 1233–1235 (1990).
 To our knowledge, first study to provide definitive proof that *H. pylori* eradication cures peptic ulcer disease.
- Van der Hulst, R. W. et al. Prevention of ulcer recurrence after eradication of Helicobacter pylori: a prospective long-term follow-up study. Gastroenterology 113, 1082–1086 (1997).
- Malfertheiner, P., Chan, F. K. & McColl, K. E. Peptic ulcer disease. Lancet 374, 1449–1461 (2009).
- 15. Pincock, S. Nobel Prize winners Robin Warren and Barry Marshall. Lancet 366, 1429 (2005).
- Hooi, J. K. Y. et al. Global prevalence of *Helicobacter pylori* infection: systematic review and meta-analysis. Gastroenterology **153**, 420–429 (2017).
- Liou, J. M. et al. Screening and eradication of *Helicobacter pylori* for gastric cancer prevention: the Taipei global consensus. *Gut* 69, 2093–2112 (2020).
- Plummer, M., Franceschi, S., Vignat, J., Forman, D. & de Martel, C. Global burden of gastric cancer attributable to *Helicobacter pylori*. Int. J. Cancer **136**, 487–490 (2015).

- Malaty, H. M. et al. Age at acquisition of *Helicobacter pylori* infection: a follow-up study from infancy to adulthood. *Lancet* 359, 931–935 (2002).
- Gao, L., Weck, M. N., Nieters, A. & Brenner, H. Inverse association between a proinflammatory genetic profile and *Helicobacter pylori* seropositivity among patients with chronic atrophic gastritis: enhanced elimination of the infection during disease progression? *Eur. J. Cancer* 45, 2860–2866 (2009).
- Parsonnet, J. et al. Helicobacter pylori infection and the risk of gastric carcinoma. N. Engl. J. Med. 325, 1127–1131 (1991).
- Eusebi, L. H., Zagari, R. M. & Bazzoli, F. Epidemiology of Helicobacter pylori infection. Helicobacter 19, 1–5 (2014).
- Park, J. S., Jun, J. S., Seo, J. H., Youn, H. S. & Rhee, K. H. Changing prevalence of Helicobacter pylori infection in children and adolescents. Clin. Exp. Pediatr. 64, 21–25 (2021).
- Yuan, C. et al. The global prevalence of and factors associated with Helicobacter pylori infection in children: a systematic review and meta-analysis. Lancet Child. Adolesc. Health 6, 185–194 (2022).
- Syam, A. F. et al. *Helicobacter pylori* in the Indonesian Malay's descendants might be imported from other ethnicities. *Gut Pathog.* 13, 36 (2021).
- 26. Liou, J. M. et al. IL-1B-511 C->T polymorphism is associated with increased host
- susceptibility to *Helicobacter pylori* infection in Chinese. *Helicobacter* 12, 142–149 (2007).
 Mayerle, J. et al. Identification of genetic loci associated with *Helicobacter pylori* serologic status. *JAMA* 309, 1912–1920 (2013).
- Lam, S. Y. et al. Toll-like receptor 1 locus re-examined in a genome-wide association study update on anti-*Helicobacter pylori* IgG titers. *Gastroenterology* **162**, 1705–1715 (2022).
- El-Omar, E. M. Genetic predisposition for *Helicobacter pylori* infection-the jury is still out! Gastroenterology 162, 1591–1593 (2022).
- Kayali, S. et al. *Helicobacter pylori*, transmission routes and recurrence of infection: state of the art. Acta Biomed. 89, 72–76 (2018).
- Brown, L. M. Helicobacter pylori: epidemiology and routes of transmission. Epidemiol. Rev. 22, 283–297 (2000).
- Fox, J. G. Non-human reservoirs of Helicobacter pylori. Aliment. Pharmacol. Ther. 9, 93–103 (1995).
- Parsonnet, J., Shmuely, H. & Haggerty, T. Fecal and oral shedding of *Helicobacter pylori* from healthy infected adults. JAMA 282, 2240–2245 (1999).
- Weyermann, M., Rothenbacher, D. & Brenner, H. Acquisition of *Helicobacter pylori* infection in early childhood: independent contributions of infected mothers, fathers, and siblings. *Am. J. Gastroenterol.* **104**, 182–189 (2009).
- Kivi, M. et al. Concordance of Helicobacter pylori strains within families. J. Clin. Microbiol. 41, 5604–5608 (2003).
- Georgopoulos, S. D. et al. *Helicobacter pylori* infection in spouses of patients with duodenal ulcers and comparison of ribosomal RNA gene patterns. *Gut* **39**, 634–638 (1996).
- Luman, W., Zhao, Y., Ng, H. S. & Ling, K. L. Helicobacter pylori infection is unlikely to be transmitted between partners: evidence from genotypic study in partners of infected patients. Eur. J. Gastroenterol. Hepatol. 14, 521–528 (2002).
- Schwarz, S. et al. Horizontal versus familial transmission of *Helicobacter pylori*. PLoS Pathog. 4, e1000180 (2008).
- Didelot, X. et al. Genomic evolution and transmission of Helicobacter pylori in two South African families. Proc. Natl Acad. Sci. USA 110, 13880–13885 (2013).
- Liou, J. M. et al. Long-term changes of gut microbiota, antibiotic resistance, and metabolic parameters after *Helicobacter pylori* eradication: a multicentre, open-label, randomised trial. *Lancet Infect. Dis.* 19, 1109–1120 (2019).
- Zhao, J. B. et al. Whole family-based *Helicobacter pylori* eradication is a superior strategy to single-infected patient treatment approach: a systematic review and meta-analysis. *Helicobacter* 26, e12793 (2021).
- Malfertheiner, P. et al. Management of *Helicobacter pylori* infection the Maastricht V/Florence consensus report. Gut 66, 6–30 (2017).
- Kuipers, E. J., Thijs, J. C. & Festen, H. P. The prevalence of Helicobacter pylori in peptic ulcer disease. Aliment. Pharmacol. Ther. 9, 59–69 (1995).
- 44. Lanas, A. & Chan, F. K. L. Peptic ulcer disease. Lancet 390, 613-624 (2017).
- Sipponen, P. et al. Cumulative 10-year risk of symptomatic duodenal and gastric ulcer in patients with or without chronic gastritis. A clinical follow-up study of 454 outpatients. Scand. J. Gastroenterol. 25, 966–973 (1990).
- Schottker, B., Adamu, M. A., Weck, M. N. & Brenner, H. Helicobacter pylori infection is strongly associated with gastric and duodenal ulcers in a large prospective study. *Clin. Gastroenterol. Hepatol.* **10**, 487–493.e1 (2012).
- Xia, B. et al. Trends in the prevalence of peptic ulcer disease and *Helicobacter pylori* infection in family physician-referred uninvestigated dyspeptic patients in Hong Kong. *Aliment. Pharmacol. Ther.* 22, 243–249 (2005).
- Perez-Aisa, M. A., Del Pino, D., Siles, M. & Lanas, A. Clinical trends in ulcer diagnosis in a population with high prevalence of *Helicobacter pylori* infection. *Aliment. Pharmacol. Ther.* 21, 65–72 (2005).
- Leow, A. H., Lim, Y. Y., Liew, W. C. & Goh, K. L. Time trends in upper gastrointestinal diseases and *Helicobacter pylori* infection in a multiracial Asian population — a 20-year experience over three time periods. *Aliment. Pharmacol. Ther.* 43, 831–837 (2016).
- Azhari, H. et al. The global incidence of peptic ulcer disease is decreasing since the turn of the 21st century: a study of the Organisation for Economic Co-operation and Development (OECD). Am. J. Gastroenterol. **117**, 1419–1427 (2022).

- Yamamichi, N. et al. Inverse time trends of peptic ulcer and reflux esophagitis show significant association with reduced prevalence of *Helicobacter pylori* infection. *Ann. Med.* 52, 506–514 (2020).
- Jiang, J. X. et al. Downward trend in the prevalence of *Helicobacter pylori* infections and corresponding frequent upper gastrointestinal diseases profile changes in Southeastern China between 2003 and 2012. Springerplus 5, 1601 (2016).
- Xie, X., Ren, K., Zhou, Z., Dang, C. & Zhang, H. The global, regional and national burden of peptic ulcer disease from 1990 to 2019: a population-based study. *BMC Gastroenterol.* 22, 58 (2022).
- Malfertheiner, P. & Schulz.C. Peptic ulcer: chapter closed? Dig. Dis. https://doi.org/10.1159/ 000505367 (2020).
- Huang, J. Q., Sridhar, S. & Hunt, R. H. Role of *Helicobacter pylori* infection and nonsteroidal anti-inflammatory drugs in peptic-ulcer disease: a meta-analysis. *Lancet* 359, 14–22 (2002).
- Venerito, M. et al. Contribution of *Helicobacter pylori* infection to the risk of peptic ulcer bleeding in patients on nonsteroidal anti-inflammatory drugs, antiplatelet agents, anticoagulants, corticosteroids and selective serotonin reuptake inhibitors. *Aliment. Pharmacol. Ther.* 47, 1464–1471 (2018).
- Rosenstock, S., Jorgensen, T., Bonnevie, O. & Andersen, L. Risk factors for peptic ulcer disease: a population based prospective cohort study comprising 2416 Danish adults. *Gut* 52, 186–193 (2003).
- Zagari, R. M. et al. Prevalence of upper gastrointestinal endoscopic findings in the community: a systematic review of studies in unselected samples of subjects. J. Gastroenterol. Hepatol. 31, 1527–1538 (2016).
- 59. Eslick, G. et al. Clinical and economic impact of "triple therapy" for *Helicobacter pylori* eradication on peptic ulcer disease in Australia. *Helicobacter* **25**, e12751 (2020).
- Sung, J., Kuipers, E. & El-Serag, H. Systematic review: the global incidence and prevalence of peptic ulcer disease. Aliment. Pharmacol. Ther. 29, 938–946 (2009).
- Moss, S. F. The clinical evidence linking Helicobacter pylori to gastric cancer. Cell Mol. Gastroenterol. Hepatol. 3, 183–191 (2017).
- de Martel, C., Georges, D., Bray, F., Ferlay, J. & Clifford, G. M. Global burden of cancer attributable to infections in 2018: a worldwide incidence analysis. *Lancet Glob. Health* 8, e180–e190 (2020).
- Fann, J. C. et al. Personalized risk assessment for dynamic transition of gastric neoplasms. J. Biomed. Sci. 25, 84 (2018).
- Arnold, M. et al. The burden of stomach cancer in indigenous populations: a systematic review and global assessment. Gut 63, 64–71 (2014).
- Kumar, S., Metz, D. C., Ellenberg, S., Kaplan, D. E. & Goldberg, D. S. Risk factors and incidence of gastric cancer after detection of *Helicobacter pylori* infection: a large cohort study. *Gastroenterology* 158, 527–536.e7 (2020).
- Gonzalez, C. A. & Lopez-Carrillo, L. Helicobacter pylori, nutrition and smoking interactions: their impact in gastric carcinogenesis. Scand. J. Gastroenterol. 45, 6–14 (2010).
- Venneman, K. et al. The epidemiology of *Helicobacter pylori* infection in Europe and the impact of lifestyle on its natural evolution toward stomach cancer after infection: a systematic review. *Helicobacter* 23, e12483 (2018).
- Wong, F., Rayner-Hartley, E. & Byrne, M. F. Extraintestinal manifestations of Helicobacter pylori: a concise review. World J. Gastroenterol. 20, 11950–11961 (2014).
- 69. Takeuchi, H. & Okamoto, A. *Helicobacter pylori* infection and chronic immune thrombocytopenia. *J. Clin. Med.* **11**, 4822 (2022).
- Malfertheiner, P. et al. Management of *Helicobacter pylori* infection the Maastricht IV/Florence Consensus Report. Gut 61, 646–664 (2012).
- Gasbarrini, A. et al. Regression of autoimmune thrombocytopenia after eradication of *Helicobacter pylori*. Lancet **352**, 878 (1998).
- Figura, N. et al. Extragastric manifestations of *Helicobacter pylori* infection. *Helicobacter* 15, 60–68 (2010).
- Franceschi, F., Zuccala, G., Roccarina, D. & Gasbarrini, A. Clinical effects of Helicobacter pylori outside the stomach. Nat. Rev. Gastroenterol. Hepatol. 11, 234–242 (2014).
- Gravina, A. G. et al. Extra-gastric manifestations of *Helicobacter pylori* infection. J. Clin. Med. 9, 3887 (2020).
- McCune, A. et al. Reduced risk of atopic disorders in adults with *Helicobacter pylori* infection. *Eur. J. Gastroenterol. Hepatol.* **15**, 637–640 (2003).
- Chen, Y. & Blaser, M. J. Inverse associations of Helicobacter pylori with asthma and allergy. Arch. Intern. Med. 167, 821–827 (2007).
- Blaser, M. J., Chen, Y. & Reibman, J. Does *Helicobacter pylori* protect against asthma and allergy? Gut 57, 561–567 (2008).
- Alvarez, C. S. et al. Associations of *Helicobacter pylori* and hepatitis A seropositivity with asthma in the Hispanic Community Health Study/Study of Latinos (HCHS/SOL): addressing the hygiene hypothesis. *Allergy Asthma Clin. Immunol.* **17**, 120 (2021).
- Rokkas, T., Pistiolas, D., Sechopoulos, P., Robotis, I. & Margantinis, G. Relationship between *Helicobacter pylori* infection and esophageal neoplasia: a meta-analysis. *Clin. Gastroenterol. Hepatol.* 5, 1413–1417 (2007).
- Fischbach, L. A. et al. The association between Barrett's esophagus and Helicobacter pylori infection: a meta-analysis. Helicobacter 17, 163–175 (2012).
- Rubenstein, J. H. et al. Association between *Helicobacter pylori* and Barrett's esophagus, erosive esophagitis, and gastroesophageal reflux symptoms. *Clin. Gastroenterol. Hepatol.* 12, 239–245 (2014).
- Vicari, J. J. et al. The seroprevalence of cagA-positive Helicobacter pylori strains in the spectrum of gastroesophageal reflux disease. Gastroenterology 115, 50–57 (1998).

- Doorakkers, E., Lagergren, J., Santoni, G., Engstrand, L. & Brusselaers, N. Helicobacter pylori eradication treatment and the risk of Barrett's esophagus and esophageal adenocarcinoma. Helicobacter 25, e12688 (2020).
- Wang, Z. et al. *Helicobacter pylori* infection is associated with reduced risk of Barrett's esophagus: an analysis of the Barrett's and esophageal adenocarcinoma consortium. *Arn. J. Gastroenterol.* **113**, 1148–1155 (2018).
- Zamani, M., Alizadeh-Tabari, S., Hasanpour, A. H., Eusebi, L. H. & Ford, A. C. Systematic review with meta-analysis: association of *Helicobacter pylori* infection with gastrooesophageal reflux and its complications. *Aliment. Pharmacol. Ther.* 54, 988–998 (2021).
- Malfertheiner, P. et al. Management of Helicobacter pylori infection: the Maastricht VI/ Florence consensus report. Gut https://doi.org/10.1136/gutinl-2022-327745 (2022).
- Tomb, J. F. et al. The complete genome sequence of the gastric pathogen Helicobacter pylori. Nature 388, 539–547 (1997); erratum 389, 412 (1997).
- Alm, R. A. et al. Genomic-sequence comparison of two unrelated isolates of the human gastric pathogen *Helicobacter pylori*. *Nature* **397**, 176–180 (1999).
- Gressmann, H. et al. Gain and loss of multiple genes during the evolution of *Helicobacter* pylori. PLoS Genet. 1, e43 (2005).
- Krebes, J. et al. The complex methylome of the human gastric pathogen Helicobacter pylori. Nucleic Acids Res. 42, 2415–2432 (2014).
- Ailloud, F. et al. Within-host evolution of *Helicobacter pylori* shaped by niche-specific adaptation, intragastric migrations and selective sweeps. *Nat. Commun.* 10, 2273 (2019).
- Jackson, L. K. et al. *Helicobacter pylori* diversification during chronic infection within a single host generates sub-populations with distinct phenotypes. *PLoS Pathog.* 16, e1008686 (2020).
- Suerbaum, S. & Josenhans, C. Helicobacter pylori evolution and phenotypic diversification in a changing host. Nat. Rev. Microbiol. 5, 441–452 (2007).
- Kang, J. & Blaser, M. J. Bacterial populations as perfect gases: genomic integrity and diversification tensions in *Helicobacter pylori*. Nat. Rev. Microbiol. 4, 826–836 (2006).
- Garcia-Ortiz, M. V. et al. Unexpected role for *Helicobacter pylori* DNA polymerase I as a source of genetic variability. *PLoS Genet.* 7, e1002152 (2011).
- Hofreuter, D., Odenbreit, S. & Haas, R. Natural transformation competence in Helicobacter pylori is mediated by the basic components of a type IV secretion system. Mol. Microbiol. 41, 379–391 (2001).
- Stingl, K., Muller, S., Scheidgen-Kleyboldt, G., Clausen, M. & Maier, B. Composite system mediates two-step DNA uptake into *Helicobacter pylori*. Proc. Natl Acad. Sci. USA 107, 1184–1189 (2009).
- Suerbaum, S. et al. Free recombination within Helicobacter pylori. Proc. Natl Acad. Sci. USA 95, 12619–12624 (1998).
- Kennemann, L. et al. *Helicobacter pylori* genome evolution during human infection. Proc. Natl Acad. Sci. USA 108, 5033–5038 (2011).
- Bubendorfer, S. et al. Genome-wide analysis of chromosomal import patterns after natural transformation of *Helicobacter pylori*. Nat. Commun. 7, 11995 (2016).
- Falush, D. et al. Traces of human migrations in *Helicobacter pylori* populations. Science 299, 1582–1585 (2003).
- 102. Linz, B. et al. An African origin for the intimate association between humans and Helicobacter pylori. Nature **445**, 915–918 (2007).
- Moodley, Y. et al. Age of the association between *Helicobacter pylori* and man. *PLoS Pathog.* 8, e1002693 (2012).
- 104. Ailloud, F., Estibariz, I. & Suerbaum, S. Evolved to vary: genome and epigenome variation in the human pathogen *Helicobacter pylori*. *FEMS Microbiol. Rev.* 45, fuaa042 (2021).
- Moodley, Y. et al. The peopling of the Pacific from a bacterial perspective. Science 323, 527–530 (2009).
- 106. Censini, S. et al. cag, a pathogenicity island of *Helicobacter pylori*, encodes type I-specific and disease-associated virulence factors. *Proc. Natl Acad. Sci. USA* 93, 14648–14653 (1996).
- Olbermann, P. et al. A global overview of the genetic and functional diversity in the Helicobacter pylori cag pathogenicity island. PLoS Genet. 6, e1001069 (2010).
- Johnson, K. S. & Otteman, K. M. Colonization, localization, and inflammation: the roles of H. pylori chemotaxis in vivo. Curr. Opin. Microbiol. 41, 51–57 (2018).
- 109. Josenhans, C., Labigne, A. & Suerbaum, S. Comparative ultrastructural and functional studies of *Helicobacter pylori* and *Helicobacter mustelae* flagellin mutants: both flagellin subunits, FlaA and FlaB, are necessary for full motility in *Helicobacter* species. *J. Bacteriol.* **177**, 3010–3020 (1995).
- Lee, S. K. et al. *Helicobacter pylori* flagellins have very low intrinsic activity to stimulate human gastric epithelial cells via TLR5. *Microbes Infect.* 5, 1345–1356 (2003).
- Andersen-Nissen, E. et al. Evasion of Toll-like receptor 5 by flagellated bacteria. Proc. Natl Acad. Sci. USA 102, 9247–9252 (2005).
- Schreiber, S. et al. The spatial orientation of *Helicobacter pylori* in the gastric mucus. Proc. Natl Acad. Sci. USA 101, 5024–5029 (2004).
- Suerbaum, S. et al. Identification of antimotilins, novel inhibitors of *Helicobacter pylori* flagellar motility that inhibit stomach colonization in a mouse model. *mBio* 13, e0375521 (2022).
- Mobley, H. L. in *Helicobacter pylori: Molecular and Cellular Biology* (eds Achtman, M. & Suerbaum, S.) (Horizon Scientific Press, 2001).
- Weeks, D. L., Eskandari, S., Scott, D. R. & Sachs, G. A H⁺-gated urea channel: the link between *Helicobacter pylori* urease and gastric colonization. *Science* 287, 482–485 (2000).
- Eaton, K. A., Brooks, C. L., Morgan, D. R. & Krakowka, S. Essential role of urease in pathogenesis of gastritis induced by *Helicobacter pylori* in gnotobiotic piglets. *Infect. Immun.* 59, 2470–2475 (1991).

- Kirschner, D. E. & Blaser, M. J. The dynamics of *Helicobacter pylori* infection of the human stomach. J. Theor. Biol. **176**, 281–290 (1995).
- Borén, T., Falk, P., Roth, K. A., Larson, G. & Normark, S. Attachment of *Helicobacter pylori* to human gastric epithelium mediated by blood group antigens. *Science* 262, 1892–1895 (1993).
- Mahdavi, J. et al. *Helicobacter pylori* SabA adhesin in persistent infection and chronic inflammation. Science **297**, 573–578 (2002).
- Javaheri, A. et al. *Helicobacter pylori* adhesin HopQ engages in a virulence-enhancing interaction with human CEACAMs. *Nat. Microbiol.* 2, 16189 (2016).
- Koniger, V. et al. *Helicobacter pylori* exploits human CEACAMs via HopQ for adherence and translocation of CagA. *Nat. Microbiol.* 2, 16188 (2016).
- Senkovich, O. A. et al. *Helicobacter pylori* AlpA and AlpB bind host laminin and influence gastric inflammation in gerbils. *Infect. Immun.* 79, 3106–3116 (2011).
- Odenbreit, S. et al. Translocation of *Helicobacter pylori* CagA into gastric epithelial cells by type IV secretion. Science 287, 1497–1500 (2000).
- 124. Stein, S. C. et al. *Helicobacter pylori* modulates host cell responses by CagT4SSdependent translocation of an intermediate metabolite of LPS inner core heptose biosynthesis. *PLoS Pathog.* **13**, e1006514 (2017).
- Pfannkuch, L. et al. ADP heptose, a novel pathogen-associated molecular pattern identified in Helicobacter pylori. FASEB J. 33, 9087–9099 (2019).
- Viala, J. et al. Nod1 responds to peptidoglycan delivered by the *Helicobacter pylori cag* pathogenicity island. *Nat. Immunol.* 5, 1166–1174 (2004).
- Varga, M. G. et al. Pathogenic Helicobacter pylori strains translocate DNA and activate TLR9 via the cancer-associated cag type IV secretion system. Oncogene 35, 6262–6269 (2016).
- Asahi, M. et al. *Helicobacter pylori* CagA protein can be tyrosine phosphorylated in gastric epithelial cells. J. Exp. Med. **191**, 593–602 (2000).
- Higashi, H. et al. SHP-2 tyrosine phosphatase as an intracellular target of *Helicobacter* pylori CagA protein. Science 295, 683–686 (2002).
- Saadat, I. et al. Helicobacter pylori CagA targets PAR1/MARK kinase to disrupt epithelial cell polarity. Nature 447, 330–333 (2007).
- Nesic, D. et al. Helicobacter pylori CagA inhibits PARI-MARK family kinases by mimicking host substrates. Nat. Struct. Mol. Biol. 17, 130–132 (2010).
- Buti, L. et al. *Helicobacter pylori* cytotoxin-associated gene A (CagA) subverts the apoptosis-stimulating protein of p53 (ASPP2) tumor suppressor pathway of the host. *Proc. Natl Acad. Sci. USA* **108**, 9238–9243 (2011).
- Ding, S. Z., Goldberg, J. B. & Hatakeyama, M. Helicobacter pylori infection, oncogenic pathways and epigenetic mechanisms in gastric carcinogenesis. *Future Oncol.* 6, 851–862 (2010).
- Bauer, M. et al. The ALPK1/TIFA/NF-kB axis links a bacterial carcinogen to R-loop-induced replication stress. Nat. Commun. 11, 5117 (2020).
- Fass, L. et al. Contribution of heptose metabolites and the cag pathogenicity island to the activation of monocytes/macrophages by *Helicobacter pylori*. Front. Immunol. 12, 632154 (2021).
- Coletta, S. et al. ADP-heptose enables *Helicobacter pylori* to exploit macrophages as a survival niche by suppressing antigen-presenting HLA-II expression. *FEBS Lett.* 595, 2160–2168 (2021).
- Cover, T. L. & Blaser, M. J. Purification and characterization of the vacuolating toxin from Helicobacter pylori. J. Biol. Chem. 267, 10570–10575 (1992).
- Cover, T. L. & Blanke, S. R. Helicobacter pylori VacA, a paradigm for toxin multifunctionality. Nat. Rev. Microbiol. 3, 320–332 (2005).
- Foegeding, N. J., Caston, R. R., McClain, M. S., Ohi, M. D. & Cover, T. L. An overview of *Helicobacter pylori* VacA toxin biology. *Toxins* 8, 173 (2016).
- Altobelli, A., Bauer, M., Velez, K., Cover, T. L. & Muller, A. Helicobacter pylori VacA targets myeloid cells in the gastric lamina propria to promote peripherally induced regulatory T-cell differentiation and persistent infection. mBio 10, e00261-19 (2019).
- Zhang, X., Arnold, I. C. & Muller, A. Mechanisms of persistence, innate immune activation and immunomodulation by the gastric pathogen *Helicobacter pylori. Curr. Opin. Microbiol.* 54, 1–10 (2020).
- Gobert, A. P. & Wilson, K. T. Induction and regulation of the innate immune response in Helicobacter pylori infection. Cell Mol. Gastroenterol. Hepatol. 13, 1347–1363 (2022).
- 143. Faass, L., Hauke, M., Stein, S. C. & Josenhans, C. Innate immune activation and modulatory factors of *Helicobacter pylori* towards phagocytic and nonphagocytic cells. *Curr. Opin. Immunol.* 82, 102301 (2023).
- de Bernard, M. & Josenhans, C. Pathogenesis of Helicobacter pylori infection. Helicobacter 19, 11–18 (2014).
- Crabtree, J. E. et al. Interleukin-8 expression in *Helicobacter pylori* infected, normal, and neoplastic gastroduodenal mucosa. J. Clin. Pathol. 47, 61–66 (1994).
- 146. Sharma, S. A., Tummuru, M. K., Blaser, M. J. & Kerr, L. D. Activation of IL-8 gene expression by *Helicobacter pylori* is regulated by transcription factor nuclear factor-kappa B in gastric epithelial cells. *J. Immunol.* **160**, 2401–2407 (1998).
- Maubach, G., Vieth, M., Boccellato, F. & Naumann, M. Helicobacter pylori-induced NF-κB: trailblazer for gastric pathophysiology. Trends Mol. Med. 28, 210–222 (2022).
- Rugge, M., Savarino, E., Sbaraglia, M., Bricca, L. & Malfertheiner, P. Gastritis: the clinico-pathological spectrum. *Dig. Liver Dis.* 53, 1237–1246 (2021).
- 149. Sipponen, P., Kekki, M. & Siurala, M. The Sydney System: epidemiology and natural history of chronic gastritis. J. Gastroenterol. Hepatol. 6, 244–251 (1991).
- Oertli, M. et al. DC-derived IL-18 drives Treg differentiation, murine *Helicobacter pylori*specific immune tolerance, and asthma protection. J. Clin. Invest. **122**, 1082–1096 (2012).

- Arshad, U., Sarkar, S., Alipour Talesh, G. & Sutton, P. A lack of role for antibodies in regulating *Helicobacter pylori* colonization and associated gastritis. *Helicobacter* 25, e12681 (2020).
- 152. Ermak, T. H. et al. Immunization of mice with urease vaccine affords protection against Helicobacter pylori infection in the absence of antibodies and is mediated by MHC class II-restricted responses. J. Exp. Med. 188, 2277–2288 (1998).
- D'Elios, M. M. & Czinn, S. J. Immunity, inflammation, and vaccines for Helicobacter pylori. Helicobacter 19, 19–26 (2014).
- Arnold, I. C. et al. *Helicobacter pylori* infection prevents allergic asthma in mouse models through the induction of regulatory T cells. *J. Clin. Invest.* **121**, 3088–3093 (2011).
- 155. Kyburz, A. et al. Transmaternal *Helicobacter pylori* exposure reduces allergic airway inflammation in offspring through regulatory T cells. J. Allergy Clin. Immunol. **143**, 1496–1512.e11 (2019).
- Cook, K. W. et al. CCL20/CCR6-mediated migration of regulatory T cells to the Helicobacter pylori-infected human gastric mucosa. Gut 63, 1550–1559 (2014).
 Robinson K. et al. Helicobacter pylori-induced pentic ulcer disease is associated
- Robinson, K. et al. *Helicobacter pylori*-induced peptic ulcer disease is associated with inadequate regulatory T cell responses. *Gut* 57, 1375–1385 (2008).
 Backert, S., Haas, R., Gerhard, M. & Naumann, M. The *Helicobacter pylori* type IV
- secretion system encoded by the cag pathogenicity island: architecture, function, and signaling. *Curr. Top. Microbiol. Immunol.* **413**, 187–220 (2017).
- Correa, P. Human gastric carcinogenesis: a multistep and multifactorial process. First American Cancer Society award lecture on cancer epidemiology and prevention. *Cancer Res.* 52, 6735 (1992).
- Ferreira, R. M. et al. Gastric microbial community profiling reveals a dysbiotic cancer-associated microbiota. Gut 67, 226–236 (2018).
- 161. Guo, Y. et al. Effect of *Helicobacter pylori* on gastrointestinal microbiota: a population-based study in Linqu, a high-risk area of gastric cancer. Gut 69, 1598–1607 (2020).
- Kwon, S. K. et al. Human gastric microbiota transplantation recapitulates premalignant lesions in germ-free mice. Gut 71, 1266–1276 (2022).
- 163. Pereira-Marques, J., Ferreira, R. M., Machado, J. C. & Figueiredo, C. The influence of the gastric microbiota in gastric cancer development. *Best Pract. Res. Clin. Gastroenterol.* 50–51, 101734 (2021).
- Bayerdorffer, E. et al. Regression of primary gastric lymphoma of mucosa-associated lymphoid tissue type after cure of *Helicobacter pylori* infection. MALT Lymphoma Study Group. *Lancet* 345, 1591–1594 (1995).
- 165. Hunt, R. H. et al. The stomach in health and disease. Gut 64, 1650-1668 (2015).
- El-Omar, E. M. et al. *Helicobacter pylori* infection and chronic gastric acid hyposecretion. Gastroenterology **113**, 15–24 (1997).
- Amieva, M. & Peek, R. M. Jr. Pathobiology of *Helicobacter pylori*-induced gastric cancer. Gastroenterology 150, 64–78 (2016).
- Kwon, S. K. et al. Human gastric microbiota transplantation recapitulates premalignant lesions in germ-free mice. Gut 71, 1266–1276 (2021).
- Jones, N. L. et al. Joint ESPGHAN/NASPGHAN guidelines for the management of Helicobacter pylori in children and adolescents (Update 2016). J. Pediatr. Gastroenterol. Nutr. 64, 991–1003 (2017).
- Sobala, G. M. et al. Acute *Helicobacter pylori* infection: clinical features, local and systemic immune response, gastric mucosal histology, and gastric juice ascorbic acid concentrations. *Gut* **32**, 1415–1418 (1991).
- 171. Graham, D. Y. et al. Challenge model for *Helicobacter pylori* infection in human volunteers. *Gut* **53**, 1235–1243 (2004).
- Malfertheiner, P. et al. Efficacy, immunogenicity, and safety of a parenteral vaccine against *Helicobacter pylori* in healthy volunteers challenged with a Cag-positive strain: a randomised, placebo-controlled phase 1/2 study. *Lancet Gastroenterol. Hepatol.* 3, 698–707 (2018).
- Spee, L. A., Madderom, M. B., Pijpers, M., van Leeuwen, Y. & Berger, M. Y. Association between *Helicobacter pylori* and gastrointestinal symptoms in children. *Pediatrics* 125, e651–e669 (2010).
- 174. Fischbach, W., Goebeler-Kolve, M. E., Dragosics, B., Greiner, A. & Stolte, M. Long term outcome of patients with gastric marginal zone B cell lymphoma of mucosa associated lymphoid tissue (MALT) following exclusive *Helicobacter pylori* eradication therapy: experience from a large prospective series. *Gut* 53, 34–37 (2004).
- Malfertheiner, P. Diagnostic methods for H. pylori infection: choices, opportunities and pitfalls. United European Gastroenterol. J. 3, 429–431 (2015).
- Pilotto, A. & Franceschi, M. Helicobacter pylori infection in older people. World J. Gastroenterol. 20, 6364–6373 (2014).
- Uotani, T. & Graham, D. Y. Diagnosis of *Helicobacter pylori* using the rapid urease test. Ann. Transl Med. 3, 9 (2015).
- Smith, S. I. et al. Helicobacter pylori infection in Africa: update of the current situation and challenges. Dig. Dis. 40, 535–544 (2021).
- Bordin, D. S., Voynovan, I. N. & Andreev, D. N. Maev IV. Current Helicobacter pylori diagnostics. Diagnostics 11, 1458 (2021).
- Miftahussurur, M. & Yamaoka, Y. Diagnostic methods of *Helicobacter pylori* infection for epidemiological studies: critical importance of indirect test validation. *Biomed. Res. Int.* 2016, 4819423 (2016).
- Talebi Bezmin Abadi, A. Diagnosis of *Helicobacter pylori* using invasive and noninvasive approaches. J. Pathog. **2018**, 9064952 (2018).

- Dixon, M. F., Genta, R. M., Yardley, J. H. & Correa, P. Classification and grading of gastritis. The updated Sydney System. International Workshop on the Histopathology of Gastritis, Houston 1994. Am. J. Surg. Pathol. 20, 1161–1181 (1996).
- Rugge, M. et al. Gastritis staging in clinical practice: the OLGA staging system. Gut 56, 631–636 (2007).
- Capelle, L. G. et al. The staging of gastritis with the OLGA system by using intestinal metaplasia as an accurate alternative for atrophic gastritis. *Gastrointest. Endosc.* 71, 1150–1158 (2010).
- Ajayi, A., Jolaiya, T. & Smith, S. I. Direct detection of *Helicobacter pylori* from biopsies of patients in Lagos, Nigeria using real-time PCR-a pilot study. *BMC Res. Notes* 14, 90 (2021).
- Moss, S. F. et al. Comparable results of *Helicobacter pylori* antibiotic resistance testing of stools vs gastric biopsies using next-generation sequencing. *Gastroenterology* 162, 2095–2097.e2 (2022).
- Schulz, C., Kalali, B., Link, A., Gerhard, M. & Malfertheiner, P. New rapid Helicobacter pylori blood test based on dual detection of FliD and CagA antibodies for on-site testing. *Clin. Gastroenterol. Hepatol.* 21, 229–231.e1 (2021).
- Megraud, F. et al. Helicobacter pylori resistance to antibiotics in Europe in 2018 and its relationship to antibiotic consumption in the community. Gut 70, 1815–1822 (2021).
- 189. Savoldi, A., Carrara, E., Graham, D. Y., Conti, M. & Tacconelli, E. Prevalence of antibiotic resistance in *Helicobacter pylori*: a systematic review and meta-analysis in World Health Organization Regions. *Gastroenterology* **155**, 1372–1382.e17 (2018).
- Hulten, K. G. et al. Comparison of culture with antibiogram to next-generation sequencing using bacterial isolates and formalin-fixed, paraffin-embedded gastric biopsies. Gastroenterology 161, 1433–1442.e2 (2021).
- Argueta, A. E., Alsamman, M. A., Moss, S. F. & D'Agata, E. M. C. Impact of antimicrobial resistance rates on eradication of *Helicobacter pylori* in a US population. *Gastroenterology* **160**, 2181–2183.e1 (2021).
- David, Y. G. & Steven, F. M. Antimicrobial susceptibility testing for *Helicobacter pylori* is now widely available: when, how, why. *Am. J. Gastroenterol.* 117, 524–528 (2022).
- Hu, Y., Zhang, M., Lu, B. & Dai, J. Helicobacter pylori and antibiotic resistance, a continuing and intractable problem. Helicobacter 21, 349–363 (2016).
- 194. Egli, K. et al. Comparison of the diagnostic performance of qPCR, sanger sequencing, and whole-genome sequencing in determining clarithromycin and levofloxacin resistance in *Helicobacter pylori. Front. Cell Infect. Microbiol.* **10**, 596371 (2020).
- Zamani, M., Rahbar, A. & Shokri-Shirvani, J. Resistance of Helicobacter pylori to furazolidone and levofloxacin: a viewpoint. World J. Gastroenterol. 23, 6920–6922 (2017).
- Wang, Y. H. et al. A systematic review and meta-analysis of genotypic methods for detecting antibiotic resistance in *Helicobacter pylori*. *Helicobacter* 23, e12467 (2018).
- Li, Y. et al. Detection of clarithromycin resistance in *Helicobacter pylori* following noncryogenic storage of rapid urease tests for 30 days. J. Dig. Dis. 13, 54–59 (2012).
- Chung, W. C. et al. Dual-priming oligonucleotide-based multiplex PCR using tissue samples in rapid urease test in the detection of *Helicobacter pylori* infection. *World J. Gastroenterol.* 20, 6547–6553 (2014).
- Chung, W. C. et al. Dual-priming oligonucleotide-based multiplex PCR using tissue samples from the rapid urease test kit for the detection of *Helicobacter pylori* in bleeding peptic ulcers. *Dig. Liver Dis.* 48, 899–903 (2016).
- 200. Chen, T., Meng, X., Zhang, H., Tsang, R. W. & Tsang, T. K. Comparing multiplex PCR and rapid urease test in the detection of *H. pylori* in patients on proton pump inhibitors. *Gastroenterol. Res. Pract.* **2012**, 898276 (2012).
- Goji, S. et al. Helicobacter suis-infected nodular gastritis and a review of diagnostic sensitivity for *Helicobacter heilmannii*-like organisms. *Case Rep. Gastroenterol.* 9, 179–187 (2015).
- Kobayashi, M. et al. Helicobacter heilmannii-like organisms in parietal cells: a diagnostic pitfall. Pathol. Int. 66, 120–122 (2016).
- De Witte, C., Schulz, C., Smet, A., Malfertheiner, P. & Haesebrouck, F. Other Helicobacters and gastric microbiota. *Helicobacter* 21 (Suppl. 1), 62–68 (2016).
- Seiichi, K. et al. The updated JSPGHAN guidelines for the management of Helicobacter pylori infection in childhood. Pediatr. Int. 62, 1315–1331 (2020).
- Moayyedi, P. et al. Guideline: management of dyspepsia. Am. J. Gastroenterol. 112, 988–1013 (2017).
- Chey, W. D., Leontiadis, G. I., Howden, C. W. & Moss, S. F. ACG clinical guideline: treatment of *Helicobacter pylori* infection. Am. J. Gastroenterol. **112**, 212–239 (2017).
- Talley, N. J. How to manage the difficult-to-treat dyspeptic patient. Nat. Clin. Pract. Gastroenterol. Hepatol. 4, 35–42 (2007).
- Koletzko, L., Macke, L., Schulz, C. & Malfertheiner, P. Helicobacter pylori eradication in dyspepsia: new evidence for symptomatic benefit. Best Pract. Res. Clin. Gastroenterol. 40-41, 101637 (2019).
- 209. Mahadeva, S., Chia, Y. C., Vinothini, A., Mohazmi, M. & Goh, K. L. Cost-effectiveness of and satisfaction with a *Helicobacter pylori* "test and treat" strategy compared with prompt endoscopy in young Asians with dyspepsia. Gut **57**, 1214–1220 (2008).
- Malfertheiner, P. et al. *Helicobacter pylori* eradication is beneficial in the treatment of functional dyspepsia. *Aliment. Pharmacol. Ther.* 18, 615–625 (2003).
- Hawkey, C. et al. *Helicobacter pylori* eradication for primary prevention of peptic ulcer bleeding in older patients prescribed aspirin in primary care (HEAT): a randomised, double-blind, placebo-controlled trial. *Lancet* **400**, 1597–1606 (2022).
- Ford, A. C., Yuan, Y., Forman, D., Hunt, R. & Moayyedi, P. Helicobacter pylori eradication for the prevention of gastric neoplasia. Cochrane Database Syst. Rev. 7, CD005583 (2020).

- Ford, A. C., Yuan, Y. & Moayyedi, P. Helicobacter pylori eradication therapy to prevent gastric cancer: systematic review and meta-analysis. Gut 69, 2113–2121 (2020).
- Choi, I. J. et al. Family history of gastric cancer and Helicobacter pylori treatment. N. Engl. J. Med. 382, 427–436 (2020).
- Rokkas, T., Rokka, A. & Portincasa, P. A systematic review and meta-analysis of the role of *Helicobacter pylori* eradication in preventing gastric cancer. *Ann. Gastroenterol.* 30, 414–423 (2017).
- Khan, M. Y. et al. Effectiveness of *Helicobacter pylori* eradication in preventing metachronous gastric cancer and preneoplastic lesions. A systematic review and meta-analysis. *Eur. J. Gastroenterol. Hepatol.* **32**, 686–694 (2020).
- Zhao, B. et al. Does *Helicobacter pylori* eradication reduce the incidence of metachronous gastric cancer after curative endoscopic resection of early gastric cancer: a systematic review and meta-analysis. J. Clin. Gastroenterol. 54, 235–241 (2020).
- Fan, F., Wang, Z., Li, B. & Zhang, H. Effects of eradicating *Helicobacter pylori* on metachronous gastric cancer prevention: a systematic review and meta-analysis. *J. Eval. Clin. Pract.* 26, 308–315 (2020).
- Choi, I. J. et al. Helicobacter pylori therapy for the prevention of metachronous gastric cancer. N. Engl. J. Med. 378, 1085–1095 (2018).
- Malfertheiner, P. Helicobacter pylori treatment for gastric cancer prevention. N. Engl. J. Med. 378, 1154–1156 (2018).
- Lauren, P. The two histological main types of gastric carcinoma: diffuse and so-called intestinal-type carcinoma. An attempt at a histo-clinical classification. Acta Pathol. Microbiol. Scand. 64, 31–49 (1965).
- 222. Ma, J., Shen, H., Kapesa, L. & Zeng, S. Lauren classification and individualized chemotherapy in gastric cancer. Oncol. Lett. **11**, 2959–2964 (2016).
- 223. Pan, K. F. et al. A large randomised controlled intervention trial to prevent gastric cancer by eradication of *Helicobacter pylori* in Linqu County, China: baseline results and factors affecting the eradication. Gut 65, 9–18 (2016).
- Herrero, R., Park, J. Y. & Forman, D. The fight against gastric cancer the IARC Working Group report. Best Pract. Res. Clin. Gastroenterol. 28, 1107–1114 (2014).
- Leja, M. et al. Multicentric randomised study of *Helicobacter pylori* eradication and pepsinogen testing for prevention of gastric cancer mortality: the GISTAR study. *BMJ Open* 7, e016999 (2017).
- 226. Ford, A. C., Tsipotis, E., Yuan, Y., Leontiadis, G. I. & Moayyedi, P. Efficacy of *Helicobacter pylori* eradication therapy for functional dyspepsia: updated systematic review and meta-analysis. *Gut* https://doi.org/10.1136/gutjnl-2021-326583 (2022).
- Kim, B. J., Kim, H. S., Jang, H. J. & Kim, J. H. *Helicobacter pylori* eradication in idiopathic thrombocytopenic purpura: a meta-analysis of randomized trials. *Gastroenterol. Res. Pract.* 2018, 6090878 (2018).
- Hudak, L., Jaraisy, A., Haj, S. & Muhsen, K. An updated systematic review and metaanalysis on the association between *Helicobacter pylori* infection and iron deficiency anemia. *Helicobacter* 22, 12330 (2017).
- Malfertheiner, P., Selgrad, M. & Bornschein, J. Helicobacter pylori: clinical management. Curr. Opin. Gastroenterol. 28, 608–614 (2012).
- Ferreri, A. J., Govi, S. & Ponzoni, M. The role of *Helicobacter pylori* eradication in the treatment of diffuse large B-cell and marginal zone lymphomas of the stomach. *Curr. Opin. Oncol.* 25, 470–479 (2013).
- Salar, A. Gastric MALT lymphoma and Helicobacter pylori. Med. Clin. 152, 65–71 (2019).
 Wundisch, T. et al. Long-term follow-up of gastric MALT lymphoma after Helicobacter
- pylori eradication. J. Clin. Oncol. 23, 8018–8024 (2005). 233. Miki, K. Gastric cancer screening using the serum pepsinogen test method. Gastric Cancer 9, 245–253 (2006).
- Sui, Z. et al. Risk for gastric cancer in patients with gastric atrophy: a systematic review and meta-analysis. Transl Cancer Res. 9, 1618–1624 (2020).
- Yoshida, T. et al. Cancer development based on chronic active gastritis and resulting gastric atrophy as assessed by serum levels of pepsinogen and *Helicobacter pylori* antibody titer. Int. J. Cancer **134**, 1445–1457 (2014).
- Miki, K. Gastric cancer screening by combined assay for serum anti-Helicobacter pylori IgG antibody and serum pepsinogen levels — "ABC method". Proc. Jpn Acad. Ser. B Phys. Biol. Sci. 87, 405–414 (2011).
- Miki, K., Fujishiro, M., Kodashima, S. & Yahagi, N. Long-term results of gastric cancer screening using the serum pepsinogen test method among an asymptomatic middle-aged Japanese population. *Dig. Endosc.* 21, 78–81 (2009).
- Sung, H. et al. Global Cancer Statistics 2020: GLOBOCAN estimates of incidence and mortality worldwide for 36 cancers in 185 countries. CA Cancer J. Clin. 71, 209–249 (2021).
- Mabe, K. et al. Endoscopic screening for gastric cancer in Japan: current status and future perspectives. *Dig. Endosc.* 34, 412–419 (2022).
 Device the status of the status
- 240. Ryu, J. E. et al. Trends in the performance of the Korean National Cancer Screening Program for Gastric Cancer from 2007 to 2016. Cancer Res. Treat. 54, 842–849 (2022).
- Pimentel-Nunes, P. et al. Management of epithelial precancerous conditions and lesions in the stomach (MAPS II): European Society of Gastrointestinal Endoscopy (ESGE), European Helicobacter and Microbiota Study Group (EHMSG), European Society of Pathology (ESP), and Sociedade Portuguesa de Endoscopia Digestiva (SPED) guideline update 2019. Endoscopy 51, 365–388 (2019).
- 242. Gupta, S. et al. AGA clinical practice guidelines on management of gastric intestinal metaplasia. *Gastroenterology* **158**, 693–702 (2020).
- Lee, Y. C. et al. Association between *Helicobacter pylori* eradication and gastric cancer incidence: a systematic review and meta-analysis. *Gastroenterology* **150**, 1113–1124.e5 (2016).

- Fallone, C. A., Moss, S. F. & Malfertheiner, P. Reconciliation of recent *Helicobacter pylori* treatment guidelines in a time of increasing resistance to antibiotics. *Gastroenterology* 157, 44–53 (2019).
- El-Serag, H. B. et al. Houston consensus conference on testing for *Helicobacter pylori* infection in the United States. *Clin. Gastroenterol. Hepatol.* 16, 992–1002.e6 (2018).
- Shah, S. C., Iyer, P. G. & Moss, S. F. AGA clinical practice update on the management of refractory *Helicobacter pylori* infection: expert review. *Gastroenterology* 160, 1831–1841 (2021).
- Fallone, C. A. et al. The Toronto consensus for the treatment of *Helicobacter pylori* infection in adults. *Gastroenterology* 151, 51–69.e14 (2016).
- Lind, T. et al. The MACH2 study: role of omeprazole in eradication of *Helicobacter pylori* with 1-week triple therapies. *Gastroenterology* **116**, 248–253 (1999).
- Bazzoli, F. et al. Evaluation of short-term low-dose triple therapy for the eradication of Helicobacter pylori by factorial design in a randomized, double-blind, controlled study. Aliment. Pharmacol. Ther. 12, 439–445 (1998).
- Mahachai, V. et al. Helicobacter pylori management in ASEAN: the Bangkok consensus report. J. Gastroenterol. Hepatol. 33, 37–56 (2018).
- Boyanova, L., Hadzhiyski, P., Gergova, R. & Markovska, R. Evolution of Helicobacter pylori resistance to antibiotics: a topic of increasing concern. Antibiotics 12, 332 (2023).
- Furuta, T. et al. Dual therapy with vonoprazan and amoxicillin is as effective as triple therapy with vonoprazan, amoxicillin and clarithromycin for eradication of *Helicobacter pylori*. *Digestion* **101**, 743–751 (2020).
- Lima, J. J. et al. Clinical pharmacogenetics implementation consortium (CPIC) guideline for CYP2C19 and proton pump inhibitor dosing. *Clin. Pharmacol. Ther.* **109**, 1417–1423 (2021).
- 254. Erah, P. O., Goddard, A. F., Barrett, D. A., Shaw, P. N. & Spiller, R. C. The stability of amoxycillin, clarithromycin and metronidazole in gastric juice: relevance to the treatment of *Helicobacter pylori* infection. J. Antimicrob. Chemother. **39**, 5–12 (1997).
- Furuta, T. & Graham, D. Y. Pharmacologic aspects of eradication therapy for Helicobacter pylori infection. Gastroenterol. Clin. North Am. 39, 465–480 (2010).
- McNicholl, A. G., Linares, P. M., Nyssen, O. P., Calvet, X. & Gisbert, J. P. Meta-analysis: esomeprazole or rabeprazole vs. first-generation pump inhibitors in the treatment of *Helicobacter pylori* infection. *Aliment. Pharmacol. Ther.* **36**, 414–425 (2012).
- Gisbert, J. P. Potent gastric acid inhibition in *Helicobacter pylori* eradication. Drugs 65, 83–96 (2005).
- Villoria, A., Garcia, P., Calvet, X., Gisbert, J. P. & Vergara, M. Meta-analysis: high-dose proton pump inhibitors vs. standard dose in triple therapy for *Helicobacter pylori* eradication. *Aliment. Pharmacol. Ther.* 28, 868–877 (2008).
- Miehlke, S. et al. An increasing dose of omeprazole combined with amoxycillin cures Helicobacter pylori infection more effectively. *Aliment. Pharmacol. Ther.* **11**, 323–329 (1997).
- 260. Gao, W., Zhang, X., Yin, Y., Yu, S. & Wang, L. Different dose of new generation proton pump inhibitors for the treatment of *Helicobacter pylori* infection: a meta-analysis. *Int. J. Immunopathol. Pharmacol.* **35**, 20587384211030397 (2021).
- 261. Nyssen, O. P. et al. European Registry on *Helicobacter pylori* management (Hp-EuReg): patterns and trends in first-line empirical eradication prescription and outcomes of 5 years and 21 533 patients. Gut **70**, 40–54 (2021).
- Scordo, M. G., Caputi, A. P., D'Arrigo, C., Fava, G. & Spina, E. Allele and genotype frequencies of CYP2C9, CYP2C19 and CYP2D6 in an Italian population. *Pharmacol. Res.* 50, 195–200 (2004).
- El Rouby, N., Lima, J. J. & Johnson, J. A. Proton pump inhibitors: from CYP2C19 pharmacogenetics to precision medicine. *Expert Opin. Drug Metab. Toxicol.* 14, 447–460 (2018).
- 264. Strom, C. M. et al. Testing for variants in CYP2C19: population frequencies and testing experience in a clinical laboratory. *Genet. Med.* **14**, 95–100 (2012).
- Sugimoto, K., Uno, T., Yamazaki, H. & Tateishi, T. Limited frequency of the CYP2C19*17 allele and its minor role in a Japanese population. *Br. J. Clin. Pharmacol.* 65, 437–439 (2008).
- Yusuf, I. et al. Ethnic and geographical distributions of CYP2C19 alleles in the populations of Southeast Asia. Adv. Exp. Med. Biol. 531, 37–46 (2003).
- Xie, H. G. Genetic variations of S-mephenytoin 4'-hydroxylase (CYP2C19) in the Chinese population. *Life Sci.* 66, PL175–PL181 (2000).
- Morino, Y. et al. Influence of cytochrome P450 2C19 genotype on Helicobacter pylori proton pump inhibitor-amoxicillin-clarithromycin eradication therapy: a meta-analysis. Front. Pharmacol. 12, 759249 (2021).
- Sugimoto, M. & Furuta, T. Efficacy of tailored *Helicobacter pylori* eradication therapy based on antibiotic susceptibility and CYP2C19 genotype. *World J. Gastroenterol.* 20, 6400–6411 (2014).
- Zhang, H. J. et al. Effects of genetic polymorphisms on the pharmacokinetics and pharmacodynamics of proton pump inhibitors. *Pharmacol. Res.* 152, 104606 (2020).
- Sugimoto, M. et al. Rabeprazole 10 mg q.d.s. decreases 24-h intragastric acidity significantly more than rabeprazole 20 mg b.d. or 40 mg o.m., overcoming CYP2C19 genotype. Aliment. Pharmacol. Ther. 36, 627–634 (2012).
- Saitoh, T. et al. Effects of rabeprazole, lansoprazole and omeprazole on intragastric pH in CYP2C19 extensive metabolizers. *Aliment. Pharmacol. Ther.* 16, 1811–1817 (2002).
- 273. Sahara, S. et al. Twice-daily dosing of esomeprazole effectively inhibits acid secretion in CYP2C19 rapid metabolisers compared with twice-daily omeprazole, rabeprazole or lansoprazole. *Aliment. Pharmacol. Ther.* **38**, 1129–1137 (2013).

- Graham, D. Y. et al. Factors influencing the eradication of *Helicobacter pylori* with triple therapy. *Gastroenterology* **102**, 493–496 (1992).
- Zhou, B. G. et al. Effect of enhanced patient instructions on *Helicobacter pylori* eradication: A systematic review and meta-analysis of randomized controlled trials. *Helicobacter* 27, e12869 (2022).
- Graham, D. Y. & Fischbach, L. *Helicobacter pylori* treatment in the era of increasing antibiotic resistance. *Gut* 59, 1143–1153 (2010).
- Meyer, J. M. et al. Risk factors for *Helicobacter pylori* resistance in the United States: the surveillance of *H. pylori* antimicrobial resistance partnership (SHARP) study, 1993-1999.
 Ann. Intern. Med. **136**, 13–24 (2002).
- Bujanda, L. et al. Antibiotic resistance prevalence and trends in patients infected with Helicobacter pylori in the period 2013-2020: results of the European Registry on H. pylori management (Hp-EuReg). Antibiotics 10, 1058 (2021).
- Camargo, M. C. et al. The problem of *Helicobacter pylori* resistance to antibiotics: a systematic review in Latin America. *Am. J. Gastroenterol.* **109**, 485–495 (2014).
- Tacconelli, E. et al. Discovery, research, and development of new antibiotics: the WHO priority list of antibiotic-resistant bacteria and tuberculosis. *Lancet Infect. Dis.* 18, 318–327 (2018).
- Molina-Infante, J. et al. Optimised empiric triple and concomitant therapy for *Helicobacter pylori* eradication in clinical practice: the OPTRICON study. *Aliment. Pharmacol. Ther.* 41, 581–589 (2015).
- Liou, J. M. et al. Concomitant, bismuth quadruple, and 14-day triple therapy in the first-line treatment of *Helicobacter pylori*: a multicentre, open-label, randomised trial. *Lancet* 388, 2355–2365 (2016).
- 283. Crowe, S. E. Helicobacter pylori infection. N. Engl. J. Med. 380, 1158–1165 (2019).
- Romano, M. et al. Empirical levofloxacin-containing versus clarithromycin-containing sequential therapy for *Helicobacter pylori* eradication: a randomised trial. *Gut* 59, 1465–1470 (2010).
- Federico, A. et al. Efficacy of 5-day levofloxacin-containing concomitant therapy in eradication of *Helicobacter pylori* infection. *Gastroenterology* 143, 55–61.e1 (2012).
- Gisbert, J. P. Optimization strategies aimed to increase the efficacy of Helicobacter pylori eradication therapies with quinolones. Molecules 25, 5084 (2020).
- Kuo, Y. T. et al. Primary antibiotic resistance in *Helicobacter pylori* in the Asia-Pacific region: a systematic review and meta-analysis. *Lancet Gastroenterol. Hepatol.* 2, 707–715 (2017).
- 288. An, Y. et al. Fourth-generation quinolones in the treatment of *Helicobacter pylori* infection: a meta-analysis. *World J. Gastroenterol.* **24**, 3302–3312 (2018).
- Sugimoto, M. et al. High Helicobacter pylori cure rate with sitafloxacin-based triple therapy. Aliment. Pharmacol. Ther. 42, 477–483 (2015).
- Megraud, F. Antibiotic resistance is the key element in treatment of *Helicobacter pylori* infection. Gastroenterology 155, 1300–1302 (2018).
- 291. Malfertheiner, P. et al. *Helicobacter pylori* eradication with a capsule containing bismuth subcitrate potassium, metronidazole, and tetracycline given with omeprazole versus clarithromycin-based triple therapy: a randomised, open-label, non-inferiority, phase 3 trial. *Lancet* **377**, 905–913 (2011).
- 292. Gisbert, J. P. Rifabutin for the treatment of *Helicobacter pylori* infection: a review. *Pathogens* **10**, 15 (2020).
- Graham, D. Y. et al. Rifabutin-based triple therapy (RHB-105) for *Helicobacter pylori* eradication: a double-blind, randomized, controlled trial. *Ann. Intern. Med.* **172**, 795–802 (2020).
- FDA. TALICIA (Omeprazole Magnesium, Amoxicillin and Rifabutin) Delayed-release Capsules [Package Insert]. Raleigh, NC: RedHill Biopharma Inc. https://www.accessdata. fda.gov/drugsatfda_docs/label/2019/213004lbl.pdf (2019).
- 295. Ji, C. R. et al. Safety of furazolidone-containing regimen in *Helicobacter pylori* infection: a systematic review and meta-analysis. *BMJ Open* **10**, e037375 (2020).
- Treiber, G., Ammon, S., Malfertheiner, P. & Klotz, U. Impact of furazolidone-based quadruple therapy for eradication of *Helicobacter pylori* after previous treatment failures. *Helicobacter* 7, 225–231 (2002).
- Tshibangu-Kabamba, E. & Yamaoka, Y. Helicobacter pylori infection and antibiotic resistance - from biology to clinical implications. Nat. Rev. Gastroenterol. Hepatol. 18, 613–629 (2021).
- Wang, Y. H. et al. Characteristics of *Helicobacter pylori* heteroresistance in gastric biopsies and its clinical Relevance. *Front. Cell Infect. Microbiol.* **11**, 819506 (2021).
- Selgrad, M. et al. Different antibiotic susceptibility between antrum and corpus of the stomach, a possible reason for treatment failure of *Helicobacter pylori* infection. *World J. Gastroenterol.* 20, 16245–16251 (2014).
- Malfertheiner, P. Infection: bismuth improves PPI-based triple therapy for H. pylori eradication. Nat. Rev. Gastroenterol. Hepatol. 7, 538–539 (2010).
- Dore, M. P., Lu, H. & Graham, D. Y. Role of bismuth in improving *Helicobacter pylori* eradication with triple therapy. *Gut* 65, 870–878 (2016).
- 302. Marcus, E. A., Sachs, G. & Scott, D. R. Colloidal bismuth subcitrate impedes proton entry into *Helicobacter pylori* and increases the efficacy of growth-dependent antibiotics. *Aliment. Pharmacol. Ther.* 42, 922–933 (2015).
- 303. Nyssen, O. P. et al. European registry on *Helicobacter pylori* management: single-capsule bismuth quadruple therapy is effective in real-world clinical practice. *United European Gastroenterol. J.* 9, 38–46 (2021).
- 304. Zagari, R. M. et al. The "three-in-one" formulation of bismuth quadruple therapy for Helicobacter pylori eradication with or without probiotics supplementation: efficacy and safety in daily clinical practice. *Helicobacter* 23, e12502 (2018).

- Zhang, W. et al. Bismuth, lansoprazole, amoxicillin and metronidazole or clarithromycin as first-line *Helicobacter pylori* therapy. Gut 64, 1715–1720 (2015).
- Bang, C. S. et al. Amoxicillin or tetracycline in bismuth-containing quadruple therapy as first-line treatment for *Helicobacter pylori* infection. Gut Microbes 11, 1314–1323 (2020).
- 307. Delchier, J. C., Malfertheiner, P. & Thieroff-Ekerdt, R. Use of a combination formulation of bismuth, metronidazole and tetracycline with omeprazole as a rescue therapy
- for eradication of *Helicobacter pylori*. Aliment. Pharmacol. Ther. **40**, 171–177 (2014).
 308. Chen, Q. et al. Rescue therapy for *Helicobacter pylori* eradication: a randomized non-inferiority trial of amoxicillin or tetracycline in bismuth quadruple therapy. Am. J. Gastroenterol. **111**, 1736–1742 (2016).
- Hunt, R. H. & Scarpignato, C. Potent acid suppression with PPIs and P-CABs: what's new? Curr. Treat. Options Gastroenterol. 16, 570–590 (2018).
- Abdel-Aziz, Y., Metz, D. C. & Howden, C. W. Review article: potassium-competitive acid blockers for the treatment of acid-related disorders. *Aliment. Pharmacol. Ther.* 53, 794–809 (2021).
- Murakami, K. et al. Vonoprazan, a novel potassium-competitive acid blocker, as a component of first-line and second-line triple therapy for *Helicobacter pylori* eradication: a phase III, randomised, double-blind study. *Gut* 65, 1439–1446 (2016).
- Chey, W. D. et al. S1382 Vonoprazan dual and triple therapy for *Helicobacter pylori* eradication. J. Am. Coll. Gastroenterol. 116, S634 (2021).
- Rokkas, T. et al. Comparative effectiveness of multiple different first-line treatment regimens for *Helicobacter pylori* infection: a network meta-analysis. *Gastroenterology* 161, 495–507.e4 (2021).
- Malfertheiner, P. et al. Potassium-competitive acid blocker and proton pump inhibitorbased regimens for first-line *Helicobacter pylori* eradication: a network meta-analysis. *Gastro. Hep. Adv.* 1, 824–834 (2022).
- Liou, J. M. et al. Efficacies of genotypic resistance-guided vs empirical therapy for refractory *Helicobacter pylori* infection. *Gastroenterology* 155, 1109–1119 (2018).
- Yu, L. et al. Susceptibility-guided therapy for Helicobacter pylori infection treatment failures. Ther. Adv. Gastroenterol. 12, 1756284819874922 (2019).
- 317. Chen, P. Y. et al. Systematic review with meta-analysis: the efficacy of levofloxacin triple therapy as the first- or second-line treatments of *Helicobacter pylori* infection. *Aliment. Pharmacol. Ther.* 44, 427–437 (2016).
- Gao, C. P. et al. PPI-amoxicillin dual therapy for *Helicobacter pylori* infection: an update based on a systematic review and meta-analysis. *Helicobacter* 25, e12692 (2020).
- 319. Gao, W. et al. Eradication rate and safety of a "simplified rescue therapy": 14-day vonoprazan and amoxicillin dual regimen as rescue therapy on treatment of *Helicobacter pylori* infection previously failed in eradication: a real-world, retrospective clinical study in China. *Helicobacter* 27, e12918 (2022).
- Shiotani, A., Roy, P., Lu, H. & Graham, D. Y. Helicobacter pylori diagnosis and therapy in the era of antimicrobial stewardship. Ther. Adv. Gastroenterol. 14, 17562848211064080 (2021).
- Graham, D. Y. & Liou, J. M. Primer for development of guidelines for *Helicobacter pylori* therapy using antimicrobial stewardship. *Clin. Gastroenterol. Hepatol.* 20, 973–983.e1 (2021).
- McFarland, L. V., Huang, Y., Wang, L. & Malfertheiner, P. Systematic review and metaanalysis: Multi-strain probiotics as adjunct therapy for *Helicobacter pylori* eradication and prevention of adverse events. *United European Gastroenterol. J.* 4, 546–561 (2016).
- 323. Fernandez-Salazar, L. et al. Effectiveness and safety of high-dose dual therapy: results of the European Registry on the management of *Helicobacter pylori* infection (Hp-EuReg). *J. Clin. Med.* **11**, 3544 (2022).
- 324. Liou, J. M. et al. Second-line levofloxacin-based quadruple therapy versus bismuth-based quadruple therapy for *Helicobacter pylori* eradication and long-term changes to the gut microbiota and antibiotic resistome: a multicentre, open-label, randomised controlled trial. *Lancet Gastroenterol. Hepatol.* 8, 228–241 (2023).
- 325. Guillemard, E. et al. A randomised, controlled trial: effect of a multi-strain fermented milk on the gut microbiota recovery after *Helicobacter pylori* therapy. *Nutrients* 13, 3171 (2021).
- 326. Ford, A. C. et al. Adverse events with bismuth salts for *Helicobacter pylori* eradication: systematic review and meta-analysis. *World J. Gastroenterol.* **14**, 7361–7370 (2008).
- Kumar, S., Metz, D. C., Kaplan, D. E. & Goldberg, D. S. Treatment of *Helicobacter pylori* is not associated with future clostridium difficile infection. *Am. J. Gastroenterol.* **115**, 716–722 (2020).
- Lu, M. et al. Efficacy of probiotic supplementation therapy for *Helicobacter pylori* eradication: a meta-analysis of randomized controlled trials. *PLoS ONE* 11, e0163743 (2016).
- 329. Szajewska, H., Horvath, A. & Kolodziej, M. Systematic review with meta-analysis: Saccharomyces boulardii supplementation and eradication of *Helicobacter pylori* infection. *Aliment. Pharmacol. Ther.* **41**, 1237–1245 (2015).
- Viazis, N. et al. A four-probiotics regimen combined with a standard Helicobacter pylorieradication treatment reduces side effects and increases eradication rates. Nutrients 14, 632 (2022).
- 331. Zhao, Y. et al. Saccharomyces boulardii combined with quadruple therapy for Helicobacter pylori eradication decreased the duration and severity of diarrhea: a multi-center prospective randomized controlled trial. Front. Med. 8, 776955 (2021).
- Malfertheiner, P. et al. Helicobacter pylori eradication and gastric ulcer healing comparison of three pantoprazole-based triple therapies. Aliment. Pharmacol. Ther. 17, 1125–1135 (2003).

- Gralnek, I. M. et al. Endoscopic diagnosis and management of nonvariceal upper gastrointestinal hemorrhage (NVUGIH): European Society of Gastrointestinal Endoscopy (ESGE) Guideline — Update 2021. Endoscopy 53, 300–332 (2021).
- Barkun, A. N. et al. Management of nonvariceal upper gastrointestinal bleeding: guideline recommendations from the International Consensus Group. Ann. Intern. Med. 171, 805–822 (2019).
- 335. Sostres, C. et al. Peptic ulcer bleeding risk. The role of Helicobacter pylori infection in NSAID/low-dose aspirin users. Am. J. Gastroenterol. 110, 684–689 (2015).
- 336. Malfertheiner, P. et al. Current concepts in the management of *Helicobacter pylori* infection: the Maastricht III Consensus Report. Gut 56, 772–781 (2007).
- Ruskone-Fourmestraux, A. et al. EGILS consensus report. Gastric extranodal marginal zone B-cell lymphoma of MALT. Gut 60, 747–758 (2011).
- Raderer, M., Kiesewetter, B. & Ferreri, A. J. Clinicopathologic characteristics and treatment of marginal zone lymphoma of mucosa-associated lymphoid tissue (MALT lymphoma). CA Cancer J. Clin. 66, 153–171 (2016).
- 339. Jung, K., Kim, D. H., Seo, H. I., Gong, E. J. & Bang, C. S. Efficacy of eradication therapy in *Helicobacter pylori*-negative gastric mucosa-associated lymphoid tissue lymphoma: a meta-analysis. *Helicobacter* 26, e12774 (2021).
- Laine, L. & Dhir, V. Helicobacter pylori eradication does not worsen quality of life related to reflux symptoms: a prospective trial. Aliment. Pharmacol. Ther. 16, 1143–1148 (2002).
- Hirata, K. et al. Improvement of reflux symptom related quality of life after Helicobacter pylori eradication therapy. J. Clin. Biochem. Nutr. 52, 172–178 (2013).
- Taguchi, H. et al. *Helicobacter pylori* eradication improves the quality of life regardless of the treatment outcome: a multicenter prospective cohort study. *Medicine* **96**, e9507 (2017).
- Piriyapong, K., Tangaroonsanti, A., Mahachai, V. & Vilaichone, R. K. Helicobacter pylori infection impacts on functional dyspepsia in Thailand. Asian Pac. J. Cancer Prev. 15, 10887-10891 (2014).
- 344. Moayyedi, P. et al. Effect of population screening and treatment for *Helicobacter pylori* on dyspepsia and quality of life in the community: a randomised controlled trial. Leeds HELP Study Group. *Lancet* 355, 1665–1669 (2000).
- Bektas, M., Soykan, I., Altan, M., Alkan, M. & Ozden, A. The effect of *Helicobacter pylori* eradication on dyspeptic symptoms, acid reflux and quality of life in patients with functional dyspepsia. *Eur. J. Intern. Med.* **20**, 419–423 (2009).
- 346. Buzas, G. M. Quality of life in patients with functional dyspepsia: short- and long-term effect of *Helicobacter pylori* eradication with pantoprazole, amoxicillin, and clarithromycin or cisapride therapy: a prospective, parallel-group study. *Curr. Ther. Res. Clin. Exp.* **67**, 305–320 (2006).
- Mestrovic, A., Bozic, J., Vukojevic, K. & Tonkic, A. Impact of different *Helicobacter pylori* eradication therapies on gastrointestinal symptoms. *Medicina* 57, 803 (2021).
- Bitwayiki, R. et al. Dyspepsia prevalence and impact on quality of life among Rwandan healthcare workers: a cross-sectional survey. S. Afr. Med. J. 105, 1064–1069 (2015).
- 349. Kabakambira, J. D. et al. Efficacy of Helicobacter pylori eradication regimens in Rwanda: a randomized controlled trial. BMC Gastroenterol. 18, 134 (2018).
- Labenz, J. et al. Curing Helicobacter pylori infection in patients with duodenal ulcer may provoke reflux esophagitis. Gastroenterology 112, 1442–1447 (1997).
- 351. Zamani, M. et al. Systematic review with meta-analysis: the worldwide prevalence of Helicobacter pylori infection. Aliment. Pharmacol. Ther. 47, 868–876 (2018).
- Ding, S. Z. et al. Chinese consensus report on family-based *Helicobacter pylori* infection control and management (2021 edition). *Gut* **71**, 238–253 (2022).
 Chiang, T. H. et al. Bismuth salts with versus without acid suppression for *Helicobacter*
- pylori infection: a transmission electron microscope study. *Helicobacter* 26, e12801 (2021).
- 354. Kafarski, P. & Talma, M. Recent advances in design of new urease inhibitors: a review. J. Adv. Res. 13, 101–112 (2018).
- 355. Strugatsky, D. et al. Structure of the proton-gated urea channel from the gastric pathogen *Helicobacter pylori*. *Nature* **493**, 255–258 (2013).
- Chu, J. K. et al. Loss of a cardiolipin synthase in *Helicobacter pylori* G27 blocks flagellum assembly. J. Bacteriol. **201**, e00372-19 (2019).
- 357. Doohan, D., Rezkitha, Y. A. A., Waskito, L. A., Yamaoka, Y. & Miftahussurur, M. Helicobacter pylori BabA-SabA key roles in the adherence phase: the synergic mechanism for successful colonization and disease development. *Toxins* **13**, 485 (2021).
- Zhang, Y. et al. Inhibition of pathogen adhesion by bacterial outer membrane-coated nanoparticles. Angew. Chem. Int. Ed. Engl. 58, 11404–11408 (2019).
- 359. Ensign, L. M., Cone, R. & Hanes, J. Oral drug delivery with polymeric nanoparticles: the gastrointestinal mucus barriers. Adv. Drug Deliv. Rev. 64, 557–570 (2012).
- 360. Yang, C. et al. Effects of non-viable Lactobacillus reuteri combining with 14-day standard triple therapy on *Helicobacter pylori* eradication: a randomized double-blind placebo-controlled trial. *Helicobacter* 26, e12856 (2021).
- Liang, B. et al. Current and future perspectives for *Helicobacter pylori* treatment and management: from antibiotics to probiotics. *Front. Cell Infect. Microbiol.* **12**, 1042070 (2022).
- Yang, Y. J. & Sheu, B. S. Metabolic interaction of *Helicobacter pylori* infection and gut microbiota. *Microorganisms* 4, 15 (2016).
- 363. Yang, I., Nell, S. & Suerbaum, S. Survival in hostile territory: the microbiota of the stomach. FEMS Microbiol. Rev. 37, 736–761 (2013).
- Chen, C. C. et al. The interplay between *Helicobacter pylori* and gastrointestinal microbiota. *Gut Microbes* 13, 1–22 (2021).

- 365. Schulz, C. et al. The active bacterial assemblages of the upper GI tract in individuals with and without *Helicobacter* infection. Gut 67, 216–225 (2018).
- 366. Sun, Q. H. et al. Microbiome changes in the gastric mucosa and gastric juice in different histological stages of *Helicobacter pylori-negative gastric cancers*. *World J. Gastroenterol.* 28, 365–380 (2022).
- Sung, J. J. Y. et al. Gastric microbes associated with gastric inflammation, atrophy and intestinal metaplasia 1 year after *Helicobacter pylori* eradication. *Gut* 69, 1572–1580 (2020).
- Guo, Y., Cao, X. S., Zhou, M. G. & Yu, B. Gastric microbiota in gastric cancer: different roles of *Helicobacter pylori* and other microbes. *Front. Cell Infect. Microbiol.* 12, 1105811 (2022).
- Coker, O. O. et al. Mucosal microbiome dysbiosis in gastric carcinogenesis. Gut 67, 1024–1032 (2018).
- Vaillant, L., Oster, P., McMillan, B., Orozco Fernandez, E. & Velin, D. GM-CSF is key in the efficacy of vaccine-induced reduction of *Helicobacter pylori* infection. *Helicobacter* 27, e12875 (2022).
- Oster, P. et al. *Helicobacter pylori* infection has a detrimental impact on the efficacy of cancer immunotherapies. *Gut* 71, 457–466 (2022).
- Oster, P., Vaillant, L., McMillan, B. & Velin, D. The efficacy of cancer immunotherapies is compromised by *Helicobacter pylori* infection. *Front. Immunol.* 13, 899161 (2022).
- Abadi, A. T. & Kusters, J. G. Management of *Helicobacter pylori* infections. *BMC Gastroenterol.* 16, 94 (2016).
- Celiberto, F. et al. The state of the art of molecular fecal investigations for Helicobacter pylori (H. pylori) antibiotic resistances. Int. J. Mol. Sci. 24, 4361 (2023).
- Ranjbar, R., Ebrahimi, A. & Sahebkar, A. Helicobacter pylor infection: conventional and molecular strategies for bacterial diagnosis and antibiotic resistance testing. *Curr. Pharm. Biotechnol.* https://doi.org/10.2174/1389201023666220920094342 (2022).
- Best, L. M. et al. Non-invasive diagnostic tests for Helicobacter pylori infection. Cochrane Database Syst. Rev. 3, CD012080 (2018).
- No authors listed. Schistosomes, liver flukes and Helicobacter pylori. IARC Monogr. Eval. Carcinog. Risks Hum. 61, 1–241 (1994).
- Wotherspoon, A. C. et al. Regression of primary low-grade B-cell gastric lymphoma of mucosa-associated lymphoid tissue type after eradication of *Helicobacter pylori*. *Lancet* 342, 575–577 (1993).
- 379. Rugge, M. et al. OLGA staging for gastritis: a tutorial. *Dig. Liver Dis.* **40**, 650–658 (2008).
- Chiang, T. H. et al. Mass eradication of *Helicobacter pylori* to reduce gastric cancer incidence and mortality: a long-term cohort study on Matsu Islands. *Gut* 70, 243–250 (2021).
- Du, M. Q. & Isaccson, P. G. Gastric MALT lymphoma: from aetiology to treatment. Lancet Oncol. 3, 97–104 (2002).
- Glupczynski, Y., Megraud, F., Lopez-Brea, M. & Andersen, L. P. European multicentre survey of in vitro antimicrobial resistance in *Helicobacter pylori. Eur. J. Clin. Microbiol. Infect. Dis.* 20, 820–823 (2001).
- Li, W. Q. et al. Effects of *Helicobacter pylori* treatment and vitamin and garlic supplementation on gastric cancer incidence and mortality: follow-up of a randomized intervention trial. *BMJ* 366, I5016 (2019).
- Wang, Z. et al. Changes of the gastric mucosal microbiome associated with histological stages of gastric carcinogenesis. Front. Microbiol. 11, 997 (2020).
- Song, P., Wu, L. & Guan, W. Dietary nitrates, nitrites, and nitrosamines intake and the risk of gastric cancer: a meta-analysis. *Nutrients* 7, 9872–9895 (2015).
- Lucker, S. et al. A Nitrospira metagenome illuminates the physiology and evolution of globally important nitrite-oxidizing bacteria. Proc. Natl Acad. Sci. USA 107, 13479–13484 (2010).
- Rudnicka, K., Backert, S. & Chmiela, M. Genetic polymorphisms in inflammatory and other regulators in gastric cancer: risks and clinical consequences. *Curr. Top. Microbiol. Immunol.* 421, 53–76 (2019).
- 388. Ford, A. C., Marwaha, A., Lim, A. & Moayyedi, P. What is the prevalence of clinically significant endoscopic findings in subjects with dyspepsia? Systematic review and meta-analysis. *Clin. Gastroenterol. Hepatol.* 8, 830–837 (2010).
- Gisbert, J. P. & Calvet, X. Helicobacter pylori "test-and-treat" strategy for management of dyspepsia: a comprehensive review. Clin. Transl Gastroenterol. 4, e32 (2013).
- 390. Beresniak, A. et al. *Helicobacter pylori* "test-and-treat" strategy with urea breath test: a cost-effective strategy for the management of dyspepsia and the prevention of ulcer and gastric cancer in Spain — results of the Hp-Breath initiative. *Helicobacter* 25, e12693 (2020).
- Pritchard, D. M. et al. Cost-effectiveness modelling of use of urea breath test for the management of *Helicobacter pylori*-related dyspepsia and peptic ulcer in the UK. *BMJ Open Gastroenterol.* 8, e000685 (2021).
- Kawasaki, K. et al. Low-dose aspirin and non-steroidal anti-inflammatory drugs increase the risk of bleeding in patients with gastroduodenal ulcer. *Dig. Dis. Sci.* 60, 1010–1015 (2015).
- Eusebi, L. H., Black, C. J., Howden, C. W. & Ford, A. C. Effectiveness of management strategies for uninvestigated dyspepsia: systematic review and network meta-analysis. *BMJ* 367, I6483 (2019).
- Wu, R. et al. Prevalence of gastric cancer precursors in gastroscopy-screened adults by family history of gastric cancer and of cancers other than gastric. *BMC Cancer* 20, 1110 (2020).

- 395. De, R. V. et al. Pepsinogens to distinguish patients with gastric intestinal metaplasia and Helicobacter pylori infection among populations at risk for gastric cancer. Clin. Transl Gastroenterol. 7, e183 (2016).
- 396. Zagari, R. M. et al. Systematic review with meta-analysis: diagnostic performance of the combination of pepsinogen, gastrin-17 and anti-*Helicobacter pylori* antibodies serum assays for the diagnosis of atrophic gastritis. *Aliment. Pharmacol. Ther.* 46, 657–667 (2017).

Acknowledgements

The authors thank Y.-C. Chen and H.-T. Yu for help with figure design.

Author contributions

Introduction (P.M.); Epidemiology (M.C.C., J.-M.L. and S.S.); Mechanisms/pathophysiology (S.S., E.E.-O. and S.S.); Diagnosis, screening and prevention (M.C.C., R.P., C.S. and S.I.S.); Management (P.M., E.E.-O. and J.-M.L.); Quality of life (M.C.C., R.P. and S.I.S.); Outlook (P.M. and J.-M.L.); Overview of the Primer (P.M.).

Competing interests

P.M. has consulted for Aboca, Bayer Healthcare, Cinclus, Imevax, Menarini Foundation and Phatom. P.M. has received honoraria for lectures from Allergosan, Biohit, Biocodex and Malesci. S.I.S. has received scientific support from Richen. C.S. has received speaker fees from Imevax, Falk Foundation and Lilly. S.S. is listed as an inventor on a patent application related to the use of bacterial motility inhibitors as potential treatment for *Helicobacter pylori* infection.

Additional information

Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41572-023-00431-8.

Peer review information Nature Reviews Disease Primers thanks F. Carneiro, K. McColl, Y. Yamaoka and the other, anonymous, reviewer(s) for their contribution to the peer review of this work.

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