

MEDICAL POLICY

POLICY TITLE	GENETIC TESTING FOR <i>HELICOBACTER PYLORI</i> TREATMENT
POLICY NUMBER	MP 2.308

CLINICAL BENEFIT	☐ MINIMIZE SAFETY RISK OR CONCERN. ☒ MINIMIZE HARMFUL OR INEFFECTIVE INTERVENTIONS. ☐ ASSURE APPROPRIATE LEVEL OF CARE. ☐ ASSURE APPROPRIATE DURATION OF SERVICE FOR INTERVENTIONS. ☐ ASSURE THAT RECOMMENDED MEDICAL PREREQUISITES HAVE BEEN MET. ☐ ASSURE APPROPRIATE SITE OF TREATMENT OR SERVICE.
Effective Date:	RETIRED 7/1/2026

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I. POLICY

Genotyping to determine cytochrome p450 (CYP2C19) genetic polymorphisms is considered **investigational** for the purpose of managing the treatment of *H. pylori* infection. There is insufficient evidence to support a general conclusion concerning the health outcomes or benefits associated with this procedure.

II. PRODUCT VARIATIONS

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This policy is only applicable to certain programs and products administered by Capital Blue Cross and subject to benefit variations as discussed in Section VI. Please see additional information below.

FEP PPO - Refer to FEP Medical Policy Manual. The FEP Medical Policy manual can be found at:

<https://www.fepblue.org/benefit-plans/medical-policies-and-utilization-management-guidelines/medical-policies>

III. DESCRIPTION/BACKGROUND

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Helicobacter pylori infection of the gastrointestinal (GI) tract is treated with a combination of antibiotics and proton-pump inhibitors (PPI). Genetic factors may influence the success of *H. pylori* treatment through effects on PPI metabolism. Individuals with polymorphisms in the *CYP2C19* gene metabolize PPIs more rapidly than normal and may have a reduced therapeutic effect. Therefore, individualized treatment regimens based on genetic testing may improve eradication rates.

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Helicobacter pylori (*H pylori*) is a bacterium associated with a range of gastrointestinal (GI) disorders, such as peptic ulcer disease, chronic gastritis, and gastric malignancy. Eradication of *H pylori* has been proven beneficial for a number of indications.

Currently, multiple regimens are available for treating *H pylori* infection. These include proton pump inhibitors (PPI), as well as similar medication(s), to suppress acid production in combination with antibiotic treatment, consisting of one or more agents such as amoxicillin, clarithromycin, or metronidazole. These first-line regimens generally achieve eradication rates in the 70–90% range. Differences in eradication rates are dependent on the regimen used and the population being treated. Treatment failures are most often attributed to antibiotic resistance or poor patient compliance. Resistance to clarithromycin is an important factor associated with treatment failure, with high rates of treatment failure for standard first-line regimens in patients infected with clarithromycin-resistant strains of *H pylori*. A 2002 survey from the U.S. estimated that 13% of *H pylori* strains are resistant to clarithromycin and that the rate of resistance was rising in comparison to earlier studies.

Genetic factors may influence the success of *H pylori* treatment through effects on PPI metabolism. Individuals with polymorphisms in the *CYP2C19* gene, a component of the cytochrome p450 (CYP450) system, metabolize PPIs more slowly than normal. Genetic variation in the CYP450 enzyme system is one of the most extensively studied in the field of pharmacogenomics. This family of enzymes is found in the liver and is important for metabolizing and eliminating a large number of pharmacologic agents. Differences in PPI metabolism lead to variability in gastric acid suppression, with associated variability in gastric pH and potential impact on the efficacy of *H pylori* treatment. Observational research suggests that patients who are extensive metabolizers of PPIs have lower eradication rates following standard treatment for *H pylori*, compared with poor metabolizers.

Three major *CYP2C19* alleles determine enzymatic activity, as shown in Table 1. The *1 allele is the wild type found in most individuals, while the *2 and *3 alleles are the most common polymorphisms that are known to impact enzymatic activity. Both the *2 and *3 alleles are examples of “null” alleles, which have no enzymatic activity. Each null allele is caused by a single nucleotide change that results in a splice defect or a stop codon. (1)

Table 1. CYP2C19 polymorphisms**

Allele	Nucleotide Change	Predicted Enzyme Activity
*1	None	Normal
*2	681G>A	None
*3	636G>A	None

Table 2. CYP2C19 phenotypes**

Allele	1	2	3
1	EM	IM	IM
2		PM	PM
3			PM

EM: extensive metabolizers; IM: intermediate metabolizers; PM: poor metabolizers.

** Adapted from AmpliChip package insert

Polymorphisms of the *CYP2C19* gene are relatively common and vary by ethnicity. Patients with no polymorphisms of *CYP2C19* have 2 wild-type alleles and no reduction in their ability to metabolize PPIs. These patients are typically called extensive metabolizers (EM) (Table 2).

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Heterozygous polymorphisms are found in 27–37% of the Caucasian population and 46–50% of the Asian population. These patients have a minor reduction in their ability to eliminate PPIs and are called intermediate metabolizers (IM). Homozygous polymorphisms of the *CYP2C19* gene are found in 3–6% of Caucasians and in 12–20% of Asians. These patients eliminate PPIs from the circulation substantially more slowly than unaffected patients and are termed poor metabolizers (PM).

In patients treated with PPIs, intragastric pH has been shown to correlate with *CYP2C19* status. Patients homozygous for a *CYP2C19* mutation (PM) exhibit a less acidic pH when compared to patients without a *CYP2C19* mutation, with heterozygous patients exhibiting intermediate values. Intragastric pH has important implications for treating *H. pylori*. *H. pylori* is more sensitive to antibiotics at less acidic pH levels. Less acidic pH levels also lead to greater stability and bioavailability of antibiotics. Therefore, it is expected that treatment of *H. pylori* will be more successful if there is maximal suppression of gastric acid production and higher intragastric pH levels.

Therefore, it has been proposed that a pharmacogenomics-based treatment regimen individualized by *CYP2C19* status may improve the success rate of treatment for *H. pylori*. If *CYP2C19* status is known prior to treatment, adjustments can be made in the selection of PPI and/or the dosing schedule to achieve optimal acid suppression in all patients. Improved eradication rates for *H. pylori* could lead to improved health outcomes by reducing the need for retreatment following treatment failure, reducing recurrences of *H. pylori*-associated disorders and reducing the morbidity and mortality associated with disease recurrence.

The American Gastroenterological Association published a “Clinical Practice Update on the Management of Refractory *Helicobacter pylori* Infection: Expert Review” in January of 2021. The update outlines the importance of *CYP2C19* polymorphisms in metabolism of PPIs and failure of eradication of *H. pylori*. Studies have been completed in Asian-Pacific populations, however the literature highlights that the lack of study in the US population is important, given the “substantive racial and ethnic differences in the prevalence of *CYP2C19* variant alleles and genotypes in the United States.” The guideline goes on to state that “current data are insufficient to support genetic polymorphism testing for guiding therapeutic selection in refractory (or primary) eradication therapy. Given the high population prevalence of metabolism-enhancing phenotypes of *CYP2C19* at least in non-Asian groups, empiric selection of strategies that achieve greater intragastric acid suppression might be reasonable in the management of refractory *H. pylori* infection.”

At least one commercially available genetic test, the Roche AmpliChip Cytochrome P450® Genotyping test, has been approved by the U.S. Food and Drug Administration (FDA) as a class II medical device. This test examines polymorphisms in *CYP2D6* and *CYP2C19* isoenzymes of the cytochrome p450 enzyme system. Approval for this device was originally granted in December 2004 as an aid in determining treatment choice and individualizing treatment dose for therapeutics that are primarily metabolized by the *CYP2D6* enzyme. The use of information on *CYP2C19* polymorphisms was not addressed as part of the FDA approval process.

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IV. RATIONALE

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Summary

The scientific evidence does not permit conclusions on whether the use of a pharmacogenomics-based treatment regimen for *H pylori* improves eradication rates. In general, eradication rates of *H pylori* vary by CYP2C19 status, with the highest rates found in patients who are poor metabolizers of PPIs. In the single randomized controlled trial comparing a pharmacogenomics-based treatment regimen with a standard regimen, eradication rates after first-line treatment were higher for the pharmacogenomics group compared with the standard treatment group. However, because of numerous variations in treatment protocol within the pharmacogenomics group, it is not possible to determine whether the improvement resulted from the tailored PPI dosages according to CYP2C19 genetic status or due to other variations in the treatment protocol unrelated to CYP2C19 status. It is possible that other clinical factors, such as clarithromycin resistance, or other treatment factors, such as length of antibiotic treatment, may have influenced eradication rates. The use of a PPI that is less susceptible to CYP2C19 status, such as rabeprazole, has been associated with higher eradication rates compared to other PPIs. Therefore, additional trials are needed to address the issues noted above, including alternative treatment regimens, before conclusions can be made on whether a pharmacogenomics-based treatment regimen improves *H pylori* eradication rates compared to a standard treatment regimen. Therefore, the use of genetic testing for *Helicobacter pylori* treatment is considered investigational.

V. DEFINITIONS

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GENOTYPE refers to the pair of genes present for a particular characteristic or protein.

POLYMORPHISM refers to the state or quality of existing or occurring in several different forms.

VII. DISCLAIMER

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Capital Blue Cross' medical policies are used to determine coverage for specific medical technologies, procedures, equipment, and services. These medical policies do not constitute medical advice and are subject to change as required by law or applicable clinical evidence from independent treatment guidelines. Treating providers are solely responsible for medical advice and treatment of members. These policies are not a guarantee of coverage or payment. Payment of claims is subject to a determination regarding the member's benefit program and eligibility on the date of service, and a determination that the services are medically necessary and appropriate. Final processing of a claim is based upon the terms of contract that applies to the members' benefit program, including benefit limitations and exclusions. If a provider or a member has a question concerning this medical policy, please contact Capital Blue Cross' Provider Services or Member Services.

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VIII. CODING INFORMATION

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Note: This list of codes may not be all-inclusive, and codes are subject to change at any time. The identification of a code in this section does not denote coverage as coverage is determined by the terms of member benefit information. In addition, not all covered services are eligible for separate reimbursement.

Investigational; therefore, not covered, genetic testing for the purpose of managing the treatment of helicobacter pylori infection:

Procedure Codes							
81225							

IX. REFERENCES

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X. POLICY HISTORY

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MP 2.308	05/22/2020 Consensus Review. No change to policy statement. Variations updated. References reviewed with no changes. Coding reviewed with no changes.
	04/14/2021 Consensus Review. No change to policy statement. References updated and coding reviewed.
	07/29/2022 Consensus Review. No change to policy statement. FEP, references updated, coding reviewed.
	07/31/2023 Consensus Review. No change to policy statement. Updated background, references. Coding reviewed, no changes.
	01/19/2024 Administrative Update. Clinical benefit added.
	09/03/2024 Consensus Review. No change to policy statement. Updated references. Coding reviewed, no changes.
	08/08/2025 Consensus Review. No change to policy statement. Coding reviewed, no changes.
	09/23/2025 Administrative Update. Removed Benefit Variations Section and updated Disclaimer.
	03/09/2026 Retirement Review. Services managed by EviCore.

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