Pharmaceuticals

INNOVATION IN GI MEDICINE

NOVEMBER 2019

Safe Harbor Statement

This presentation contains forward-looking statements. All statements other than statements of historical facts contained in this presentation, including statements regarding our future results of operations and financial position, business strategy, prospective products, product approvals, research and development costs, timing and likelihood of success, plans and objectives of management for future operations, and future results of current and anticipated products, are forward-looking statements.

In some cases, you can identify forward-looking statements by terms such as "may," "will," "should," "expect," "plan," "anticipate," "could," "intend," "target," "project," "contemplates," "believes," "estimates," "predicts," "potential" or "continue" or the negative of these terms or other similar expressions. These statements involve known and unknown risks, uncertainties and other important factors that may cause our actual results, performance or achievements to be materially different from any future results, performance or achievements expressed or implied by the forward-looking statements. These risks, uncertainties and other factors include, without limitation: potential delays in the commencement, enrollment and completion of clinical trials; our dependence on third parties in connection with product manufacturing, research and preclinical and clinical testing; the success of our clinical trials of vonoprazan, and the results of prior clinical trials and other investigator-initiated clinical trials of vonoprazan are not necessarily predictive of our future results and the FDA and comparable foreign regulatory authorities may not accept the data from such prior trials to support approval; regulatory developments in the United States and foreign countries; unexpected adverse side effects or inadequate efficacy of vonoprazan that may limit its development, regulatory approval and/or commercialization, or may result in recalls or product liability claims; our ability to obtain and maintain intellectual property protection for vonoprazan; our ability to comply with our license agreement with Takeda; and other risks described in our filings with the Securities and Exchange Commission (SEC), including our Form S-1 and related prospectus, an any subsequent filings with the SEC. You are cautioned to place undue reliance on these forward-looking statements, which speak only as of the date hereof, and we undertake no obligation to revise or update this presentation to reflect events or circumstances after the date hereof. All forward-looking statements are qualified in their entirety by this cautionary statement, which is made under the safe harbor provisions of the Private Securities Litigation Reform Act of 1995.

This presentation also contains estimates and other statistical data made by independent parties and by us relating to market size and growth and other data about our industry. This data involves a number of assumptions and limitations, and you are cautioned not to give undue weight to such estimates. In addition, projections, assumptions, and estimates of our future performance and the future performance of the markets in which we operate are necessarily subject to a high degree of uncertainty and risk.

Executive leadership team



Tachi Yamada, MD Chairman

CMO & CSO, Takeda President Global Health, Gates Foundation

Chairman R&D, GSK

President, American Gastroenterology Association



David Socks

Venture Partner, Frazier CEO, Outpost Medicine COO, Incline Therapeutics SVP, Cadence Pharmaceuticals



Terrie Curran Incoming CEO*

President I&I Franchise, Celgene Led OTEZLA business from US launch through pending \$13b sale SVP Global Women's Health, Merck



Azmi Nabulsi, MD

Deputy CMO & CSO, Takeda Global Head Development, Takeda Division VP, Abbott



Aditya Kohli, PhD CBO

Vice President, Frazier

VP Business Development, Scout Bio

Engagement Manager, McKinsey



Tom Harris SVP, Regulatory & Quality

SVP/Head of Global Regulatory, Takeda

VP US Regulatory, Takeda Humira Global Project Head, Abbott

*We expect Ms. Curran will succeed Mr. Socks as CEO effective upon the sale of Celgene Corporation to Bristol Myers Squibb Company



POTENTIAL BREAKTHROUGH for acid-related disorders



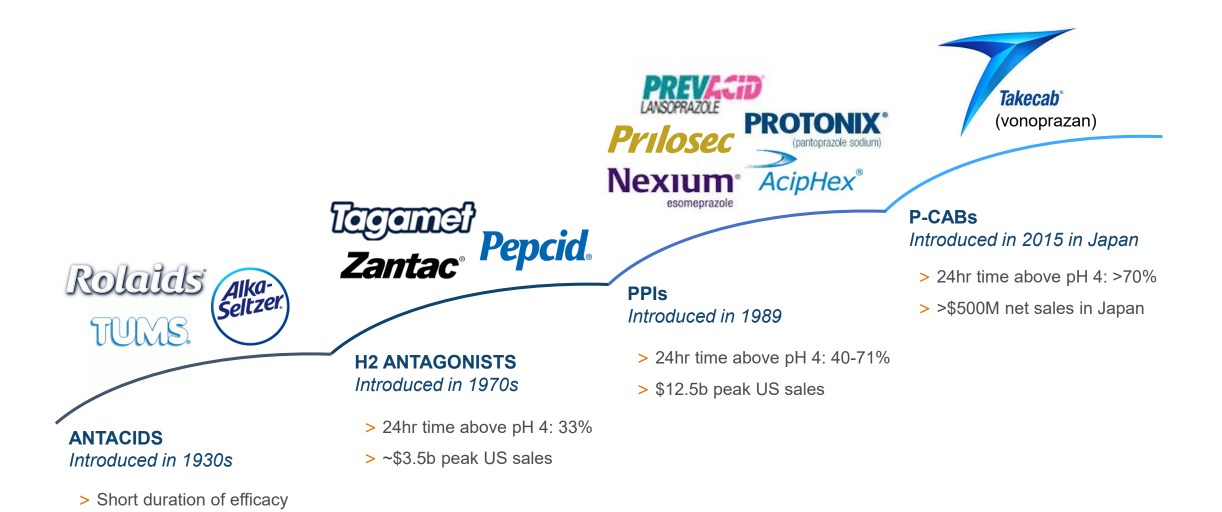
VONOPRAZAN



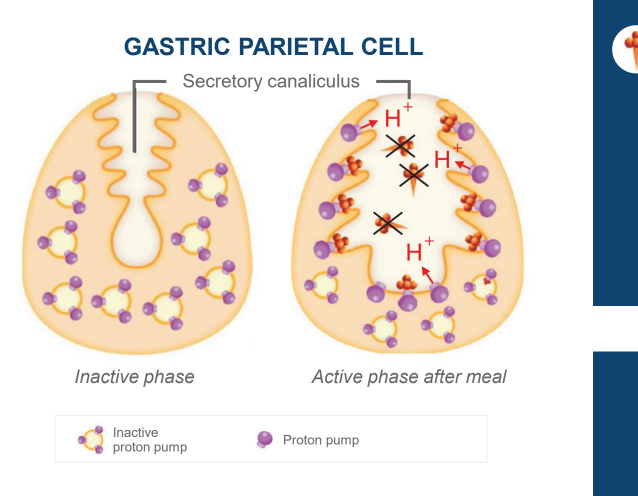
- > Potassium competitive acid blocker (P-CAB)
- > Potentially first-in-class in US, Europe, and Canada
- > US/EU/Canada rights licensed from Takeda
- > 17 Phase 3 studies completed by Takeda in >6,000 subjects
- > Approved in 9 countries across Asia and Latin America
- > >\$500M net sales in Japan in fourth full year on the market



P-CABs: next generation of acid-control therapeutics



PPIs: mechanism limits effectiveness



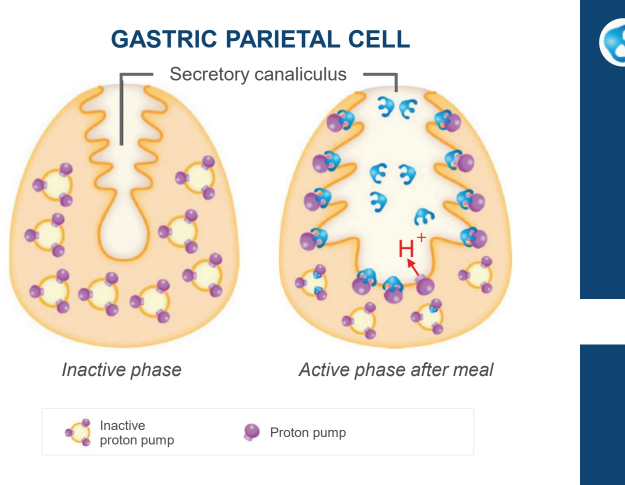
PPI: COVALENTLY BINDING PRODRUG

Acid needed for activation but unstable in presence of acid Meal required to stimulate pumps Short plasma half-life of 1 to 2 hours Primarily metabolized via CYP2C19

Slow onset of action
 Limited potency
 Limited duration of activity

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Vonoprazan: distinct mechanism designed to address PPI shortcomings



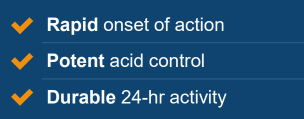
VONOPRAZAN: COMPETITIVE ENZYME INHIBITOR

Stable in acid

Binds with slow dissociation rate

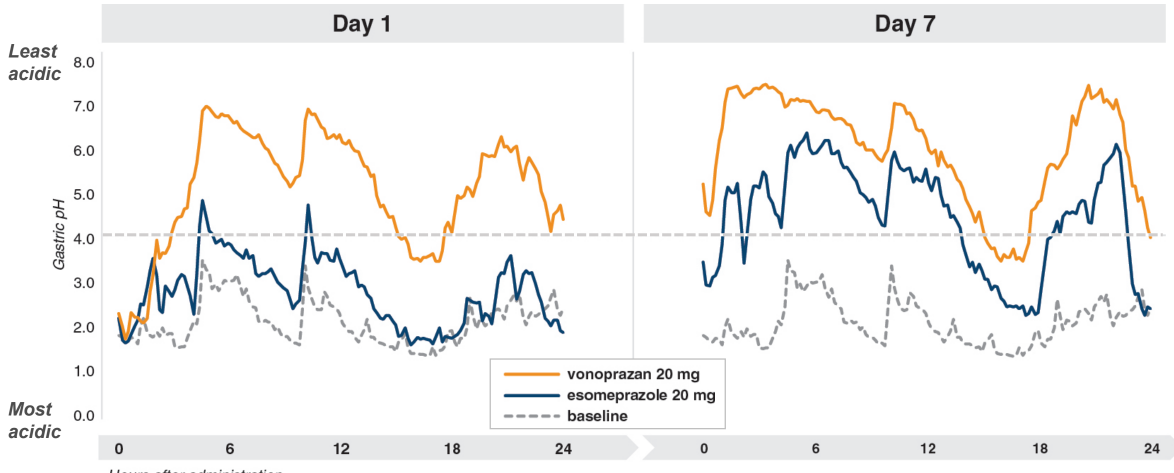
Long plasma half-life of 7 hours

Primarily metabolized via CYP3A4/5





Vonoprazan demonstrated faster and more potent acid control vs. PPI



Hours after administration

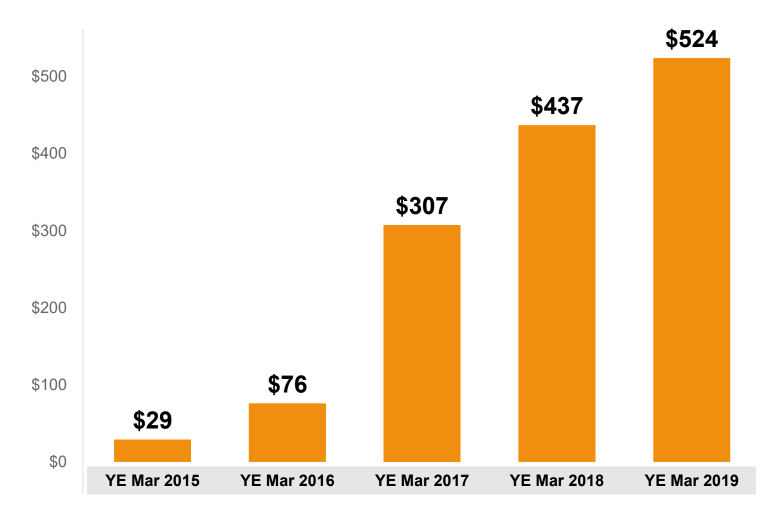


Vonoprazan achieved RAPID ADOPTION and strong sales growth in Japan

TAKECAB[®] (VONOPRAZAN) JAPAN LAUNCH FEBRUARY 2015

Takecab® is a registered trademark of Takeda Pharmaceutical Co. Ltd.

VONOPRAZAN NET SALES, US\$ MILLION*



Note: vonoprazan net sales of approximately \$165M for the three months ended June 30, 2019

*U.S. dollars based on the June 30, 2019 conversion rate of 0.009 yen to one dollar

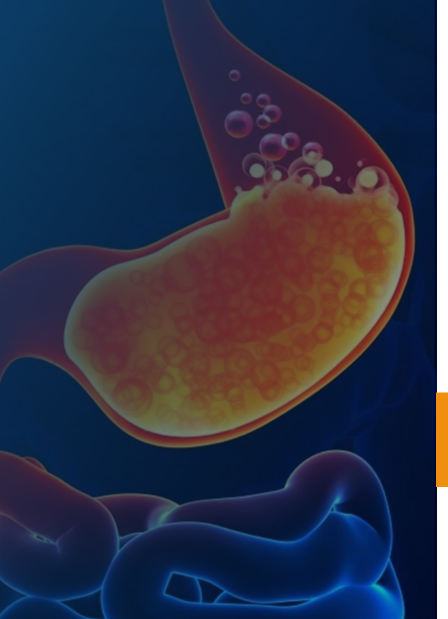


Phathom pipeline

	TARGET INDICATION	PHASE 1*	PHASE 2*	PHASE 3	EXPECTED MILESTONES
Vonoprazan	GERD Healing of erosive esophagitis and relief of heartburn Maintenance of healing of erosive esophagitis and relief of heartburn				Initiate Phase 3 trial 4Q19 Phase 3 results 2021
Vonoprazan + antibiotics	<i>H. pylori</i> treatment Dual therapy (vonoprazan + amoxicillin) Triple therapy (vonoprazan + amoxicillin + clarithromycin)				Initiate Phase 3 trial 4Q19 Phase 3 results 2021



Vonoprazan for GERD



~65M US and ~50M EU5 patients with GERD

6.1 billion PPI doses prescribed in US for the 12 months ended May 31, 2019

~15-45% inadequately treated with PPIs

Many patients experience breakthrough heartburn and recurrence of erosions while on PPIs

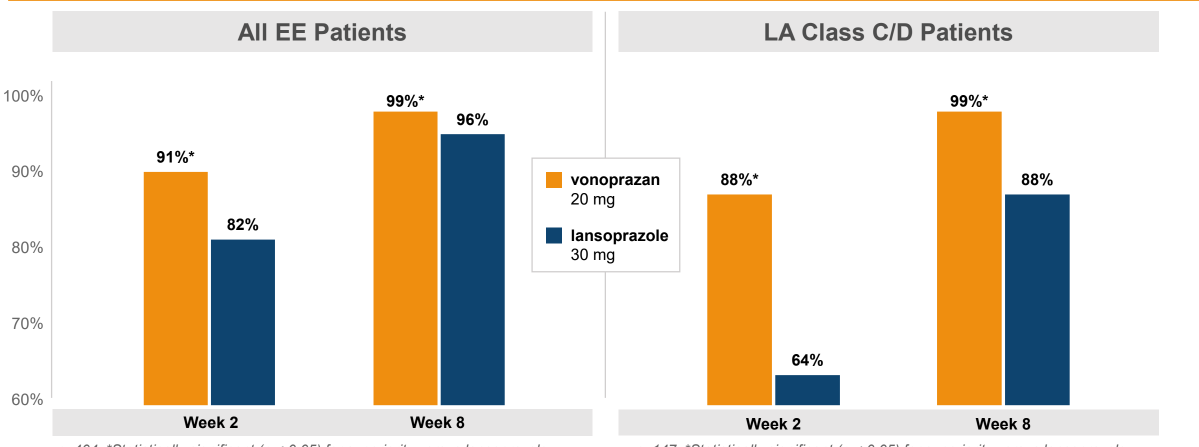
Vonoprazan may offer more rapid, potent, and durable healing and symptom control

El-Serag APT 2010; El-Serag Gut 2014; IQVIA data July 2019



Japan erosive esophagitis (EE) Phase 3: demonstrated faster and improved healing vs. PPI

Patients with healed EE, %



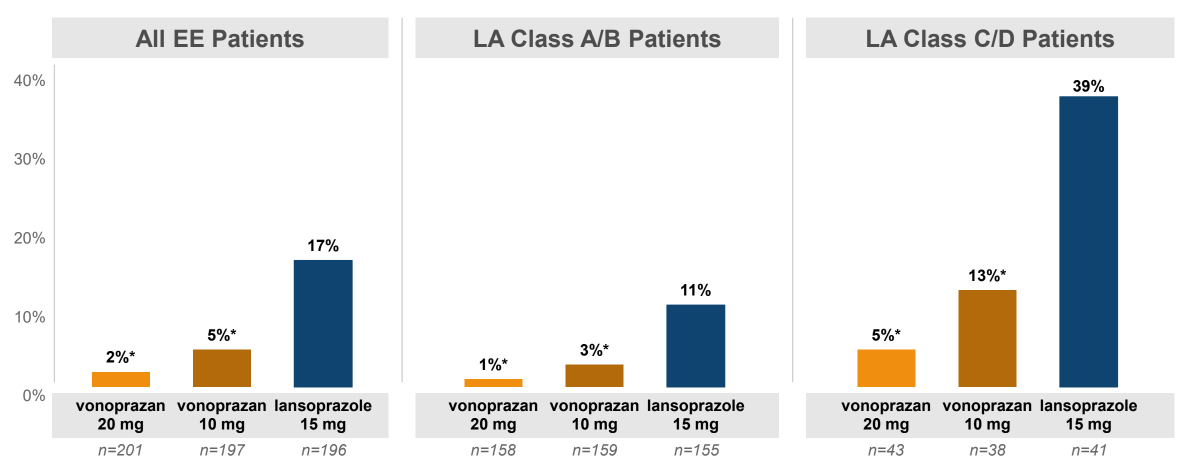
n=404, *Statistically significant (p < 0.05) for superiority versus lansoprazole

n=147, *Statistically significant (p < 0.05) for superiority versus lansoprazole



Japan erosive esophagitis (EE) Phase 3: demonstrated lower recurrence rates vs. PPI

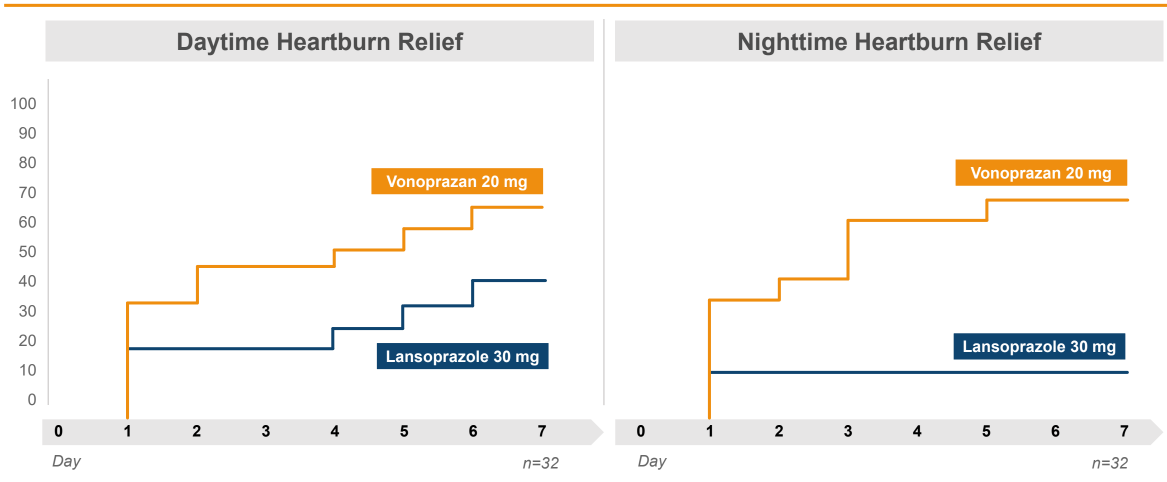
Patients with recurrence of EE at 6 months, %



* p < 0.05 for superiority of vonoprazan 20 mg and vonoprazan 10 mg vs. lansoprazole

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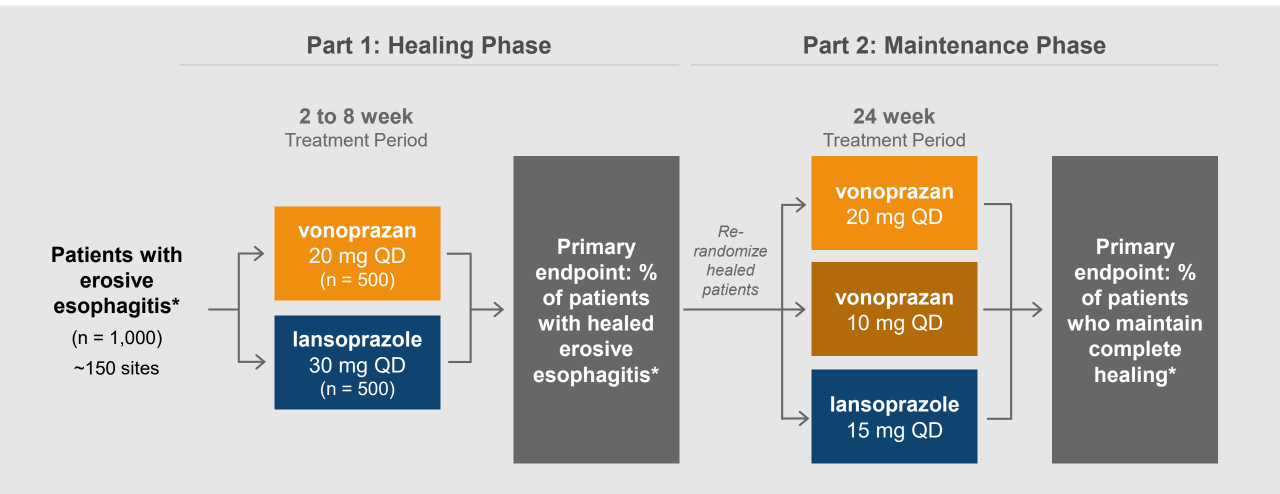
Faster and more complete heartburn relief of vonoprazan vs. PPI



Patients with complete symptom relief, %



Phathom US/Europe EE Phase 3 study design





Vonoprazan for *H. pylori* infection



~2.5M US patients treated annually

H. pylori designated as a Class I carcinogen by WHO and Qualifying Pathogen under FDA GAIN Act

Eradication rates have fallen to <80% due to increasing antibiotic resistance

Antibiotic potency increases at higher pH

Vonoprazan-based regimens may restore eradication rates above 90% in the US and Europe

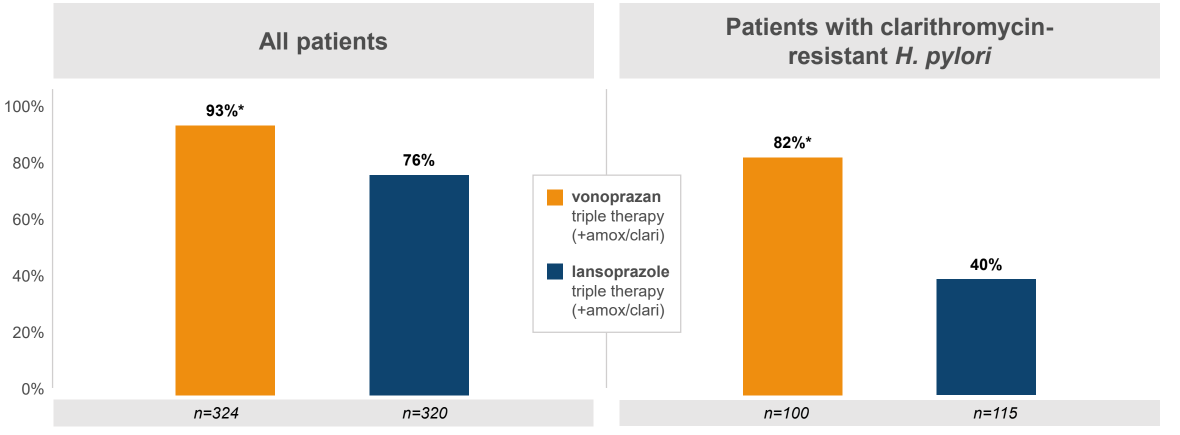
Hooi Gastroenterology 2017; Graham et al 2018; Erah et al 1997



Japan H. pylori Phase 3: vonoprazan triple therapy demonstrated superiority to PPI therapy

First-line triple therapy eradication rates of H. pylori

(combo with amoxicillin/clarithromycin), %



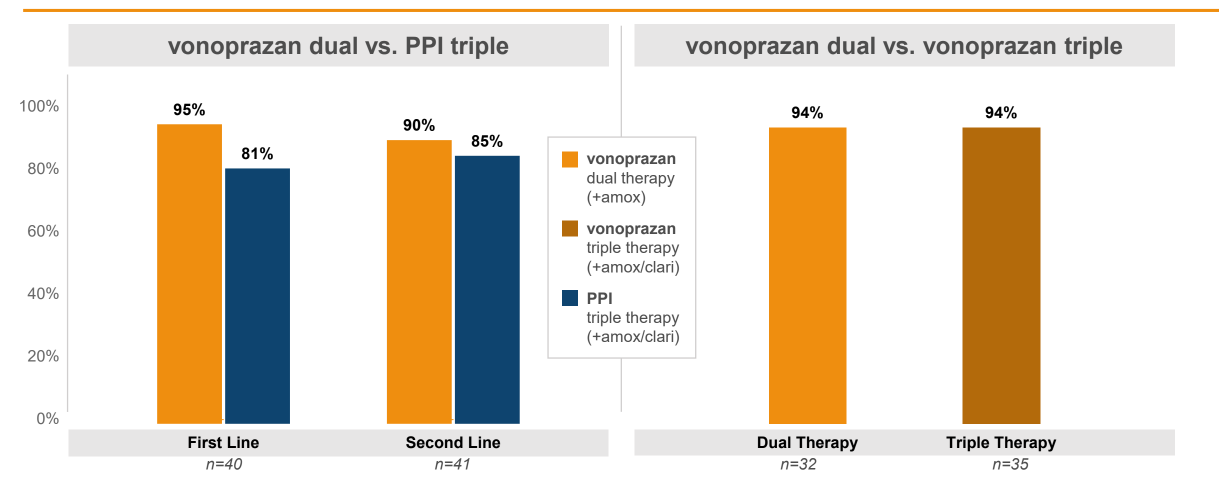
* *p* < 0.0001 for superiority of vonoprazan-based triple therapy to lansoprazole-based triple therapy

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Vonoprazan dual therapy also demonstrated >90% H. pylori eradication

Eradication rates of *H. pylori* (dual or triple therapy)

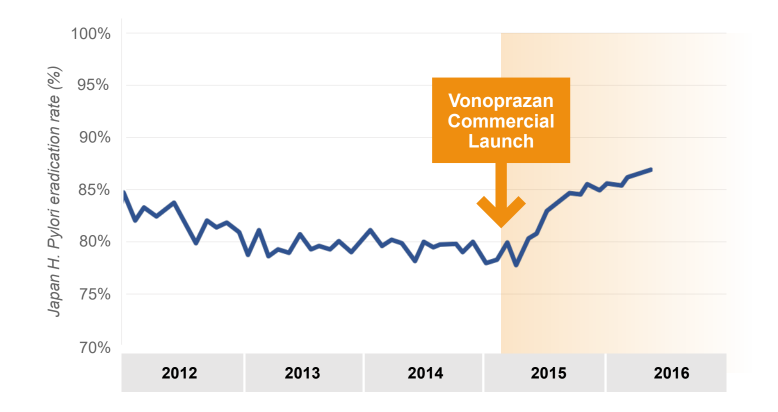
(combo with antibiotics), %





H. pylori eradication rates in Japan have increased since the launch of vonoprazan

VONOPRAZAN-BASED REGIMENS ACHIEVED ~80% SHARE IN JAPAN BY 2016





Deguchi et al Digestion 2019

Phathom US/Europe H. pylori Phase 3 study design



1. vonoprazan 20 mg BID + amoxicillin 1 g BID + clarithromycin 500 mg BID

2. vonoprazan 20 mg BID + amoxicillin 1 g TID (partially blinded)

3. lansoprazole 30 mg BID + amoxicillin 1 g BID + clarithromycin 500 mg BID

Vonoprazan safety profile SIMILAR TO PPIs

6,683 subjects received vonoprazan in clinical studies

No dose-related increase in adverse events observed

>23 million patients received vonoprazan since launch

ADVERSE EVENTS REFLECTED IN JAPANESE PRESCRIBING INFORMATION

Incidence of 0.1-5.0%

Diarrhea ¹	Elevated liver enzymes
Constipation	Rash
Nausea	Eosinophilia

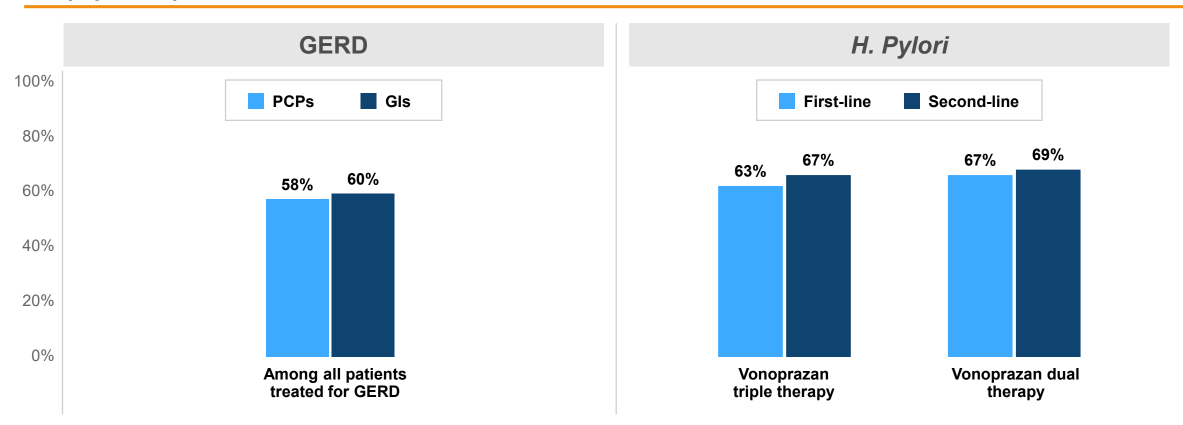
HEPATIC EVENTS OF SPECIFIC INTEREST IN LIGHT OF FIRST-GENERATION PCABs

Pooled data across head-to-head Phase 2 and 3 studies	vonoprazan 10 and 20mg	lansoprazole 15 and 30mg
ALT or AST > 3X ULN or Bilirubin >2X ULN	1.0%	0.8%

1. 10.6% in combination with antibiotics for treatment of H. pylori



US physicians have strong preference to prescribe vonoprazan

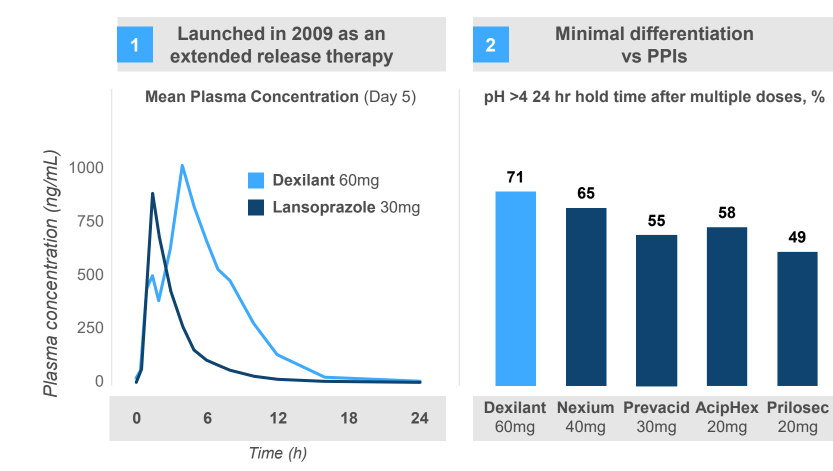


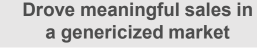
US physician preference share, %

2019 US survey of 100 gastroenterologists and 100 primary care physicians

Dexilant case study: last of the branded PPIs



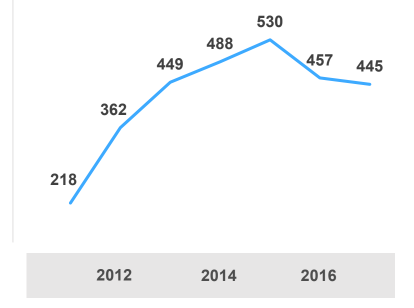




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Dexilant case study: market access

~\$9/dose US WAC1

~90% of commercial and ~80% of Medicare covered lives have access to Dexilant²

65% of commercial covered lives have unrestricted access without step edits or prior authorization²

35% of commercial covered lives have access at the lowest branded cost tier²

FORMULARY STATUS AMONG TOP 5 PLANS By covered lives³ **HEALTH PLAN** COVERAGE Aetna Self-Insured Tier 2 Preferred **Cigna Standard 3-Tier Tier 2 Preferred** (National) **CVS Caremark Advanced Tier 2 Preferred Control Specialty Express Scripts Tier 3 Non-Preferred National Preferred UnitedHealthcare Tier 3 Non-Preferred Advantage 3-Tier NO STEP-EDITS OR PRIOR AUTHORIZATION**



Financial highlights

Cash and Cash Equivalents (As of 9/30/2019) Note: excludes net proceeds from IPO of \$191.5M on October 29, 2019	\$75M
Debt ¹	\$25M
Common Shares Outstanding (As of 11/12/2019)	24,526,537

¹ Silicon Valley Bank Term Loan.
\$25M drawn as of 9/30/19.
Additional \$25M will be available through 3/31/20 subject to certain conditions.



NASDAQ: PHAT

Pharmaceuticals

- Significant unmet medical need
- Established safety and efficacy in Japan
- ✓ Late-stage US/EU program
- Large commercial opportunity
- Seasoned team and investors

