

Statistical Analysis Plan Version 3 J2A-MC-GZGT

A Phase 3, Randomized, Double-Blind Study to Investigate the Efficacy and Safety of
Once Daily Oral LY3502970 Compared With Placebo in Adult Participants With Type 2
Diabetes and Inadequate Glycemic Control With Diet and Exercise Alone

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Title Page

Protocol Title: A Phase 3, Randomized, Double-Blind Study to Investigate the Efficacy and Safety of Once Daily Oral LY3502970 Compared with Placebo in Adult Participants with Type 2 Diabetes and Inadequate Glycemic Control with Diet and Exercise Alone (ACHIEVE-1)

Protocol Number: J2A-MC-GZGT

Compound Number: Orforglipron (LY3502970)

Short Title: Effect of Orforglipron Compared with Placebo in Participants with Type 2 Diabetes Inadequately Controlled with Diet and Exercise Alone (ACHIEVE-1)

Acronym: ACHIEVE-1

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Version History

The original Statistical Analysis Plan (SAP) for Study J2A-MC-GZGT (GZGT) was approved on 25-Jan-2024 and was written based on Protocol GZGT amendment (b), dated 06-Sep-2023.

SAP GZGT version 2 SAP was updated based on regulatory feedback and was approved on 06-Nov-2024. SAP GZGT version 3 SAP was updated to add the final graphical approach and further clarification. The major updates in this amendment are listed in the table below.

SAP Version History Summary

SAP Version	Approval Date	Change Section	Change	Rationale
1	25 January 2024		Not Applicable	Original version
2	06 November 2024	1.1	Added clarification to the definition of estimands population	Clarification
		2.1	Added details of graphical approach	Graphical approach was added to control Type 1 error
		4.1	Added clarification for different types of time-to-event analyses	Clarification
		4.1.2.1.3	Further clarified the definition of Scenario 5. Removed the direct imputation method Added more details regarding the imputation methods. Added footnote in Table GZGT.4.4 for protocol deviation, inadvertent enrollment, and emergency unblinding	The direct imputation method is not fully aligned with the primary imputation strategy. Used a different method to handle Scenario 5B, based on simulation results from Liu et al. (2024) Clarified the distinction of imputing discontinuation due to protocol deviation, inadvertent enrolment
		4.1.2.3	Updated MMRM model for safety analysis	To be consistent with the PSAP

			Clarified the definition of start time for different analyses	
		4.2	Updated Table GZGT.4.5	To be consistent with other studies in the program
		4.3.1	Added note that for missing or canceled HbA1c measurement at endpoint visit, retests may be used	Minimize the missing HbA1c in the primary endpoint analysis
		4.4.2	Added definition of valid SMBG profile	Clarification
		4.6.3	Updated Table GZGT.4.9 by adding details on MACE, an additional table for pancreatic events, and an additional table for hepatic events	To be consistent with the PSAP
		4.7.1	Updated the testing of subgroup effects. Added details for the Bayesian shrinkage method	Clarification
		7	Reference section was updated	To add extra references
3	See date on page 1	2.1	Added the final graphical approach	Adding details to the graphical testing scheme
		4.1	Updated wording on which analyses will adjust for strata	Clarification
		4.1.2.1.2	Updated logistic regression section to include interaction terms in model	Consistency with other analyses
		4.1.2.1.3	Updated imputation procedure to placebo washout if not enough retrieved dropouts. Further clarified the imputation strategy for all scenarios	Clarification

			Updated study discontinuation reason categories for imputation purpose	
		4.1.2.1.4	Removed modified multiple imputation section	Clarification
		4.6.2	Clarified sorting order of maximum severity TEAE table Added table to summarize dose reductions due to AE	Clarification
		4.6.3	Removed redundant TE NVDC by Max Severity table Added shift tables for hepatic parameters	Clarification
		4.7.1	Further clarified subgroup analysis section.	Clarification
		6.4	Updated time period of analysis for initiation of additional antihyperglycemic medication to “during treatment period”	Clarification

Abbreviation: ANCOVA = analysis of covariance; HbA1c = hemoglobin A1c, MMRM = mixed model repeated measure, PSAP= program safety analysis plan; SAP = statistical analysis plan; SMBG= self-monitoring of blood glucose.

1. Introduction

Changes to the protocol-planned analyses are described in Section 4.9.

1.1. Objectives, Endpoints, and Estimands

Objectives	Endpoints
Primary	
To demonstrate that orforglipron \blacksquare mg, \blacksquare mg, and/or \blacksquare mg are superior to placebo in glycemic control	<ul style="list-style-type: none"> Change from baseline in HbA1c at Week 40
Key Secondary	
To demonstrate that orforglipron \blacksquare mg, \blacksquare mg, and/or \blacksquare mg are superior to placebo in additional measures of glycemic control	<ul style="list-style-type: none"> Proportion of participants with an HbA1c target value of <7.0% (53 mmol/mol) at Week 40 Proportion of participants with an HbA1c target value of \leq6.5% (48 mmol/mol) at Week 40 Change from baseline in fasting serum glucose (central laboratory) at Week 40
To demonstrate that orforglipron \blacksquare mg, \blacksquare mg, and/or \blacksquare mg are superior to placebo in weight management	<ul style="list-style-type: none"> Percentage change from baseline in body weight at Week 40 Change from baseline in body weight at Week 40
To demonstrate that orforglipron \blacksquare mg and/or \blacksquare mg are superior to placebo on lipid parameters	<ul style="list-style-type: none"> Percentage change from baseline in non-HDL cholesterol at Week 40 Percentage change from baseline in triglycerides at Week 40
Other Secondary	
To compare orforglipron \blacksquare mg, \blacksquare mg, and \blacksquare mg with placebo on additional measures of glycemic control	<ul style="list-style-type: none"> Proportion of participants with an HbA1c target value of <5.7% (39 mmol/mol) at Week 40 Change from baseline in daily average 7-point SMBG at Week 40
To compare orforglipron \blacksquare mg, \blacksquare mg, and \blacksquare mg QD with placebo on additional measurements of weight management	<ul style="list-style-type: none"> Proportion of participants who achieved weight loss of \geq5%, \geq10%, and \geq15% from baseline at Week 40 Change from baseline in waist circumference at Week 40 Change from baseline in BMI at Week 40
To compare orforglipron \blacksquare mg, \blacksquare mg, and \blacksquare mg QD with placebo on measures of blood pressure	<ul style="list-style-type: none"> Change from baseline at Week 40 in: <ul style="list-style-type: none"> systolic blood pressure diastolic blood pressure
To compare orforglipron \blacksquare mg, \blacksquare mg, and \blacksquare mg QD with placebo on lipid parameters	<ul style="list-style-type: none"> Percentage change from baseline in lipid parameters at Week 40 <ul style="list-style-type: none"> total cholesterol HDL-cholesterol LDL-cholesterol VLDL-cholesterol

Objectives	Endpoints
To compare orforglipron CC mg with placebo on lipid parameters	<ul style="list-style-type: none"> Percentage change from baseline in lipid parameters at Week 40 <ul style="list-style-type: none"> non-HDL cholesterol triglycerides
To compare orforglipron CC mg, CC mg, and CC mg QD with placebo on patient-reported outcomes	<ul style="list-style-type: none"> Change from baseline at Week 40 in: <ul style="list-style-type: none"> SF-36v2 Acute Form domain scores EQ-5D-5L health state utilities and VAS APPADL scores IW-SP score DTSQs scores DTSQc scores at Week 40
To describe the safety of orforglipron in participants with T2D	<p>Summary of safety data, including number and incidence of:</p> <ul style="list-style-type: none"> SAEs TEAEs discontinuations from study intervention or study due to AEs

Abbreviations: AE = adverse event; APPADL = Ability to Perform Physical Activities of Daily Living; BMI = body mass index; DTSQc = Diabetes Treatment Satisfaction Questionnaire-Change Version; DTSQs = Diabetes Treatment Satisfaction Questionnaire-Status Version;; HbA1c = hemoglobin A1c; HDL = high-density lipoprotein; IW-SP = Impact of Weight on Self-Perception; LDL = low-density lipoprotein; QD = once daily; SAE = serious adverse event; SF-36v2, Acute = Short Form-36 version 2 Health Survey acute form; SMBG = self-monitored blood glucose; T2D = type 2 diabetes; TEAE = treatment-emergent adverse event; **CC**; VLDL = very low-density lipoprotein.

Primary estimands

There will be 2 estimands for the primary objective planned in Study GZGT. Details on plan for use of each estimand can be found in Section 2.1. The 2 estimands, treatment regimen estimand and efficacy estimand, address intercurrent events (ICEs) using the treatment policy strategy and the hypothetical strategy (FDA 2021), respectively.

Treatment regimen estimand

The clinical question of interest is: *What is the treatment difference in the change in HbA1c (hemoglobin A1c) from baseline at Week 40 between orforglipron **CC** mg, **CC** mg, and/or **CC** mg, compared with placebo in individuals who meet eligibility criteria regardless of adherence to study intervention or initiation of additional antihyperglycemic medications?*

Treatment policy strategy

The occurrence of the ICE is considered irrelevant in defining the treatment effect of interest; the values for the variable of interest are used regardless of whether the ICE occurs.

Rationale for the treatment regimen estimand

The estimand aims to evaluate the efficacy of orforglipron that reflects the real-life behavior of the target population.

Treatment regimen estimand attributes

- *Population*: individuals who meet the eligibility criteria. This represents the target population identified through study inclusion/exclusion criteria (ICH E9). For the analysis, intent-to-treatment principle will be applied. Details are defined in Section 3.
- *Endpoint*: change from baseline in HbA1c at Week 40.
- *Treatment condition*: the randomized treatment with allowance for potential dose interruptions and dose modification regardless of adherence to study intervention or initiation of additional antihyperglycemic medications.
- *Intercurrent events*: no ICEs are defined because treatment adherence and the initiation of additional antihyperglycemic medications are a part of the treatment condition.
- *Population-level summary and treatment effect of interest*: difference in mean changes in HbA1c from baseline at Week 40 between orforglipron ■ mg, ■ mg, and/or ■ mg compared with placebo.

Efficacy estimand

The clinical question of interest is: *What is the treatment difference in the change in HbA1c from baseline at Week 40 between orforglipron ■ mg, ■ mg, and/or ■ mg compared with placebo in individuals who meet the eligibility criteria if they would remain on their randomly assigned study intervention for ■ weeks and would not initiate additional antihyperglycemic medications?*

Hypothetical strategy

A scenario is envisaged in which the ICE would not occur. The value of the variable to reflect the clinical question of interest is the value that the variable would have taken in the hypothetical scenario defined.

Rationale for the efficacy estimand

This estimand aims to evaluate the efficacy of orforglipron under the ideal condition that all participants would adhere to the randomly assigned study intervention without being confounded by the initiation of additional antihyperglycemic medications.

Efficacy estimand attributes

- *Population*: individuals who meet the eligibility criteria. This represents the target population identified through study inclusion/exclusion criteria (ICH E9). For the analysis, intent-to-treatment principle will be applied. Details are defined in Section 3.
- *Endpoint*: change from baseline in HbA1c at Week 40.
- *Treatment condition*: the randomized treatment with allowance for potential dose interruptions and dose modification.
- *Intercurrent events*: ICEs include permanent discontinuation of study intervention and initiation of additional antihyperglycemic medications and will be handled by the hypothetical strategy. The potential outcome of interest is the response in the efficacy measurement if participants would remain on their randomly assigned study intervention for ■ weeks and would not initiate additional antihyperglycemic medications. Dose

modification and interruption will not be considered an ICE because they are part of the treatment condition.

- *Population-level summary and treatment effect of interest:* difference in mean changes in HbA1c from baseline at Week 40 between orforglipron cc mg, cc mg, and/or cc mg compared with placebo.

The same strategies will also be applied to the estimands for the key secondary endpoints as specified in Section 4.

1.2. Study Design

ACHIEVE-1 (Study GZGT) is a Phase 3, multicenter, randomized, parallel-arm, double-blind, placebo-controlled study to compare the effect of orforglipron (LY3502970) with that of placebo on the glycemic control in individuals with type 2 diabetes (T2D) who have inadequate glycemic control with diet and exercise alone, are naïve to insulin use and have not been treated with any antihyperglycemic medication during the 90 days preceding Visit 1.

This study will consist of 3 periods (Figure GZGT.1.1):

- a screening and lead-in period: up to 4 weeks



Figure GZGT.1.1. Illustration of study design for Protocol GZGT.

The treatment duration will be up to cc weeks for participants from non-Japan sites. For participants from Japan sites, treatment duration will be up to cc weeks (Section 6.6). The

treatment will be double-blind, which means neither the participants nor the researchers will know which study intervention participants are receiving until the study is over.

The visit frequency is listed in [Table GZGT.1.1](#).

Table GZGT.1.1. Visit Frequency

	Visit Number	From Visit 3 (Randomization)
Screening and lead-in period	601 ^a	CCI
	1	
	2	
Treatment period	3	
	4	
	5	
	6	
	7	
	8	
	9	
	9T ^b	
	10	
	10T ^b	
	11	
	12 ^c	
	ED	
Safety follow-up period	801	12 weeks post end of treatment period

Abbreviations: ED = early discontinuation; T = telephone visit.

^a Optional prescreening visit.

^b Only for participants from China sites (Section 6.7).

^c Only for participants from Japan sites (Section 6.6).

Participants will be randomly assigned in a **CCI** ratio to a once daily dose of orforglipron **CC** mg, **CCI** mg, **CCI** mg, or placebo.

Participants will be stratified based on country, HbA1c ($\leq 8.0\%$, $> 8.0\%$) at baseline, and prior use of any antihyperglycemic medication (Yes or No). Stratification factors will be called strata.

2. Statistical Hypotheses

The null hypotheses corresponding to the primary objective are

- $H_{1,36,0}$, $H_{1,12,0}$, and $H_{1,3,0}$: No difference in orforglipron [redacted] mg, [redacted] mg, and [redacted] mg compared to placebo with respect to the mean change from baseline in HbA1c at Week 40, respectively.

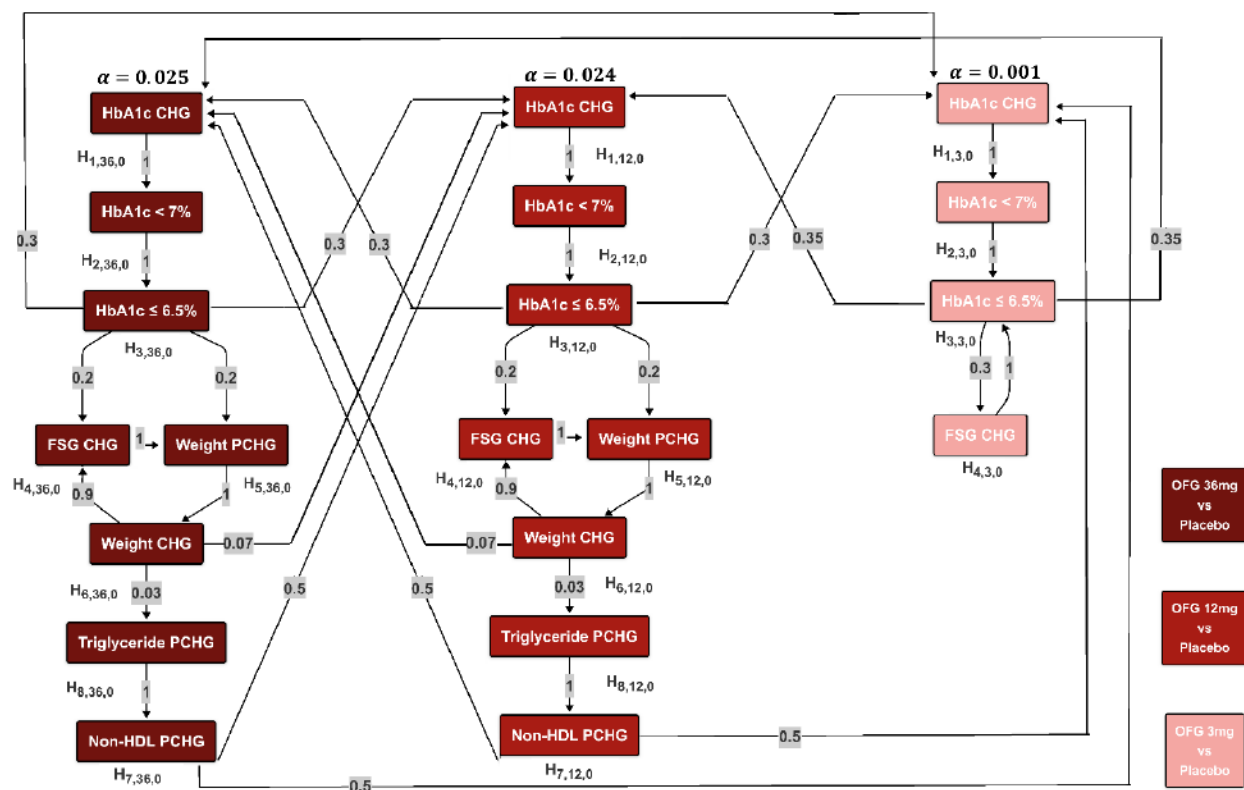
The null hypotheses corresponding to the key secondary objectives are

- $H_{2,36,0}$, $H_{2,12,0}$, and $H_{2,3,0}$: No difference in orforglipron [redacted] mg, [redacted] mg, and [redacted] mg compared to placebo with respect to the proportion of participants who achieved HbA1c <7.0% at Week 40, respectively.
- $H_{3,36,0}$, $H_{3,12,0}$, and $H_{3,3,0}$: No difference in orforglipron [redacted] mg, [redacted] mg, and [redacted] mg compared to placebo with respect to the proportion of participants who achieved HbA1c ≤6.5% at Week 40, respectively.
- $H_{4,36,0}$, $H_{4,12,0}$, and $H_{4,3,0}$: No difference in orforglipron [redacted] mg, [redacted] mg, and [redacted] mg compared to placebo with respect to the mean change from baseline in fasting serum glucose at Week 40, respectively.
- $H_{5,36,0}$, $H_{5,12,0}$, and $H_{5,3,0}$: No difference in orforglipron [redacted] mg, [redacted] mg, and [redacted] mg compared to placebo with respect to the mean percentage change from baseline in body weight at Week 40, respectively.
- $H_{6,36,0}$, $H_{6,12,0}$, and $H_{6,3,0}$: No difference in orforglipron [redacted] mg, [redacted] mg, and [redacted] mg compared to placebo with respect to the mean change from baseline in body weight at Week 40, respectively.
- $H_{7,36,0}$ and $H_{7,12,0}$: No difference in orforglipron [redacted] mg and [redacted] mg compared to placebo with respect to the mean percentage change from baseline in non-high-density lipoprotein (HDL) cholesterol at Week 40, respectively.
- $H_{8,36,0}$ and $H_{8,12,0}$: No difference in orforglipron [redacted] mg and [redacted] mg compared to placebo with respect to the mean percentage change from baseline in triglycerides at Week 40, respectively.

2.1. Multiplicity Adjustment

Multiplicity adjusted analyses will be performed on the primary and key secondary objectives listed in [Figure GZGT 2.1](#) in order to control the overall family-wise Type I error rate at a 2-sided alpha level of 0.05. The graphical multiple testing procedure described in Bretz et al. (2009, 2011) will be used. This approach is a closed testing procedure; hence, it strongly controls the family-wise Type I error rate across all hypotheses (Alosh et al. 2014).

[Figure GZGT 2.1](#) illustrates the current graphical testing procedure. Since more than one types of estimands are used to assess primary and key secondary objectives and are intended for separate and independent purposes, no multiplicity adjustment will be made for conducting separate analyses on the same objectives. Treatment regimen estimand will be the primary estimand, with the efficacy estimand considered supportive. There will be no adjustment for multiple comparisons for any other analyses beyond the primary and key secondary objectives.



Abbreviations: CHG = change from baseline; FSG = fasting serum glucose; HbA1c = hemoglobin A1c; Non-HDL = non-HDL cholesterol; PCHG = percent change from baseline..

Figure GZGT 2.1. Type 1 error control strategy for primary and key secondary efficacy objectives.

3. Analysis Sets

The participant analysis sets are defined as

Participant Analysis Set	Description
Entered participants	All participants who sign informed consent ^a .
Randomized participants	All participants who are randomly assigned a study intervention.
Safety participants	All participants who are randomly assigned a study intervention and who take at least 1 dose of study intervention.

^a Refers to the informed consent for the study.

The data points sets are defined as

Data Points Sets	Description
Treatment regimen estimand data points set	All data points obtained during the treatment period ^a defined as at or after baseline and up to the last visit within the treatment period, regardless of study intervention discontinuation or initiation of additional antihyperglycemic medications ^b .
Safety data points set	All data points obtained during the treatment period ^a and the follow-up period ^a defined as at or after baseline and up to the date of study withdrawal or study completion, including the follow-up period regardless of study intervention discontinuation or initiation of additional antihyperglycemic medications ^b .
Efficacy estimand data points set	All data points obtained during the treatment period ^a defined as at or after baseline and up to the earliest date of discontinuation of study intervention or initiation of additional antihyperglycemic medications ^b .

^a Data in participants from Japan sites who continued beyond the Week 40 visit will be excluded. Subgroup analyses for participants from Japan sites are separately defined in Appendix 6 (Section 6.6).

^b “Additional antihyperglycemic medications” refers to any antihyperglycemic therapy that is used for more than 2 weeks (14 days).

4. Statistical Analyses

4.1. General Considerations

Statistical analysis will be the responsibility of Eli Lilly and Company (Lilly) or its designee. Some analyses and summaries described in this analysis plan may not be conducted if warranted by data (that is, too few events to justify conducting an analysis). Additional analyses of the data may be conducted as deemed appropriate without further changes made to the protocol or SAP, even after the final database lock.

Unless otherwise noted, all tests of treatment effects will be conducted at the 2-sided alpha level of 0.05, and the confidence intervals (CIs) will be calculated at 95%, 2-sided.

Participants will be analyzed according to the treatment group to which they were randomly assigned.

Change from baseline will be calculated as the value of interest at the visit minus the baseline value. In general, percent change from baseline will be calculated as the value of change from baseline divided by the baseline value in 100% scale. Some specific predefined parameters may be log-transformed before statistical analysis, if deemed necessary. If baseline value is missing for a particular variable, then the change from baseline and percent change from baseline will not be calculated when deriving the summary statistics.

For continuous measures, summary statistics will include sample size, mean, standard deviation (SD), median, minimum, and maximum for the actual value, the change from baseline and if applicable, the percent change from baseline measurements as well. Least squares (LS) means and standard errors derived from the analysis models will also be displayed for the actual value and the change or percent change from baseline measurements. For analysis of log-transformed parameters, model estimated means and standard errors on the original scale will be derived through back-transformation using the delta method from the LS means and standard errors on the natural log-scale. Treatment comparisons will be displayed showing the LS means, the 95% CIs, and the p-values for the treatment differences. All baseline measures will be analyzed using an analysis of variance (ANOVA) model that has treatment group as the fixed effect.

For categorical measures, summary statistics will include sample size, frequency, and percentages. For safety categorical measures, risk difference and its 95% CI will be provided wherever applicable. If applicable, Fisher's exact test or Pearson's chi-squared test will be used for treatment comparisons.

For time-to-event measures, participants without an event will be censored, and time-to-event will be the number of days between the start date and the date of the participant's end of follow-up (depending on the estimand definition) + 1 day. For participants experiencing the event, "time-to-first-event" will be the time (in days) from start date to the first occurrence of the event + 1 day. For safety and study intervention discontinuation analyses, the date of first dose of study intervention will be used as the start date. For other analyses (for example, study discontinuation), the randomization date will be used as the start date (in rare cases where randomization occurred prior to Visit 3 date, Visit 3 date will be used instead).

Data may exist at postbaseline visits where a variable is not scheduled to be collected. Data from unscheduled visits and from early discontinuation (ED) visits that do not correspond to a

scheduled visit will be excluded from mixed-model for repeated measures (MMRM) analysis, analysis of covariance (ANCOVA), or logistic regression analysis, unless otherwise specified.

For stratification factors, countries with fewer than 10 randomized participants will be pooled into 1 category (pooled country). A single strata variable will be created for other stratification factors (HbA1c at baseline [$\leq 8.0\%$, $> 8.0\%$], and prior use of any antihyperglycemic medication [Yes or No]). Unless otherwise specified, country and this strata variable will be used in the analysis models including MMRM, ANCOVA, and logistic regression analysis.

No data collected at the optional prescreening visit (Visit 601), except adverse events and demographic information, will be included in statistical analyses, given that the purpose of prescreening visit is not for data collection (Protocol GZGT, Section 1.3) but for operational efficiency.

Details about the analyses regarding demographic and baseline characteristics, historical illnesses and preexisting conditions, treatment compliance, concomitant medications, and important protocol deviations can be found in Appendices 1 through 5 (Section 6.1 through Section 6.5), respectively.

A subset of the planned efficacy, health outcomes, and safety analyses at Week 40 will be reproduced based on participants from Japan sites to confirm consistency of the Japanese population in support of the regulatory submission in Japan. Efficacy and safety analyses for participants from Japan sites will also be analyzed at Week 52. See Appendix 6 (Section 6.6) for details.

A subset of the planned efficacy, health outcomes, and safety analyses will also be reproduced based on participants enrolled in China and East Asian countries. See Appendix 7 (Section 6.7) for details.

4.1.1. Baseline Definition

Unless otherwise specified, the baseline for efficacy assessments is defined as the last available non-missing measurement prior to the first dose of study intervention, which in most cases will be the measurement recorded at Week 0 (Visit 3). If there are no doses of study intervention, the baseline will be defined as the last available non-missing measure on or prior to randomization. In cases where the measurement is taken on the same day (where the time is not collected or not reliable) as the first dose, this measurement will be used as the baseline value for data analysis. For patient-reported outcome measures, data obtained at Visit 3, regardless of the timing relative to first dose, will serve as the baseline. For the safety related parameters, the definition of the baseline and postbaseline are specified in [Table GZGT.4.1](#).

Table GZGT.4.1. Baseline and Postbaseline Definitions for Safety Analyses

Analysis Type	Participant Analysis Set/Data Points Set	Baseline Period	Postbaseline Period^a
Treatment-emergent adverse events	Safety participants/safety data points set	Start of screening ^b and ends prior to the first dose of study intervention (typically at Week 0).	Starts after the first dose of study intervention and ends at the end of the safety follow-up period or the date of study withdrawal ^c .
Treatment-emergent abnormal laboratory values, vital signs, and ECGs	Safety participants/safety data points set	Start of screening ^b and ends prior to the first dose of study intervention (typically at Week 0). All scheduled and unscheduled measurements will be included.	Starts after the first dose and ends at the end of the safety follow-up period or the date of study withdrawal. All scheduled and unscheduled measurements will be included.
Change from last baseline to each postbaseline week and to last postbaseline for laboratory values, vital signs, and ECGs	Safety participants/safety data points set	When the term “last baseline” is used, starts from informed consent date and ends prior to the first dose (typically at Week 0).	Starts after the first dose and ends at the end of the safety follow-up period or the date of study withdrawal. Only scheduled and early termination visits will be included.

Abbreviations: ECG = electrocardiogram; PMDA = Pharmaceuticals and Medical Devices Agency.

- ^a The safety analysis for treatment effects will exclude any data that are collected after Visit 11 (Week 40) for participants from Japan sites; additional analysis (Appendix 6, Section 6.6) will be performed to these additional safety data in Japanese participants for PMDA submission.
- ^b Adverse event data collected between Visit 601 and Visit 1 will be included as part of screening as well.
- ^c For events occurring on the day of first dose, the case report form collected information will be used to determine whether the event was pre- versus post-treatment if available. If the relevant information is not available, then the events will be counted as postbaseline.

4.1.2. Analysis Methods

The primary and key secondary objectives, as well as blood pressure, lipid parameters and patient-reported outcomes, will be evaluated based on both the treatment regimen estimand and efficacy estimand. All other secondary endpoints (except the summary of safety data) will be guided by the efficacy estimand. The assessments guided by the treatment regimen estimand will be conducted using the randomized participants analysis set and the treatment regimen estimand data points set. The assessments guided by the efficacy estimand will be conducted using the randomized participants analysis set and the efficacy estimand data points set (Section 3).

Unless otherwise specified, safety and tolerability assessments will be guided by an estimand comparing safety of orforglipron treatment groups with placebo for the entire study period (the treatment period plus safety follow-up) irrespective of adherence to study intervention or initiation of additional anti-hyperglycemic medications for all study population. It is noted that for some safety endpoints, such as treatment discontinuation due to adverse events and

laboratory values that are planned to be measured only during the treatment period, the analyses will be based on data observed during the treatment period only.

The estimands, participant analysis sets, analysis data points sets, and analysis models for efficacy and safety endpoints are summarized in [Table GZGT.4.2](#).

Table GZGT.4.2. Estimands and Analysis Models for All Endpoints

	Treatment Regimen Estimand	Efficacy Estimand	Estimand Guiding Safety
Participants	Randomized	Randomized	Safety
Data points set	Treatment regimen	Efficacy estimand	Safety
Primary and key secondary endpoints	X	X	
Selected other secondary endpoints			
<ul style="list-style-type: none"> - blood pressure - lipid parameters - patient-reported outcomes 			
Other secondary endpoints		X	
<ul style="list-style-type: none"> - additional measures of glycemic control - additional measures of weight management 			
Subgroup analyses	X		
Safety endpoints (not including lipid parameters)			X
Analysis model	<ul style="list-style-type: none"> • Continuous: ANCOVA with missing endpoints imputed^a • Binary: logistic regression with missing endpoints imputed^a 	<ul style="list-style-type: none"> • Continuous: MMRM (if there are multiple longitudinal postbaseline measurements) or ANCOVA (if there is only 1 postbaseline measurement) with missing endpoints imputed^b • Binary: logistic regression with missing endpoints imputed^b 	<ul style="list-style-type: none"> • Continuous: MMRM (if there are multiple longitudinal postbaseline measurements) or ANCOVA with no imputation to be conducted (if there is only 1 postbaseline measurement) • Binary: risk difference (95% CI) will be presented; p-value from Fisher's exact or Pearson's chi-squared test if applicable; no imputation will be conducted • Time-to-event: Cox Regression and Kaplan-Meier

Abbreviations: ANCOVA = analysis of covariance; CI = confidence interval; MMRM = mixed model for repeated measures.

^a The treatment regimen estimand imputation method is described in Section 4.1.2.1.3.

^b The efficacy estimand imputation method is described in Section 4.1.2.2.3.

4.1.2.1. Analysis Methods for Treatment Regimen Estimand

4.1.2.1.1. Method for Continuous Variables

For the assessment of continuous efficacy measures guided by the treatment regimen estimand, data for participants with missing values at Week 40 will be imputed based on methods described in Section 4.1.2.1.3, and then analyzed using an ANCOVA with a variance estimator that is robust to model misspecification and heteroscedasticity (Ye et al. 2022, FDA 2023). The model will include

- treatment group as a factor variable
- pooled country as a factor variable
- a single strata variable for other stratification factors (HbA1c at baseline [$\leq 8.0\%$, $>8.0\%$], and prior use of any antihyperglycemic medication [Yes or No]), as a factor variable
- baseline value (of the dependent variable)
- interaction between treatment group and strata variable, and
- interaction between treatment group and baseline value.

Statistical comparisons will be performed between each dose of orforglipron and placebo. If the model fails to converge, the interaction terms will be removed. The addition of interaction terms is not intended to estimate the heterogeneity effect but to provide robustness and efficiency for the estimate of treatment comparisons on the unconditional effect. The final inference will be derived using Rubin's rule which combines estimates from multiple imputed datasets. The imputation procedure for creating a single imputed dataset is detailed in Table GZGT.4.3.

Some parameters may be log-transformed before fitting the ANCOVA as specified in Section 4.4, Section 4.5, and Section 4.6, with the associated log-transformed baseline value as a covariate. In these cases, LS means and 95% CIs for each treatment group and treatment difference will be back-transformed and presented as mean percent change from baseline and as relative treatment difference to placebo in percent change.

4.1.2.1.2. Method for Binary Variables

For binary efficacy measures derived from a continuous variable and guided by the treatment regimen estimand, the missing value in the underlying continuous variable will be imputed first based on methods described in Section 4.1.2.1.3 and then the corresponding binary variable will be derived.

For the assessment of binary efficacy measures, a logistic regression model will be used to examine the treatment difference with missing values at Week 40 imputed based on methods described in Section 4.1.2.1.3. The model will include:

- treatment group as a factor variable
- pooled country as a factor variable
- a single strata variable for other stratification factors (HbA1c at baseline [$\leq 8.0\%$, $>8.0\%$], and prior use of any antihyperglycemic medication [Yes or No]), as a factor variable
- baseline value (of the dependent variable)
- interaction between treatment group and strata variable, and

- interaction between treatment group and baseline value.

The unconditional treatment group effect will be assessed by risk difference and relative risk based on the delta-method using the formula provided (Ye et al. 2023). The final inference will be derived using Rubin's rule by combining estimates from multiple imputed datasets, with the imputation procedure for creating a single imputed dataset detailed in [Table GZGT.4.3](#). The estimated treatment group-specific risk, risk difference, relative risk, p-value and 95% CI will be presented.

4.1.2.1.3. Primary Multiple Imputation Strategy

Discontinuation is expected to be uncommon. Participants who discontinue study intervention (that is, discontinue study treatment) will be encouraged to continue in Study GZGT for the treatment period and follow-up period (note, the case report forms [CRFs] use "phase," which is interchangeable with the "period" in SAP). In this section, study discontinuation refers to treatment phase discontinuation captured in CRF.

If there are occurrences of missing data despite all reasonable precautions, missing data will be imputed in a manner consistent with what the values would likely have been, had they been collected.

In general, 5 scenarios are considered regarding the treatment journey of a trial participant relative to the primary time point (the Week 40 visit) shown in [Figure GZGT.4.1](#).

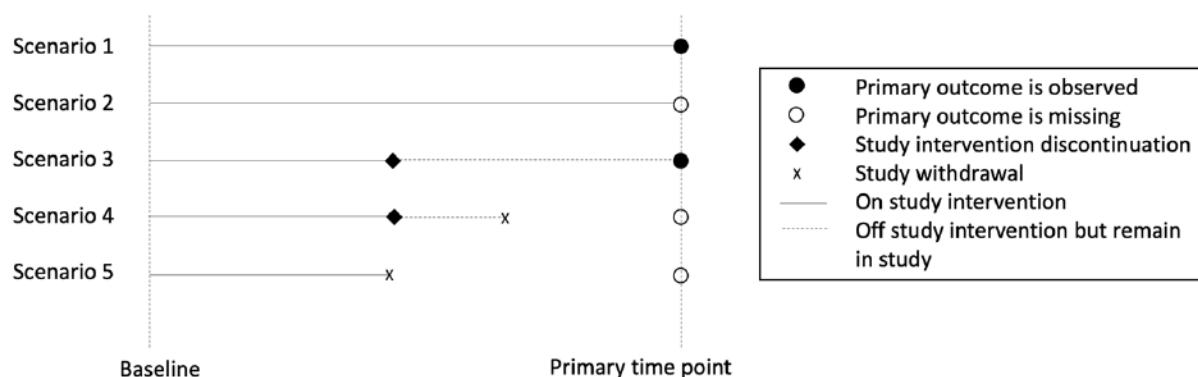


Figure GZGT.4.1. Treatment journey of a trial participant in relation to permanent discontinuation of study intervention or study.

Scenario 5 also covers situations where a participant discontinues from the study intervention and then comes back for a study discontinuation visit for the same discontinuation reason on or prior to a scheduled visit.

In principle, missing data due to permanent discontinuation of the study intervention (Scenario 4 and 5A) will be imputed by treatment group using retrieved dropouts (MI-RD), namely using multiple imputation based on data retrieved from participants who permanently discontinued the study intervention but continued in Study GZGT with non-missing measurements from the same treatment group. For scenarios 4 and 5A, if there are not enough retrieved dropouts to provide a reliable imputation model (that is, the model implemented does not converge), an alternative

multiple imputation method (placebo-washout method) (Wang 2023) will be used. Any missing values at baseline will be imputed in the same model when imputing the endpoint.

Table GZGT.4.3. Imputation Procedure

Scenario (Study Intervention/Study Discontinuation; Missingness at Endpoint)	Methods to Handle Missing Values at Endpoint
1. No study intervention discontinuation; no missing value	N/A
2. No study intervention discontinuation; with missing value	Use observed data, including baseline and all post-baseline visits, across all time points from Scenario 1 and 2 to impute under the MAR assumption by treatment group. There should be very few such cases in a clinical trial. No additional covariates will be included in the imputation model.
3. Study intervention discontinuation; no missing value	N/A
4. Study intervention discontinuation with study discontinuation at a subsequent visit or completion of treatment period; with missing value	Missing values at the endpoint visit will be imputed through MCMC (predictive mean matching method), using baseline and endpoint visit data from Scenario 3 (MI-RD) and impute missing values by treatment group. No additional covariates will be included in the imputation model.
5. Study discontinuation resulting in study intervention discontinuation; with missing value	<p>5A: If the study discontinuation is possibly related to study intervention (see study discontinuation reasons classified as Category 5A in Table GZGT.4.4), missing values at the endpoint visit will be imputed in the same way as Scenario 4.</p> <p>5B: If the study discontinuation is clearly due to administrative reasons not related to study intervention (see study discontinuation reasons classified as Category 5B in Table GZGT.4.4), missing values at the endpoint visit will be imputed in the same way as Scenario 2.</p>

Abbreviations: MAR = missing at random; MCMC = Markov chain Monte Carlo; MI-RD = multiple imputation using retrieved dropouts; N/A = not applicable.

Table GZGT.4.4. Study Discontinuation Reasons

Disposition Reason	Associated Sub-categories	Category
Adverse event		5A
Death		5A
Protocol deviation ^a		5A
Pregnancy		5A
Non-compliance with study drug		5A
Lack of efficacy		5A
Withdrawal by subject	Concern about study procedures/perceived risks	5A
	Scheduling conflicts	5A
	Subject is moving or has moved	5A
	Personal issue unrelated to trial	5A
	Due to epidemic/pandemic	5B
	Other (option to include a specify field)	5A
Physician decision	Concern about study procedures/perceived risks	5A
	Due to epidemic/pandemic	5B
	Other (option to include a specify field)	5A
Study terminated by sponsor		5B
Site terminated by sponsor		5B
Study terminated by IRB/ERB		5B
Lost to follow-up		5A
Other		5A

Abbreviations: ERB = ethical review board; IRB = institutional review board.

Note: For participants that discontinued the study due to emergency unblinding, the study disposition will be classified based upon the reason leading to emergency unblinding (for example, Adverse Event [5A]).

4.1.2.1.4. Multiple Imputation Based Tipping-Point Analysis

A multiple imputation-based tipping-point (MI-TP) analysis is planned as a sensitivity analysis to explore how different patterns of HbA1c change post-treatment period discontinuation by different treatment groups could impact the treatment comparisons.

To start with, missing data are imputed according to the imputation procedure as shown in [Table GZGT.4.3](#). A penalty is then added to those imputed values at the Week 40 visit. The analysis is intended to vary the magnitude of the penalties added for all treatment groups under comparison and evaluate the impact these would have on the study conclusion. A 2-dimensional space of penalties will be assessed for each dose of orforglipron and placebo. MI-TP aims to evaluate the robustness of the superiority claim to the assumptions of using the observed data to impute the missing HbA1c in all treatment groups.

4.1.2.2. Analysis Methods for Efficacy Estimand

4.1.2.2.1. Method for Continuous Variables

For the assessment of continuous efficacy measures guided by the efficacy estimand, a restricted maximum likelihood-based MMRM (Wang and Du 2023) analysis will be used. All the longitudinal observations at each scheduled postbaseline visit will be included in the analysis. The MMRM model will include these terms:

- treatment group as a factor variable
- visit as a factor variable
- pooled country as a factor variable
- a single strata variable for other stratification factors (HbA1c at baseline [$\leq 8.0\%$, $> 8.0\%$]), and prior use of any antihyperglycemic medication [Yes or No]), as a factor variable
- baseline value (of the dependent variable)
- interaction between treatment group, visit and strata variable, and
- interaction between treatment group, visit and baseline value.

The estimated treatment group effect and comparison between each of the orforglipron treatment groups and placebo at the scheduled visits will be reported together with the variability estimated using the robust inference as provided in Wang and Du (2023). The associated 2-sided 95% CI and corresponding p-values will also be reported. The sandwich estimator (Diggle et al. 1994) for the variance-covariance matrix will be used. An unstructured covariance matrix by treatment groups will be used to model the within-participant errors, assuming heteroscedasticity (variant covariance structure across treatment groups) and the measurements for different participants are independent. If the model fails to converge, the model will be simplified to include the following terms, removing 3-way interactions and the heteroscedasticity assumption:

- treatment group as a factor variable
- visit as a factor variable
- pooled country as a factor variable
- a single strata variable for other stratification factors (HbA1c at baseline [$\leq 8.0\%$, $> 8.0\%$]), and prior use of any antihyperglycemic medication [Yes or No]), as a factor variable
- baseline value (of the dependent variable)
- interaction between visit and treatment group
- interaction between visit and strata variable, and
- interaction between visit and baseline value.

If the model still fails to converge, the following covariance structures will be tested in this order for the simplified model:

- unstructured covariance matrix
- heterogeneous Toeplitz
- heterogeneous autoregressive
- heterogeneous compound symmetry
- homogeneous Toeplitz

- homogeneous autoregressive, and
- homogeneous compound symmetry.

The first covariance structure that converges will be used.

Some parameters may be log-transformed before fitting the MMRM as specified in Section 4.4, Section 4.5, and Section 4.6. In these cases, LS means and 95% CIs for each treatment group and treatment difference will be back-transformed and presented as mean percent change from baseline and as relative treatment difference to placebo in percent change.

If the data does not warrant the use of an MMRM model, then an ANCOVA model will be conducted, as described in Section 4.1.2.1.1. Missing values will be imputed based on the methods described in Section 4.1.2.2.3.

4.1.2.2.2. Method for Binary Variables

For binary efficacy measures derived from a continuous variable and guided by the efficacy estimand, the missing value in the underlying continuous variable will be imputed first based on methods described in Section 4.1.2.2.3 and then the corresponding binary variable will be derived.

For the assessment of binary efficacy objectives, the logistic regression model described in Section 4.1.2.1.2 will be used after imputation and estimation of treatment group specific risk, risk difference, relative risk, p-value, and 95% CI will be presented.

4.1.2.2.3. Handling of Missing Data

Missing data should be minimized at the best precaution. For continuous efficacy analyses guided by efficacy estimand, the hypothetical strategy will be used to handle the ICEs. For the MMRM analysis, only data collected before the occurrence of any ICEs will be used. Through the MMRM, the potential efficacy measures (after the ICEs) will be implicitly imputed as if participants did not have ICEs. For the ANCOVA analysis, missing values at the last visit will be imputed through multiple imputation using all non-missing data (excluding data collected after ICEs) from the same treatment group under the missing at random (MAR) assumption. The same method will also be used to impute the missing value in the underlying continuous variables of binary outcomes.

4.1.2.3. Analysis Methods for Safety

For categorical safety measures, treatment group differences of percentages will be conducted using Fisher's exact test or Pearson's chi-squared test if applicable, unless otherwise specified. Risk difference and the 95% CI will be provided.

For selected continuous safety measures, unless otherwise specified, treatment group differences of mean change or mean percent change from baseline at all scheduled visits will be assessed via an MMRM using restricted maximum likelihood (REML), which will include the following terms:

- treatment group as a factor variable
- visit as a factor variable
- baseline value (of the dependent variable), and
- interaction between visit and treatment group.

If the data does not warrant the MMRM model, then an ANCOVA model with treatment group as a fixed effect and the continuous baseline value as a covariate will be used.

No explicit imputation will be conducted for safety measures. Some parameters (such as urinary albumin to creatinine ratio [UACR], p-amylase, and lipase) may be log-transformed before fitting the MMRM as specified in Section 4.6. In these cases, LS means and 95% CIs for each treatment group and treatment difference will be back-transformed and presented as mean percent change from baseline and as relative treatment difference of orforglipron treatment groups compared to placebo in percent change.

For selected safety parameters, time-to-first-event analysis via the Cox proportional hazards model may be conducted. A log-rank test will be used to calculate p-values. The Kaplan-Meier method will be used to estimate the cumulative event curve over time. Counts and proportions of participants who experience the event will be calculated by treatment group.

Where necessary, the rate of events will be analyzed using a generalized linear mixed -effects model, assuming the number of events follows a negative binomial distribution with the mean modeled using treatment as a fixed effect. The logarithm of days during analysis interval will be adjusted as an offset to account for possible unequal duration of follow-up between participants.

Some safety analyses may be conducted excluding data after the initiation of new anti-hyperglycemic medications.

4.2. Participant Dispositions

The participant dispositions for the screening period, the study intervention/treatment, the treatment period and/or the follow-up period will be collected in CRFs with the corresponding primary reason. The study completion for a participant is defined as the participant completing both the treatment period and the follow-up period, regardless of completion of study intervention.

The planned listings and summary tables for dispositions are provided in [Table GZGT.4.5](#).

Table GZGT.4.5. Listings and Summary Tables Related to Dispositions

Analysis	Population/Period
Summary of disposition prior to randomization	Entered
Summary of study and study intervention disposition	Randomized/TP + FP (TP for study intervention disposition)
Participant allocation by region, country, and center/site	Entered
Kaplan-Meier plot of time to study discontinuation	Randomized/TP + FP
Kaplan-Meier plot of time to study intervention discontinuation	Randomized/TP
Kaplan-Meier plot of time to study intervention discontinuation due to AE	Randomized/TP
Listing of randomization	Randomized
Listing of randomized participants who were discontinued from the study intervention due to inadvertent enrollment	Randomized
Listing of study and study intervention disposition	Randomized

Abbreviations: AE = adverse event; FP = follow-up period; TP = treatment period.

4.3. Primary Endpoint/Estimand Analysis

The primary study objective is to demonstrate that orforglipron \blacksquare mg, \blacksquare mg, and/or \blacksquare mg are superior to placebo with respect to the mean change from baseline in HbA1c at Week 40.

4.3.1. Definition of Endpoint

The primary efficacy measure is change from baseline in HbA1c at Week 40. The change from baseline in HbA1c for each participant at each nominal visit is defined as postbaseline HbA1c at the nominal visit minus baseline, where baseline is defined as in Section 4.1.

In addition, if HbA1c measurement is missing/canceled at the primary endpoint visit (Week 40), retests conducted within 7 days of Week 40 will be used as primary endpoint in model analyses.

4.3.2. Main Analytical Approach

The analytical approaches are specified for the treatment regimen estimand in Section 4.1.2.1.1 (ANCOVA) and the efficacy estimand in Section 4.1.2.2.1 (MMRM).

For each hypothesis, orforglipron treatment groups will be declared superior to placebo in controlling HbA1c if the p-value is less than the alpha level allocated to the hypothesis according to the graphical approach (Section 2.1).

4.3.3. Sensitivity Analysis

The sensitivity analyses are specified in Section 4.1.2.1.4 and Section 4.1.2.1.4 (MI-TP).

4.4. Secondary Endpoints/Estimands Analysis

4.4.1. Key Secondary Endpoints/Estimands

The key secondary study objectives are listed in Section 1.1.

4.4.1.1. Definition of Endpoints

The definition of endpoints are:

- incidence of participants (proportion at the population-level) achieving an HbA1c target value of <7.0% (53 mmol/mol) at Week 40 (Yes or No)
- incidence of participants (proportion at the population-level) achieving an HbA1c target value of $\leq 6.5\%$ (48 mmol/mol) at Week 40 (Yes or No)
- change from baseline in fasting serum glucose at Week 40
- percentage change from baseline in body weight at Week 40
- change from baseline in body weight at Week 40
- percentage change from baseline in non-HDL cholesterol at Week 40 (CC mg and CC mg only), and
- percentage change from baseline in triglycerides at Week 40 (CC mg and CC mg only).

4.4.1.2. Main Analytical Approach

The analytical approaches are specified for the treatment regimen estimand in Section 4.1.2.1 and for the efficacy estimand in Section 4.1.2.2.

For each hypothesis, orforglipron treatment groups will be declared superior to placebo if the p-value is less than the alpha level allocated to the hypothesis according to the graphical approach (Section 2.1).

4.4.2. Other Secondary Endpoints/Estimands

The other secondary efficacy endpoints will be guided by the efficacy estimand as shown:

- incidence of participants (proportion at the population-level) achieving an HbA1c target value of <5.7% (39 mmol/mol) at Week 40 (Yes or No)
- change from baseline in daily average 7-point self-monitored blood glucose (SMBG) at Week 40
- incidence of participants (proportion at the population-level) achieving a weight loss of $\geq 5\%$, $\geq 10\%$, and $\geq 15\%$ from baseline at Week 40
- change from baseline in waist circumference at Week 40, and
- change from baseline in body mass index (BMI) at Week 40.

For SMBG, valid SMBG profiles will be used for analysis, defined as having non-missing values at ≥ 4 time points (including at least one post-meal point) among the 7 pre-specified time points and being collected within 2 weeks prior to the given visit. For each time point, the average of the corresponding SMBG values from the valid SMBG profiles will be used for analysis. In addition to the analyses of the mean of all 7-point measurements, similar analyses will be performed for each of the 7 points, the 2-hour morning, midday, and evening meal excursions, the mean of all meals' 2-hour excursion, the mean of all pre-meal measurements, and the mean of all 2-hour postprandial measurements.

The other secondary efficacy endpoints will be guided by the treatment regimen estimand and the efficacy estimand as shown:

- change from baseline at Week 40 in:

- systolic blood pressure
- diastolic blood pressure
- percentage change from baseline in lipid parameters at Week 40 (log-transformed data)
 - total cholesterol
 - HDL-cholesterol
 - low-density lipoprotein (LDL)-cholesterol
 - very low-density lipoprotein (VLDL)-cholesterol
 - non-HDL cholesterol (■ mg only), and
 - triglycerides (■ mg only).
- change from baseline at Week 40 in:
 - Short Form-36 version 2 Health Survey (SF-36v2) Acute Form domain scores
 - EQ-5D-5L health state utilities and CCI ■■■■■■
 - Ability to Perform Physical Activities of Daily Living (APPADL) scores
 - Impact of Weight on Self-Perception (IW-SP) scores
 - Diabetes Treatment Satisfaction Questionnaire-Status (DTSQs) scores, and
- score of the Diabetes Treatment Satisfaction Questionnaire-Change (DTSQc) at Week 40.

The analytical approaches for the other secondary efficacy endpoints are specified for the treatment regimen estimand in Section 4.1.2.1 and the efficacy estimand in Section 4.1.2.2, unless otherwise stated. The analytical approaches for the other secondary safety endpoints are specified in Section 4.1.2.3.

No multiplicity adjustment will be conducted for non-key secondary endpoints.

4.4.2.1. Health Outcomes

The patient-reported outcome questionnaires will be completed by the participants at baseline and postbaseline (Week 40 or early discontinuation visit). Guided by the treatment regimen estimand and efficacy estimand, the analysis will be conducted using an ANCOVA model as described in Section 4.1.2.1.1, with missing values at Week 40 imputed based on the methods described in Section 4.1.2.1.3 and Section 4.1.2.2.3, respectively, for the two estimands.

4.4.2.1.1. Short-Form-36 Health Survey Version 2, Acute Form

Per copyright owner, the PRO CoRE 2.0 Smart Measurement® System will be used to derive the following 8 domains and 2 component scores:

- Physical Functioning domain
- Role-Physical domain
- Bodily Pain domain
- General Health domain
- Vitality domain
- Social Functioning domain
- Role-Emotional domain
- Mental Health domain
- Mental component score, and

- Physical component score.

Each domain is scored individually and information from these 8 domains is further aggregated into the 2 health component scores. Items are answered on Likert scales of varying lengths (3-point, 5-point, or 6-point scales). Scoring of each domain and both component scores are norm-based and presented in the form of T-scores, with a mean of 50 and an SD of 10; higher scores indicate better levels of function and/or better health (Maruish 2011). The following analyses for the actual values and change from baseline values for the 8 domains and the two component scores will be conducted at Week 40:

- descriptive summaries by treatment group, and
- ANCOVA as described in Section 4.4.2.1.

4.4.2.1.2. EQ-5D-5L

The EQ-5D-5L is a standardized 5-item instrument that assesses 5 dimensions of health. These are:

- mobility
- self-care
- usual activities
- pain/discomfort, and
- anxiety/depression.

Each dimension has 5 levels:

- no problems
- slight problems
- moderate problems
- severe problems, and
- extreme problems.

The EQ-5D-5L also has a vertical visual analog scale, called the EQ VAS, which records the respondent's self-rated health with endpoints labeled as "best imaginable health state" (100) and "worst imaginable health state" (0). The EQ-5D-5L health states are converted into a single index "utility" score using a scoring algorithm based on public preferences. The single index "utility" score will be based on the appropriate United Kingdom scoring algorithm (Devlin et al. 2018), where higher scores indicate better overall health related quality of life.

Each item will be summarized descriptively by treatment group at each scheduled visit at which the EQ-5D-5L is administered. The changes from baseline to Week 40 in the index, and the VAS scores, will be analyzed using an ANCOVA as described in Section 4.4.2.1.

4.4.2.1.3. Ability to Perform Physical Activities of Daily Living

The APPADL contains 7 items that assess how difficult it is for participants to engage in certain activities considered to be integral to normal daily life, such as walking, standing, and climbing stairs (Hayes et al. 2012). Items are scored on a 5-point numeric rating scale, where 5 = "not at all difficult" and 1 = "unable to do." A raw overall score is calculated by simply summing the scores of the 7 items, and a transformed overall score is obtained by linearly transforming the

raw overall score to a 0 to 100 scale. A higher raw overall score and a higher transformed overall score are indicative of better ability to perform activities of daily living.

Descriptive summaries by treatment group at each scheduled visit at which the APPADL is administered will be presented for each item. Treatment comparison in the transformed overall APPADL score change from baseline to Week 40 will be analyzed using an ANCOVA as described in Section 4.4.2.1.

4.4.2.1.4. Impact of Weight on Self-perceptions Questionnaire

The IW-SP contains 3 items that assess how often a participant's body weight affects how happy they are with their appearance and how often they feel self-conscious when out in public (Hayes and DeLozier, 2015). Each item is rated on a 5-point scale ranging from 1 = "always" to 5 = "never." Total scores for the IW-SP are derived by summing the item scores and dividing by the number of items. The score can also be transformed to a range from 0 to 100. Higher IW-SP scores correspond to better self-perception.

Descriptive summaries by treatment group at each scheduled visit at which the IW-SP is administered will be presented for each item. Treatment comparison in the transformed overall IW-SP total score change from baseline to Week 40 will be analyzed using an ANCOVA as described in Section 4.4.2.1.

4.4.2.1.5. Diabetes Treatment Satisfaction Questionnaire

The Diabetes Treatment Satisfaction Questionnaire (DTSQ) contains 8 items (conceptually the same items in the status [DTSQs] and change [DTSQc] versions) (Bradley 1994).

- For DTSQs, 6 items (1, and 4 through 8) are rated from 0 to 6 and item scores are summed to produce a measure of treatment satisfaction. Greater scores indicate greater treatment satisfaction. The 2 remaining items (2 and 3) are treated individually to assess, respectively, the perceived frequency of hyperglycemia and hypoglycemia.
- For DTSQc, 6 items (1, and 4 through 8) are scored on a scale of -3 to +3. Item scores are summed to produce a measure of change in treatment satisfaction. For all items except item 2 (perceived frequency of hyperglycemia) and item 3 (perceived frequency of hypoglycemia):
 - the higher the score, the greater the improvement in treatment satisfaction
 - the lower the score, the greater the deterioration in treatment satisfaction
 - a score of 0 represents no change.
 - For items 2 and 3: the lower the score, the better the glucose control.

Descriptive summaries will be provided at baseline (DTSQs only) and at Week 40 (DTSQs and DTSQc) for the perceived hyperglycemia item, perceived hypoglycemia item, and the 6-item overall satisfaction score.

Treatment comparison in the change from baseline in DTSQs scores and in the actual responses to DTSQc at Week 40 will be analyzed using an ANCOVA as described in Section 4.4.2.1, except that baseline will not be used as a covariate in DTSQc. The analyses will be conducted for the perceived hyperglycemia item, perceived hypoglycemia item, and the 6-item overall satisfaction score.

4.5. Tertiary and Exploratory Endpoints Analysis

4.5.1. Pharmacokinetic and Pharmacokinetic/Pharmacodynamic Methods

Pharmacokinetic (PK), pharmacodynamic (PD), and PK/PD analysis are the responsibility of Lilly's PK/PD group. Please refer to the Population PK Analysis plan.

4.5.2. Exploratory Endpoints Analysis

The exploratory endpoints and analysis are specified in [Table GZGT.4.6](#) and [Table GZGT.4.7](#). Additional exploratory analysis may be conducted as deemed necessary.

Table GZGT.4.6. Description and Derivation of Exploratory Analysis

Measure	Description	Variable	Derivation/Comment	Handling Missing Components
Homeostasis Model Assessment	The homeostasis model assessment 2 B (HOMA2 B) estimates steady state beta cell function	HOMA2 B (insulin)	Calculated from the link ^a using fasting glucose and insulin laboratory values	Missing if fasting glucose or insulin is missing
		Change from baseline in HOMA2 B (insulin)	Calculated as: postbaseline HOMA2 B (insulin) – baseline HOMA2 B (insulin)	Missing if baseline or postbaseline value is missing
		Percent change from baseline in HOMA2 B (insulin)	Calculated as: [postbaseline HOMA2 B (insulin) / baseline HOMA2 B (insulin) – 1] × 100%	Missing if baseline or postbaseline value is missing
		HOMA2 B (C-peptide)	Calculated from the link ^a using fasting glucose and C-peptide laboratory values	Missing if fasting glucose or C-peptide is missing
		Change from baseline in HOMA2 B (C-peptide)	Calculated as: postbaseline HOMA2 B (C-peptide) – baseline HOMA2 B (C-peptide)	Missing if baseline or postbaseline value is missing
		Percent change from baseline in HOMA2 B (C-peptide)	Calculated as: [postbaseline HOMA2 B (C-peptide) / baseline HOMA2 B (C-peptide) – 1] × 100%	Missing if baseline or postbaseline value is missing
	The homeostasis model assessment 2 IR (HOMA2 IR) estimates insulin resistance	HOMA2 IR (insulin)	Calculated from the link ^a using fasting glucose and insulin laboratory values	Missing if fasting glucose or insulin is missing
		Change from baseline in HOMA2 IR (insulin)	Calculated as: postbaseline HOMA2 IR (insulin) – baseline HOMA2 IR (insulin)	Missing if baseline or postbaseline value is missing
		Percent change from baseline in HOMA2 IR (insulin)	Calculated as: [postbaseline HOMA2 IR (insulin) / baseline HOMA2 IR (insulin) – 1] × 100%	Missing if baseline or postbaseline value is missing
		HOMA2 IR (C-peptide)	Calculated from the link ^a using fasting glucose and C-peptide laboratory values	Missing if fasting glucose or C-peptide is missing
		Change from baseline in HOMA2 IR (C-peptide)	Calculated as: postbaseline HOMA2 IR (C-peptide) – baseline HOMA2 IR (C-peptide)	Missing if baseline or postbaseline value is missing

		Percent change from baseline in HOMA2 IR (C-peptide)	Calculated as: [postbaseline HOMA2 IR (C-peptide) / baseline HOMA2 IR (C-peptide) - 1] × 100%	Missing if baseline or postbaseline value is missing
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Abbreviations: HOMA2 B = homeostasis model assessment 2 B; HOMA2 IR = homeostasis model assessment 2 IR.

^a Derive HOMA2 B and HOMA2 IR using the calculator at <https://www.rdm.ox.ac.uk/about/our-clinical-facilities-and-mrc-units/DTU/software/homa>.

Table GZGT.4.7. Description of Exploratory Analysis

Measure	Variable	Estimand Data Points Set/Population ^a	Analysis Method	Time Point ^b
HOMA	Change from baseline in HOMA2 B (insulin)	Efficacy	MMRM	Week 40 visit
	Percent change from baseline in HOMA2 B (insulin)	Efficacy	MMRM	Week 40 visit
	Change from baseline in HOMA2 IR (insulin)	Efficacy	MMRM	Week 40 visit
	Percent change from baseline in HOMA2 IR (insulin)	Efficacy	MMRM	Week 40 visit
	Change from baseline in HOMA2 B (C-peptide)	Efficacy	MMRM	Week 40 visit
	Percent change from baseline in HOMA2 B (C-peptide)	Efficacy	MMRM	Week 40 visit
	Change from baseline in HOMA2 IR (C-peptide)	Efficacy	MMRM	Week 40 visit
	Percent change from baseline in HOMA2 IR (C-peptide)	Efficacy	MMRM	Week 40 visit

Abbreviations: HOMA = homeostasis model assessment; HOMA2 B = homeostasis model assessment 2 B; HOMA2 IR = homeostasis model assessment 2 IR; MMRM = mixed-effects model for repeated measures.

^a Population is displayed if different from randomized participants.

^b Assessments collected at multiple postbaseline visits will be analyzed at those scheduled visits using MMRM as supplemental analyses, in addition to the primary time point listed in the table.

4.6. Safety Analyses

The planned safety analyses are consistent with compound-level safety standards, which are based on various sources, including company standards, internal and external subject matter experts, publications from cross-industry initiatives (for example, PHUSE 2013, PHUSE 2015, PHUSE 2017, PHUSE 2018, PHUSE 2022), and publications from regulatory agencies (for example, FDA 2010, EMA 2014, CDER/BIRRS 2022, 2023). Descriptions of the safety analyses are provided in SAP GZGT, but some details are found in the compound-level safety standards.

Unless otherwise specified, safety analyses will be conducted using the safety participants and the safety data points set (Table GZGT.4.1). It is noted that for some safety endpoints, such as study intervention discontinuation due to an AE and such, the analyses will be based on data observed during the treatment period only. The analytical approaches for safety analyses are specified in Section 4.1.2.3. Summary tables with risk difference will be sorted by decreasing order of risk difference. Not all displays will necessarily be created as a static display. Some displays will be incorporated into interactive display tools instead of, or in addition to, a static display.

4.6.1. Extent of Exposure

Duration of exposure to study intervention will be summarized by treatment group for safety participants. Exposure will be summarized and calculated for the treatment period being

considered as the date of last dose of study intervention minus the date of first dose of study intervention plus 1 day using safety participants. Duration on study (date of end of study participation – date of randomization + 1) will also be summarized by treatment group.

Descriptive statistics (including n, mean, SD, minimum, first quartile, median, third quartile, and maximum) will be provided and the frequency of participants falling into different exposure ranges will be summarized. Overall exposure will be summarized in total participant-year (PY) of exposure, derived in the following manner:

$$\text{total PY of exposure} = \text{sum of duration of exposure in days (for all participants in treatment group)} / 365.25$$

The frequency and percentages of participants falling into the following ranges will be summarized by planned treatment group as well:

- >0
- ≥ 4 weeks
- ≥ 8 weeks
- ≥ 12 weeks
- ≥ 16 weeks
- ≥ 20 weeks
- ≥ 24 weeks
- ≥ 32 weeks
- ≥ 40 weeks, and
- ≥ 52 weeks.

In addition, the frequency and percentages of participants falling into the following exposure ranges for study and study intervention may be summarized by planned treatment group:

- 0 weeks
- >0 to <4 weeks
- ≥ 4 to <8 weeks
- ≥ 8 to <12 weeks
- ≥ 12 to <16 weeks
- ≥ 16 to <20 weeks
- ≥ 20 to <24 weeks
- ≥ 24 to <32 weeks
- ≥ 32 to <40 weeks
- ≥ 40 to <52 weeks, and
- ≥ 52 weeks.

No p-values will be reported in these summaries as they are intended to describe the study populations rather than test hypotheses about said populations.

4.6.2. Adverse Events

Planned summary tables and listings related to the adverse events are provided in [Table GZGT.4.8](#), and details are described in the compound-level safety standards.

Table GZGT.4.8. Listings and Summary Tables Related to Adverse Events

Analysis	Method	Population/Period
Overview of AEs, including <ul style="list-style-type: none"> • TEAE • SAE • death, and • permanent discontinuation from study intervention due to an AE. 	Fisher's exact	Safety/TP + FP (TP for study intervention discontinuation due to an AE)
TEAEs by PT within SOC	Fisher's exact	Safety/TP + FP
Maximum Severity TEAEs		Safety/TP + FP
TEAEs with incidence $\geq 5\%$ by PT	Fisher's exact	Safety/TP + FP
SAEs by PT within SOC	Fisher's exact	Safety/TP + FP
Primary AEs leading to permanent discontinuation of study intervention by PT within SOC	Fisher's exact	Safety/TP
Primary AEs leading to permanent discontinuation of study by PT within SOC	Fisher's exact	Safety/TP
AEs leading to study intervention interruption by PT within SOC	Fisher's exact	Safety/TP
AEs leading to study intervention reduction by PT within SOC	Fisher's exact	Safety/TP
Listing of SAEs		Safety/TP + FP
Listing of primary AEs leading to permanent discontinuation of study intervention		Safety/TP
Listing of primary AEs leading to permanent discontinuation of study		Safety/TP + FP
Listing of deaths		Safety/TP + FP
Listing of AEs of suspected overdosing of orforglipron		Safety/TP
Listing of participants with at least 1 notable event		Randomized/TP + FP

Abbreviations: AE = adverse event; eCRF = electronic case report form; FP = follow-up period; PT = preferred term; SAE = serious adverse event; SOC = system organ class; TEAE = treatment-emergent adverse event; TP = treatment period.

4.6.3. Safety Topics of Interest

This section includes safety topics of interest whether due to observed safety findings, potential findings based on drug class, or safety topics anticipated to be requested by a regulatory agency for any reason. In general, safety topics of interest will be identified by 1 or more standardized Medical Dictionary for Regulatory Activities (MedDRA) query(ies) (SMQs), system organ class (SOC), high-level term (HLT), FDA Medical Query (FMQ), or by a Lilly defined MedDRA preferred term (PT) listing based upon the review of the most current MedDRA Version, or by relevant laboratory changes. Search criteria are detailed in the compound-level safety standards.

The planned analyses for safety topics of interest are provided in [Table GZGT.4.9](#) and are described more fully in the compound-level safety standards.

Table GZGT.4.9. Description and Analyses of Safety Topics of Interest

Safety Topic of Interest	Short Description	Analysis	Method	Population/ Period
MACE	Death and nonfatal CV AEs will be adjudicated by a committee of physicians external to Lilly with cardiology expertise: a CEC.	Positively adjudicated MACE by category/subcategory and PT (if necessary). MACE reported by investigators may be summarized	Fisher's exact	Safety/TP + FP
		Listing of all MACE reported by investigator (whether or not positively adjudicated) (if necessary).		Safety/TP + FP
Arrhythmias and cardiac conduction disorders	The TE arrhythmias and cardiac conduction disorders events will be derived using the MedDRA PTs contained in certain SMQs.	TE arrhythmias and cardiac conduction disorders by PT nested within SMQ and HLT ^a .	Fisher's exact	Safety/TP + FP
Hypotension, orthostatic hypotension, and syncope	The AE database will be searched using predefined PTs.	TE hypotension, orthostatic hypotension and syncope disorders by PT ^a .	Fisher's exact	Safety/TP + FP
Hypoglycemia	The 2023 American Diabetes Association position statement on glycemic targets ^b will be used to define: <ul style="list-style-type: none"> • Level 1 hypoglycemia • Level 2 hypoglycemia • Level 3 hypoglycemia (severe/serious hypoglycemia) 	Incidence (percent of participants experiencing 1 or more episodes) and the rate (episodes/participant/year) of level 3 (severe/serious) hypoglycemia.	Logistic regression (for incidence) and negative binomial regression (for rate; the logarithm of days during the analysis interval will be adjusted as an offset).	Primary analysis: Safety/TP + FP excluding events after initiation of new antihyperglycemic therapy (regardless of how many days of use). Supportive analysis: Safety/TP + FP regardless of initiation of new antihyperglycemic therapy.
		Incidence and rate of level 2 or level 3 hypoglycemia.		
		Incidence and rate of level 1 hypoglycemia (if warranted by data).		
		Listing of level 2 or level 3 hypoglycemia events.		Safety/TP + FP

Safety Topic of Interest	Short Description	Analysis	Method	Population/ Period
Severe persistent hyperglycemia	Initiation of new antihyperglycemic therapy taken for severe persistent hyperglycemia	Rescue therapy taken for severe persistent hyperglycemia (if warranted by data) ^a .	Fisher's exact	Safety/TP + FP
GI adverse events	GI AEs using gastrointestinal disorders SOC will be captured.	Severe or serious TE gastrointestinal events by PT.	Fisher's exact	Safety/TP + FP
		Plots of prevalence and incidence over time for TE nausea, vomiting, diarrhea, constipation, and NVD by maximum severity.		Safety/TP + FP
		Summary of prevalence and incidence over time for TE nausea, vomiting, diarrhea, constipation and NVD.		Safety/TP + FP
		Plot of time to onset of TE nausea, vomiting, diarrhea, constipation and NVD.	Kaplan-Meier	Safety/TP + FP
		Study intervention discontinuation due to GI.	Fisher's exact	Safety/TP + FP
Renal safety	Laboratory measures related to renal safety will be analyzed. Renal events including acute renal failure and chronic renal failure exacerbation will be captured using SMQs. Dehydration events will also be identified using SMQs.	Shift of minimum-to-minimum for eGFR as estimated by the CKD-EPI equation ^c .		Safety/TP + FP
		Shift of maximum-to-maximum for UACR.		Safety/TP + FP
		MMRM analyses for eGFR ^c .	MMRM	Safety/TP + FP
		MMRM analyses for UACR (log-transformation).	MMRM	Safety/TP + FP
		TE renal events by PT nested within SMQs ^a .	Fisher's exact	Safety/TP + FP
		TE dehydration events by PT ^a .	Fisher's exact	Safety/TP + FP
Pancreatitis		Shift of maximum-to-maximum for pancreatic enzyme.		Safety/TP + FP

Safety Topic of Interest	Short Description	Analysis	Method	Population/ Period
	The pancreatic enzyme data (p-amylase and lipase) will be observed through laboratory testing. All suspected cases of acute or chronic pancreatitis and AEs of severe or serious abdominal pain of unknown etiology will be sent for adjudication by an independent CEC.	MMRM analysis for pancreatic enzymes: p-amylase and lipase (log-transformation).	MMRM	Safety/TP + FP
		Investigator-reported events and positively adjudicated pancreatic events, respectively	Fisher's exact	Safety/TP + FP
		TE pancreatic events by "Acute pancreatitis" SMQ and "Chronic pancreatitis" PT	Fisher's exact	Safety/TP + FP
		Listing of adjudicated and investigator-reported pancreatitis.		Safety/TP + FP
Thyroid malignancies and C-cell hyperplasia	TE thyroid malignancies and C-cell hyperplasia will be identified using MedDRA HLTs and PTs. The purpose of calcitonin measurements is to assess the potential effect of orforglipron on thyroid C-cell function.	TE thyroid C-cell hyperplasia and malignancies by PT.	Fisher's exact	Safety/TP + FP
		Shift of maximum-to-maximum for calcitonin value in the thresholds.		Safety/TP + FP
		MMRM analysis for calcitonin (log-transformation)	MMRM	Safety/TP + FP
		Postbaseline calcitonin value of ≥ 35 ng/mL that has increased at least 50% over baseline ^a .	Fisher's exact	Safety/TP + FP
Malignancies	The malignancy events will be derived using the MedDRA PTs contained in certain SMQs.	TE malignancy by PT nested within SMQ ^a .	Fisher's exact	Safety/TP + FP
Hepatic safety	Hepatic labs include ALT, AST, serum ALP, TBL, DBL, INR, and GGT.	Abnormal postbaseline categories for hepatic safety parameters: ALT, AST, ALP, TBL, DBL, and GGT.		Safety/TP + FP
		TE potentially drug-related hepatic disorders by PT nested within SMQ	Fisher's exact	Safety/TP + FP
		Hepatocellular drug-induced liver injury screening plot (TBL vs ALT or AST).		Safety/TP + FP

Safety Topic of Interest	Short Description	Analysis	Method	Population/ Period
	When criteria are met for hepatic evaluations, investigators will conduct close monitoring of hepatic symptoms and liver tests, perform a comprehensive evaluation for alternative causes of abnormal liver tests, and complete follow-up hepatic safety eCRFs. Potentially drug-related hepatic disorders will be identified using SMQs.	Hepatocellular drug-induced liver injury screening table.		Safety/TP + FP
		Listing of participants with ALT or AST $\geq 3 \times$ ULN.		Safety/TP + FP
		Cholestatic drug-induced liver injury screening plot (TBL vs ALP).		Safety/TP + FP
		Cholestatic drug-induced liver injury screening table.		Safety/TP + FP
		Listing of participants with ALP or TBL $\geq 2 \times$ ULN.		Safety/TP + FP
		Participant profiles for participants meeting criteria for a comprehensive hepatic evaluation (as defined in Protocol GZGT).		Safety/TP + FP
		Shift of maximum-to-maximum for ALT, AST, ALP, TBL		Safety/TP + FP
Gallbladder and biliary tract disorders	All events of TEAE biliary colic, cholecystitis, or other suspected events related to gallbladder disease will be identified by using certain SMQs.	TE gallbladder and biliary tract disorders by PT nested within SMQ ^a .	Fisher's exact	Safety/TP + FP
Hypersensitivity reactions	AEs will be searched using MedDRA PTs from narrow scope FMQ or SMQ.	TE allergic reactions and hypersensitivities by PT nested within applicable FMQ or SMQ ^a .	Fisher's exact	Safety/TP + FP
Major depression, suicidal ideation, and behavior	AEs will be searched using MedDRA PTs that satisfies the search criteria.	TE depression, suicidal ideation, and behavior events by PT nested within applicable SMQ or FMQ ^a .	Fisher's exact	Safety/TP + FP
Abuse potential	AEs will be searched using a modified abuse potential FMQ.	TE abuse potential events by PT ^a .	Fisher's exact	Safety/TP + FP

Safety Topic of Interest	Short Description	Analysis	Method	Population/ Period
Diabetic retinopathy	Diabetic retinopathy and macular edema will be monitored using retinal fundus photography and additional ophthalmological follow-up as warranted throughout the study for participants with T2D. Diabetic retinopathy complication will be searched using MedDRA PTs.	Baseline and postbaseline diabetic retinopathy exam by treatment group and by eye (that is, any eye, left eye and right eye).		Safety/TP + FP
		TE diabetic retinopathy, diabetic macular edema and related complications by PTs.	Fisher's exact	Safety/TP + FP
		Shift of baseline diabetic retinopathy and macular edema status to most severe postbaseline diabetic retinopathy and macular edema status.		Safety/TP + FP

Abbreviations: AE = adverse event; ALT = alanine aminotransferase; ALP = alkaline phosphatase; AST = aspartate aminotransferase; CEC = clinical endpoint committee; CKD-EPI = Chronic Kidney Disease Epidemiology; CV = cardiovascular; DBL = direct bilirubin; eCRF = electronic case report form; eGFR = estimated glomerular filtration rate; FDA = United States Food and Drug Administration; FMQ = FDA Medical Query; FP = follow-up period; GGT = gamma-glutamyltransferase; GI = gastrointestinal; HLT = high-level term; INR = international normalized ratio; MACE = major adverse cardiovascular event; MedDRA = Medical Dictionary for Regulatory Activities; MMRM = mixed model for repeated measures; NVD = nausea, vomiting, diarrhea; PT = preferred term; SMQ = standardized MedDRA query; T2D = type 2 diabetes; TE = treatment-emergent; TEAE = treatment-emergent adverse event; TBL = total bilirubin; TP = treatment period; UACR = urinary albumin to creatinine ratio.

^a For these tables, if the number of events is less than 10, a listing will be provided instead.

^b ElSayed et al. 2023.

^c eGFR will be calculated by Chronic Kidney Disease-Epidemiology cystatin-c equation (Inker et al. 2012).

4.6.4. Additional Safety Assessments

Actual and change from baseline to postbaseline values of central laboratory measures, vital signs, and selected electrocardiogram (ECG) parameters will be summarized at each scheduled visit. Continuous variables, as well as the change from baseline for these variables, will be analyzed by MMRM or ANCOVA models as described in Section 4.1.2.3.

4.6.4.1. Central Laboratory Measures

All laboratory data will be reported in the International System of Units (SI). Selected laboratory measures will also be reported using conventional units.

The planned summaries for clinical laboratory evaluations are provided in Table GZGT.4.10 and are described more fully in the compound-level safety standards.

Table GZGT.4.10. Summary Tables Related to Clinical Laboratory Evaluations

Analysis	Method	Population/Period
Box plots and means/SDs (or 95% CIs) for observed values by visit	Descriptive statistics	Safety/TP + FP
Box plots and means/SDs (or 95% CIs) for change from baseline values by visit	Descriptive statistics	Safety/TP + FP
Summary of participants with elevated or low values meeting specified levels	Descriptive statistics	Safety/TP + FP
Listing of abnormal laboratory findings		Safety/TP + FP

Abbreviations: CI = confidence interval; FP = follow-up period; SD = standard deviation; TP = treatment period.

4.6.4.2. Vital Signs

Measurements taken in triplicate for a particular visit will be averaged before any analysis or summary outlined in this section.

The planned summaries for vital signs (systolic blood pressure, diastolic blood pressure, and pulse rate) are provided in Table GZGT.4.11, and are described more fully in the compound-level safety standards.

Table GZGT.4.11. Summary Tables Related to Vital Signs

Analysis	Method	Population/Period
Box plots and means/SDs (or 95% CIs) for observed values by visit	Descriptive statistics	Safety/TP + FP
Box plots and means/SDs (or 95% CIs) for change from baseline values by visit	Descriptive statistics	Safety/TP + FP
Analysis of pulse rate and change from baseline	MMRM	Safety/TP + FP
Summary of participants meeting specific blood pressure and pulse rate levels	Descriptive statistics	Safety/TP + FP
Shift of maximum-to-maximum for pulse rate		Safety/TP + FP

Abbreviations: CI = confidence interval; FP = follow-up period; MMRM = mixed model for repeated measures; SD = standard deviation; TP = treatment period.

4.6.4.3. Electrocardiograms

Measurements taken in triplicate for a particular visit will be averaged before any analysis or summary outlined in this section. The planned summaries for ECG parameters are provided in [Table GZGT.4.12](#) and are described more fully in the compound-level safety standards.

Table GZGT.4.12. Summary Tables Related to Electrocardiogram Parameters

Analysis	Method	Population/Period
Box plot and mean/SD (or 95% CI) for observed values by visit	Descriptive statistics	Safety/TP + FP
Box plot and mean/SD (or 95% CI) for change from baseline values by visit	Descriptive statistics	Safety/TP + FP
Analysis of change from baseline in ECG parameters (heart rate, PR interval)	MMRM	Safety/TP + FP
Heart rate and PR interval in specified categories	Descriptive statistics	Safety/TP + FP

Abbreviations: CI = confidence interval; ECG = electrocardiogram; FP = follow-up period; MMRM = mixed model for repeated measures; PR = pulse rate; SD = standard deviation; TP = treatment period.

4.7. Other Analyses

4.7.1. Subgroup Analyses

Subgroup analyses will be conducted for the primary endpoint with the treatment-regimen estimand:

- change from baseline in HbA1c at Week 40 visit.

The analyses will be conducted across these subgroups (including but not limited to)

- age group (<65, ≥65 years)
- age group (<75, ≥75 years)
- sex (female, male)
- baseline HbA1c stratum (≤8.0%, >8.0%)
- baseline BMI (<30, ≥30 and <35, ≥35 kg/m²)
- baseline eGFR calculated by CKD-EPI cystatin-c equation (<60 and ≥60 mL/min/1.73 m²)
- duration of T2D (<median, ≥median)

- duration of T2D (≤ 5 , > 5 and ≤ 10 , > 10 years)
- race
- ethnicity
- geographic region (United States versus outside of the United States), and
- country.

The ANCOVA model specified in Section 4.1.2.1.1 will be fitted separately within each category of subgroup. The LS means, LS means difference, SE, and 95% CI will be presented. The LS means and variance-covariance estimates from these separate models will be used to test the treatment-by-subgroup interaction at a significance level of 0.10.

If any category within the subgroup (for example, status of yes or no) is $< 10\%$ of the total population, only descriptive statistics will be provided for that category (that is, no inferential testing will be performed within the subgroup category).

A Bayesian shrinkage method will be used to provide adjusted estimates and inferences from subgroup analyses to potentially account for multiplicity. Following the paper by Wang et.al (2024), for a given baseline characteristic with k subgroups, let Y_i ($i = 1, \dots, k$) be the observed sample estimate of the treatment effect difference in subgroup i . The following hierarchical model will be used:

$$\begin{aligned} Y_i &\sim N(\mu_i, \sigma_i^2), \\ \mu_i &\sim N(\mu, \tau^2), \\ \mu &\sim N(0, 5^2), \\ \tau &\sim N^+(1), \end{aligned}$$

where σ_i^2 is set to the observed variance for sample estimate, $N^+(1)$ is the half-normal distribution with a scaled parameter of 1. A weakly informative prior, as suggested by Röver et al. (2021) and specified above, will be applied to μ and τ . Of note, a standard deviation of 5% is chosen for the centrality parameter μ , so that its standard deviation is approximately twice the unit information standard deviation (UISD). A scale parameter of 1 is chosen for the half-normal distribution such that the median heterogeneity in HbA1c reduction is approximately 0.67%, which is a reasonably large difference.

Additional subgroup evaluations may be conducted as exploratory analyses.

4.8. Interim Analyses

No interim analyses are planned for this study. If an unplanned interim analysis is deemed necessary for reasons other than a safety concern, the protocol must be amended.

4.8.1. Data Monitoring Committee

No data monitoring committee (DMC) is planned for this study.

An independent DMC will be established for interim safety monitoring of Study J2A-MC-GZGS (ACHIEVE-4) (GZGS), “A Phase 3, Open Label, Study of Once-Daily LY3502970 Compared with Insulin Glargine in Adult Participants with Type 2 Diabetes and Obesity or Overweight at Increased Cardiovascular Risk”, a study within the orforglipron Phase 3 program for T2D. The

DMC will have the responsibility to review unblinded interim safety analysis results from that study. The DMC may be asked to review unblinded safety data from Study GZGT if a need arises following the blinded trial-level safety reviews (TLSRs) conducted by the sponsor. If needed, permanent data transfers will occur for the purposes of supporting the DMC. Only the statisticians from the statistical analysis center (SAC) will have access to the unblinded data that are presented to the DMC. The SAC and members of the DMC will abide by the principles and responsibilities described in the Study GZGS DMC charter, which includes keeping all unblinded information confidential until the planned unblinding of the trial.

4.9. Changesto Protocol-planned Analyses

The objective to demonstrate that orforglipron 120 mg is superior to placebo in weight management was excluded from the graphical multiple testing procedure and is not included in Figure GZGT.2.1.

5. Sample Size Determination

Approximately [REDACTED] participants will be randomly assigned in a [REDACTED] ratio to orforglipron [REDACTED] mg once daily (QD), [REDACTED] mg QD, [REDACTED] mg QD, or placebo, respectively.

This trial is powered at over 90% to assess superiority of orforglipron [REDACTED] mg, [REDACTED] mg, and [REDACTED] mg, each tested in parallel, against placebo at a 2-sided significance level of 0.0167, relative to the primary endpoint (mean change in HbA1c from baseline to [REDACTED] weeks), under the following assumptions:

- use of a 2-sample t-test to compare treatment means using HbA1c data for the treatment regimen estimand
- 0.6% greater mean reduction in HbA1c from baseline for [REDACTED] mg, [REDACTED] mg, and [REDACTED] mg of orforglipron compared with placebo, and
- common SD of 1.2%.

These assumptions are considered to ensure the power for the primary endpoint for both the efficacy and treatment regimen estimands.

The power calculations were performed with nQuery version 9.1.0.0.

6. Supporting Documentation

6.1. Appendix 1: Demographic and Baseline Characteristics

Demographics and other baseline characteristics will be summarized by treatment group for randomly assigned participants. The listing of basic demographic characteristics (that is, age, sex, ethnicity, race, country, baseline body weight, and so on) for randomly assigned participants will be provided.

Continuous variables will be summarized using descriptive statistics; categorical variables will be summarized using frequency counts and percentages.

[Table GZGT.6.1](#) describes the specific variables and how they will be summarized. Additional variables may also be included in the final study report.

Table GZGT.6.1. Demographics and Baseline Characteristics

Variable	Quantitative Summary	Categorical Summary
Age ^a	Yes	<65, ≥65 years <75, ≥75 years
Sex	No	Male, Female
Ethnicity	No	Hispanic/Latino, Non-Hispanic/Non-Latino
Race	No	American Indian/Alaska Native, Asian, Black/African American, Native Hawaiian or other Pacific Islander, White, or Multiple
Country	No	By each country
Geographic Region	No	US, Latin America, East Asian, India
Height (cm)	Yes	
Baseline tobacco use	No	Never, Current, Former
Baseline waist circumference (cm)	Yes	
Baseline body weight (kg)	Yes	
Baseline BMI (kg/m ²)	Yes	<30, ≥30 and <35, ≥35 kg/m ²
Duration of T2D (years)	Yes	≤5, >5 and ≤10, >10 years
Baseline systolic blood pressure (mmHg)	Yes	
Baseline diastolic blood pressure (mmHg)	Yes	
Baseline pulse rate (beats/min)	Yes	
Baseline eGFR (CKD-EPI) (mL/min/1.73m ²) ^b	Yes	≥15 to <30, ≥30 to <45, ≥45 to <60, ≥60 to <90, ≥90 mL/min/1.73 m ²
Baseline UACR (g/kg)	Yes	<30, ≥30 and ≤300, >300 g/kg
Baseline fasting insulin (pmol/L)	Yes	
Baseline HbA1c (%)	Yes	≤8.0%, >8.0%
Baseline HbA1c (mmol/mol)	Yes	
Baseline fasting serum glucose (mg/dL)	Yes	
Baseline fasting serum glucose (mmol/L)	Yes	
Baseline fasting lipid parameters (SI)	Yes	
Baseline fasting lipid parameters (CN)	Yes	

Abbreviations: BMI = body mass index; CKD-EPI = Chronic Kidney Disease Epidemiology; eCRF = electronic case report form; eGFR = estimated glomerular filtration rate; HbA1c = hemoglobin A1c; T2D = type 2 diabetes; UACR = urinary albumin to creatinine ratio.

^a Age in years will be calculated as length of the time interval from the imputed date of birth (01 July in the year of birth collected in the eCRF) to the informed consent date.

^b eGFR will be calculated by Chronic Kidney Disease-Epidemiology cystatin-c equation (Inker et al. 2012).

6.2. Appendix 2: Historical Illnesses and Preexisting Conditions

The count and percentages of participants with historical illnesses and preexisting conditions will be summarized by treatment group using the MedDRA PTs nested within SOC. The SOC will be in alphabetical order. Events will be ordered by decreasing frequency. Conditions (that is, PTs) will be ordered by decreasing frequency within SOC. This will be summarized for all randomized participants. Historical illnesses are illnesses that end prior to informed consent and

preexisting conditions are conditions that are still ongoing at inform consent. No statistical comparisons between treatment groups will be performed.

6.3. Appendix 3: Treatment Compliance

Treatment compliance is defined as taking at least 75% of scheduled study intervention during the treatment period. Compliance will be calculated by

$$([\text{total number of doses dispensed} - \text{total number of doses returned}] / \text{total number of doses expected to be administered}) \times 100\%.$$

If the data warrant, frequency counts and percentages of participants who are non-compliant to treatment and/or have dose interruptions/modifications will be summarized by treatment group for safety participants. Listings of such participants may also be provided.

6.4. Appendix 4: Prior/Concomitant Medications

Medications that start before or at the last date of the treatment period or follow-up period, and are ongoing or ended during the treatment period or follow-up period, will be classified as concomitant medication. Medications that start and end before first dose date of study intervention will be classified as prior therapy.

Baseline is defined as the corresponding medication taken on the day of the first dose of study intervention. If there are no doses of study intervention, the randomization date will be used instead of the first dose date.

Concomitant medication will be summarized by PTs, by treatment group and by decreasing frequency for randomized participants. Additionally, the following prior/concomitant medications of interest will be summarized by treatment group:

- prior antihyperglycemic use prior to study entry
- baseline antihypertensive therapy, by type
- baseline lipid lowering therapy, by type
- utilization of the following medication during treatment period
 - antihypertensive therapy
 - lipid lowering therapy
- initiation of additional antihyperglycemic medication during treatment period
- initiation of following medications during treatment period and safety follow-up period
 - antidiarrheal medication
 - antiemetic medication, and
 - constipation medication.

6.5. Appendix 5: Important Protocol Deviations

Important protocol deviations are defined in the Trial Issues Management Plan. A listing and a summary of important protocol deviations by treatment group will be provided at the end of the study.

6.6. Appendix 6: Statistical Analysis for Japan

For the Pharmaceuticals and Medical Devices Agency (PMDA), there are 2 sets of analyses.

The first set of analyses is a subset of the analyses described in the main part of SAP GZGT, which will be reproduced for participants from all investigative sites in Japan. This set of analyses aims to evaluate the efficacy, health outcomes, and safety at Week 40 for participants from all investigative sites in Japan. If there is not a sufficient number of participants in this subpopulation, summary statistics will be provided. The analyses to be included will be documented in a separate list of analyses which should include disposition, demographics, and selected efficacy and safety endpoints.

The other set of analyses will assess the safety and tolerability of orforglipron administered QD during the 52-week treatment period for participants from all investigative sites in Japan, since the Protocol GZGT addendum extends the study intervention for 12 weeks for participants from Japan sites. Additional details for the 52-week analysis are in Section 6.6.1, Section 6.6.2, and Section 6.6.3.

6.6.1. Objectives and Endpoints

Objectives	Endpoints
Primary	
To assess safety and tolerability of orforglipron (■, ■■■, or ■■■■mg) administered QD during 52-week treatment as monotherapy in the treatment of T2D in participants from Japan	<ul style="list-style-type: none"> Incidence of TEAEs
Secondary	
To assess safety and tolerability of orforglipron (■ mg, ■■■ mg, or ■■■■ mg) administered QD during 52-week treatment as monotherapy in the treatment of T2D in participants from Japan	<p>Summary of safety data, including the number and incidence of</p> <ul style="list-style-type: none"> SAEs discontinuations from study intervention or study due to AEs AEs of special interest
To assess the efficacy of orforglipron (■ mg, ■■■ mg, or ■■■■ mg) administered QD during 52-week treatment as monotherapy in the treatment of T2D in participants from Japan	<ul style="list-style-type: none"> Change from baseline in HbA1c at Week 52 Proportion of participants with an HbA1c target value of <7.0% (53 mmol/mol) at Week 52 Proportion of participants with an HbA1c target value of ≤6.5% (48 mmol/mol) at Week 52 Proportion of participants with an HbA1c target value of <5.7% (39 mmol/mol) at Week 52 Change from baseline in fasting serum glucose (central laboratory) at Week 52 Change from baseline in body weight at Week 52 Percentage change from baseline in body weight at Week 52 Proportion of participants who achieved weight loss of ≥5%, ≥10%, and ≥15% at Week 52 Change from baseline in waist circumference at Week 52 Change from baseline in BMI at Week 52
Tertiary	
To assess the efficacy of orforglipron (■ mg, ■■■ mg, or ■■■■ mg) administered QD during 52-week treatment as monotherapy in the treatment of T2D in participants from Japan.	<ul style="list-style-type: none"> Change from baseline in blood pressure at Week 52 <ul style="list-style-type: none"> systolic blood pressure diastolic blood pressure Percentage change from baseline in lipid parameters at Week 52 <ul style="list-style-type: none"> total cholesterol HDL-cholesterol LDL-cholesterol VLDL-cholesterol non-HDL cholesterol triglycerides Percentage change from baseline in liver enzyme at Week 52

Objectives	Endpoints
	<ul style="list-style-type: none"> ○ ALT ○ AST

Abbreviations: AE = adverse event; ALT = alanine aminotransferase; AST = aspartate aminotransferase; HbA1c = hemoglobin A1c; HDL = high-density lipoprotein; LDL = low-density lipoprotein; QD = once daily; SAE = serious adverse event; T2D = type 2 diabetes; TEAE = treatment-emergent adverse event; VLDL = very low-density lipoprotein.

6.6.2. Analysis Sets

The following participant analysis sets and data point sets are defined for participants from Japan sites only, where data from the 52-week treatment period will be included.

Participant Analysis Set	Description
Entered participants – Japan	All participants from Japan sites who sign informed consent ^a .
Randomized participants – Japan	All participants from Japan sites who are randomly assigned a study intervention.
Safety participants – Japan	All participants from Japan sites who are randomly assigned a study intervention and who take at least 1 dose of study intervention.

^a Refers to the informed consent for the study.

Data Points Sets	Description
Treatment regimen estimand data points set - Japan	All data points obtained during the treatment period (Weeks 0- 52), defined as at or after baseline and up to the last visit within the treatment period, regardless of study intervention discontinuation or initiation of additional antihyperglycemic medications ^a .
Safety data points set – Japan	All data points obtained during the treatment period (Weeks 0-52) and the follow-up period defined as after baseline and up to the date of study withdrawal/study completion including the follow-up period regardless of study intervention discontinuation or initiation of additional antihyperglycemic medications ^a .
Efficacy estimand data points set – Japan	All data points obtained during the treatment period (Weeks 0-52) defined as after baseline and up to the earliest date of discontinuation of study intervention or initiation of additional antihyperglycemic medications ^a .

^a “Additional antihyperglycemic medications” refers to any antihyperglycemic therapy that is used for more than 2 weeks (14 days).

6.6.3. Statistical Analyses

Participants will be analyzed according to the treatment group to which they were randomly assigned.

Unless otherwise specified, safety and tolerability assessments will be guided by an estimand comparing safety of orforglipron treatment groups with placebo for the entire study period (the treatment period [Weeks 0-52] plus safety follow-up) irrespective of adherence to study intervention or initiation of additional anti-hyperglycemic medications for all study population.

Thus, safety and tolerability assessments will use the safety participants analysis set – Japan and the safety data points set – Japan.

Efficacy assessments will be guided by the treatment regimen estimand using the randomized participants analysis set – Japan, the treatment regimen estimand data points set – Japan, and by the efficacy estimand using the randomized participants analysis set – Japan and the efficacy estimand data points set – Japan.

6.6.3.1. Primary Analysis

The primary objective is to assess safety and tolerability of orforglipron (■ mg, ■■ mg, or ■■ mg) administered QD during 52-week treatment as monotherapy in the treatment of T2D in participants from all investigative sites in Japan measured by the incidence of TEAEs.

Counts and proportions of participants experiencing TEAEs will be reported for each treatment group, and Fisher's exact test or Pearson's chi-squared test, if applicable, will be used to compare the treatment groups. Risk difference and the 95% CI will be provided.

6.6.3.2. Secondary Analyses

A secondary objective is to assess safety and tolerability of orforglipron (■ mg, ■■ mg, or ■■ mg) administered QD during 52-week treatment as monotherapy in the treatment of T2D in Japanese participants as measured by the incidence of the following:

- SAEs
- discontinuations from study intervention or study due to AEs, and
- AEs of special interest.

Counts and proportions of participants experiencing these events will be reported for each treatment group, and Fisher's exact test or Pearson's chi-squared test, if applicable, will be used to compare the treatment groups. Risk difference and the 95% CI will be provided.

The following secondary efficacy endpoints will be guided by the treatment regimen and efficacy estimand:

- change from baseline in HbA1c at Week 52
- incidence of participants (proportion at the population-level) achieving an HbA1c target value of <7.0% (53 mmol/mol) at Week 52
- incidence of participants (proportion at the population-level) achieving an HbA1c target value of ≤6.5% (48 mmol/mol) at Week 52
- change from baseline in fasting serum glucose at Week 52
- change from baseline in body weight at Week 52, and
- percentage change from baseline in body weight at Week 52.

The following secondary efficacy endpoints will be guided by the efficacy estimand:

- incidence of participants (proportion at the population-level) achieving an HbA1c target value of <5.7% (39 mmol/mol) at Week 52
- incidence of participants (proportion at the population-level) achieving weight loss of ≥5%, ≥10%, and ≥15% from baseline at Week 52
- change from baseline in waist circumference at Week 52, and

- change from baseline in BMI at Week 52.

The analytical approaches for the secondary efficacy endpoints are specified for the treatment regimen estimand in Section 4.1.2.1 and the efficacy estimand in Section 4.1.2.2.

6.6.3.3. Tertiary Analyses

The following tertiary efficacy endpoints will be guided by the treatment regimen and efficacy estimand:

- change from baseline at Week 52 in
 - systolic blood pressure
 - diastolic blood pressure
- percentage change from baseline in lipid parameters at Week 52 (log-transformed data)
 - total cholesterol
 - HDL-cholesterol
 - LDL-cholesterol
 - VLDL-cholesterol
 - non-HDL cholesterol
 - triglycerides
- percentage change from baseline in liver enzyme at Week 52 (log transformed data)
 - ALT, and
 - AST.

The analytical approaches for the tertiary efficacy endpoints are specified for the treatment regimen estimand in Section 4.1.2.1 and the efficacy estimand in Section 4.1.2.2.

6.7. Appendix 7: Statistical Analysis for China

In accordance with an authority requirement in the People's Republic of China, telephone visits between Visit 9 and Visit 10, and Visit 10 and Visit 11, were added at all investigative sites in China to assess information related to conditions/symptoms and/or concomitant medications.

Analyses will be performed for the following subpopulations:

- participants enrolled in East Asian Countries (China and Japan), and
- participants enrolled in China.

The analysis methods for these subgroups will be similar to those described for the main part of this SAP, which excludes data in participants from Japan sites who continued beyond the Week 40 visit. If there is not a sufficient number of participants in the subpopulation, summary statistics will be provided.

The analyses to be included will be documented in a separate list of analyses which should include disposition, demographics, and selected efficacy and safety endpoints.

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