#### **Supplement to:**

## Clinical Pharmacogenetics Implementation Consortium (CPIC) Guideline for CYP2C19 and Proton Pump Inhibitor Dosing

John J. Lima<sup>1</sup>, Cameron D. Thomas<sup>2</sup>, Julia Barbarino<sup>3</sup>, Zeruesenay Desta<sup>4</sup>, Sara L. Van Driest<sup>5</sup>, Nihal El Rouby<sup>2,6</sup>, Julie A. Johnson<sup>2</sup>, Larisa H. Cavallari<sup>2</sup>, Valentina Shakhnovich<sup>7,8,9</sup>, David L. Thacker<sup>10,11</sup>, Stuart A. Scott<sup>12,13</sup>, Matthias Schwab<sup>14,15,16</sup>, Chakradhara Rao S Uppugunduri<sup>17,18</sup>, Christine M. Formea<sup>19</sup>, James P. Franciosi<sup>20,21</sup>, Katrin Sangkuhl<sup>3</sup>, Andrea Gaedigk<sup>7</sup>, Teri E. Klein<sup>3</sup>, Roseann S. Gammal<sup>22,23</sup>, Takahisa Furuta<sup>24</sup>

- <sup>1</sup> Center for Pharmacogenomics and Translational Research, Nemours Children's Health, Jacksonville, FL, USA
- <sup>2</sup> Department of Pharmacotherapy and Translational Research, and Center for Pharmacogenomics and Precision Medicine, College of Pharmacy, University of Florida, Gainesville, FL, USA
- <sup>3</sup> Department of Biomedical Data Science, Stanford University, Stanford, CA, USA
- <sup>4</sup> Department of Medicine, Division of Clinical Pharmacology, Indiana University School of Medicine, Indianapolis, IN, USA
- <sup>5</sup> Departments of Pediatrics and Medicine, Vanderbilt University School of Medicine, Nashville, TN, USA
- <sup>6</sup> Division of Pharmacy Practice & Administrative Sciences, University of Cincinnati James Winkle College of Pharmacy, Cincinnati, OH, USA
- <sup>7</sup> Division of Clinical Pharmacology, Toxicology & Therapeutic Innovation, Children's Mercy Kansas City and University of Missouri Kansas City School of Medicine, Kansas City, MO, USA
- <sup>8</sup> Division of Gastroenterology, Hepatology, and Nutrition, Children's Mercy Kansas City, Kansas City, MO, USA
- <sup>9</sup> Center for Children's Healthy Lifestyles & Nutrition, Kansas City, MO, USA
- <sup>10</sup> Department of Clinical Pharmacology, Indiana University School of Medicine, Indianapolis, IN, USA
- <sup>11</sup> Translational Software, Bellevue, WA, USA
- <sup>12</sup> Department of Genetics and Genomic Sciences, Icahn School of Medicine at Mount Sinai, New York, NY, USA
- <sup>13</sup> Sema4, Stamford, CT, USA
- <sup>14</sup> Dr Margarete Fischer-Bosch-Institute of Clinical Pharmacology, Stuttgart, Germany
- <sup>15</sup> Department of Clinical Pharmacology, University Hospital, Tuebingen, Germany
- <sup>16</sup> Department of Pharmacy and Biochemistry, University of Tuebingen, Tuebingen, Germany
- <sup>17</sup> CANSEARCH Research Laboratory, Department of Paediatrics, Gynaecology and Obstetrics, Faculty of Medicine, University of Geneva, Geneva, Switzerland
- <sup>18</sup> Oncology-Hematology Unit, Department of Paediatrics, Gynaecology and Obstetrics, Geneva University Hospital, Geneva, Switzerland
- <sup>19</sup> Department of Pharmacy Services and Intermountain Precision Genomics, Intermountain Healthcare, Salt Lake City, UT, USA
- $^{20}\,\mathrm{Division}$  of Gastroenterology, Hepatology and Nutrition, Nemours Children's Hospital, Orlando, FL

<sup>&</sup>lt;sup>21</sup> Department of Pediatrics, University of Central Florida College of Medicine, Orlando, FL, USA

<sup>&</sup>lt;sup>22</sup> Department of Pharmacy Practice, MCPHS University School of Pharmacy, Boston, MA, USA

<sup>&</sup>lt;sup>23</sup> Department of Pharmaceutical Sciences, St. Jude Children's Research Hospital, Memphis, TN, USA

<sup>&</sup>lt;sup>24</sup> Center for Clinical Research, Hamamatsu University School of Medicine, Hamamatsu, Shizuoka, Japan

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#### **GUIDELINE UPDATES**

The Clinical Pharmacogenetics Implementation Consortium (CPIC) guideline for *CYP2C19* and proton pump inhibitor (PPI) dosing is published in full on the CPIC website (1). Relevant information will be reviewed periodically and updated guidelines published online.

#### LITERATURE REVIEW

The PubMed® database (1966 to April 2018) was searched for the following keywords: (CYP2C19 OR cytochrome P450 2C19) AND (proton pump inhibitor OR PPI OR \*omeprazole OR \*lansoprazole OR pantoprazole OR rabeprazole). The search was limited to studies conducted in humans and written in the English language, and review articles were excluded. Using these search terms, 831 publications were identified. Study inclusion criteria included publications that incorporated analyses for the association between *CYP2C19* genotype and PPI pharmacokinetic parameters or PPI-related clinical outcomes. Following the application of these criteria, 244 publications were reviewed and included in the evidence tables (**Tables S1-S7**).

#### **GENETIC TEST INTERPRETATION**

The haplotype, or star (\*) allele name, is determined by a specific single nucleotide polymorphism (SNP) or a combination of SNPs that are interrogated in the genotyping analysis. Numerous deletion and duplication events affecting the *CYP2C* gene locus have been described (see Botton et al for a comprehensive summary (2) and the PharmVar Structural Variation document at <a href="https://www.pharmvar.org/gene/CYP2C19">https://www.pharmvar.org/gene/CYP2C19</a>). Many of the gene deletion and duplication events involve more than one of the *CYP2C* genes and can even encompass a large number of genes within this chromosomal region. To date, PharmVar has defined deletion events

encompassing the entire *CYP2C19* gene under the *CYP2C19\*36* designation and those with partial *CYP2C19* gene deletion events (that include at least exon 1) as *CYP2C19\*37* (2). *CYP2C* copy number variants appear to be rare and are typically not part of pharmacogenetic testing.

The genotypes that constitute the haplotype, or star (\*) alleles for *CYP2C19*, and the rs# for each of the specific genomic nucleotide alterations that define the alleles, are described in the *CYP2C19* Allele Definition Table online (1, 3). The genotype results are generally reported as a diplotype, which includes one maternal and one paternal star allele (e.g., \*1/\*2). The *CYP2C19* function associated with each of the common star alleles is summarized in the *CYP2C19* Allele Functionality Table online (1, 3).

#### **AVAILABLE GENETIC TEST OPTIONS**

Registry provides a central location for voluntary submission of genetic test information by providers and is available at <a href="http://www.ncbi.nlm.nih.gov/gtr">http://www.ncbi.nlm.nih.gov/gtr</a>. Desirable characteristics of pharmacogenetic tests, including naming of alleles and test report contents, have been extensively reviewed by an international group, including CPIC members (4). CPIC recommends that clinical laboratories adhere to these test reporting standards. CPIC gene-specific tables adhere to these allele nomenclature standards. Moreover, these tables (*CYP2C19* Allele Definition Table, *CYP2C19* Allele Functionality Table, and *CYP2C19* Allele Frequency Table) may be used to assemble lists of known functional and actionable pharmacogenetic variants and their population frequencies, which may inform decisions as to whether tests are adequately comprehensive in interrogations of alleles (1, 3). Furthermore, the Association for Molecular Pathology has published a recommendation for the key attributes of alleles

recommended for clinical testing and a minimum set of variants that should be included in clinical genotyping assays for *CYP2C19* (5).

#### LEVELS OF EVIDENCE LINKING GENOTYPE TO PHENOTYPE

The evidence summarized in **Tables S1-S7** is graded on a scale of high, moderate, and weak (6) based upon the level of evidence:

High: Evidence includes consistent results from well-designed, well-conducted studies.

**Moderate:** Evidence is sufficient to determine effects, but the strength of the evidence is limited by the number, quality or consistency of the individual studies, generalizability to routine practice, or the indirect nature of the evidence.

**Weak:** Evidence is insufficient to assess the effects on health outcomes because of limited number or power of studies, important flaws in their design or conduct, gaps in the chain of evidence, or lack of information.

To determine these levels of evidence for major finding statements, individual studies are first evaluated using a structured framework:

- 1. Quality elements for individual studies are evaluated as yes, no, partially, unclear, or not relevant:
  - Confounders and use of concomitant medications with possible drug interactions
    are reported and potential impact on the major finding are analyzed and reported.
  - Phenotype assignments (when comparing phenotype groups) are based on CPIC phenotype assignment or similar.
  - Reported data are based on steady-state kinetics where appropriate.

- Sample size adequate to assess difference between genotype/phenotype groups, especially for negative findings.
- Adequate phenotyping or genotyping methods:
  - o States all genetic variants screened
  - o Alleles tested are adequate to determine "wild-type" genotype
  - Adequate phenotyping or genotyping method used
  - Appropriate attainment of samples
  - o Defines how \* alleles are defined, if applicable
  - o Clearly states which genotypes were found in the study
- Race and/or ancestry is discussed and appropriately considered.
- Outcome definition clearly defined and measured.
- Appropriate statistics performed.
- 2. The individual study is rated with respect to how well it supports the major finding statement:
  - First, it is determined whether the study supports the major finding statement or does not support it.
  - Second, a qualifier is added to the statement (if needed) based on the quality elements listed above:
    - Some study quality flaws: Enough of the items in step 1 are rated "partially," "unclear," or "no" to introduce some uncertainty about the validity of the conclusions.

- o **Major study quality flaws:** Enough of the items in step 1 are rated "partially," "unclear," or "no" to introduce serious uncertainty about the validity of the conclusions.
- No qualification on statement: If few items in step 1 are rated as "partially," "unclear," or "no."
- There are six possible ratings for individual studies:
  - Supports the statement
  - o Supports the statement but with some quality flaws
  - o Supports the statement but with major quality flaws
  - Does not support the statement
  - o Does not support the statement but with some quality flaws
  - o Does not support the statement but with major quality flaws

#### STRENGTH OF RECOMMENDATIONS

CPIC's therapeutic recommendations are based on weighing the evidence from a combination of preclinical functional and clinical data, as well as on some existing disease-specific consensus guidelines. Some of the factors that are taken into account in evaluating the evidence supporting therapeutic recommendations include *in vivo* pharmacokinetic and pharmacodynamic data, *in vitro* enzyme activity of tissues expressing wild-type/reference or variant-containing CYP2C19, *in vitro* CYP2C19 enzyme activity from tissues isolated from individuals of known *CYP2C19* genotypes, and *in vivo* pre-clinical and clinical pharmacokinetic and pharmacodynamic studies.

Overall, the therapeutic recommendations are simplified to allow rapid interpretation by clinicians. CPIC uses a slight modification of a transparent and simple system for recommendations adopted from the rating scale for evidence-based guidelines on the use of antiretroviral agents (7):

- **Strong** recommendation for the statement: The evidence is high quality and the desirable effects clearly outweigh the undesirable effects.
- Moderate recommendation for the statement: There is a close or uncertain balance as to
  whether the evidence is high quality and the desirable effects clearly outweigh the
  undesirable effects.
- **Optional** recommendation for the statement: The desirable effects are closely balanced with undesirable effects, or the evidence is weak or based on extrapolations. There is room for differences in opinion as to the need for the recommended course of action.
- **No recommendation**: There is insufficient evidence, confidence, or agreement to provide a recommendation to guide clinical practice at this time.

# RESOURCES TO INCORPORATE PHARMACOGENETICS INTO AN ELECTRONIC HEALTH RECORD WITH CLINICAL DECISION SUPPORT

Clinical decision support (CDS) tools integrated within electronic health records (EHRs) can help guide clinical pharmacogenetics at the point of care (8-13). See <a href="https://cpicpgx.org/guidelines/cpic-guideline-for-proton-pump-inhibitors-and-cyp2c19/">https://cpicpgx.org/guidelines/cpic-guideline-for-proton-pump-inhibitors-and-cyp2c19/</a> for resources to support the adoption of CPIC guidelines within an EHR (1, 14). Based on the capabilities of various EHRs and local preferences, we recognize that approaches may vary across organizations. Our intent is to synthesize foundational knowledge that provides a common

starting point for incorporating *CYP2C19* genotype results in an EHR to guide proton pump inhibitor dosing.

Effective incorporation of pharmacogenetic information into an EHR to optimize drug therapy should have some key attributes. Pharmacogenetic results, an interpreted phenotype, and a concise interpretation or summary of the result must be documented in the EHR (15). To incorporate a phenotype in the EHR in a standardized manner, genotype test results provided by the laboratory must be consistently translated into an interpreted phenotype (**Table 1, main manuscript**; *CYP2C19* **Diplotype to Phenotype Table** (1, 3)). Because clinicians must be able to easily find the information, the interpreted phenotype may be documented as a problem list entry or in a patient's summary section; these phenotypes are best stored in the EHR at the "person level" rather than at the date-centric "encounter level". Additionally, results should be entered as standardized and discrete terms to facilitate using them to provide point-of-care CDS (see **PPI Pre- and Post-Test Alerts and Flow Chart** for example CDS alerts; <a href="https://epicpgx.org/guidelines/cpic-guideline-for-proton-pump-inhibitors-and-cyp2c19/">https://epicpgx.org/guidelines/cpic-guideline-for-proton-pump-inhibitors-and-cyp2c19/</a>) (16, 17).

Because pharmacogenetic results have lifetime implications and clinical significance, results should be placed into a section of the EHR that is accessible independent of the test result date to allow clinicians to quickly find the result at any time after it is initially placed in the EHR. To facilitate this process, CPIC is providing gene-specific information figures and tables that include complete diplotype to phenotype translation tables, diagram(s) that illustrate how CYP2C19 pharmacogenetic test results could be entered into an EHR, example EHR consultation/genetic test interpretation language and widely used nomenclature systems (see <a href="https://cpicpgx.org/guidelines/cpic-guideline-for-proton-pump-inhibitors-and-cyp2c19">https://cpicpgx.org/guidelines/cpic-guideline-for-proton-pump-inhibitors-and-cyp2c19</a>) (1, 18).

Point-of-care CDS should be designed to effectively notify clinicians of prescribing implications at any time after the test result is entered into the EHR. CPIC is also providing gene-drug specific tables that provide guidance to achieve these objectives with diagrams that illustrate how point-of-care CDS should be entered into the EHR, example pre- and post-test alert language, and widely used nomenclature systems for relevant drugs (see <a href="https://cpicpgx.org/guidelines/cpic-guideline-for-proton-pump-inhibitors-and-cyp2c19">https://cpicpgx.org/guidelines/cpic-guideline-for-proton-pump-inhibitors-and-cyp2c19</a> (1)).

There are some unique implementation challenges associated with CYP2C19/PPIs given the indication-specific recommendations (increased dose for *H. pylori* infection and erosive esophagitis) for CYP2C19 RMs and NMs and time-dependent recommendations (short-term vs long-term use >12 weeks) for CYP2C19 IMs, CYP2C19 likely IMs, CYP2C19 PMs, and CYP2C19 likely PMs. CDS post-test alert language is provided for all CYP2C19 phenotypes; however, the possibility of alert fatigue must be considered given the frequency with which PPIs are prescribed. Considering that for most initial PPI prescriptions the recommendation will be to initiate standard dosing, pre-test alerts are not recommended to fire for all PPI orders. Given these challenges, implementation will ultimately be institution-specific and may require creative solutions. Alternative CDS solutions include incorporating an option to order CYP2C19 genotype into disease-specific order sets (e.g., for *H. pylori* infection and/or erosive esophagitis); custom-built order sets that include PPIs for specific indications (e.g., H. pylori infection and/or erosive esophagitis) that account for known CYP2C19 results; limit pre-test and/or post-test alerts to providers within a specific specialty area (e.g., gastrointestinal specialists); and include pre-test and/or post-test alerts when all the required elements for a specific indication are present (e.g., alert will fire only when all medications for the treatment of *H. pylori* are added to the

patient's active medication list or *H. pylori* infection and/or erosive esophagitis are in the patient's problem list). See the pre- and post-test alert tables for examples of CDS alerts (1).

#### PEDIATRIC CONSIDERATIONS

PPIs are some of the most commonly prescribed drugs for pediatric populations, and prescription rates continue to rise (19-21). PPIs are available over-the-counter (without prescriptions) in some countries. In children younger than 18 years of age, PPIs currently have U.S. Food and Drug Administration (FDA)-approved indications for the short-term treatment of symptomatic gastroesophageal reflux disease (GERD), healing of erosive esophagitis, treatment of peptic ulcer disease, and for eradication of *H. pylori* as part of a multi-drug regimen (22-24). Despite current lack of FDA approval for the indication of eosinophilic esophagitis, PPI therapy is now considered standard of care for this condition in North America and Europe (25). Offlabel use of long-term PPI therapy in children is common, and there are increasing concerns that PPIs are over-utilized in pediatric populations (26, 27). Esomeprazole has been FDA-approved for infants as young as one month of age only for confirmed erosive esophagitis, yet many PPIs are frequently prescribed incorrectly for symptoms suspected to be secondary to GERD without proven benefit. PPI therapy has been studied in children with respiratory symptoms, sleep disorders, and excessive crying, with minimal benefit demonstrated (28-30).

There are emerging concerns that PPI therapy use (and/or misuse) is associated with numerous side effects including, but not limited to, gastrointestinal and respiratory infections, malabsorption of vitamins and minerals, bone fracture, and possible association to chronic diseases such as celiac disease and chronic kidney disease (31-36). PPI therapy did not show

benefit in children with asthma in terms of lung function but did demonstrate increased rates of respiratory infections (37), highlighting this specific side effect.

CYP2C19 enzyme function is reported to be very low during early fetal life, consistent with very low apparent clearance of PPIs in preterm neonates and term infants less than 2-3 months of age, but clearance is consistent with adult values after that age (38-40). There is emerging evidence for the influence of CYP2C19 function on PPI pharmacokinetics and response in children (41-46). CYP2C19 RM or UM phenotypes have been associated with decreased therapeutic efficacy compared to PM and NM phenotypes when treating children with GERD and eosinophilic esophagitis (47-50). The CYP2C19 PM phenotype has been associated with higher rates of pediatric respiratory and gastrointestinal infections than NM, RM, or UM phenotypes (51). There is one report of increased infection events in infants and young children treated with PPIs who are CYP2C19 NMs versus UMs (52). A recent pilot study of CYP2C19 genotype-guided dosing of PPI medications in children has been promising, and additional studies are ongoing (53, 54). These data support optimization of PPIs therapy in infants and children over one year of age based on CYP2C19 genotype data, with weaker evidence to support genotype-guided dosing in pre-term infants or infants less than 2-3 months of age (55).

TABLE S1. EVIDENCE LINKING CYP2C19 TO OMEPRAZOLE PHENOTYPE

Type of experimental model	Major findings	References	Level of evidence <sup>a</sup>
Metabolism			
Clinical	CYP2C19 PMs are associated with decreased metabolism of omeprazole as compared to IMs.	Ieiri, et al. 1996 (56) Herrlin, et al. 1998 (57) Furuta, et al. 1999 (58) Furuta, et al. 1999 (59) Sakai, et al. 2001 (60) Shirai, et al. 2003 (62) Yin, et al. 2004 (63) Rosemary, et al. 2005 (64) Ohnishi, et al. 2005 (65) Qiao, et al. 2006 (66) Shimizu, et al. 2006 (67) Uno, et al. 2007 (68) Wang, et al. 2007 (70) Wang, et al. 2010 (71) Shiohira, et al. 2011 (72) Shiohira, et al. 2012 (73) Yamada, et al. 2014 (75) Park, et al. 2017 (76)	High
Clinical	CYP2C19 PMs (as determined by phenotyping) are associated with decreased metabolism of omeprazole as compared to IMs.	Chang, et al. 1995 (77)	High
Clinical	CYP2C19 PMs are associated with decreased metabolism of omeprazole as compared to NMs.	Rost and Roots 1996 (78) Ieiri, <i>et al.</i> 1996 (56)	High

			-
		Furuta, <i>et al.</i> 1999 (58)	
		Furuta, <i>et al</i> . 1999 (59)	
		Zhou, et al. 1999 (79)	
		Sakai, et al. 2001 (60)	
		Shirai, et al. 2001 (61)	
		Kita, et al. 2002 (80)	
		He, et al. 2003 (62)	
		Yin, et al. 2004 (63)	
		Rosemary, et al. 2005 (64)	
		Ohnishi, et al. 2005 (65)	
		Ieiri, et al. 2005 (81)	
		Qiao, et al. 2006 (66)	
		Shimizu, et al. 2006 (67)	
		Uno, et al. 2007 (68)	
		Wang, et al. 2007 (69)	
		Niioka, et al. 2007 (82)	
		Hu, et al. 2007 (70)	
		Uno, et al. 2008 (83)	
		Chen, et al. 2009 (84)	
		Chaudhry, et al. 2009 (85)	
		Wang, et al. 2010 (71)	
		Shiohira, <i>et al.</i> 2011 (72)	
		Shiohira, et al. 2012 (73)	
		Yamada, et al. 2013 (74)	
		Payan, et al. 2014 (75)	
		Park, et al. 2017 (76)	
Clinical	CYP2C19 PMs (as determined by phenotyping) are associated with	Rost, et al. 1992 (86)	High
	decreased metabolism of omeprazole as compared to NMs.	Yasuda, et al. 1995 (87)	
		Tybring, et al. 1997 (88)	
		Bottiger, et al. 1997 (89)	
		Andersson, et al. 1998 (90)	
		Mihara, <i>et al</i> . 1999 (91)	
		Tassaneeyakul, et al. 2000 (92)	

Clinical	CYP2C19 PMs are associated with decreased metabolism of omeprazole as compared to RMs.	Payan, et al. 2014 (75)	Moderate
Clinical	CYP2C19 PMs are associated with decreased metabolism of omeprazole as compared to UMs.	Payan, et al. 2014 (75)	Moderate
Clinical	CYP2C19 IMs are associated with decreased metabolism of omeprazole as compared to NMs.	Chang, et al. 1995 (93) Marinac, et al. 1996 (94) Herrlin, et al. 1998 (57) Furuta, et al. 1999 (59) Furuta, et al. 1999 (58) Shu, et al. 2000 (95) Sakai, et al. 2001 (60) Kim, et al. 2002 (96) He, et al. 2003 (62) Kearns, et al. 2003 (97) Yin, et al. 2004 (63) Rosemary, et al. 2005 (64) Shimizu, et al. 2006 (67) Uno, et al. 2007 (68) Wang, et al. 2007 (69) Niioka, et al. 2007 (82) Hunfeld, et al. 2008 (98) Yamada, et al. 2014 (75)	Moderate
Clinical	CYP2C19 IMs (as determined by phenotyping) are associated with decreased metabolism of omeprazole as compared to NMs.	Chang, et al. 1995 (77)	Moderate
Clinical	CYP2C19 IMs are associated with decreased metabolism of omeprazole as compared to RMs.	Roman, et al. 2014 (99) Payan, et al. 2014 (75)	Moderate
Clinical	CYP2C19 IMs are associated with decreased metabolism of omeprazole as compared to UMs.	Payan, et al. 2014 (75)	Moderate
Clinical	CYP2C19 NMs are associated with decreased metabolism of omeprazole as compared to RMs.	Sim, et al. 2006 (100) Hunfeld, et al. 2008 (98) Roman, et al. 2014 (99)	Weak

		Payan, et al. 2014 (75)	
Clinical	CYP2C19 NMs are associated with decreased metabolism of omeprazole as compared to UMs.	Sim, et al. 2006 (100) Baldwin, et al. 2008 (101) Payan, et al. 2014 (75)	Moderate
Clinical	CYP2C19 RMs are associated with decreased metabolism of omeprazole as compared to UMs.	Payan, et al. 2014 (75)	Moderate
Clinical	CYP2C19 PMs are associated with decreased metabolism of omeprazole as compared to IMs+NMs.	Chang, et al. 1995 (93) Ishizawa, et al. 2005 (102) Kuhlmann, et al. 2014 (103) Nazir, et al. 2015 (104) Nazir, et al. 2016 (105)	Moderate
Clinical	CYP2C19 PMs have decreased metabolism of omeprazole as compared to NMs+RMs+UMs.	Zhao, et al. 2018 (46)	Moderate
Clinical	CYP2C19 PMs+IMs are not associated with altered metabolism of omeprazole as compared to NMs.	Denisenko, et al. 2017 (106)	Weak
Clinical	CYP2C19 PMs+IMs are not associated with altered metabolism of omeprazole as compared to UMs.	Denisenko, et al. 2017 (106)	Weak
Clinical	CYP2C19 IMs have decreased metabolism of omeprazole as compared to NMs+RMs+UMs.	Zhao, et al. 2018 (46)	Moderate
Clinical	CYP2C19 NMs are associated with decreased metabolism of omeprazole as compared to RMs+UMs.	Denisenko, et al. 2017 (106)	Weak
Clinical	CYP2C19 is associated with omeprazole metabolism when comparing PMs vs IMs vs NMs.	Roh, et al. 1996 (107) Sagar, et al. 1998 (108) Furuta, et al. 1999 (58) Furuta, et al. 1999 (59) Roh, et al. 2004 (109) Isaza, et al. 2007 (110)	Moderate
Clinical	CYP2C19 is related to omeprazole metabolism when comparing PMs vs IMs vs NMs vs RMs.	Koukoula, et al. 2017 (111)	Weak
Clinical	CYP2C19 is associated with omeprazole metabolism when comparing PMs vs IMs vs NMs vs RMs+UMs.	Xavier, et al. 2016 (112)	Weak

Clinical	CYP2C19 is not associated with omeprazole metabolism when	Kearns, et al. 2010 (113)	Weak
	comparing PMs vs IMs vs NMs vs RMs vs UMs.		
Clinical	CYP2C19 is not associated with omeprazole metabolism when comparing IMs vs NMs vs RMs.	Chwiesko, et al. 2016 (114)	Weak
Clinical	CYP2C19 is not associated with omeprazole metabolism when comparing IMs vs NMs vs RMs+UMs.	Chwiesko, et al. 2016 (114)	Weak
In vitro	CYP2C19*9, *10, *16, *19 and *26 are associated with decreased metabolism of omeprazole as compared to CYP2C19*1.	Hanioka, et al. 2008 (115) Lee, et al. 2009 (116) Wang, et al. 2011 (117) Langaee, et al. 2014 (118)	Weak
In vitro	CYP2C19*1B, *11, *13, *15 and *18 are not associated with altered metabolism of omeprazole as compared to CYP2C19*1.	Hanioka, et al. 2008 (115) Wang, et al. 2011 (117)	Weak
In vitro	CYP2C19*3, *5, *6, and *24 are associated with absent metabolism of omeprazole as compared to CYP2C19*1.	Wang, et al. 2011 (117) Dai, et al. 2015 (119) Lau, et al. 2017 (120)	Weak
In vitro	CYP2C19*8 is associated with decreased metabolism of omeprazole as compared to CYP2C19*1.	Wang, et al. 2011 (117)	Weak
In vitro	CYP2C19*14 and *32 (H99R) are not associated with altered metabolism of omeprazole as compared to CYP2C19*1.	Wang, et al. 2011 (117) Dai, et al. 2015 (119)	Weak
In vitro	CYP2C19*23, *29 (K28I), *30, *31 (H78Y) and *33 (D188N) are associated with decreased metabolism of omeprazole as compared to CYP2C19*1.	Dai, et al. 2015 (119) Lau, et al. 2017 (120)	Weak
In vitro	CYP2C19*2B (E92D), *2C (A161P), *2E (M271I), *2F (D341N), *2G (D360V), *2H (H396D), *2J (K421Q), *3B (D360N) and *3C (M136K) are associated with decreased metabolism of omeprazole as compared to CYP2C19*1.	Wang, et al. 2011 (117) Dai, et al. 2015 (119)	Weak
Efficacy			
Remission of			
Clinical	CYP2C19 IMs are associated with increased remission of reflux when treated with omeprazole as compared to NMs.	Zendehdel, et al. 2010 (121)	Weak
Clinical	CYP2C19 IMs are associated with increased healing rate of ulcers when treated with omeprazole as compared to NMs.	Ando, et al. 2005 (122) Ando, et al. 2008 (123)	Weak

Clinical	CYP2C19 NMs have a decreased healing rate of ulcers when	Hizawa, et al. 2006 (124)	Weak
	treated with omeprazole.		
Clinical	CYP2C19 IMs have a decreased healing rate of ulcers when treated	Okamura, et al. 2013 (125)	Weak
	with omeprazole compared to CYP2C19 NMs.		
Clinical	CYP2C19 RMs have a decreased healing rate of reflux when	Fukaya, et al. 2016 (126)	Weak
	treated with omeprazole compared to CYP2C19 NMs.		
Clinical	CYP2C19 PMs are not associated with healing rate of ulcers when	Ji, et al. 2006 (127)	Weak
	treated with omeprazole as compared to IMs+NMs.		
Clinical	CYP2C19 is not associated with healing rate of reflux when treated	Ohkusa, et al. 2005 (128)	Weak
	with omeprazole when comparing PMs vs IMs vs NMs.		
<i>H. pylori</i> er			
Clinical	CYP2C19 PMs are associated with an increased <i>H. pylori</i>	Furuta, et al. 1998 (129)	Weak
	eradication rate when treated with omeprazole as compared to IMs.	Lin, et al. 2017 (130)	
Clinical	CYP2C19 PMs are associated with an increased <i>H. pylori</i>	Furuta, et al. 1998 (129)	Moderate
	eradication rate when treated with omeprazole as compared to NMs.	Chaudhry, et al. 2009 (85)	
		Lin, et al. 2017 (130)	
Clinical	CYP2C19 IMs are associated with an increased <i>H. pylori</i>	Furuta, et al. 1998 (129)	Weak
	eradication rate when treated with omeprazole as compared to NMs	Sapone, et al. 2003 (131)	
		Miehlke, et al. 2006 (132)	
		Sezgin, et al. 2014 (133)	
		Lin, et al. 2017 (130)	
Clinical	CYP2C19 PMs are associated with increased <i>H. pylori</i> eradication	Tanigawara, et al. 1999 (134)	Weak
	rate when treated with omeprazole as compared to IMs+NMs.	Miwa, et al. 2001 (135)	
		Yang, et al. 2011 (136)	
Clinical	CYP2C19 PMs+IMs are associated with increased <i>H. pylori</i>	Sheu, et al. 2005 (137)	Moderate
	eradication rate when treated with omeprazole as compared to NMs.	Hong, et al. 2016 (138)	
Clinical	CYP2C19 is associated with H. pylori eradication rate when treated	Furuta, et al. 1998 (129)	Weak
	with omeprazole when comparing PMs vs IMs vs NMs.	Dojo, et al. 2001 (139)	
		Inaba, et al. 2002 (140)	
		Higuchi, et al. 2006 (141)	
		Zhang, et al. 2010 (142)	
		Nabinger, et al. 2016 (143)	
Acid secret	ion indices		

Clinical	CYP2C19 PMs are associated with better acid secretion indices when treated with omeprazole as compared to IMs.	Furuta, et al. 1999 (58) Shirai, et al. 2001 (61) Shimatani, et al. 2003 (144) Hu, et al. 2007 (70) Wang, et al. 2010 (71) Sahara, et al. 2013 (145) Sugimoto, et al. 2014 (146) Park, et al. 2017 (76)	Moderate
Clinical	CYP2C19 PMs are associated with better acid secretion indices when treated with omeprazole as compared to NMs.	Furuta, et al. 1999 (58) Shirai, et al. 2001 (61) Shimatani, et al. 2003 (144) Sugimoto, et al. 2006 (147) Hu, et al. 2007 (70) Wang, et al. 2010 (71) Furuta, et al. 2010 (148) Sahara, et al. 2013 (145) Sugimoto, et al. 2014 (146) Park, et al. 2017 (76)	Moderate
Clinical	CYP2C19 IMs are associated with better acid secretion indices when treated with omeprazole as compared to NMs.	Furuta, et al. 1999 (58) Adachi, et al. 2000 (149) Sagar, et al. 2000 (150) Sagar, et al. 2000 (151) Sagar, et al. 2000 (150) Shirai, et al. 2001 (61) Sugimoto, et al. 2006 (147) Hu, et al. 2007 (70) Hunfeld, et al. 2008 (98) Sahara, et al. 2013 (145) Sugimoto, et al. 2014 (146)	Moderate
Clinical	CYP2C19 NMs are not associated with altered acid secretion indices when treated with omeprazole as compared to RMs	Hunfeld, et al.2008 (98) Chwiesko, et al. 2016 (114)	Weak
Clinical	CYP2C19 PMs have better acid secretion indices when treated with omeprazole as compared to IMs+NMs.	Yang, et al. 2011 (136)	Weak

Clinical	CYP2C19 NMs are not associated with altered acid secretion indices when treated with omeprazole as compared to RMs+UMs.	Chwiesko, et al. 2016 (114)	Weak
Clinical	CYP2C19 is associated with acid secretion indices when treated	Furuta, et al. 1999 (58)	Weak
Toxicity	with omeprazole when comparing PMs vs IMs vs NMs.	Roh, et al. 2004 (109)	
Clinical	CYP2C19 PMs are not associated with risk for adverse events when treated with omeprazole as compared to IMs.	Ohkusa, et al. 2005 (128)	Weak
Clinical	CYP2C19 PMs are not associated with risk for adverse events when treated with omeprazole as compared to NMs.	Ohkusa, et al. 2005 (128)	Weak
Clinical	CYP2C19 PMs do not have an altered risk for acute interstitial nephritis when treated with omeprazole as compared to NMs.	Helsby, et al. 2010 (152)	Weak
Clinical	CYP2C19 UMs have an increased risk for agranulocytosis when treated with omeprazole.	Dury, et al. 2012 (153)	Weak
Clinical	CYP2C19 is not associated with risk for visual disorders when treated with omeprazole when comparing PMs vs IMs vs NMs.	Lutz, et al. 2002 (154)	Weak

<sup>a</sup>Rating scheme described in the **Supplemental Material**IM, intermediate metabolizer; NM, normal metabolizer; PM, poor metabolizer; RM, rapid metabolizer; UM, ultrarapid metabolizer

TABLE S2. EVIDENCE LINKING CYP2C19 TO LANSOPRAZOLE PHENOTYPE

Type of	Major findings	References	Level of
experimental			evidence <sup>a</sup>
model			
Metabolism			
Clinical	CYP2C19 PMs are associated with decreased metabolism of	Sakai, et al. 2001 (60)	High
	lansoprazole as compared to IMs.	Ieiri, et al. 2001(155)	
		Furuta, et al. 2001 (156)	
		Shirai, et al. 2002 (157)	
		Furuta, et al. 2002 (158)	
		Schwab, et al. 2004 (159)	
		Miura, et al. 2004 (160)	
		Uno, et al. 2005 (161)	
		Saito, et al. 2005 (162)	
		Uno, et al. 2005 (163)	
		Qiao, et al. 2006 (66)	
		Xu, et al. 2010 (164)	
		Zhang, et al. 2011 (165)	
		Gumus, et al. 2012 (166)	
Clinical	CYP2C19 PMs are associated with decreased metabolism of	Li, et al. 2014 (167)	TT: ~1.
Clinical		Ko, et al. 1999 (168)	High
	lansoprazole as compared to NMs.	Sakai, <i>et al.</i> 2001 (60) Ieiri, <i>et al.</i> 2001 (155)	
		Furuta, et al. 2001 (156)	
		Shirai, et al. 2001 (130)	
		Kim, et al. 2002 (157)	
		Furuta, et al. 2002 (158)	
		Hu, et al. 2004 (170)	
		Schwab, et al. 2004 (179)	
		Miura, et al. 2004 (160)	
		Uno, et al. 2005 (161)	
		Saito, et al. 2005 (162)	

Clinical	CYP2C19 PMs (as determined by phenotyping) are associated with	Uno, et al. 2005 (163) Qiao, et al. 2006 (66) Niioka, et al. 2008 (171) Miura, et al. 2010 (172) Xu, et al. 2010 (164) Zhang, et al. 2011 (165) Gumus, et al. 2012 (166) Li, et al. 2014 (167) Sohn, et al. 1997 (173)	Moderate
	decreased metabolism of lansoprazole as compared to NMs.	Andersson, et al. 1998 (90)	
Clinical	CYP2C19 IMs are associated with decreased metabolism of lansoprazole as compared to NMs.	Ieiri, et al. 2001 (155) Sakai, et al. 2001 (60) Furuta, et al. 2001 (156) Schwab, et al. 2004 (159) Howden, et al. 2006 (174) Miura, et al. 2006 (175) Hunfeld, et al. 2008 (98) Zhang, et al. 2010 (172) Xu, et al. 2010 (164) Zhang, et al. 2011 (165) Gumus, et al. 2012 (166) Zalloum, et al. 2012 (176) Li, et al. 2014 (167) Zhang, et al. 2014 (177)	Moderate
Clinical	CYP2C19 NMs are not associated with altered metabolism of lansoprazole as compared to RMs.	Gumus, et al. 2012 (166)	Weak
Clinical	CYP2C19 NMs are associated with decreased metabolism of lansoprazole as compared to UMs,	Gumus, et al. 2012 (166)	Weak
Clinical	CYP2C19 PMs are associated with decreased metabolism of lansoprazole as compared to IMs+NMs.	Katsuki, et al. 1997 (178) Ko, et al. 1999 (168) Miura, et al. 2010 (179) Zhang, et al. 2012 (180)	High

		Zhang, et al. 2013 (181)	
Clinical	CYP2C19 PMs+IMs are associated with decreased metabolism of lansoprazole as compared to NMs.	Lima, et al. 2013 (51)	Weak
Clinical	CYP2C19 is associated with lansoprazole metabolism when comparing PMs vs IMs vs NMs.	Yasui-Furukori, et al. 2004 (182) Niioka, et al. 2006 (183) Sakurai, et al. 2007 (184)	Moderate
Efficacy			
Remission of		1 2005 (105)	1361
Clinical	CYP2C19 PMs are associated with increased remission of reflux when treated with lansoprazole as compared to IMs.	Kawamura, et al. 2007 (185)	Moderate
Clinical	CYP2C19 PMs are associated with increased remission and healing rate of reflux when treated with lansoprazole as compared to NMs.	Furuta, et al. 2002 (158) Kawamura, et al. 2007 (185) Furuta, et al. 2009 (186)	High
Clinical	CYP2C19 PMs are not associated with altered healing rate of ulcers when treated with lansoprazole as compared to NMs.	Yoshizawa, et al. 2016 (187)	Weak
Clinical	CYP2C19 IMs are associated with increased remission and healing rate of reflux when treated with lansoprazole as compared to NMs.	Furuta, et al. 2002 (158) Kawamura, et al. 2003 (188) Furuta, et al. 2009 (186)	High
Clinical	CYP2C19 IMs are not associated with healing rate of ulcers when treated with lansoprazole as compared to NMs.	Yoshizawa, et al. 2016 (187)	Weak
Clinical	CYP2C19 IMs have a decreased healing rate of ulcers when treated with lansoprazole.	Okamura, et al. 2013 (125)	Weak
H. pylori era	adication		
Clinical	CYP2C19 PMs are not associated with altered <i>H. pylori</i> eradication rate when treated with lansoprazole as compared to IMs.	Okudaira, et al. 2005 (189)	Weak
Clinical	CYP2C19 PMs are associated with increased <i>H. pylori</i> eradication rate when treated with lansoprazole as compared to NMs.	Isomoto, et al. 2003 (190) Okudaira, et al. 2005 (189) Furuta, et al. 2005 (191) Furuta, et al. 2007 (192) Suzuki, et al. 2007 (193) Settin, et al. 2014 (194)	Moderate

Clinical	CYP2C19 IMs are associated with increased <i>H. pylori</i> eradication	Isomoto, et al. 2003 (190)	Moderate
	rate when treated with lansoprazole as compared to NMs.	Okudaira, et al. 2005 (189)	
		Furuta, et al. 2005 (191)	
		Furuta, et al. 2007 (192)	
		Suzuki, et al. 2007 (193)	
		Ozdil, et al. 2010 (195)	
		Settin, et al. 2014 (194)	
Clinical	CYP2C19 PMs are associated with increased <i>H. pylori</i> eradication	Schwab, et al. 2004 (159)	Weak
	rate when treated with lansoprazole as compared to IMs+NMs.	, , ,	
Clinical	CYP2C19 PMs+IMs are associated with increased <i>H. pylori</i>	Kawabata, et al. 2003 (196)	High
	eradication rate when treated with lansoprazole as compared to	Liou, et al. 2016 (197)	
	NMs.	Liou, et al. 2016 (197)	
		Liou, et al. 2016 (198)	
Clinical	CYP2C19 is not associated with H. pylori eradication rate when	Inaba, et al. 2002 (140)	Weak
	treated with lansoprazole when comparing PMs vs IMs vs NMs.	Hagiwara, et al. 2007 (199)	
		Sugimoto, et al. 2007 (200)	
		Srinarong, et al. 2014 (201)	
Acid secret			
Clinical	CYP2C19 PMs are associated with better acid secretion indices	Adachi, et al. 2000 (149)	High
	when treated with lansoprazole as compared to IMs.	Furuta, et al. 2001 (156)	
		Furuta, et al. 2005 (202)	
		Sugimoto, et al. 2007 (200)	
		Furuta, et al. 2009 (203)	
		Nishino, et al. 2011 (204)	
		Sahara, et al. 2013 (145)	
		Sugimoto, et al. 2014 (146)	
Clinical	CYP2C19 PMs are associated with better acid secretion indices	Adachi, et al. 2000 (149)	High
	when treated with lansoprazole as compared to NMs.	Ieiri, et al. 2001 (155)	
		Furuta, et al. 2001 (156)	
		Shirai, et al. 2002 (157)	
		Furuta, et al. 2005 (202)	
		Sugimoto, et al. 2007 (200)	
		Furuta, et al. 2009 (203)	

	infection or sore throat when treated with lansoprazole as compared to NMs.		
Clinical	CYP2C19 PMs+IMs have an increased risk for upper respiratory	Lima, et al. 2013 (51)	Weak
Toxicity			
	when treated with lansoprazole as compared to NMs.		
Clinical	CYP2C19 PMs+IMs are associated with decreased asthma control	Lang, et al. 2015 (206)	Weak
Other	•		
	when treated with lansoprazole as compared to IMs+NMs.		
Clinical	CYP2C19 PMs are associated with better acid secretion indices	Hata, et al. 2013 (205)	Moderate
		Sugimoto, et al. 2014 (146)	
		Sahara, et al. 2013 (145)	
		Hunfeld, et al. 2008 (98)	
		Howden, et al. 2006 (174)	
		Furuta, et al. 2005 (202)	
	when treated with lansoprazole as compared to NMs.	Shirai, et al. 2002 (157)	
Clinical	CYP2C19 IMs are associated with better acid secretion indices	Furuta, et al. 2001 (156)	High
		Sugimoto, et al. 2014 (146)	
		Sahara, et al. 2013 (145)	
		Nishino, et al. 2011 (204)	

<sup>&</sup>lt;sup>a</sup>Rating scheme described in the **Supplemental Material** IM, intermediate metabolizer; NM, normal metabolizer; PM, poor metabolizer; RM, rapid metabolizer; UM, ultrarapid metabolizer

TABLE S3. EVIDENCE LINKING CYP2C19 TO PANTOPRAZOLE PHENOTYPE

Type of experimental model	Major findings	References	Level of evidence <sup>a</sup>
Metabolism			
Clinical	CYP2C19 PMs are associated with decreased metabolism of pantoprazole as compared to IMs.	Desta, et al. 2009 (207) Furuta, et al. 2009 (203) Furuta, et al. 2010 (208) Thacker, et al. 2011 (209) Thacker, et al. 2013 (210)	High
Clinical	CYP2C19 PMs are associated with decreased metabolism of pantoprazole as compared to NMs.	Choi, et al. 2009 (211) Shao, et al. 2009 (212) Furuta, et al. 2009 (203) Furuta, et al. 2010 (208) Ward, et al. 2010 (213)	High
Clinical	CYP2C19 PMs (as determined by phenotyping) have decreased metabolism of pantoprazole as compared to NMs.	Andersson, et al. 1998 (90)	Moderate
Clinical	CYP2C19 IMs are associated with decreased metabolism of pantoprazole as compared to NMs.	Kearns, et al. 2008 (44) Furuta, et al. 2009 (203) Furuta, et al. 2010 (208) Hunfeld, et al. 2010 (214) Gawronska-Szklarz, et al. 2012 (215) Roman, et al. 2014 (99)	High
Clinical	CYP2C19 IMs are associated with decreased metabolism of pantoprazole as compared to RMs.	Roman, et al. 2014 (99)	Moderate
Clinical	CYP2C19 IMs are associated with decreased metabolism of pantoprazole as compared to UMs.	Roman, et al. 2014 (99)	Moderate
Clinical	CYP2C19 NMs are not associated with altered metabolism of pantoprazole as compared to RMs.	Hunfeld, et al. 2008 (98) Roman, et al. 2014 (99)	Weak
Clinical	CYP2C19 NMs are associated with decreased metabolism of pantoprazole as compared to UMs.	Gawronska-Szklarz, et al. 2012 (215)	Weak

		Roman, et al. 2014 (99)	
Clinical	CYP2C19 PMs are associated with decreased metabolism of	Tanaka, et al. 1997 (216)	Moderate
	pantoprazole as compared to IMs+NMs.	Tanaka, et al. 2001 (217)	
Clinical	CYP2C19 PMs are associated with decreased metabolism of	Desta, et al. 2009 (207)	Moderate
	pantoprazole as compared to NMs+RMs.	Thacker, et al. 2011 (209)	
		Thacker, et al. 2013 (210)	
Clinical	CYP2C19 IMs are associated with decreased metabolism of	Desta, et al. 2009 (207)	Moderate
	pantoprazole as compared to NMs+RMs.	Thacker, et al. 2011 (209)	
		Thacker, et al. 2013 (210)	
		Shakhnovich, et al. 2018 (218)	
		Shakhnovich, et al. 2018 (43)	
Clinical	CYP2C19 PMs+IMs+NMs are associated with decreased	Karaca, et al. 2017 (219)	Moderate
	metabolism of pantoprazole as compared to RMs+UMs.		
Clinical	CYP2C19 PMs+IMs are associated with decreased metabolism of	Kearns, et al. 2010 (113)	Weak
	pantoprazole as compared to NMs+RMs.		
Clinical	CYP2C19 is associated with metabolism of pantoprazole when	Gawronska-Szklarz, et al. 2010	Weak
	comparing IMs vs NMs vs RMs vs UMs.	(220)	
Efficacy			
Remission of			
Clinical	CYP2C19 PMs+IMs are associated with increased healing rate of	Sheu, et al. 2012 (221)	High
	reflux when treated with pantoprazole as compared to NMs.		
Clinical	CYP2C19 is associated with remission of reflux when treated with	Chen, et al. 2010 (222)	Moderate
	pantoprazole when comparing PMs vs IMs vs NMs.		
H. pylori er			
Clinical	CYP2C19 PMs are not associated with altered <i>H. pylori</i> eradication	Oh, et al. 2009 (223)	Weak
	rate when treated with pantoprazole as compared to IMs.		
Clinical	CYP2C19 PMs are not associated with altered <i>H. pylori</i> eradication	Oh, et al. 2009 (223)	Weak
	rate when treated with pantoprazole as compared to NMs.		
Clinical	CYP2C19 NMs are not associated with altered <i>H. pylori</i> eradication	Kurzawski, et al. 2006 (224)	Weak
	rate when treated with pantoprazole as compared to RMs.		
Clinical	CYP2C19 NMs are not associated with altered <i>H. pylori</i> eradication	Kurzawski, et al. 2006 (224)	Weak
	rate when treated with pantoprazole as compared to UMs.		

Clinical	CYP2C19 PMs are not associated with altered <i>H. pylori</i> eradication	Kang, et al. 2008 (225)	Weak
	rate when treated with pantoprazole as compared to IMs+NMs.	Lee, et al. 2014 (226)	
Clinical	CYP2C19 PMs+IMs are associated with increased <i>H. pylori</i>	Kurzawski, et al. 2006 (224)	Moderate
	eradication rate when treated with pantoprazole as compared to	Ormeci, et al. 2016 (227)	
	NMs.		
Clinical	CYP2C19 PMs+IMs are not associated with altered <i>H. pylori</i>	Gawronska-Szklarz, et al. 2010	Weak
	eradication rate when treated with pantoprazole as compared to	(220)	
	NMs+RMs+UMs.		
Clinical	CYP2C19 is <b>not</b> associated with H. pylori eradication rate when	Hsu, et al. 2015 (228)	Moderate
	treated with pantoprazole when comparing PMs vs IMs vs NMs.		
Clinical	CYP2C19 is not associated with H. pylori eradication rate when	Karaca, et al. 2017 (219)	Weak
	treated with pantoprazole when comparing PMs+IMs vs NMs vs		
	RMs+UMs.		
Acid secret	ion indices		
Clinical	CYP2C19 PMs are associated with better acid secretion indices	Oh, et al. 2007 (229)	Weak
	when treated with pantoprazole as compared to IMs.		
Clinical	CYP2C19 PMs are associated with better acid secretion indices	Oh, et al. 2007 (229)	Weak
	when treated with pantoprazole as compared to NMs.		
Clinical	CYP2C19 IMs are associated with better acid secretion indices	Oh, et al. 2007 (229)	Moderate
	when treated with pantoprazole as compared to NMs.	Hunfeld, et al. 2010 (214)	
Clinical	CYP2C19 NMs are not associated with altered acid secretion	Hunfeld, et al. 2008 (98)	Weak
	indices when treated with pantoprazole as compared to RMs.		
Other			
Clinical	CYP2C19 PMs are associated with a decreased number of required	Sheu, et al. 2012 (221)	Moderate
	pantoprazole tablets in patients with reflux in remission as		
	compared to IMs+NMs.		

<sup>a</sup>Rating scheme described in the **Supplemental Material**IM, intermediate metabolizer; NM, normal metabolizer; PM, poor metabolizer; RM, rapid metabolizer; UM, ultrarapid metabolizer

TABLE S4. EVIDENCE LINKING CYP2C19 TO DEXLANSOPRAZOLE PHENOTYPE

Type of experimental model	Major findings	References	Level of evidence <sup>a</sup>
Metabolism			
Clinical	CYP2C19 PMs are associated with decreased metabolism of dexlansoprazole as compared to IMs.	Sun, et al. 2017 (230)	Moderate
Clinical	CYP2C19 PMs are associated with decreased metabolism of dexlansoprazole as compare to NMs.	Sun, et al. 2017 (230)	Moderate
Clinical	CYP2C19 IMs are associated with decreased metabolism of dexlansoprazole as compared to NMs.	Sun, et al. 2017 (230)	Moderate
Clinical	CYP2C19 PMs have decreased metabolism of dexlansoprazole as compared to IMs+NMs.	Grabowski and Lee, <i>et al.</i> 2012 (231)	Weak
Efficacy			_
Clinical	CYP2C19 PMs are not associated with altered acid secretion indices when treated with dexlansoprazole as compared to IMs.	Sun, et al. 2017 (230)	Moderate
Clinical	CYP2C19 PMs are associated with better acid secretion indices when treated with dexlansoprazole as compared to NMs.	Sun, et al. 2017 (230)	Moderate
Clinical	CYP2C19 IMs are not associated with altered acid secretion indices when treated with dexlansoprazole as compared to NMs.	Sun, et al. 2017 (230)	Weak

<sup>&</sup>lt;sup>a</sup>Rating scheme described in the **Supplemental Material** 

IM, intermediate metabolizer; NM, normal metabolizer; PM, poor metabolizer; RM, rapid metabolizer; UM, ultrarapid metabolizer

TABLE S5. EVIDENCE LINKING CYP2C19 TO ESOMEPRAZOLE PHENOTYPE

Type of experimental model	Major findings	References	Level of evidence <sup>a</sup>
Metabolism			
Clinical	CYP2C19 PMs are associated with decreased metabolism of esomeprazole as compared to IMs.	Lou, et al. 2009 (232)	Weak
Clinical	CYP2C19 PMs are associated with decreased metabolism of esomeprazole as compare to NMs.	Lou, et al. 2009 (232) Liu, et al. 2009 (232)	Weak
Clinical	CYP2C19 IMs are associated with decreased metabolism of esomeprazole as compared to NMs.	Liu, et al. 2009 (232) Hunfeld, et al. 2009 (232) Hunfeld, et al. 2012 (233)	Weak
Clinical	CYP2C19 is associated with metabolism of esomeprazole when comparing PMs vs IMs vs NMs.	Yi, et al. 2017 (234)	Moderate
Efficacy			
Remission or			
Clinical	CYP2C19 is associated with remission of reflux when treated with esomeprazole when comparing PMs vs IMs vs NMs.	Sheu, et al. 2008 (235)	Weak
Clinical	CYP2C19 is not associated with healing rate of reflux when treated with esomeprazole when comparing PMs vs IMs vs NMs.	Schwab, et al. 2005 (236) Sheu, et al. 2008 (235)	Moderate
H. pylori erad	ication		
Clinical	CYP2C19 PMs are not associated with altered <i>H. pylori</i> eradication rate when treated with esomeprazole as compared to IMs+NMs.	Kang, et al. 2008 (225) Lee, et al. 2014 (226) Su, et al. 2017 (237)	High
Clinical	CYP2C19 PMs+IMs are associated with increased <i>H. pylori</i> eradication rate when treated with esomeprazole as compared to NMs.	Miehlke, et al. 2006 (132) Miehlke, et al. 2008 (238) Kuo, et al. 2009 (239) Kuo, et al. 2013 (240) Saito, et al. 2015 (241) Hong, et al. 2016 (138) Shimoyama, et al. 2017 (242)	Moderate

Clinical	CYP2C19 is not associated with H. pylori eradication rate when	Sheu, et al. 2005 (137)	High
	treated with esomeprazole when comparing PMs vs IMs vs NMs.	Lee, et al. 2010 (243)	
		Pan, et al. 2010 (244)	
		Wu, et al. 2011 (245)	
		Liou, et al. 2011 (246)	
		Song, et al. 2016 (247)	
		Song, et al. 2016 (248)	
Acid secreti	on indices		
Clinical	CYP2C19 PMs are associated with better acid secretion indices	Sahara, et al. 2013 (145)	Moderate
	when treated with esomeprazole as compared to IMs.	Sahara, et al. 2015 (249)	
Clinical	CYP2C19 PMs are associated with better acid secretion indices	Sahara, et al. 2013 (145)	Moderate
	when treated with esomeprazole as compared to NMs.	Sahara, et al. 2015 (249)	
Clinical	CYP2C19 IMs are associated with better acid secretion indices	Hunfeld, et al. 2010 (214)	Moderate
	when treated with esomeprazole as compared to NMs.	Hunfeld, et al. 2012 (233)	
		Kagami, et al. 2016 (250)	
		Yi, et al. 2017 (234)	
Clinical	CYP2C19 PMs are not associated with altered acid secretion indices when treated with esomeprazole as compared to IMs+NMs.	Li, et al. 2007 (251)	Weak
Clinical	CYP2C19 is not associated with acid secretion indices when treated	Kagami, et al. 2016 (250)	Weak
	with esomeprazole when comparing PMs vs IMs vs NMs.	Yi, et al. 2017 (234)	
Toxicity			
Clinical	CYP2C19 UMs have an increased risk for agranulocytosis when	Dury, et al. 2012 (153)	Weak
	treated with esomeprazole.		
Clinical	CYP2C19 is not associated with risk for adverse events when	Miehlke, et al. 2008 (238)	Moderate
	treated with esomeprazole when comparing PMs vs IMs vs NMs.	, ,	

<sup>a</sup>Rating scheme described in the **Supplemental Material**IM, intermediate metabolizer; NM, normal metabolizer; PM, poor metabolizer; RM, rapid metabolizer; UM, ultrarapid metabolizer

TABLE S6. EVIDENCE LINKING CYP2C19 TO RABEPRAZOLE PHENOTYPE

Type of experimental model	Major findings	References	Level of evidence <sup>a</sup>
Metabolism	CVP2C10 PM	11 1 2001 (252)	TT' 1
Clinical	CYP2C19 PMs are associated with decreased metabolism of rabeprazole as compared to IMs.	Horai, et al. 2001 (252) Ieiri, et al. 2001 (155) Sugimoto, et al. 2004 (253) Hu, et al. 2005 (254) Hu, et al. 2006 (255) Qiao, et al. 2006 (66) Uno, et al. 2006 (256) Miura, et al. 2006 (257) Shimizu, et al. 2006 (258)	High
		Yamano, et al. 2008 (259) Hayato, et al. 2012 (260)	
Clinical	CYP2C19 PMs are associated with decreased metabolism of rabeprazole as compared to NMs.	Horai, et al. 2001 (252) Ieiri, et al. 2001 (155) Shirai, et al. 2001 (61) Lin, et al. 2003 (261) Sugimoto, et al. 2004 (253) Hu, et al. 2005 (254) Qiao, et al. 2006 (66) Uno, et al. 2006 (256) Miura, et al. 2006 (257) Shimizu, et al. 2006 (258) Niioka, et al. 2006 (262) Hu, et al. 2006 (255) Yamano, et al. 2008 (259) Sheng, et al. 2010 (263) Hayato, et al. 2012 (260)	High

Clinical	CYP2C19 PMs (as determined by phenotyping) are associated with	Yasuda, et al. 1995 (87)	Moderate
	decreased metabolism of rabeprazole as compared to NMs.		
Clinical	CYP2C19 IMs are associated with decreased metabolism of rabeprazole as compared to NMs.	Horai, et al. 2001 (252) Ieiri, et al. 2001 (155) Sugimoto, et al. 2004 (253)	Weak
		Hu, et al. 2005 (254) Hu, et al. 2006 (255) Yamano, et al. 2008 (259)	
		Sheng, et al. 2010 (263) Hunfeld, et al. 2012 (233)	
		Hayato, et al. 2012 (260) Roman, et al. 2014 (99)	
Clinical	CYP2C19 IMs are associated with decreased metabolism of rabeprazole as compared to RMs.	Roman, et al. 2014 (99)	Weak
Clinical	CYP2C19 NMs are not associated with altered metabolism of rabeprazole as compared to RMs.	Roman, et al. 2014 (99)	Weak
Clinical	CYP2C19 PMs are associated with decreased metabolism of rabeprazole as compared to IMs+NMs.	Yang, et al. 2009 (264)	Moderate
Clinical	CYP2C19 is associated with rabeprazole metabolism when comparing PMs vs IMs vs NMs.	Toda, et al. 2018 (265)	Weak
Efficacy			
Remission of			
Clinical	CYP2C19 PMs are not associated with healing rate of ulcers when treated with rabeprazole as compared to IMs+NMs.	Ji, et al. 2006 (127)	Moderate
Clinical	CYP2C19 is not associated with remission of reflux when treated with rabeprazole when comparing PMs vs IMs vs NMs.	Kinoshita, et al. 2011 (266)	Moderate
Clinical	CYP2C19 is not associated with healing rate of ulcers when treated with rabeprazole when comparing PMs vs IMs vs NMs.	Ando, et al. 2005 (122) Ando, et al. 2008 (123) Nakamura, et al. 2016 (267)	Weak
Clinical	CYP2C19 does not alter healing rate of erosive lesions when treated with rabeprazole when comparing PMs vs IMs vs NMs.	Yamano, et al. 2008 (259)	Weak
Clinical	CYP2C19 is not associated with healing rate of reflux when treated with rabeprazole when comparing PMs vs IMs vs NMs.	Ariizumi, et al. 2006 (268) Kinoshita, et al. 2018 (269)	Moderate

H. pylori er	adication		
Clinical	CYP2C19 PMs are associated with an increased <i>H. pylori</i> eradication rate when treated with rabeprazole as compared to IMs.	Furuta, et al. 2001 (270) Lay, et al. 2010 (271) Lin, et al. 2017 (130)	Weak
Clinical	CYP2C19 PMs are associated with an increased <i>H. pylori</i> eradication rate when treated with rabeprazole as compared to NMs.	Furuta, et al. 2001 (270) Hokari, et al. 2001 (272) Lay, et al. 2010 (271) Lin, et al. 2017 (130)	Weak
Clinical	CYP2C19 IMs are associated with an increased <i>H. pylori</i> eradication rate when treated with rabeprazole as compared to NMs.	Furuta, et al. 2001 (270) Inaba, et al. 2002 (140) Hsu, et al. 2008 (273) Lay, et al. 2010 (271) Lin, et al. 2017 (130)	Weak
Clinical	CYP2C19 PMs+IMs are associated with increased <i>H. pylori</i> eradication rate when treated with rabeprazole as compared to NMs.	Ormeci, et al. 2016 (227) Sugimoto, et al. 2017 (274) Shimoyama, et al. 2017 (242)	Weak
Clinical	CYP2C19 PMs are not related to altered <i>H. pylori</i> eradication rate when treated with rabeprazole as compared to IMs+NMs.	Kawabata, et al. 2003 (196)	Weak
Clinical	CYP2C19 is not associated with H. pylori eradication rate when treated with rabeprazole when comparing PMs vs IMs vs NMs.	Dojo, et al. 2001 (139) Isomoto, et al. 2003 (275) Lee, et al. 2003 (276) Kuwayama, et al. 2007 (277) Yang, et al. 2009 (264) Lee, et al. 2010 (243) Zhang, et al. 2010 (142) Pan, et al. 2010 (244) Sugimoto, et al. 2014 (278) Sugimoto, et al. 2015 (279) Yang, et al. 2015 (280)	High
Clinical	Acid secretion indices  CYP2C19 PMs are associated with better acid secretion indices when treated with rabeprazole as compared to IMs.	Horai, et al. 2001 (252) Shirai, et al. 2001 (61) Sugimoto, et al. 2004 (253)	Weak

		G : 1 2005 (201)	
		Sugimoto, et al. 2005 (281)	
		Yamano, et al. 2008 (259)	
		Sugimoto, et al. 2010 (282)	
		Nishino, et al. 2010 (283)	
		Sugimoto, et al. 2012 (284)	
		Hayato, et al. 2012 (260)	
		Sugimoto, et al. 2014 (146)	
Clinical	CYP2C19 PMs are associated with better acid secretion indices	Horai, et al. 2001 (252)	High
	when treated with rabeprazole as compared to NMs.	Ieiri, et al. 2001 (155)	
		Shirai, et al. 2001 (61)	
		Lin, et al. 2003 (261)	
		Shimatani, et al. 2004 (285)	
		Sugimoto, et al. 2004 (253)	
		Sugimoto, et al. 2005 (281)	
		Yamano, et al. 2008 (259)	
		Sugimoto, et al. 2010 (282)	
		Nishino, et al. 2010 (283)	
		Hayato, et al. 2012 (260)	
		Sugimoto, et al. 2012 (284)	
		Sahara, et al. 2013 (145)	
		Sugimoto, et al. 2014 (146)	
		Kagami, et al. 2015 (286)	
Clinical	CYP2C19 IMs are associated with better acid secretion indices	Horai, et al. 2001 (252)	Weak
	when treated with rabeprazole as compared to NMs.	Shirai, et al. 2001 (61)	
		Sugimoto, et al. 2005 (281)	
		Hayato, et al. 2012 (260)	
		Hunfeld, et al. 2012 (233)	
		Sugimoto, et al. 2012 (284)	
		Sugimoto, et al. 2014 (146)	
		Kagami, et al. 2015 (286)	
Clinical	CYP2C19 PMs are associated with better acid secretion indices	Hata, et al. 2013 (205)	Weak
	when treated with rabeprazole as compared to IMs+NMs.	Toda, et al. 2018 (265)	
		Kinoshita, et al. 2018 (269)	

Clinical	CYP2C19 is associated with acid secretion indices when treated with rabeprazole when comparing PMs vs IMs vs NMs.	Adachi, et al. 2000 (149) Hu, et al. 2005 (254) Hu, et al. 2006 (255) Li, et al. 2007 (251) Nishino, et al. 2010 (283) Sugimoto, et al. 2010 (282) Furuta, et al. 2010 (148) Sheng, et al. 2010 (263) Sugimoto, et al. 2010 (282)	Weak
	Toxicity		
Clinical	CYP2C19 PMs are associated with increased risk for celecoxib-induced small bowl injury when treated with rabeprazole as compared to IMs+NMs.	Nuki, et al. 2017 (287)	Weak
Clinical	CYP2C19 PMs are not associated with risk for adverse events when treated with rabeprazole as compared to NMs.	Hokari, et al. 2001 (272)	Weak

<sup>&</sup>lt;sup>a</sup>Rating scheme described in the **Supplemental Material**IM, intermediate metabolizer; NM, normal metabolizer; PM, poor metabolizer; RM, rapid metabolizer; UM, ultrarapid metabolizer

TABLE S7. EVIDENCE LINKING *CYP2C19* TO PROTON PUMP INHIBITOR PHENOTYPE (MIXED COHORT)

Type of experimental model	Major findings	References	Level of evidence <sup>a</sup>
Metabolism			
Clinical	CYP2C19 PMs have decreased metabolism of lansoprazole and pantoprazole as compared to NMs.	Metz, et al. 2006 (288)	Weak
Clinical	CYP2C19 IMs have decreased metabolism of lansoprazole and pantoprazole as compared to NMs.	Metz, et al. 2006 (288)	Weak
Efficacy			
Remission or	Healing		
Clinical	CYP2C19 PMs+IMs are associated with increased esophageal eosinophilia remission rate when treated with esomeprazole, lansoprazole, omeprazole, pantoprazole and rabeprazole as compared to NMs+RMs+UMs.	Molina-Infante, et al. 2015 (49)	Moderate
Clinical	CYP2C19 PMs+IMs are associated with increased healing rate of erosive esophagitis when treated with lansoprazole, omeprazole and rabeprazole as compared to NMs.	Kawara, et al. 2017 (289)	Weak
Clinical	CYP2C19 PMs+IMs are associated with decreased healing rate of nonerosive reflux disease when treated with lansoprazole, omeprazole and rabeprazole as compared to NMs.	Kawara, et al. 2017 (289)	Weak
H. pylori erad	ication		
Clinical	CYP2C19 PMs are not associated with altered <i>H. pylori</i> eradication rate when treated with lansoprazole and omeprazole as compared to IMs.	Furuta, et al. 2001 (290)	Weak
Clinical	CYP2C19 PMs are not associated with altered <i>H. pylori</i> eradication rate when treated with lansoprazole, omeprazole and rabeprazole as compared to IMs.	Take, et al. 2003 (291)	Weak
Clinical	CYP2C19 PMs are associated with an increased <i>H. pylori</i> eradication rate when treated with lansoprazole and omeprazole as compare to NMs.	Furuta, et al. 2001 (290) Furuta, et al. 2004 (292)	Moderate

Clinical	CYP2C19 PMs are not associated with altered <i>H. pylori</i> eradication rate when treated with lansoprazole and rabeprazole as compared to NMs.	Miki, et al. 2003 (293)	Weak
Clinical	CYP2C19 PMs are associated with increased <i>H. pylori</i> eradication rate when treated with lansoprazole, omeprazole and rabeprazole as compared to NMs.	Take, et al. 2003 (291) Sugimoto, et al. 2006 (294)	Moderate
Clinical	CYP2C19 IMs are associated with an increased <i>H. pylori</i> eradication rate when treated with lansoprazole and omeprazole as compared to NMs.	Furuta, et al. 2001 (290) Furuta, et al. 2004 (292)	Moderate
Clinical	CYP2C19 IMs are associated with an increased <i>H. pylori</i> eradication rate when treated with omeprazole and pantoprazole as compared to NMs.	Gawronska-Szklarz, et al. 2005 (295)	Weak
Clinical	CYP2C19 IMs are not associated with altered <i>H. pylori</i> eradication rate when treated with lansoprazole and rabeprazole as compared to NMs.	Miki, et al. 2003 (293)	Weak
Clinical	CYP2C19 IMs are associated with increased <i>H. pylori</i> eradication rate when treated with lansoprazole, omeprazole and rabepazole as compared to NMs.	Sugimoto, et al. 2006 (294)	Weak
Clinical	CYP2C19 PMs are associated with an increased <i>H. pylori</i> eradication rate when treated with esomeprazole and pantoprazole as compared to IMs+NMs.	Kang, et al. 2008 (225)	Weak
Clinical	CYP2C19 PMs are not associated with altered <i>H. pylori</i> eradication rate when treated with esomeprazole, lansoprazole, omeprazole, pantoprazole and rabeprazole as compared to IMs+NMs.	Lee, et al. 2014 (226)	Weak
Clinical	CYP2C19 PMs+IMs are associated with increased <i>H. pylori</i> eradication rate when treated with pantoprazole and rabeprazole as compared to NMs	Ormeci, et al. 2016 (227)	Weak
Clinical	CYP2C19 PMs+IMs are associated with increased <i>H. pylori</i> eradication rate when treated with esomeprazole and omeprazole as compared to NMs.	Hong, et al. 2016 (138)	Weak
Clinical	CYP2C19 PMs+IMs are associated with increased <i>H. pylori</i> eradication rate when treated with esomeprazole and rabeprazole as compared to NMs.	Kuo, et al. 2010 (296) Shimoyama, et al. 2017 (242)	Weak

Clinical	CYP2C19 is not associated with H. pylori eradication rate when	Miyoshi, et al. 2001 (297)	Weak
	treated with omeprazole and rabeprazole when comparing PMs vs		
	IMs vs NMs.		
Clinical	CYP2C19 is not associated with H. pylori eradication rate when	Pan, et al. 2010 (244)	Weak
	treated with esomeprazole or rabeprazole when comparing PMs vs		
	IMs vs NMs.		
Acid secreti	on indices		
Clinical	CYP2C19 PMs+IMs are associated with better acid secretion	Egan, et al. 2003 (298)	Weak
	indices when treated with esomeprazole, lansoprazole and		
	omeprazole as compared to NMs.		
Clinical	CYP2C19 IMs+NMs are associated with better acid secretion	Franciosi, et al. 2018 (47)	Weak
	indices when treated with esomeprazole, lansoprazole, omeprazole		
	and pantoprazole as compared to RMs+UMs.		
Clinical	CYP2C19 is not associated with acid secretion indices when treated	Shiotani, et al. 2018 (299)	Weak
	with esomeprazole, lansoprazole and rabeprazole when comparing		
	PMs vs IMs vs NMs.		
Other			
Clinical	CYP2C19 PMs+IMs are associated with decreased resistance to	Franciosi, et al. 2018 (48)	Weak
	treatment with esomeprazole, lansoprazole, omeprazole and		
	pantoprazole in patients with reflux as compared to RMs+UMs.		
Clinical	CYP2C19 PMs+IMs are associated with decreased resistance to	Franciosi, et al. 2018 (48)	Weak
	treatment with esomeprazole, lansoprazole, omeprazole and		
	pantoprazole in patients with reflux as compared to NMs.		
Clinical	CYP2C19 is not associated with resistance to treatment with PPIs in	Wada, et al. 2002 (300)	Weak
	patients with ulcers when comparing PMs vs IMs vs NMs.		

<sup>&</sup>lt;sup>a</sup>Rating scheme described in the **Supplemental Material** 

IM, intermediate metabolizer; NM, normal metabolizer; PM, poor metabolizer; RM, rapid metabolizer; UM, ultrarapid metabolizer

TABLE S8. DOSING RECOMMENDATIONS FOR ESOMEPRAZOLE AND RABEPRAZOLE BASED ON CYP2C19 PHENOTYPE

CYP2C19 Phenotype <sup>a</sup>	Implications for Phenotypic Measures	Therapeutic Recommendation	Classification of Recommendation <sup>b</sup>
CYP2C19 ultrarapid metabolizer	Inconsistent findings regarding the effect of <i>CYP2C19</i> genotype on pharmacokinetics and therapeutic	No recommendation	No recommendation
	response	1000mmonduvion	1000mmondavion
CYP2C19 rapid metabolizer	Inconsistent findings regarding the effect of CYP2C19	No	No
	genotype on pharmacokinetics and therapeutic response	recommendation	recommendation
CYP2C19 normal metabolizer	Inconsistent findings regarding the effect of CYP2C19	No	No
	genotype on pharmacokinetics and therapeutic	recommendation	recommendation
	response		
CYP2C19 likely intermediate	Inconsistent findings regarding the effect of CYP2C19	No	No
metabolizer	genotype on pharmacokinetics and therapeutic	recommendation	recommendation
	response		
CYP2C19 intermediate	Inconsistent findings regarding the effect of CYP2C19	No	No
metabolizer	genotype on pharmacokinetics and therapeutic	recommendation	recommendation
	response		
CYP2C19 likely poor	Inconsistent findings regarding the effect of CYP2C19	No	No
metabolizer	genotype on pharmacokinetics and therapeutic	recommendation	recommendation
	response		
CYP2C19 poor metabolizer	Inconsistent findings regarding the effect of CYP2C19	No	No
	genotype on pharmacokinetics and therapeutic	recommendation	recommendation
	response		

<sup>&</sup>lt;sup>a</sup>The online *CYP2C19* Frequency Table provides phenotype frequencies for major race/ethnic groups, and the online *CYP2C19* Diplotype-Phenotype Table provides a complete list of possible diplotypes and phenotype assignments.

<sup>&</sup>lt;sup>b</sup>Rating scheme described in the **Supplemental Material** 

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