

Title: A Randomized Double-Blind, Double-Dummy, Phase 3 Study to Evaluate the Efficacy and Safety of Oral TAK-438 20 mg Compared to Lansoprazole 30 mg Once- or Twice-Daily in the Treatment of Endoscopically Confirmed Gastric Ulcer Subjects With or Without Helicobacter pylori Infection

NCT Number: NCT03050307

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STATISTICAL ANALYSIS PLAN

STUDY NUMBER: TAK-438 302

Applicable terms of Use A Randomized Double-Blind, Double-Dummy, Phase 3 Study to Evaluate the Efficacy and Safety of Oral TAK-438 20 mg Compared to Lansoprazole 30 mgOnce- or Twice-Daily in the Treatment of Endoscopically Confirmed Gastric Ulcer Subjects With or Without Helicobacter pylori Infection

PHASE 3 Version: 3.0 Date: 05 Jun 2020 **Prepared by:** PPD Based on: Protocol Version: Amendment 02 Property of Take Protocol Date: 21 December 2016

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Approval Signatures 1.1

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3.0 LIST OF ABBREVIATIONS

	3.0 LIST OF ABI	BREVIATIONS
	¹³ C-UBT	¹³ C Urea Breath Test
	AE	adverse event
	ALT	alanine aminotransferase
	AST	aspartate aminotransferase
	BID	twice daily
	CI	confidence interval
	СМН	Cochran–Mantel–Haenszel
	ECG	electrocardiogram
	eCRF	electronic case report form
	EQ-5D-5L	Euro Quality of Life-5D-5L
	F-1	Follow-up 2 weeks posttreatment
	F-2	Follow-up 4 weeks posttreatment
	FAS	full analysis set
	hCG	human chorionic gonadotropin
	HP	Helicobacter pylori
	HP+	Helicobacter pylori infected
	HP-	non-Helicobacter pylori infected
	LLN	lower limit of normal
	MedDRA	Medical Dictionary for Regulatory Activities
	PPI	proton pump inhibitor
	PPS	per protocol set
	PTE	pretreatment event
	Q1	25 th percentile
	Q3	75 th percentile
	QD	oncedaily
	QOL	quality of life
	SAF	safety analysis set
	SD	standard deviation
	TEAE	Treatment-emergent adverse event
	ULN SO	upper limit of normal
	VAS	visual analogue scale
	WHO Drug	World Health Organization Drug Dictionary
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4.0 **OBJECTIVES**

4.1 **Primary Objectives**

Jerms of Use To demonstrate the noninferior efficacy of TAK-438 versus lansoprazole in the treatment of subjects with gastric ulcer.

4.2 **Secondary Objectives**

To demonstrate the noninferiority of Helicobacter pylori (HP) eradication with TAK-438 versus lansoprazole.

To compare the safety of TAK-438 versus lansoprazole in subjects with gastric ulcer.

4.3 **Study Design**

This is a phase 3, multicenter, randomized, double-blind, double-dummy, parallel-group, noninferiority study of TAK-438 versus lansoprazole in subjects who have endoscopic evidence of gastric ulcer with or without *H pylori* infection.

Approximately 830 subjects across approximately 60 sites in Asia will be randomized in a 1:1 ratio to either TAK-438 20 mg or to lansoprazole 30 mg (415 subjects per treatment arm). The randomization will be stratified by Helicobacter pylori (H pylori) status, ie,

- TAK-438: 415 subjects.
- Lansoprazole: 415 subjects.

Enrollment of H pylori negative (ie, non-Helicobacter pylori infected) (HP-) subjects will be initiated first, until the availability of PK and safety data from a drug-drug interaction study to assess the effects of co-administration of TAK-438 with clarithromycin, amoxicillin, and bismuth as quadruple therapy in *Helicobacter pylori* infected (HP+) subjects.

Treatment Period:

A schematic of the study design is included as Figure 4.a. A schedule of assessments is listed in Table 4.a.

Dosing will begin on Day 1 after randomization (Visit 2) and last for up to 8 weeks. Study drug administration will depend on the *H pylori* infection status of the subject.

Administration of the 2 treatments will be according to the following regimens:

H pylori infected (HP+) subjects will take TAK-438 20 mg or lansoprazole 30 mg (blinded with matching placebo) twice daily (BID) in addition to bismuth-containing quadruple antibiotic therapy (amoxicillin 1 g BID, clarithromycin 500 mg BID, and bismuth potassium citrate/bismuth tripotassium dicitrate 600 mg [equivalent to 220 mg bismuth] BID) for 2 weeks. After 2 weeks of eradication therapy, subjects will be required to take only the TAK-438 or lansoprazole once daily (OD) for up to 6 weeks (ie, up to a total of 8 weeks of treatment). The subject remains in the same treatment arm throughout the study duration.

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• Non-*H pylori* infected (HP-) subjects will take TAK-438 or lansoprazole (blinded with matching placebo) QD for up to 8 weeks.

For all subjects, ulcer healing rates will be assessed by endoscopy at Weeks 4 and 8 of treatment. With or without endoscopic evidence of gastric ulcer healing, the treatment duration of the study is no more than 8 weeks.

Subjects (HP+ or HP-) with endoscopic healing of gastric ulcer at Week 4 or at Week 8 will discontinue treatment at that time and enter into the Follow-up period.

HP+ subjects with no healing at Week 8 will discontinue treatment and enter Follow-up.

HP- subjects with no ulcer healing by Week 8 of the treatment period will be considered "completed cases" and be discontinued from the study.

Follow-up Period:

The subjects to be followed up will include both HP+ and HP- subjects with confirmed ulcer healing during the treatment period. HP+ subjects with no ulcer healing during treatment will require one follow-up visit (F-2) to provide a post-study ¹³C-UBT to ascertain HP eradication status. HP- subjects with no ulcer healing are not required to be followed-up.

The recovery of serum gastrin-17 level in follow-up period will be monitored for a maximum of 4 weeks in subjects with endoscopic healing of gastric ulcer confirmed during the treatment period. The follow-up period ends when the recovery of serum gastrin-17 level is confirmed. No additional follow-up is planned for subjects without recovery during the 4-week follow-up period.

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35° Follow-up Period **Screening Period Treatment Period** Up to 8 weeks Up to 28 days Up to 4 weeks TAK-438 20 mg BID (part of HP Gastrin bismuth-containing quadruple recovery and Endoscopic eradication therapy) for 2 weeks, safety followhealing at followed by 20 mg OD up for up to 4 HP+ Week 4 or weeks Lansoprazole 30 mg BID (part of HP ···- - I. A Screening bismuth-containing quadruple Assessments No eradication therapy) for 2 weeks, Gastrin including endoscopic followed by 30 mg OD recovery and Informed healing at safety follow-Consent TAK-438 20 mg QD Q. HPup for Lansoprazole 30 mg QD maximum 4 0 weeks PLUS R HP- subjects: Endoscopy (Week 4 and Week 8) Randomization **HP+ subjects:** Completed HP status Discontinue follow-up at F-

* Enrollment of HP- subjects will be initiated first, until the availability of PK and safety data from a drug-drug interaction study to assess the effects of co-administration of TAK-438 with clarithromycin, amoxicillin, and bismuth as quadruple therapy in HP+ subjects.

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Figure 4.a **Schematic of Study Design**

	Screening Period Treatment Period (b)					Follow-up Period (d)			
Study Day/Week:	D -28 to	D1 (a, u)	W2 (D15)	W4 (D29) (u)	W6 (D43) (u)	W8 (D57) (u)	ET Visit (c)	2w Post- treatment	4w Post- treatmen
Visit Windows (Days):	D -1	-	±3d	±3d	±3d	±3d	Within 14d of last dose	D15 to 22	D29 to 3
Visit Number:	1	2	3	4	5	6	- 0	F-1	F-2
Informed consent	X								
Inclusion/exclusion criteria	Х	X				7	QX		
Demographics and medical history	X					0.			
Medication history	X					<u>n</u>			
Physical examination	X	X	X	X	XO	X	X	X	X
Vital signs	X	X	X	X	ČХ	X	X		
Weight and height	X			1	7				
Concomitant medications	X	X	XC	X	X	X	X	X(e)	X(e)
Concurrent medical conditions	X		6	1					
Hepatitis B and C tests (f)	X(h)		N.						
Clinical laboratory tests (g)	X(h)	14		X		X	X		
Liver function test (r)		0	X		X				
Serum gastrin-17 / Pepsinogen I/II levels (total and ratio)	JSE	x		x		X	X	X	X
Urine pregnancy test (hCG) (i)	· X	X	X	X	X	X	X		
FSH (j)	X								
Guidance on avoidance of pregnancy (i)	X	X	X	X	X	X	X		
ECG	X(l)			X(k)		X	X		
Endoscopy ()	X(l)			X		X	X		
¹³ C-UBT Test to confirm <i>H pylori</i> infection status	X(h)								X(0)
Investigator review of gastrointestinal symptoms		X	X	x	x	X	X		
QOL questionnaire (EQ-5D-5L) (p)	X		X	X	X	X	X		
Randomization via IWRS		X(m)							
Obtain subject number via IWRS	X								
Dispense TAK-438/ lansoprazole via IWRS		X	X	X(n)	X(n)				
Dispense companion drug via IWRS		X(q)							
Drug return/ accountability/ compliance			X	X	X	X	X		
Register subject as discontinued or completed via IWRS				X(k)		X	X		
PTE assessment	X	X							
AE assessment		X							

Table 4.aSchedule of Study Procedures

D=Day, W=Week, ET=Early Termination.

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(a) The day of first investigational drug administration for Treatment period is Day 1. The day before first investigational drug administration for Treatment period is Day -1.

(b) If and when endoscopic healing of gastric ulcer has been confirmed in the subject, he/she will complete the treatment period and move directly into the Follow-up period.

(c) Conduct Final Visit procedures for subjects discontinued from treatment early per Section 9.3.7 of the protocol. (d) The subject will move on to the Follow-up period when endoscopic healing of the gastric ulcer has been confirmed during the treatment period at either Visit 4 (Week 4) or Visit 6 (Week 8). Subjects with confirmed endoscopic healing will complete the scheduled Follow-up period visits for a maximum of 4 weeks until the recovery of serum gastrin-17 level is confirmed and the HP+ subjects will provide the ¹³C-UBT sample at F-2. However, only F-2 will be performed for HP+ subjects who do not have endoscopically confirmed healing (ie, treatment Week 8 visit).

(e) The use of gastric mucosal protective agents may be allowed during the Follow-up period.

(f) Hepatitis B and C analysis in accordance with Section 9.1.8 Table 9.b of the protocol will be conducted at the central laboratory. Hepatitis B and C tests performed as per local routine practice within 14 days prior to randomization (Day 1), before signing of informed consent form will be accepted.

(g) Hematology, serum chemistries, and urinalysis tests will be done at the central laboratory. Local laboratories may be used to perform laboratory tests only under special circumstances, on case by case basis with sponsor's agreement.

(h) Visit window: Day -14 to Day 1 (predose).

(i) Women of childbearing potential.

(i) Only if menopause is suspected.

(k) ECG and registration in IWRS of study completion to be performed at Week 4 only for those subjects in whom endoscopic healing is confirmed; thereafter those subjects will move directly into the Follow-up period.

(1) Visit window: Day -14 to Day 1 (predose). If endoscopy with associated images has been performed within 14 days prior to Day 1, the study-specific endoscopy may be waived in favor of the existing results, providing that the subject has not been using agents that affect the digestive organs since that examination. If an ECG has been performed within 14 days prior to randomization, the study-specific ECG may be waived.

(m) Subjects must not have been given any other agent affecting the digestive organs on the day of randomization and must take the first dose of study medication on the day of randomization during the Day 1 visit.

(n) To be performed as appropriate for those subjects in which endoscopic healing has not been confirmed during visits involving endoscopy

(o) 13 C-UBT should be repeated at least 4 weeks after eradication therapy for all HP+ subjects.

(p) EQ-5D-5L questionnaire shall be the first priority in all procedures (except signing the ICF) at each visit.

(q) Only for HP+ subjects.

(r) Liver function tests include ALT, AST, total bilirubin and direct bilirubin.

(s) Completed by subjects with ulcer healing regardless of HP status.

(t) Completed by subjects with ulcer healing regardless of HP status, only if no recovery at F-1 visit.

(u) The subject will be instructed to present to the clinic prior to taking the study medication at the visits during the treatment e .a anc .ct after For No Fronetty of Takeda. phase in which serum gastrin and pepsinogen I/II levels will be measured or endoscopy will be performed. The study medication

will be taken by the subject after completion of assessments on the days of those study visits.

5.0 ANALYSIS ENDPOINTS

The primary endpoint for this study is the percentage of subjects with endoscopically confirmed healing* of gastric ulcer at Week 4 or 8. *Rate of endoscopic healing

*Rate of endoscopic healing: defined as the proportion of subjects in whom the disappearance of all white coats associated with gastric ulcers has been endoscopically confirmed.

Secondary Endpoints

Secondary endpoints for this study are:

- Percentage of HP+ subjects with successful H pylori eradication after 4 or 8 weeks of treatment, as determined by ¹³C urea breath test (¹³C-UBT) at F-2.
- Percentage of subjects with endoscopically confirmed healing of gastric ulcer at Week 4.
- Percentage of subjects with posttreatment resolution of gastrointestinal symptoms associated with gastric ulcer (epigastric pain [postprandial, fasting, nocturnal], abdominal bloating, nausea/vomiting, heartburn, lack of appetite) at Weeks 2 through 8.

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Additional Endpoints

Additional efficacy endpoints include:

- QOL measurement EQ-5D-5L.
- Percentage of subjects with endoscopically confirmed healing of gastric ulcer at either Week 4 or Week 8 based on their baseline *H pylori* status.
- Percentage of subjects with endoscopically confirmed healing of gastric ulcer at Week 4 based on their baseline *H pylori* status.

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- Percentage of subjects with *H pylori* infection at Baseline who achieve endoscopically confirmed healing of gastric ulcer either at Week 4 or Week 8 based on their *H pylori* status at Follow-up visit F-2.
- Percentage of subjects with *H pylori* infection at Baseline who achieve endoscopically confirmed healing of gastric ulcer at Week 4 based on their *H pylori* status at Follow-up visit F-2.

Safety endpoints include adverse events (AEs), laboratory test values, ECG, vital signs, serum gastrin-17, and pepsinogen I/II values (total and ratio).

6.0 DETERMINATION OF SAMPLE SIZE

Assuming that the true Week 4 or Week 8 healing rate of gastric ulcers is 93.4% for both TAK-438 and lansoprazole, and assuming that the dropout rate is up to 20%, a sample size of 415 subjects per group will provide 98% power to establish noninferiority using a 2-sided 95% CI with a -8% noninferiority margin and will provide a sample size of 290 subjects per group for regulatory requirements in China. Furthermore, assuming that the true eradication rate of *H pylori* is 90% for both TAK-438 and lansoprazole bismuth-containing quadruple therapies and that the dropout rate is up to 20%, a sample size of 270 *H pylori* positive subjects per group will provide 92% power to establish noninferiority using a 2-sided 95% CI with a -10% noninferiority margin and will provide a sample size of 232 subjects per group for regulatory requirements in China. The overall power is therefore approximately 90%. Assuming that the proportion of subjects with *H pylori* infection at Baseline is 80%, a sample size of 332 subjects per group will provide 94% for noninferiority based on same effect size as overall population.

The assumptions of rates for ulcer healing and *H pylori* eradication are based on historical studies with lansoprazole, and literature on omeprazole/amoxicillin/clarithromycin/bismuth quadruple therapy (bismuth-containing) [3], respectively.

In a phase 3 study in Japan to compare the efficacy of lansoprazole and famotidine for healing of gastric ulcer, the Week 8 healing rate was 93.4% for lansoprazole and 82.5% for famotidine, corresponding to a difference of 10.9%. Additionally, in a study of lansoprazole versus placebo in US (M87-091), the Week 8 healing rate was 96.8% for lansoprazole 30 mg and 76.7% for placebo, corresponding to a difference of 20.1%. A noninferiority margin of -8% is specified for ulcer healing rate, which is less than the difference between lansoprazole and famotidine in the Japanese study and less than half of the difference between lansoprazole and placebo in the US study.

Limited data are available from clinical trials with PPI/amoxicillin/clarithromycin/bismuth quadruple therapy (bismuth-containing) [3,4]. In the past phase 3 trials, the eradication rate of *H pylori* with PPI-based triple therapy in HP+ subjects with gastric or gastric ulcers was shown to be 86.4% to 89.2% for lansoprazole/amoxicillin/clarithromycin, 78.8% to 83.0% for omeprazole/amoxicillin/ clarithromycin, and 85.7% to 91.4% for

rabeprazole/amoxicillin/clarithromycin; the range of these eradication rates (78.8% to 91.4%) corresponds to a difference of ≥10%. Hence the noninferiority margin is specified as -10%.

7.0 METHODS OF ANALYSIS AND PRESENTATION

7.1 **General Principles**

All statistical analyses will be conducted using SAS[®] Version 9.4, or higher.

IS OF USE All CIs, statistical tests, and resulting P-values will be reported as 2-sided and will be assessed at α =0.05 significance level unless otherwise stated. P-values will be rounded to 4 decimal places prior to assessment of statistical significance.

Means, 25th percentile (Q1), medians, and 75th percentile (Q3) will be presented to 1 more decimal place than the recorded data. The standard deviations (SDs) will be presented to 2 more decimal places than the recorded data. Minimum and maximum values will be presented using the same number of decimal places as the recorded data. CIs about a parameter estimate will be presented using the same number of decimal places as the parameter estimate.

Where appropriate, variables will be summarized descriptively by study visit. For the categorical variables, the count and proportions of each possible value will be tabulated by treatment group. The denominator for the proportion will be based on the number of subjects who provided non-missing responses to the categorical variable. The proportion will be reported to 1 decimal place.

For continuous variables, the number of subjects with non-missing values, mean, Q1, median, Q3, SD, minimum, and maximum values will be tabulated unless otherwise noted.

7.1.1 Study Definitions

The following definitions and calculation formulas will be used.

- Descriptive statistics: number of subjects with non-missing values, mean, SD, maximum, minimum, Q1, median, Q3.
- Percentage of subjects with endoscopically confirmed healing of gastric ulcer (ie, the rate of endoscopic healing). The proportion of subjects in whom the disappearance of all white coats associated with gastric ulcers has been endoscopically confirmed.
- Screening Failure: Subject who is not eligible prior to randomization.
- Treatment Emergent Adverse Event (TEAE): An AE whose date of onset occurs on or after the start of study drug. A TEAE whose relationship to study drug is missing will be considered drug-related. A TEAE whose intensity is missing will be considered severe.
- Significant TEAE: Any TEAE (not including serious TEAEs) that led to an intervention, including withdrawal of drug treatment, dose increase, dose reduction, dose interrupted or significant additional concomitant therapy.
- Pretreatment Events: A PTE is defined as any untoward medical occurrence in a clinical investigation subject who has signed informed consent to participate in a study but prior to administration of any study drug; it does not necessarily have to have a causal relationship with study participation.

- CI for one sample proportion: Exact (Clopper-Pearson) CI will be used.
- CI for two sample proportion difference: Wald CI will be used unless otherwise specified. •
- ermsofuse CI for two sample proportion difference for efficacy endpoints of healing rate of gastric • ulcer: Newcombe methods will be used.
- Age (years): Age was obtained at screening visit and presented in years.
- Multiracial: Any subject who selects more than one race category will be classified as 'Multiracial'.
- BMI (kg/m^2) : weight $(kg) / (height (m))^2$ (rounded to 1 decimal place).
- Time Since Onset of Current Ulcers (Days): Time since onset of current ulcers to informed consent signed.
- Time Since Onset of Recurrent Ulcers (Days): Time since onset of recurrent ulcers to informed consent signed.
- Pepsinogen I/II Ratio: Pepsinogen I (µg/L) / Pepsinogen II (µg/L) (rounded to 1 decimal ٠ place).
- QTcF interval (msec): QT interval (msec) / (RR interval (sec)) $^{0.33}$ (rounded to the nearest whole number).

Definition of Study Days and Baseline

Study Days: Study Day 1 is defined as the date on which a subject is administered their first dose of the medication ['Date of Dose' in First Dose (QD Treatment) CRF page for HPsubjects; 'Date of Dose' in First Dose (BID Treatment) for HP+ subjects]. Other study days are defined relative to the Study Day 1 with Day 1 being Study Day 1 and Day -1 being the day prior to Study Day 1.

The day after the last dose of the study medication will be defined as Follow-up Day 1. Other follow-up days are defined relative to Follow-up Day 1.

If the date of an event is on or after the date of first dose then Study Day = date of event – date of first dose + 1. If the date of an event is prior to the date of first dose then Study Day = date of event – date of first dose.

Baseline values are defined as the last observed value before the first dose of study medication on Day 1 (including a screening value or unscheduled assessment, if necessary).

Duration of Exposure

Duration of exposure to double-blind study medication (days): date of last dose of doubleblind study medication - date of first dose of double-blind study medication + 1.

Table 7.a Electronic Case Report Form (eCRF) Data to be Applied for Duration of **Exposure**

	eCRF Data to be Applied	l for Duration of Exposure
	Date of First Dose	Date of Last Dose
Double-Blind Study	y Medication	
For HP+ subjects	'Date of Dose' in First Dose (BID Treatment)	'Date of Last Dose' in End of Study
For HP- subjects	'Date of Dose' in First Dose (QD Treatment)	'Date of Last Dose' in End of Study
Each Companion D	Irug	
For <u>HP+ subjects in</u> the first 2 weeks	'Date of Dose' in First Dose (BID Treatment)	'Date of Dose End' in Last Dose (BID treatment)
		Pr
Study Drug Con	npliance	ine
• Study drug co	ompliance (%).	× *0

Study Drug Compliance

- Study drug compliance (%).
 - In the case that the date of last dose of study medication is missing, the last visit date \geq of the study will be used as last dose of study medication.
- = (number of days the study medication taken) / duration of exposure to study medication x100% (rounded to 1 decimal place)
- = (number of study medication dispensed number of study medication returned) $\div \Theta$ / (date of last dose -date of first dose +1) x 100%

Θ: Number of tablet/capsules required to be taken per day.

* Overall study drug compliance for double-blind study medication (HP+ subjects)= $\Sigma_{\ell=1}^{2}$ [(number of study medication dispensed – number of study medication returned) $\div \Theta \iota$]_{x100%} date of last dose of study medication - date of first dose of study medication + 1

 Θ_1 : Number of tablet/capsules required to be taken per day in the first 2 weeks for HP+ subjects.

 Θ_2 : Number of tablet/capsules required to be taken per day in Weeks 3-8 for HP+ subjects.

- * Overall study drug compliance for double-blind study medication (HP- subjects)=
 - (number of study medication dispensed number of study medication returned) $\div \Theta x100\%$ date of last dose of study medication - date of first dose of study medication + 1

Θ: Number of tablet/capsules required to be taken per day in the 8 weeks for HP- subjects.

* Study drug compliance for each companion drug=

(number of study medication dispensed – number of study medication returned) $\div \Theta$ x100%

date of last dose of BID treatment - date of first dose of BID treatment + 1

 Θ : Number of tablet/capsules required to be taken per day.

Table 7.b Number of Study Drug to Be Taken Per Day

Planned DoseΘ: Number of T	Tablet/Capsules Required t	o Be Take	n Per Day
Double-Blind Study Medication			, O
For <u>HP+</u> subjects in the first 2 weeks			n n n n n n n n n n n n n n n n n n n
TAK-438 20 mg (or TAK-438 20 mg matching placebo) BID		2 ta	ablets
Lansoprazole 30 mg (or lansoprazole 30 mg matching placebo) BI	D	2 c	apsules
For <u>HP+ subjects in Weeks 3-8</u> and <u>HP- subjects for 8 weeks</u>		10/0	
TAK-438 20 mg (or TAK-438 20 mg matching placebo) QD		1.0° t	ablets
Lansoprazole 30 mg (or lansoprazole 30 mg matching placebo) QI		<u>)</u> c	apsules
Each Companion Drug	Pr		
For <u>HP+</u> subjects in the first 2 weeks	'/lo		
Amoxicillin 1 g BID	×0	4 c	apsules
Clarithromycin 500 mg BID	, Cr	2 ta	ablets
Bismuth potassium citrate/bismuth tripotassium dicitrate 600 mg E	BID O	4 ta	ablets

Note: Amoxicillin 500 mg/cap, Clarithromycin 500 mg/tab, bismuth potassium citrate/bismuth tripotassium dicitrate OUNSUG 300 mg/tab.

Definition of Study Visit Windows 7.1.2

For each visit, all evaluable observation (ie, non-missing) obtained in the corresponding time interval (ie, window) will be used. This window will establish a time interval around which data will be considered for the analysis of the scheduled visit pertaining to that window. If more than one observation lies within the same visit window, the observation with the closest Study Day to the scheduled Study Day will be used. If there are two observations equidistant to the scheduled Study Day, the later observation will be used. If there are two observations with the same Study Day, the earlier observation will be used. The visit windows and applicable study day ranges are presented in Table 7.c. In the case that the date of last dose of study medication is missing, the last visit date of the study will be used as last dose of study medication.

Visit Analysis Window for Efficacy and Safety Variables Table 7.c

	, cc	<i>SO</i>	Visit Window (Days)							
06	Visit A	Scheduled Visit Day	(a) Variables Scheduled at Week 4, Week 8	(b) Variables Scheduled at Week 4, Week 8	(c) Variables Scheduled at Week 2, Week 4, Week 6, Week 8	(d) Variables Scheduled at Week 2, Week 4, Week 6, Week 8, F-1, F-2	Serum gastrin-17, Pepsinogen I/II (total and ratio) Scheduled at Week 4, Week 8, F-1, F-2	¹³ C-UBT Scheduled at F-2		
<i>S</i> ⁽⁰⁾	Baseline	1	Day -14 - <=1	Day -14 - <=1	Day -28 - <=1	Day -28 - <=1	<=1	Day -14 - <=1		
	Week 2	15		2 – 21 and <= 14 Days after the Last Dose	2 – 21 and <= 14 Days after the Last Dose	2 - 21 and <= 14 Days after the Last Dose				

				Visi	t Window (Days)		
Visit	Scheduled Visit Day	(a) Variables Scheduled at Week 4, Week 8	(b) Variables Scheduled at Week 4, Week 8	(c) Variables Scheduled at Week 2, Week 4, Week 6, Week 8	(d) Variables Scheduled at Week 2, Week 4, Week 6, Week 8, F-1, F-2	Serum gastrin-17, Pepsinogen I/II (total and ratio) Scheduled at Week 4, Week 8, F-1, F-2	¹³ C-UBT Scheduled at F-2
Week 4	29	2-35 and <= 14 Days after the Last Dose	22 – 35 and <= 14 Days after the Last Dose	22 – 35 and <= 14 Days after the Last Dose	22 - 35 and <= 14 Days after the Last Dose	2 – 35 and <= 14 Days after the Last Dose	
Week 6	43		36 - 49 and <= 14 Days after the Last Dose	36 - 49 and <= 14 Days after the Last Dose	36 - 49 and <= 14 Days after the Last Dose	the AP	
Week 8	57	36 - 63 and <= 14 Days after the Last Dose	50 - 63 and <= 14 Days after the Last Dose	50 - 63 and <= 14 Days after the Last Dose	50 - 63 and <= 14 Days after the Last Dose	36 – 63 and <= 14 Days after the Last Dose	
2 Weeks Post- Treatment (F-1)	Follow-up Day 15				Follow-up Day 15 - 28	Follow-up Day 15 - 28	
4 Weeks Post- Treatment (F-2)	Follow-up Day 29			17 31	Follow-up Day 29 - 42	Follow-up Day 29 – 42 (e)	Day 2 - (f)

Note: Baseline values are defined as the last observed value before the first dose of study medication on Day 1 (including a screening value or unscheduled assessment, if necessary). Any value collected on Day 1 but after the administration of first dose will be grouped under Week 2. Note: Week 2, Week 4, Week 6, and Week 8 values are used within 14 Days after the last does. The day after the last dose of the study medication will be defined as 1 Day after the last dose.

(a) Clinical laboratory tests including hematology, serum chemistries (variables other than ALT, AST, total bilirubin, and direct bilirubin), and urinalysis tests, Endoscopy, ECG.

(b) Variables of liver function tests including ALT, AST, total bilirubin, and direct bilirubin.

(c) Vital signs, Urine pregnancy test (hCG), Gastrointestinal symptoms, QOL questionnaire (EQ-5D-5L).

(d) Physical examination.

(e) Only if no serum gastrin-17 or pepsinogen I/II recovery at F-1 visit.

(f) 13 C-UBT should be repeated for HP+ subjects at F-2 to ascertain HP eradication status. If subjects have administered antibiotics (defined as "J01" of the second level ATC code), PPIs (defined as "A02BC" of the fourth level ATC code), or study drug within 14 days prior to 13C-UBT, the data will be excluded from the analysis.

7.1.3 Data Handling Rules

7.1.3.1 Handling of Rate of Endoscopic Healing

A subject in whom the disappearance of all white coats associated with gastric ulcers has been endoscopically confirmed at Week 4 or Week 8 will be considered as a healed subject. Subjects who do not have any endoscopy data during this period will be treated as missing. Subjects who do not meet this requirement will be considered as 'not healed'. Healing rate will be calculated from the healed and not healed subjects.

7.1.3.2 Handling of Percentage of Subjects with Posttreatment Resolution of Each Gastrointestinal Symptom Associated with Gastric Ulcer at Week 2 through 8

Each gastrointestinal symptom related to gastric ulcer (epigastric pain [postprandial, fasting/nocturnal], abdominal bloating, nausea/vomiting, heartburn, lack of appetite) will be captured at the start of the treatment period (Visit 2) in "Gastrointestinal Symptoms" eCRF page.

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At all visits from Visit 2 through the completion of the study drug administration, symptoms noted at Visit 2 will be followed up. If any symptom resolved during the treatment period, the date of resolution will be recorded in the "Gastrointestinal Symptoms" eCRF page. Subjects with symptom who do not have any information on either date of resolution or end of treatment status will be excluded from analysis for corresponding symptom. Subjects without will also be.

Percentage of subjects with posttreatment resolution of each gastrointestinal symptom associated with gastric ulcer at Week 2 through 8 = (number of subjects with date of resolution reported in "If Yes, and resolved, date of resolution" field at the specified gastrointestinal symptom) / (number of subjects with "Yes" selected for the specified gastrointestinal symptom associated with gastric ulcer at the start of treatment period) * 100 (rounded to 1 decimal place).

7.1.3.3 Handling of EQ-5D-5L

The EQ-5D-5L index values will be calculated from the EQ-5D-5L descriptive system scores based on the EQ-5D-5L Crosswalk value sets. If any of the questions are not answered, the EQ-5D-5L index value of the day will be treated as missing.

7.1.3.4 Handling of Gastrin/Pepsinogen samples had been exposed to ambient temperature

Some of gastrin samples and pepsinogen samples were identified to be kept at had been exposed to ambient temperature for more than 2 hours. As for the samples kept at ambient temperature for more than 2 hours, there is no sample stability data for Gastrin & Pepsinogen I/II assays, the testing results for these samples cannot be considered as reliable. Therefore, the affected sample results will be excluded from the analysis to ensure data integrity.

7.1.4 Conventions for Missing AE Dates

Imputing Missing or Partial Dates for AEs is described in Appendix 9.1 Flowcharts for Imputing Missing or Partial Dates for AEs and Concomitant Medications (Guideline for Data Handling Rule Version 1.0, dated of September 20, 2011).

AEs with onset occurring from the time that the subject is first administered study drug (visit 2, randomization) to the subject's last study visit (ie, early termination or follow-up visit) should be included in data listings but not summary tables.

7.1.5 Conventions for Missing Concomitant Medication Dates

Imputing Missing or Partial Dates for Concomitant Medications is described in Appendix 9.1 Flowcharts for Imputing Missing or Partial Dates for AEs and Concomitant Medications.

Analysis Sets

7.2

Subjects whose sites failed CFDA inspection will be excluded from all analysis sets.

Analysis of efficacy variables will be conducted in the full analysis set defined as all subjects who were randomized and received at least 1 dose of the study drug (ie, TAK-438 or

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lansoprazole or matching placebo). Subjects in this set will be analyzed according to the original randomization.

ofUse The primary efficacy endpoint will also be analyzed in the per-protocol analysis set (PPS) defined as all FAS subjects who did not have any of the major protocol deviations listed below (ert Analyses will be based on the randomized treatment.

- Subjects who did not meet inclusion criteria #3. •
- Subjects who met exclusion criteria #8, #11, #12, #13, #14, #15, #16, #17, #18, #19 or #20. •
- Subjects with study medication compliance of less than 70%. •
- Subjects who have been unblinded prior to database lock.
- Subjects who have violated the rules specified in section 7.3 of the protocol.

Analysis of safety variables will be conducted in the safety analysis set defined as all subjects who take at least 1 dose of study medication (ie, TAK-438 or lansoprazole or matching placebo) n, USbnewnang and will be based on the treatment received.

7.3 **Disposition of Subjects**

Study Information 7.3.1

Analysis Set:	All Subjects Who Signed the Informed Consent Form		
Analysis Variables:	Date First Subject Signed Informed Consent Form		
	Date of Last Subject's Last Visit/Contact		
	MedDRA Version		
	WHO Drug Version		
	SAS Version Used for Creating the Datasets		
Analytical Methods:	(1) Study Information		
	Study information shown in the analysis variables section will be provided.		
	~°,		

7.3.2 Screen Failures

Analysis Set: 🔊	All Subjects Who Were Not Randomized	
Analysis Variables:	Age (years) [Min<= - <65, 65<= - <75, 75<= - <=Max]	
XOL	Gender	[Male, Female]
×	Race	[American Indian or Alaska Native, Asian, Black or African American,
~		Native Hawaiian or Other Pacific Islander, White, Multiracial]
		_

Analytical Methods: (1) Screen Failures

Frequency distributions for categorical variables and descriptive statistics for continuous variables will be provided.

7.3.3 Subject Eligibility

7.5.5 Subject E	chgionity	-0
Analysis Set:	All Subjects Who Signed the Informed Consent Form	
Analysis Variables:	Is the subject eligible for randomization into the [Yes, No]	
	Treatment Phase?	S
	Primary Reason for Subject Not Being Eligible [Pretreatment Event/Adverse Ever	ıt,
	Significant Protocol Deviation,	0
	Lost to Follow-Up,	
	Voluntary Withdrawal,	
	Study Termination,	
	Did Not Meet Entrance Criteria,	
	Other]	
Analytical Methods:	(1) Eligibility for Randomization	
	Frequency distributions will be provided. When calculating the percentages	for the
	primary reasons for subject not being eligible, the total number of ineligible subject	ects will
	be used as the denominator.	
	· ©	

7.3.4 Number of Subjects Randomized by Country, Site, and Treatment Group

Analysis Set:	Randomized Set	All's
Analysis Variables:	Randomization Status	[Yes]
Stratum:	Country	[China, Korea, Philippines, Taiwan]
	Site	[Site numbers will be used as categories]
Analytical Methods:	(1) Number of Subjects Ran	domized by Country, Site, and Treatment Group
	Frequency distribution wil	l be provided for each stratum by treatment group and overall.

7.3.5 Disposition of Subjects

Analysis Set:	Randomized Set	
Analysis Variables:	Double-Blind Study Drug Administration	[No]
	Status	
	Reason for Not Being Treated	[Pretreatment Event/Adverse Event,
<pre></pre>	with Double-Blind Study Drug	Significant Protocol Deviation,
20.		Lost to Follow-Up,
		Voluntary Withdrawal, Study Termination,
XOL		Pregnancy, Lack of Efficacy, Other]
4 A	Double-Blind Study Drug Completion Statu	as [Completed Study Drug,
0*		Prematurely Discontinued Study Drug]
and the second s	Reason for Discontinuation of	[Pretreatment Event/Adverse Event,
000	Double-Blind Study Drug	Significant Protocol Deviation,
0(0)		Lost to Follow-Up,
		Voluntary Withdrawal, Study Termination,
		Pregnancy, Lack of Efficacy, Other]

Analytical Methods: (1) Disposition of Subjects

Frequency distributions will be provided for each treatment group and overall. When

calculating percentages for the reasons for not being treated, the total number of subjects not treated by the study drug will be used as the denominator. When calculating percentages for the reasons for discontinuation of study drug, the total number of subjects who prematurely discontinued the study drug will be used as the denominator. ator. policable terms

(2) Flow Chart of Subject Distribution Flow chart will be provided.

7.3.6 Protocol Deviations and Analysis Sets

7.3.6.1 Protocol Deviations

Analysis Set: Randomized Set Analysis Variables: Protocol Deviation

[Entry Criteria, Concomitant Medication, Procedure Not Performed Per Protocol, Study Medication, Withdrawal Criteria]

Analytical Methods: (1) Protocol Deviations

Frequency distribution will be provided by treatment group and overall for each deviation category. A subject who has several deviations will be counted once in each appropriate category. A subject who has several deviations that can be classified into the same category will be counted only once.

0

7.3.6.2 Analysis Sets

Analysis Set:	Randomized Set	Nº-
Analysis Variables:	Analysis Sets	
	Full Analysis Set	[Included]
	Per Protocol Set	[Included]
	Safety Analysis Set	[Included]
Analytical Methods:	(1) Analysis Sets	

Frequency distributions will be provided by treatment group and overall.

Demographic and Other Baseline Characteristics 7.4

Analysis Set:	Randomized Set	
Analysis Variables:	Country	[China, Korea, Philippines, Taiwan]
× 31	Age (years)	[Min<= - <65, 65<= - <75, 75<= - <=Max]
× `	Gender	[Male, Female]
kx l	Race	[American Indian or Alaska Native, Asian,
		Black or African American,
		Native Hawaiian or Other Pacific Islander,
Q ^(C)		White, Multiracial]
*	Height (cm)	[Min<= - <150, 150<= - <160,
		160<= - <170, 170<= - <=Max]
	Weight (kg)	[Min<= - <50, 50<= - <60,

	60<=-<70, 70<=-<80,
	80<=-<=Max]
BMI (kg/m^2)	[Min<= - <18.5, 18.5<= - <25.0,
	25.0<= - <=Max]
Smoking Classification	[The Subject Has Never Smoked,
	The Subject Is a Current Smoker
	The Subject Is an Ex-smoker]
Consumption of Alcohol	[Drink Everyday,
	Drink a Couple of Days Per Week,
	Drink a Couple of Days Per Month,
	Never Drink]
Consumption of Caffeine	[Yes (More Than 5 Times Per Week),
	No (Never Drink or Less Than 5 Times
	Per Week)]
History of H. pylori Eradication Therapy	[Yes (Therapy Received Ever),
	No (Never Receive Any Therapy)]
H. pylori Infection Status (Result of ¹³ -C Urea	[Positive, Negative]
Breath Test)	
Characteristics of Gastric Ulcers	
Location I (a)	[Cardiac Fundus, Upper Gastric Corpus,
O.	Middle Gastric Corpus, Lower Gastric
150	Corpus,
	Gastric Angle, Pyloric Antral Zone]
Location II (a)	[Anterior Wall, Lesser Curvature,
	Posterior Wall, Greater Curvature]
Number of Ulcers Found	
Ulcer Morphology (a)	[Circular, Ellipsoidal, Other]
Ulcer Size (a)	[Minuscule (<5 mm),
	Minor (>=5 mm/<10 mm),
, 7	Intermediate (>=10 mm/<=20 mm),
	Large (>20 mm/<30 mm),
	Giant (>=30 mm)]
History of Gastric Ulcers	
Time Since Onset of Current Ulcers (Days)	
Use of NSAIDs or Low-dose Aspirin (Except	[Yes, No]
Topical Preparations) at the Time of Ulcer	
Onset	
Type of Ulcers	[Primary, Recurrent]
Time Since Onset of Recurrent Ulcers (Days)	
EQ-5D-5L	
EQ-5D-5L Index Value (Screening)	
EQ Visual Analogue Scale (VAS) Score (Scree	ning)
Analytical Methods: (1) Summary of Demographics and Other Bas	eline Characteristics

as

Frequency distributions for categorical variables and descriptive statistics for continuous 15 OF US variables will be provided by treatment group and overall.

(a) In case of multiple lesions, information needs to be obtained on the largest of the lesions.

7.5 **Medical History and Concurrent Medical Conditions**

Analysis Set:	Safety Analysis Set
Analysis Variables:	Medical History
	Concurrent Medical Conditions
Analytical Methods:	(1) Medical History by System Organ Class and Preferred Term
	(2) Concurrent Medical Conditions by System Organ Class and Preferred Term
	Frequency distributions will be provided for each treatment group MedDRA dictional

Frequency distributions will be provided for each treatment group. MedDRA dictionary will be used for coding. Summaries will be provided using SOC and PT, where SOC will be sorted alphabetically and PT will be sorted in decreasing frequency.

A subject with multiple occurrences of medical history or concurrent medical condition within a SOC will be counted only once in that SOC. A subject with multiple occurrences of medical history or concurrent medical condition within a PT will be counted only once in that PT.

Medication History and Concomitant Medications 7.6

Analysis Set:	Safety Analysis Set
Analysis Variables:	Medication History
	Concomitant Medications
Analytical Methods:	(1) Medication History by Preferred Medication Name
	(2) Concomitant Medications That Started Prior to and Were Ongoing at Baseline
	well as Those That Started After Baseline by Preferred Medication Name
	Frequency distributions will be provided for each treatment group. WHO Drug

dictionary will be used for coding. Summaries will be provided using preferred medication names and sorted in decreasing frequency based on the number of reports. A subject who has been administered several medications with the same preferred medication name will be counted only once for that preferred medication name.

Study Drug Exposure and Compliance

7.7.1 TAK-438/Lansoprazole and Matching Placebo Exposure and Compliance

Analysis Set: Safety Analysis Set Analysis Variables: Duration of Exposure to Study Drug (days) $[1 \le - \le 14, 15 \le - \le 28, 29 \le - \le 42, 15 \le - \le 28, 29 \le - \le 42, 15 \le - 10, 10, 15 \le - 10, 10, 10, 10$ 43<= - <=56, 57<= - <=Max] Study Drug Compliance (%) $[Min \le -.. \le 50.0, 50.0 \le -. \le 70.0, 50.$ 70.0<= - <90.0. 90.0<= - <=Max]

Analytical Methods: (1) Study Drug Exposure and Compliance

3rms of USE Frequency distributions of TAK-438/Lansoprazole and matching placebo for categorical variables and descriptive statistics for continuous variables will be provided by treatment group and overall.

7.7.2 **Companion Drugs Exposure and Compliance**

Analysis Set:	Safety Analysis Set with HP+ Subjects at baseline		
Analysis Variables:	Duration of Exposure to Amoxicillin(days)	[1<= - <=14, 15<= - <=Max]	
	Duration of Exposure to Clarithromycin (days)	[1<= - <=14, 15<= - <=Max]	
	Duration of Exposure to Bismuth(days)	[1<= - <=14, 15<= - <=Max]	
	Amoxicillin Compliance (%)	[Min<= - <50.0, 50.0<= - <70.0,	
		70.0<= - <90.0, 90.0<= - <=Max]	
	Clarithromycin Compliance (%)	[Min<= - <50.0, 50.0<= - <70.0,	
		70.0<= - 90.0, 90.0<= - <=Max]	
	Bismuth Compliance (%)	[Min<=- <50.0, 50.0<= - <70.0,	
		70.0<= - <90.0, 90.0<= - <=Max]	
Analytical Methods:	(1) Companion Drugs Exposure and Compliance		
	Frequency distributions of each compar	ion drug (amoxicillin, clarithromycin, and	
bismuth potassium citrate/bismuth tripotassium dicitrate) for categorical variation		tassium dicitrate) for categorical variables and	

descriptive statistics for continuous variables will be provided by treatment group and overall. 121150

7.8 **Efficacy Analysis**

7.8.1 **Primary Efficacy Endpoint(s)**

7.8.1.1 Primary Analysis

Full Analysis Set Analysis Set: Analysis Variable: Percentage of subjects with endoscopically confirmed healing* of gastric ulcer at Week 4 or **(**8: *Rate of endoscopic healing: defined as the proportion of subjects in whom the disappearance of all white coats associated with gastric ulcers has been endoscopically confirmed. Analytical Methods: The healing rate of gastric ulcer at Week 4 or Week 8 and the 2-sided 95% CI will be

provided by treatment group. The healing rate difference between the TAK-438 treatment group and the lansoprazole treatment group (the TAK-438 treatment group minus the lansoprazole group) at Week 4 or Week 8 and the 2-sided 95% CIs using newcombe methods will be provided. If the lower bound of the 95% CI is \geq -8%, non-inferiority for TAK-438 relative to lansoprazole with regard to gastric ulcer healing will be declared.

7.8.1.2 Secondary Analysis

Analysis Set:

Property

Per Protocol Set

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Analysis Variable:	Percentage of subjects with endoscopically confirmed healing* of gastric ulcer at Week 4 or 8 (%)
	*Rate of endoscopic healing: defined as the proportion of subjects in whom the disappearance of all white coats associated with gastric ulcers has been endoscopically confirmed.
Analytical Methods:	A sensitivity analysis will be performed to check the robustness of the results, the same analyses as those in section 7.8.1.1 will be performed using the PPS.

	analyses as those in section 7.8.1.1 will be performed using the PPS.
7.8.2 Secondary	v Efficacy Endpoint(s)
7.8.2.1 Percentaz Week 4	ge of Subjects with Endoscopically Confirmed Healing of Gastric Ulcer at
Analysis Set:	Full Analysis Set
Analysis Variable:	Percentage of Subjects with Endoscopically Confirmed Healing of Gastric Ulcer at Week 4 (%)
Analytical Methods:	The healing rate of gastric ulcer at Week 4 and the 2-sided 95% CI will be provided by treatment group. The healing rate difference between the TAK-438 treatment group and the lansoprazole treatment group (the TAK-438 treatment group minus the lansoprazole group) at Week 4 and the 2-sided 95% CIs using newcombe methods will be provided.

7.8.2.2 Percentage of HP+ Subjects with Successful H pylori Eradication after 4 or 8 Weeks of Treatment, as Determined by ¹³C-UBT at F-2

Full Analysis Set with HP+ Subjects at baseline
Percentage of HP+ Subjects with Successful H pylori Eradication after 4 or 8 Weeks of
Treatment, as Determined by ¹³ C-UBT at F-2 (%)
Frequency distributions will be provided by treatment group along with rates and the two- sided 95% CIs. Also, the differences in the rates between the TAK-438 group and the lansoprazole group (the TAK-438 group – the lansoprazole group) and the two-sided 95% CIs will be provided. If the lower bound of the 95% CI is \geq -10% and the testing described in 7.8.1.1 is successful, noninferiority for TAK-438 relative to lansoprazole with regard to H <i>pylori</i> Eradication will be declared. Details concerning interpretation are described in section 7.8.4.5.

7.8.2.3 Percentage of Subjects with Posttreatment Resolution of Gastrointestinal Symptoms Associated with Gastric Ulcer at Weeks 2 Through 8

Analysis Set:	Full Analysis Set
Analysis Variable:	Percentage of Subjects with Posttreatment Resolution of Gastrointestinal Symptoms
a or	Associated with Gastric Ulcer at Weeks 2 Through 8 (%)
Analytical Methods:	For each symptom (epigastric pain [postprandial, fasting, nocturnal], abdominal bloating, nausea/vomiting, heartburn, lack of appetite), Frequency distributions will be provided by treatment group along with rates and the two-sided 95% CIs. Also, the differences in the rates between the TAK-438 group and the lansoprazole group (the TAK-438 group – the lansoprazole group) and the two-sided 95% CIs will be provided.

7.8.3 Additional Efficacy Endpoint(s)

7.8.3.1 QOL Measurement EQ-5D-5L

EQ-5D-5L Index Value

7.8.3 Additional Additional	Efficacy Endpoint(s) y endpoints include:
7.8.3.1 QOL Mea	usurement EQ-5D-5L
<u>EQ-5D-5L Index Val</u>	lue contra
Analysis Set: Analysis Variables: Covariates: Visit: Analytical Methods:	 Full Analysis Set EQ-5D-5L Index Value <i>H pylori</i> status (Baseline) [HP+, HP-] EQ-5D-5L Index Value (Baseline) Baseline, Week 2, Week 4, Week 6, Week 8 (1) Summary of EQ-5D-5L Index Value Descriptive statistics will be provided for each visit for the observed values and for each post-baseline visit for the changes from baseline (each post-baseline visit - baseline) by treatment group. The mean differences in the changes from baseline between the TAK-438 group and the lansoprazole group (the TAK-438 group – the lansoprazole group) and the two-sided 95% CIs will be provided for each post-baseline visit. (2) ANCOVA The changes from baseline (each post baseline visit - baseline) in the analysis variable will be analyzed using an ANCOVA model with treatment and baseline <i>H pylori Status</i> as factors and baseline EQ-5D-5L index value as a covariate. The ANCOVA analysis will be performed at each post baseline visit. LS means and the two-sided 95% CIs will be provided for each treatment group. The difference in the LS means between the TAK-438 group and the lansoprazole group (the TAK-438 group – the lansoprazole group (the TAK-438 group and the lansoprazole group (the TAK-438 group – the
<i>EQ VAS Score</i> Analysis Set: Analysis Variables: Covariates:	Full Analysis Set EQ VAS Score <i>H pylori</i> Status (Baseline) [HP+, HP-] EQ VAS Score (Baseline)
Analytical Methods:	The same analyses as those in section 7.8.3.1 "EQ-5D-5L Index Value" will be conducted for the EQ VAS score.
7.8.3.2 Percentag either Web	ge of Subjects with Endoscopically Confirmed Healing of Gastric Ulcer at ek 4 or Week 8 Based on Their Baseline H pylori Status
Analysis Set: Analysis Variable:	Full Analysis Set Percentage of Subjects with Endoscopically Confirmed Healing of Gastric Ulcer at either

	either Week 4 or Week 8 Based on Their Baseline <i>H pylori</i> Status
Analytical Methods:	(1) Percentage of Subjects with Endoscopically Confirmed Healing of Gastric Ulcer at
	Week 4 or Week 8 Based on Their Baseline H pylori Status [HP+, HP-]
Analysis Variable:	Percentage of Subjects with Endoscopically Confirmed Healing of Gastric Ulcer at either
i marysis set.	Turi Tihurysis Set

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Percentage of subjects with endoscopically confirmed healing of gastric ulcer and the 2sided 95% CI will be provided by treatment group and baseline *H pylori* status. Healing percentage difference between the TAK-438 treatment group and the lansoprazole treatment group (the TAK-438 treatment group minus the lansoprazole group) and the 2-sided 95% stratified newcombe CI with baseline *H pylori* status as a stratified factor will be provided

7.8.3.3 Percentage of Subjects with Endoscopically Confirmed Healing of Gastric Uleer at Week 4 Based on Their Baseline H pylori Status

Analysis Set:	Full Analysis Set
Analysis Variable:	Percentage of Subjects with Endoscopically Confirmed Healing of Gastric Ulcer at Week 4
	Based on Their Baseline H pylori Status [HP+, HP-]
Analytical Methods:	The same analyses as those in section 7.8.3.2 "Percentage of Subjects with Endoscopically Confirmed Healing of Gastric Ulcer at either Week 4 or Week 8 Based on Their Baseline <i>H pylori</i> Status" will be applied.

7.8.3.4 Percentage of Subjects with H pylori Infection at Baseline Who Achieve Endoscopically Confirmed Healing of Gastric Ulcer Either at Week 4 or Week 8 Based on Their H pylori Status at Follow-up Visit F-2

Analysis Set:	Full Analysis Set with HP+ Subjects at baseline		
Analysis Variable:	Percentage of Subjects with H pylori Infection at Baseline Who Achieve Endoscopically		
	Confirmed Healing of Gastric Olcer Either at Week 4 or Week 8 Based on Their H pylori		
	Status at Follow-up Visit F-2(%)		
Analytical Methods:	(1) Percentage of Subjects with Endoscopically Confirmed Healing of Gastric Ulcer at either Week 4 or Week 8 Based on Their Baseline <i>H pylori</i> Status		
	Percentage of subjects with endoscopically confirmed healing of gastric ulcer and the 2- sided 95% CI will be provided by treatment group and <i>H pylori</i> status at Follow-up Visit F2. Healing percentage difference between the TAK-438 treatment group and the lansoprazole treatment group (the TAK-438 treatment group minus the lansoprazole group) and the 2- sided 95% stratified newcombe CI with <i>H pylori</i> status at Follow-up Visit F-2 as a stratified factor will be provided.		

7.8.3.5 Percentage of Subjects with H pylori Infection at Baseline Who Achieve Endoscopically Confirmed Healing of Gastric Ulcer at Week 4 Based on Their H pylori Status at Follow-up Visit F-2

Analysis Set:	Full Analysis Set with HP+ Subjects at baseline	
Analysis Variable:	Percentage of Subjects with H pylori Infection at Baseline Who Achieve Endoscopically	
el ci	Confirmed Healing of Gastric Ulcer at Week 4 Based on Their H pylori Status at Follow-up	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Visit F-2 (%)	
Analytical Methods:	The same analyses as those in section 7.8.3.4 "Percentage of Subjects with Endoscopically Confirmed Healing of Gastric Ulcer at either Week 4 or Week 8 Based on Their <i>H pylori</i> Status at Follow-up Visit F-2" will be applied	

# 7.8.4 Statistical/Analytical Issues

## 7.8.4.1 Adjustments for Covariates

Analysis Set:	Full Analysis Set		
Allalysis vallable.	(%)		
	Percentage of subjects with endoscopically confirmed healing of gastric ulcer at Week 4 (%) Percentage of subjects with posttreatment resolution of gastrointestinal symptoms associated with gastric ulcer at Weeks 2 through 8 (%)		
Stratified Variable:	<i>H pylori</i> status (Baseline) [HP+, HP-]		
Analytical Methods:	<ul> <li>(1) CMH Test for Percentage of subjects with endoscopically confirmed healing of gastric ulcer at Week 4 or 8 (%)</li> <li>(2) CMH Test for Percentage of subjects with endoscopically confirmed healing of gastric ulcer at Week 4 (%)</li> </ul>		
	(3) CMH Test for Percentage of subjects with posttreatment resolution of gastrointestinal symptoms associated with gastric ulcer at Weeks 2 through 8 (%)		
	A Cochran–Mantel–Haenszel (CMH) test with baseline <i>H pylori</i> status as a stratification factor will be used to compare the above analysis variable between the TAK-438 group and the lansoprazole group for treatment differences. Mantel-Haenszel estimate of risk difference between the TAK-438 group and the lansoprazole group (the TAK-438 group – the lansoprazole group) and the two-sided 95% CI using the Wald method will also be provided		
	provided.		

# 7.8.4.2 Handling of Dropouts or Missing Data

For the primary endpoint "Percentage of subjects with endoscopically confirmed healing of gastric ulcer at Week 4 or 8" and the secondary efficacy endpoints "Percentage of subjects with endoscopically confirmed healing of gastric ulcer at Week 4" and "Percentage of subjects with posttreatment resolution of gastrointestinal symptoms associated with gastric ulcer (epigastric pain [postprandial, fasting, nocturnal], abdominal bloating, nausea/vomiting, heartburn, lack of appetite) at Weeks 2 through 8", missing data will be handled according to section 7.1.3.

Values below the lower limit of quantification will be treated as zero when calculating the descriptive statistics. Values above the upper limit of quantification will be treated as the upper limit values when calculating the descriptive statistics.

# 7.8.4.3 Interim Analyses and Data Monitoring

No interim analysis is planned in this study.

# 8.4.4 Multicenter Studies

Treatment-by-center interaction will not be explored in this study.

# 7.8.4.5 Multiple Comparison/Multiplicity

IS OF USE Statistical inference will be performed at 2-sided 0.05 level of significance or via 2-sided 95% CIs. Adjustment for multiplicity will be performed for the primary efficacy endpoint and the secondary efficacy endpoints in the following order under the closed testing procedure:

- The primary endpoint of healing rate of gastric ulcer during the 8-week treatment will be tested for noninferiority between the TAK-438 group and the lansoprazole group based on the FAS. If the lower bound of the 95% CI is  $\geq$ -8%, noninferiority for TAK-438 relative to lansoprazole with regard to gastric ulcer healing will be declared.
- If the previous test is successful, the secondary endpoint of eradication rate of *H pylori* will be tested for noninferiority between the TAK-438 group and the lansoprazole group based on the FAS. If the lower bound of the 95% CI of the difference is  $\geq -10\%$ , noninferiority for TAK-438 relative to lansoprazole with regard to *H pylori* eradication will be declared.
- If the previous test is successful, the primary endpoint of heating rate of gastric ulcer during the 8-week treatment will be tested for superiority between the TAK-438 group and the lansoprazole group based on the FAS. If the lower bound of the 95% CI is  $\geq 0\%$ , superiority for TAK-438 relative to lansoprazole with regard to gastric ulcer healing will be declared.
- If the previous test is successful, the secondary endpoint of eradication rate of *H pylori* will be tested for superiority between the TAK-438 group and the lansoprazole group based on the FAS. If the lower bound of the 95% Cb of the difference is  $\geq 0\%$ , superiority for TAK-438 relative to lansoprazole with regard to *Hpylori* eradication will be declared.

# 7.8.4.6 Use of an "Efficacy Subset" of Subjects

In addition to analyses on the primary efficacy endpoint using the FAS, sensitivity analyses will also be performed using the PPS to examine the robustness of the results.

# 7.8.4.7 Active-Control Studies Intended to Show Equivalence or Non-Inferiority

For the primary efficacy endpoint, non-inferiority for TAK-438 relative to lansoprazole will be confirmed in the FAS using a non-inferiority margin of 8% as described in section 7.8.1.1 "Primary Analysis".

For the secondary efficacy endpoint of eradication rate of H pylori, non-inferiority for TAK-438 relative to lansoprazole will be confirmed in the FAS using a non-inferiority margin of 10% as described in section 7.8.1.2 "Secondary Analysis".

# 7.8.4.8 Subgroup Analysis

Analysis Set: Full Analysis Set Analysis Variable: Percentage of subjects with endoscopically confirmed healing of gastric ulcer at Week 4 or 8 (%) Percentage of subjects with endoscopically confirmed healing of gastric ulcer at Week 4 (%) Percentage of subjects with posttreatment resolution of gastrointestinal symptoms associated with gastric ulcer at Weeks 2 through 8 (%)

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Subgroups:	Age (years)	[Min<= - <65, 65<= - <75, 75<= - <=Max]
	Gender	[Male, Female]
	H pylori status (Baseline)	[HP+, HP-]
Analytical Methods:	<ul> <li>For subgroup of age and gender, summaring group.</li> <li>For subgroup of <i>H pylori</i> status (Baseling group.</li> <li>(1) Percentage of subjects with endosce Week 4 or 8 (%)</li> <li>(2) Percentage of subjects with endosce Week 4 (%)</li> <li>(3) Percentage of subjects with posttreat associated with gastric ulcer at Weet The same analyses as those in section above subgroups.</li> </ul>	ries (1), (2), and (3) will be provided by treatment e), summaries (3) will be provided by treatment opically confirmed healing of gastric ulcer at opically confirmed healing of gastric ulcer at atment resolution of gastrointestinal symptoms eks 2 through 8 (%) on 7.8.1 and 7.8.2 will be performed for each of the

# Pharmacokinetic/Pharmacodynamic Analysis plicable. Pharmacokinetic Analysis plicable. 7.9

Not applicable.

# 7.9.1 Pharmacokinetic Analysis

Not applicable.

# 7.9.2 Pharmacodynamic Analysis

Not applicable.

### Other Outcomes 7.10

Not applicable.

#### Safety Analysis 7.11

# 7.11.1 Treatment-Emergent Adverse Events

7.11.1.1 Overview of Treatment-Emergent Adverse Events

Analysis Set:	Safety Analysis Set	
Analysis Variables:	TEAE	
Categories:	Relationship to TAK-438/Lansoprazole	[Yes, No]
D.CO.	Relationship to Clarithromycin	[Yes, No]
	Relationship to Amoxicillin	[Yes, No]
	Relationship to Bismuth	[Yes, No]
	Intensity	[Mild, Moderate, Severe]

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Analytical Methods: The following summaries will be provided for each treatment group. (1) Overview of Treatment-Emergent Adverse Events 1) All Treatment-Emergent Adverse Events (number of events, number and percentage of subjects) 2.1) Relationship of Treatment-Emergent Adverse Events to TAK-438/Lansoprazole. (number of events, number and percentage of subjects) 2.2) Relationship of Treatment-Emergent Adverse Events to Clarithromycin (number of events, number and percentage of subjects) 2.3) Relationship of Treatment-Emergent Adverse Events to Amoxicillin (number of events, number and percentage of subjects) 2.4) Relationship of Treatment-Emergent Adverse Events to Bismuth (number of events, number and percentage of subjects) 3) Intensity of Treatment-Emergent Adverse Events (number of events, number and percentage of subjects) 4) Treatment-Emergent Adverse Events Leading to Double-Blind Study Drug Discontinuation (number of events, number and percentage of subjects) 5) Relationship to TAK-438/Lansoprazole of Treatment-Emergent Adverse Events Leading to Double-Blind Study Drug Discontinuation (number of events, number and percentage of subjects) 6) Serious Treatment-Emergent Adverse Events (number of events, number and percentage of subjects) 7.1) Relationship of Serious Treatment-Emergent Adverse Events to TAK-438/Lansoprazole (number of events, number and percentage of subjects) 7.2) Relationship of Serious Treatment-Emergent Adverse Events to Clarithromycin (number of events, number and percentage of subjects) 7.3) Relationship of Serious Treatment-Emergent Adverse Events to Amoxicillin (number of events, number and percentage of subjects) 7.4) Relationship of Serious Treatment-Emergent Adverse Events to Bismuth (number of events, number and percentage of subjects) 8) Serious Treatment-Emergent Adverse Events Leading to Double-Blind Study Drug property of Takeda. Discontinuation (number of events, number and percentage of subjects) 9) Treatment-Emergent Adverse Events Resulting in Death (number of events, number and percentage of subjects) 10) Treatment-Emergent Adverse Events Corresponding to Liver Function Test Abnormalities (number of events, number and percentage of subjects) 11) Significant Treatment-Emergent Adverse Events (number of events, number and percentage of subjects) TEAEs will be counted according to the rules below. Number of subjects Summaries for 2), 5), 7.1), 7.2), 7.3) and 7.4)

- A subject with occurrences of TEAE in both categories (ie, Yes and No) will be counted once in the Related category.
- Summary for 3) A subject with multiple occurrences of TEAE will be counted once for the TEAE with

7.	1	1.1	1.2	D	ispla	avs	of	Treatme	ent-Emei	rgent	Adve	erse	Events
· •			•	~	ispic	vys	<i>v</i> ,	1.0000000	nit Bille	Sein	110000	100	<b>L</b> i Ciiio

7 1	1 1 2 Display	the maximum ir • Summaries othe A subject with r <u>Number of events</u> For each summary, the state of Treatment-Emerication of the second se	ntensity. er than 2), 3), 5), 7.1), 7.2), 7.3) and 7.4) nultiple occurrences of TEAE will be counted only once. ne total number of events will be calculated.
λ.n	lucic Set	Safety Analysis Set	angenii Harterise Erteniis
Ana	alysis Set.	TEAE	, ilo
Cat	egories:	Intensity	[Mild, Moderate, Severe]
	8	Time of Onset (day)	[1<= - <=14, 15<= - <=28, 29<= - <=42, 43<= - <=56, 57<= - <=Max]
Stra	atified Variable:	H pylori status	[HP+, HP-]
		(Baseline)	×0
Ana	alytical	The following summar	ries will be provided using frequency distribution for each treatment
Met	thods:	group.	
		TEAEs will be coded u	using the MedDRA and will be summarized using SOC and PT. SOC will
		soc and PT soc and	y and P1 will be sorted in decreasing frequency for tables provided by
		only or PT only	IT I will be solved in decreasing nequency for tables provided by SOC
		(1) Treatment-Emerg	ent Adverse Events by System Organ Class and Preferred Term
		(2) Treatment-Emerg	gent Adverse Events by System Organ Class
		(3) Treatment-Emerg	gent Adverse Events by Preferred Term
		(4.1) TAK-438/Lanso	prazole-Related Treatment-Emergent Adverse Events by System
		Organ Class and	Preferred Term
		(4.2) Clarithromycin-	Related Treatment-Emergent Adverse Events by System Organ
		Class and Preferr	red Term
		(4.3) Amoxicillin-Rel	ated Treatment-Emergent Adverse Events by System Organ Class
		and Preferred 1 e	rm d Tweetment Emergent Adverse Events by System Organ Class and
		(4.4) Distilutii-Keiateo Preferred Term	a Treatment-Emergent Adverse Events by System Organ Class and
	20.	(5) Intensity of Treat	ment-Emergent Adverse Events by System Organ Class and
	Xe	Preferred Term	
	£	(6) Intensity of TAK-	438/Lansoprazole-Related Treatment-Emergent Adverse Events by
1	0	System Organ Cla	ass, and Preferred Term
AN AN	<b>b</b>	(7) Treatment-Emerg	gent Adverse Events Leading to Double-Blind Study Drug
200		Discontinuation b	y System Organ Class and Preferred Term
00.		(8) TAK-438/Lansop	razole-Related Treatment-Emergent Adverse Events Leading to
		Double-Blind Stu	dy Drug Discontinuation by System Organ Class and Preferred
		1 erm	t Emorgant Advance Evants by System Owen Class and Dusferred
		Term	t-Emergent Auverse Events by System Organ Class and Preferred
		1 (1 111	

- (10) Serious TAK-438/Lansoprazole-Related Treatment-Emergent Adverse Events by System Organ Class and Preferred Term
- (11) Treatment-Emergent Adverse Events by System Organ Class and Preferred Term Over Time
- (12) Most Frequent Treatment-Emergent Adverse Events by System Organ Class and Preferred Term
- (13) Treatment-Emergent Adverse Events Corresponding to Liver Function Test Abnormalities by System Organ Class and Preferred Term
- (14) TAK-438/Lansoprazole-Related Treatment-Emergent Adverse Events Corresponding to Liver Function Test Abnormalities by System Organ Class and Preferred Term
- (15) Significant Treatment-Emergent Adverse Events by System Organ Class and Preferred Term
- (16) TAK-438/Lansoprazole-Related Significant Treatment-Emergent Adverse Events by System Organ Class and Preferred Term
- (17) Most Frequent Non-Serious Treatment-Emergent Adverse Events by System Organ Class and Preferred Term
- (18) Treatment-Emergent Adverse Events by System Organ Class and Preferred Term stratified by H pylori status at Baseline

The frequency distribution will be provided according to the rules below. <u>Number of subjects</u>

• Summary tables other than (5), (6) and (11) A subject with multiple occurrences of TEAE within a SOC will be counted only once in

that SOC. A subject with multiple occurrences of TEAE within a SOC will be counted only once in that SOC. A subject with multiple occurrences of TEAE within a PT will be counted only once in that PT. Percentages will be based on the number of subjects in the safety analysis set.

• Summary tables for (5) and (6)

A subject with multiple occurrences of TEAE within a SOC or a PT will be counted only once for the TEAE with the maximum intensity. Percentages will be based on the number of subjects in the safety analysis set.

Summary table for (11) A subject with a TEAE that occurs in more than one interval is counted in all the intervals that the TEAE occurs. For each time interval, a subject with multiple occurrences of TEAE within a SOC or a PT will be counted only once in that SOC or PT.

When calculating percentages for each time interval, the number of subjects at risk (ie, subjects who either have an exposure or have an occurrence of TEAE, during or after the corresponding time interval) will be used as the denominator. The number of subjects whose onset of any one of the TEAEs is within the time interval will be used as the numerator.

Summary table for (12)

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Most frequent TEAEs refer to PTs whose percentages are at least 2% in any one of the treatment groups.

Summary table for (17)

Most frequent non-serious TEAEs refer to PTs that are not serious whose percentages are at least 5% in any one of the treatment groups. If there are no PTs whose percentages exceed 5%, the threshold is lowered to 2%. When counting the number of "Subjects With Any TEAEs", subjects with at least one of these most frequent non-serious TEAEs will be counted.

# 7.11.2 Pretreatment Events

# 7.11.2.1 Displays of Pretreatment Events

Analysis Set: All Subjects Who Signed the Informed Consent Form

Analysis Variables: PTE

orms of Use Analytical Methods: The following summaries will be provided using frequency distribution. Pretreatment Events will be coded using the MedDRA and will be summarized using SOC and PT. SOC will be sorted alphabetically and PT will be sorted in decreasing frequency.

# (1) Pretreatment Events by System Organ Class and Preferred Term

(2) Serious pretreatment Events by System Organ Class and Preferred Term

The frequency distribution will be provided according to the rules below. Number of subjects A subject with multiple occurrences of Pretreatment Event within a SOC will be counted only once in that SOC. A subject with multiple occurrences of Pretreatment Event within a PT will

# 7.11.3 Clinical Laboratory Evaluations

	once in that SOC. A subject	with multiple occurrences of Pr	effeatment Event within a PT will
	be counted only once in that	t PT.	
7.11.3 Clinical	Laboratory Evaluation	s and	
7.11.3.1 Hemato	logy and Serum Chemist	try	
Analysis Set:	Safety Analysis Set	ON	
Analysis Variables:	Hematology	150	
	Red Blood Cells (×10 ¹² /I	L) White Blood Cells ( $\times 10^{9}/L$ )	Hemoglobin (g/L)
	Hematocrit (%)	Platelets ( $\times 10^9/L$ )	
	White Blood Cell Fraction	ons (Neutrophils (%), Eosinophil	s (%), Basophils (%),
	Monocytes (%), Lympho	ocytes (%))	
	Serum Chemistry		
	ALT (U/L)	ALP (U/L)	AST (U/L)
	GGT (U/L)	Total Bilirubin (µmol/L)	Direct Bilirubin (µmol/L)
<	LDH (U/L)	CK (CPK) (U/L)	Albumin (g/L)
20.	Total Protein (g/L)	Creatinine (µmol/L)	BUN (mmol/L)
Leor	Uric Acid (mmol/L)	Total Cholesterol (mmol/L)	Triglycerides (mmol/L)
( A DI	Glucose (mmol/L)	Potassium (mmol/L)	Sodium (mmol/L)
Ŏ	Magnesium (mmol/L)	Calcium (mmol/L)	Inorganic Phosphorus (mmol/L)
and the second s	Chloride (mmol/L)	Serum Iron (µmol/L)	Vitamin B ₁₂ (pmol/L)
21000	Note: 1) Fasting laboratory laboratory tests for hematol	test in planned per protocol. 2) T	The central laboratory will perform
X			

Variables of Liver function tests including ALT, AST, Total Bilirubin, and Direct Bilirubin: Baseline, Week 2, Week 4, Week 6, Week 8

Variables other than ALT, AST, Total Bilirubin, and Direct Bilirubin:

Visit:

Baseline, Week 4, Week 8

Analytical Methods: For each variable, summaries (1) and (2) will be provided by treatment group.

- For applicable variables, summaries (3) and (4) will be provided by treatment group. (1) Summary of Laboratory Test Results and Change from Baseline by Visit
- 150 USE Descriptive statistics for observed values and changes from baseline (each post-baseline visit - baseline) will be provided for each visit.
- (2) Summary of Shifts of Laboratory Test Results

Shift tables showing the number of subjects in each category at baseline and each postbaseline visit will be provided. For each laboratory test, the laboratory values will be classified as "Low", "Normal" or "High" relative to the normal reference range provided by the central laboratory. The shift tables will be based on these classifications.

(3) Number and Percentage of Subjects with Markedly Abnormal Values of Laboratory **Parameters** 

Overall frequency distributions of MAV during treatment phase will be provided. If a laboratory parameter has both lower and upper MAV criteria, analysis will be performed for each. Further details are given in Appendix.

(4) Number and Percentage of Subjects with Elevated Liver Enzyme Laboratory Parameters

Overall frequency distributions of elevated hepatic parameters during treatment phase will be provided. Further details are given in Appendix.

ALT=Alanine Aminotransferase, ALP=Alkaline Phosphatase, AST=Aspartate Aminotransferase, GGT= $\gamma$ -Glutamyl Transferase, LDH=Lactate Dehydrogenase, CK=Creatine Kinase, BUN=Blood Urea Nitrogen.

7.11.3.2	Urinalysis
	~

	Analysis Set:	Safety Analysis Set		
	Analysis Variables :	Urinalysis		
		Appearance	Color	рН
		Ketones	Protein	Urine Glucose
		Nitrite	Urobilinogen	Blood
		Specific Gravity		
	Visit: Kateda.	Note: For Urinalysis laboratory numeric results, the others are of Baseline, Week 4, Week 8	tests, only specific gravity vari- categoric variables with characte	able is continuous variable with er results.
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Analytical Methods:	For specific gravity variable, su For each variable other than spe group.	immaries (1) and (2) will be pro ecific gravity, summaries (2) wi	vided by treatment group. Il be provided by treatment
8		(1) Summary of Urine Labora Descriptive statistics for ob visit - baseline) will be prov	atory Test Results and Change served values and changes from vided for each visit.	e from Baseline by Visit baseline (each post-baseline

(2) Summary of Shifts of Urine Laboratory Test Results Shift tables showing the number of subjects in each category at baseline and each postbaseline visit will be provided. For specific gravity laboratory test, the laboratory values will be classified as "Low", "Normal" or "High" and for each variable other than specific gravity will be classified as "Normal" or "Abnormal" relative to the normal reference range provided by the central laboratory. The shift tables will be based on these classifications.

7.11.3.3 Serum Gastrin, Pepsinogen I/II

Analysis Set:	Safety Analysis Set
Analysis Variables :	Serum Gastrin (pmol/L)
	Pepsinogen I (µg/L)
	Pepsinogen II (µg/L)
	Pepsinogen I/II Ratio
	Note: 1) Fasting laboratory test in planned per protocol (2) The central laboratory will perform
	laboratory tests for hematology, serum chemistries, hepatitis and urinalysis.
Visit:	Baseline, Week 4, Week 8, Visit F-1, Visit F-2
Analytical Methods:	The following summaries will be provided for each treatment group.
	(1) Summary of Serum Gastrin and Pepsinogen I/II Results
	For each visit, descriptive statistics for observed values and changes from baseline (each
	post-baseline visit - baseline) will be provided.
	500
7.11.4 Vital Sign	ns
A	

Analysis Set:	Safety Analysis Set
Analysis Variables:	Body Temperature (C)
	Systolic Blood Pressure (mmHg)
	Diastolic Blood Pressure (mmHg)
	Pulse (bpm)
Visit:	Baseline, Week 2, Week 4, Week 6, Week 8
Analytical Methods:	For each variable, summaries (1) and (2) will be provided by treatment group.
	(1) Summary of Vital Signs Parameters and Change from Baseline by Visit
×Q.	Descriptive statistics for observed values and changes from baseline (each post-baseline
	visit – baseline) will be provided for each visit.
XOT	(2) Number and Percentage of Subjects with Markedly Abnormal Values of Vital Signs
	Parameters
ist O.	Overall frequency distributions of MAV during treatment phase will be provided. If a vital
	sign parameter has both lower and upper MAV criteria, analysis will be performed for
Sec.	each. Further details are given in Appendix.
OKU .	

7.11.5 12-Lead ECGs

Analysis Set: Safety Analysis Set

Analysis Variables:	Heart Rate (bpm)	01
	RR Interval (msec)	150
	PR Interval (msec)	
	QT Interval (msec)	S
	QTcF Interval (msec)	- the
	QRS Interval (msec)	$\sqrt{\circ}$
	Interpretation	["Within Normal Limits",
		"Abnormal, Not Clinically Significant",
T T 1		"Abnormal, Clinically Significant"]
Visit:	Baseline, Week 4, Week	8
Analytical Methods:	For each variable other t	han interpretations, summary (1) will be provided by treatment group.
	For applicable variables,	summary (2) will be provided by treatment group.
	For interpretation, summ	hary (3) will be provided by treatment group.
	(1) Summary of ECG I	arameters and Change from Baseline by Visit
	Descriptive statistics	for observed values and changes from baseline (each post-baseline
	visit – baseline) will	be provided for each visit.
	(2) Number and Percer	itage of Subjects with Markedly Abnormal values of ECG
	Parameters	stributions of MAR during tractment phase will be provided. If an
	ECG laboratory para	mater has both lower and upper MAV aritaria, analysis will be
	nerformed for each	Further details are given in Appendix
	(3) Summary of Shifts	of FCC Parameters
	Shift tables showing	the number of subjects in each category at baseline and each post-
	baseline visit will be	ntovided
		R. C. Lavan
7.11.6 Other Ol	oservations Related	to Safety
71161 Physica	1 Examination	
7.11.0.1 1 hysica		
Analysis Set:	Safety Analysis Set	
Analysis Variables:	Results	["Yes", "No"]
Visit:	Screening, Baseline, We	ek 2, Week 4, Week 6, Week 8, Visit F-1, Visit F-2
Analytical Methods:	Summary (1) will be pro	vided by treatment group.
	(1) Number and Percei	stributions of physical examination results during treatment phase will
<u>k</u>	be provided	surbutions of physical examination results during treatment phase with
0,	be provided.	
7.11.7 Subgrou	p Analysis for China	I
Analysis Set:	Subjects in China	
Υ.	Subjects in Countries of	her than China
Analytical Methods:	(1) The same analyses a China.	as those in section 7.3 to 7.11.6 will be conducted for the subjects in
	(2) The same analyses a	as those in section 7.3 to 7.11.6 will be conducted for the subjects in

countries other than China.

Analysis Set: Subjects in Taiwan Analytical Methods: (1) The same analyses as those in section 7.8.1.1, 7.8.1.2, 7.8.2.1, 7.8.2.2, 7.8.2.3, 7.11, 1, 1, and 7.11.3.3 will be conducted for the subjects in Taiwan.

Analysis Set: Subjects in Korea

- Analytical Methods: (1) The same analyses as those in section 7.8.1.1, 7.8.1.2, 7.8.2.1, 7.8.2.2 7.8.2.3, 7.11.1.1, and 7.11.3.3 will be conducted for the subjects in Korea.
 - (2) For primary endpoint, the forest plot was presented by study level and region (Korea, China, and Taiwan).

7.12

Not applicable.

7.13

Changes in the Statistical Analysis Plan in and Subject anges from the original SAP (Version: A Control Subject is been for The changes from the original SAP (Version: 2.0, Date: 21 April 2020) to the amended SAP were described below with a rationale for changes provided.

- It has been found that the date of tast dose of study medication for one patient. Hence, it has been described in section 7.1.1 and 7.1.2 that the end of study will be used instead to complete the missing value.
- Since missing case for compliance described in section 7.7.1 and 7.7.2 does not exist, "Missing" category has been removed.
- Some gastrin samples and pepsinogen samples were exposed to ambient temperature for more than 2 hours. As for the samples kept at ambient temperature for more than 2 hours, there is no sample stability data for Gastrin & Pepsinogen I/II assays, the testing results for these samples cannot be considered as reliable. Therefore, the section $\sqrt{7.1.3.4}$ has been added to exclude the Gastrin/Pepsinogen sample which was exposed to ambient temperature from the analysis.
- For the exclusion criteria 19 in protocol, upper GI endoscopic therapy may have the impact on efficacy/safety evaluation. Additionally, for exclusion criteria 20, Zollinger-Ellision syndrome is a disease which secrets gastrin, results in overproduction of gastric acid, and it will also impact the efficacy/safety evaluation. Therefore, to exclude the Subjects who met exclusion criteria #19 or #20 from PPS, the description has changed.
- For NDA submission of duodenal ulcer in Taiwan and Korea, subgroup analysis in each country was required. Since it is possible that same analysis would also be required for

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the study, to conduct subgroup analysis for Taiwan 7.11.9 has been added.	and Korea the section 7.11.8 and
	10 Torms
	Applicato
	ect to the
and a second	Suple
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APPENDIX 9.0



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Criteria for Markedly Abnormal Values 9.2

Hematology, Serum Chemistry, Urinalysis, Vital Signs, and 12-lead ECG (except 9.2.1 Upper MAV Criteria of QTcF Interval)

For each parameter, all evaluable data (ie, non-missing data) obtained up to Follow-up Day 14 will be classified as a MAV or not. The criteria in the table below will be used.

For each parameter and subject, classifications will be made according to the conditions i) to iii) provided below. The lower and the upper criteria will be considered separately.

- A subject with at least one evaluable data after baseline that meets the MAV criteria will Ö) be classified as a subject with MAV.
- Property ii) A subject who does not meet condition i) and has at least one evaluable data after baseline that doesn't meet the MAV criteria will be considered as a subject without MAV.
 - iii) A subject who does not meet conditions i) or ii) will be excluded from the analysis of MAV for that parameter.

Markedly Abnormal Values Criteria for Hematology Table 9.a

Table 9.a Narkedly Abnormal values <u>C</u>	<u>riteria for Hematology</u>	2
Daramatar	MAV Criteria	
Parameter	Lower Criteria	Upper Criteria
Red Blood Cells ($\times 10^{12}/L$)	<0.8×LLN	>1.2×ULN
White Blood Cells ($\times 10^9$ /L)	<0.5×LLN	>1.5×ULN
Hemoglobin (g/L)	<0.8×LLN	>1.2×ULN
Hematocrit (%)	<0.8×LLN	>1.2×ULN
Platelets ($\times 10^9$ /L)	<75	>600
Neutrophils (%)	<0.5×LLN	>13×ULN
Eosinophils (%)	-	>2×ULN
Basophils (%)	-	>3×ULN
Monocytes (%)	-	>2×ULN
Lymphocytes (%)	<0.5×LLN	>1.5×ULN

LLN=Lower limit of normal, ULN=Upper limit of normal.

Markedly Abnormal Values Criteria for Serum Chemistry Table 9.b

Deremator	MAV Criteria		
Parameter	Lower Criteria	Upper Criteria	
Alanine Aminotransferase (ALT) (U/L)	-	>3×ULN	
Albumin (g/L)	<25	-	
Alkaline Phosphatase (ALP) (U/L)	<u></u>	>3×ULN	
Aspartate Aminotransferase (AST) (U/L)	-	>3×ULN	
Total Bilirubin (µmol/L)	-	>34.2	
Direct Bilirubin (µmol/L)	-	>2×ULN	
Total Cholesterol (mmol/L)	-	>7.72	
Triglycerides (mmol/L)	-	>2.5×ULN	
Calcium (mmol/L)	<1.75	>2.88	
Chloride (mmol/L)	<75	>126	
Creatinine (µmol/L)	-	>177	
Creatine Kinase (U/L)		>5×ULN	
Blood Urea Nitrogen (BUN) (mmol/L)	-	>10.7	
γ-Glutamyl Transferase (γ-GTP)	-	>3×ULN	
Glucose (mmol/L)	<2.8	>19.4	
Inorganic Phosphorus (mmol/L)	< 0.52	>2.00	
Magnesium (mmol/L)	<0.5	>1.2	
Potassium (mmol/L)	<3.0	>6.0	
Total Protein (g/L)	<0.8×LLN	>1.2×ULN	
Sodium (mmol/L)	<130	>150	
Uric Acid (mmol/L)	-	>0.773	
Vitamin B_{12} (pmol/L)	<92	-	
Perty .			

Markedly Abnormal Values Criteria for Vital Signs Table 9.c

Table 7. What Kenty Abhot mar values <u>Cr</u>	iteria ior vitar signs		~0
Doromotor	MAV (Criteria	S
1 diameter	Lower Criteria	Upper Criteria	
Body Temperature (C)	<35.6	>37.7	
Systolic Blood Pressure (mmHg)	<85	>180	
Diastolic Blood Pressure (mmHg)	<50	>110	
Pulse (bpm)	<50	>120	
		10	
		CON CON	
Table 9.dMarkedly Abnormal Values	<u>iteria for 12-lead ECG</u>	Ollo	

Table 9.d Markedly Abnormal Values Criteria for 12-lead ECG

Darrowsstar	MAV Criteria		
Parameter	Lower Criteria	Upper Criteria	
Heart Rate (bpm)	<50	>120	
QT Interval (msec)	<=50	>=460	
QTcF Interval (msec)	<=50	See Section 9.2.2	

12-lead ECG (Upper MAV Criteria of QTcF Interval) 9.2.2

All evaluable data (ie, non-missing data) obtained up to Follow-up Day 14 will be classified as a MAV or not. The criteria in the table below will be used. Note that the observed value and the change from baseline used for classification should be measurements taken on the same day.

For each subject, classifications will be made according to the conditions i) to iii) provided below.

- i) A subject with at least one evaluable data after baseline that meets the MAV criteria will be classified as a subject with MAV.
- ii) A subject who does not meet condition i) and has at least one evaluable data after baseline that meets any of the following will be considered as a subject without MAV.
 - Observed value is less than 450 msec and not missing.
 - Change from baseline is less than 30 msec and not missing, and observed value is less • than 500 msec and not missing.
- iii) A subject who does not meet conditions i) or ii) will be excluded from the analysis of MAV.

Table 9.e Markedly Abnormal Values Criteria for 12-lead ECG (Upper MAV Criteria of QTcF Interval) X

~	Darameter	MAV Criteria		MAV Criteria	
Farameter		Lower Criteria	Upper Criteria		
RIC	QTcF Interval (msec)	-	If either of the following conditions is met: • observed value >=500 • change from baseline >= 30 and observed value >=450		

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9.3 Criteria for Elevated Liver Enzyme

All evaluable data (ie, non-missing data) obtained up to Follow-up Day 14 will be used to determine whether each criteria for elevated liver enzyme in the table below is met or not. If there is more than one parameter that need to be considered for a criteria, parameter measurements taken on the same day will be used. For each criteria and subject, classifications will be made according to the conditions i) to iii) provided below.

- i) A subject who met criteria (a) at least once after baseline will be considered to have met the criteria for elevated liver enzyme.
- ii) If condition i) is not met but if criteria (b) is met at least once after baseline, then the subject will be considered to have not met the criteria for elevated liver enzyme.
- iii) If neither i) nor ii) is met, then the subject will be excluded from the analysis for the criteria for elevated liver enzyme.

Label	Criteria for Eleva	ted Liver Enzyme	
Laber	(a) Elevated	(b) Not Elevated	
$\Delta I T > 3 \times I \parallel N$	ALT is greater than 3 times the LIL	ALT is non-missing and less than or equal	
ALT > SAOLIN	ALT is greater than 5 times the OLIV	to 3 times the ULN	
$ALT > 5 \times ULN$	ALT is greater than 5 times the ULN	ALT is non-missing and less than or equal	
		to 5 times the ULN	
$ALT > 8 \times ULN$	ALT is greater than 8 times the ULN	ALT is non-missing and less than or equal	
		to 8 times the ULN	
	ALT is greater than 3 times the ULN and	Either ALT is non-missing and less than	
ALT > 3xULN	the total bilirubin is greater than twice the	or equal to 3 times the ULN, or the total	
with $1 \text{ bill} > 2 \text{ xULN}$	ULN	bilirubin is non-missing and less than or	
		equal to twice the ULN	
AST > 3xULN	AST is greater than 3 times the ULN	AST is non-missing and less than or equal	
		AST is non-missing and loss than or equal	
AST > 5xULN	AST is greater than 5 times the ULN	to 5 times the LU N	
		AST is non-missing and less than or equal	
AST > 8xULN	AST is greater than 8 times the ULN	to 8 times the LU N	
		Fither AST is non-missing and less than	
AST > 3xUUN	AST is greater than 3 times the ULN and	or equal to 3 times the ULN or the total	
with Thili $\ge 2 \times ULN$	the total bilirubin is greater than twice the	bilirubin is non-missing and less than or	
	ULN	equal to twice the ULN	
	Either ALT or AST is greater than 3 times	Both ALT and AST are non-missing and	
AL or $AS1 > 3xULN$	the ULN	less than or equal to 3 times the ULN	
X	Either ALT or AST is greater than 5 times	Both ALT and AST are non-missing and	
ALT or AST > 5xULN	the ULN	less than or equal to 5 times the ULN	
ALT or ACT > PULL N	Either ALT or AST is greater than 8 times	Both ALT and AST are non-missing and	
$ALI \text{ OI } ASI > \delta X ULN$	the ULN	less than or equal to 8 times the ULN	
	Fither AI T or AST is greater than 3 times	If any of the following conditions is met:	
ALT or AST $>$ 3xULN	the III N and the total bilirubin is greater	- Both ALT and AST are non-missing and	
with Tbili $> 2xULN$	1 Tbili > 2xULN une ULIN and the total offit doin is greater than twice the ULIN	less than or equal to 3 times the ULN.	
		- Total bilirubin is non-missing and less	

 Table 9.f
 Markedly Abnormal Values Criteria for Elevated Liver Enzyme

Labol	Criteria for Elevated Liver Enzyme	
Lauei	(a) Elevated	(b) Not Elevated
		than or equal to twice the ULN.
ALT and AST > 3xULN	Both ALT and AST are greater than 3 times the ULN	Either ALT is non-missing and less than or equal to 3 times the ULN, or AST is non-missing and less than or equal to 3 times the ULN
ALT and AST > 5xULN	Both ALT and AST are greater than 5 times the ULN	Either ALT is non-missing and less than or equal to 5 times the ULN, or AST is non-missing and less than or equal to 5 times the ULN
ALT and AST > 8xULN	Both ALT and AST are greater than 8 times the ULN	Either ALT is non-missing and less than or equal to 8 times the ULN, or AST is non-missing and less than or equal to 8 times the ULN
ALT and AST > 3xULN with Tbili > 2xULN	Both ALT and AST are greater than 3 times the ULN and the total bilirubin is greater than twice the ULN	If any of the following conditions is met: - ALT is non-missing and less than or equal to 3 times the ULN - AST is non-missing and less than or equal to 3 times the ULN - Total bilirubin is non-missing and less than or equal to twice the ULN
ALP > 3xULN	ALP is greater than 3 times the ULN	ALP is non-missing and less than or equal to 3 times the LU N
ALP > 3xULN with ALT > 3xULN	Both ALP and ALT are greater than 3 time the ULN	Either ALP is non-missing and less than or equal to 3 times the ULN, or ALT is non-missing and less than or equal to 3 times the ULN
ALP > 3xULN with AST > 3xULN	Both ALP and AST are greater than 3 times the ULN	Either ALP is non-missing and less than or equal to 3 times the ULN, or AST is non-missing and less than or equal to 3 times the ULN
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