

## STATISTICAL ANALYSIS PLAN

### STUDY TITLE:

**An Open-label, Single-center, Three-part Study in Healthy Subjects to Investigate the Effect of Givinostat on the Pharmacokinetics of Midazolam and Dabigatran, the Effect of Clarithromycin on the Pharmacokinetics of Givinostat and the Pharmacokinetics of Single and Multiple Doses of Givinostat.**

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## 2.2. List of Abbreviations

$\alpha$	Type I Error (Significance Level)
Ab	Antibody
ADaM	Analysis Data Model
ADaMIG	Analysis Data Model Implementation Guide
ADPC	Analysis Data Pharmacokinetic Concentrations
ADPP	Analysis Data Pharmacokinetic Parameters
ADSL	Subject-Level Analysis
AE	Adverse Event
ALP	Alkaline Phosphatase
ALT	Alanine Aminotransferase
$A_{\text{mean}}$	Arithmetic Mean
ANOVA	Analysis of Variance
aPTT	Activated Partial Thromboplastin Time
AST	Aspartate Aminotransferase
ATC	Anatomic Therapeutic Chemical
AUC	Area Under the Concentration <i>Versus</i> Time Curve
$AUC_{0-t}$	AUC from Time Zero to Last Sampling Time with Quantifiable Concentrations
$AUC_{0-\infty}$	AUC from Time Zero to Infinity
$AUC_{0-t,ss}$	AUC Corresponding to the Dosing Interval, at Steady State
$AURC_{0-t}$	Area Under the Urinary Excretion Rate Curve from Time 0 to the Last Rate
$\%AUC_{\text{extrap}}$	Residual Area or Percentage of Extrapolated Part of $AUC_{0-\infty}$
BDS	Basic Data Structure
BLQ	Below the Limit of Quantification
BMI	Body Mass Index
bpm	Beats Per Minute
$C_0$	Pre-dose Concentration
CI	Confidence Interval
Cl	Total Body Clearance
CK	Creatine Kinase
Cl/F	Total Body Clearance Affected by the Bioavailability Factor
$Cl_R$	Renal Clearance
$Cl_R/F$	Renal Clearance Affected by the Bioavailability Factor
$C_{\text{last}}$	Last Quantifiable Concentration
$C_{\text{max}}$	Maximum Observed Concentration
$C_{\text{max},ss}$	Maximum Observed Plasma Concentration at Steady State
$Cr_{CL}$	Creatinine Clearance
CSP	Clinical Study Protocol
CSR	Clinical Study Report
CV%	Coefficient of Variation
CYP	Cytochrome P450
DBP	Diastolic Blood Pressure
DDI	Drug-Drug Interaction
DM	Demographics
ECG	Electrocardiogram
F	Bioavailability Factor
FDA	Food and Drug Administration
GGT	Gamma-Glutamyltransferase
GLP	Good Laboratory Practices

$G_{\text{mean}}$	Geometric Mean
GMR	Geometric Least Square Means Ratio
GCV%	Geometric Coefficient of Variation
GSD	Geometric Standard Deviation
$H_0$	Null Hypothesis
$H_1$	Alternative Hypothesis
HBsAg	Hepatitis B Surface Antigen
hCG	Human Chorionic Gonadotropin
HCVAb	Hepatitis C Virus Antibodies
HIV	Human Immunodeficiency Virus
ICH	International Conference on Harmonization
INR	International Normalized Ratio
ISCV%	Intra-Subject Coefficient of Variation
IV	Intravenous
LC-MS/MS	Liquid Chromatography with Tandem Mass Spectrometry
LDH	Lactate Dehydrogenase
LLOQ	Lower Limit of Quantification
LSmeans	Least Square Means
MCH	Mean Corpuscular Hemoglobin
MCHC	Mean Corpuscular Hemoglobin Concentration
MCV	Mean Corpuscular Volume
MedDRA	Medical Dictionary for Regulatory Activities
MSE	Mean Square Error
MW	Molecular Weight
NCA	Non-Compartmental Analysis
PC	Pharmacokinetic Concentrations
PCL	Protocol Clarification Letter
P-gp	P-Glycoprotein
PP	Pharmacokinetic Parameters
PT	MedDRA Preferred Term
QTcF	Corrected QT Interval by Fridericia
RBC	Red Blood Cell
RDW-CV	Coefficient Variation of the Red Cell Distribution Width
RSQ	Goodness of Fit for the Terminal Elimination Phase
rtf	Rich Text File
SAE	Serious Adverse Event
SAP	Statistical Analysis Plan
SAR	Serious Adverse Reaction
SARS-CoV-2	Severe Acute Respiratory Syndrome Coronavirus 2
SAS	Statistical Analysis Software
SBP	Systolic Blood Pressure
SD	Standard Deviation
SDTM	Study Data Tabulation Model
SDTMIG	Study Data Tabulation Model Implementation Guide
SOC	MedDRA System Organ Class
SOP	Standard Operating Procedure
TEAE	Treatment-Emergent Adverse Events
TFLs	Tables, Figures and Listings
$T_{\text{max}}$	Time to Maximum Observed Concentration
$T_{\text{max,ss}}$	Time to Maximum Observed Concentration at Steady State



TOST	Two One-Sided T-Tests
TSH	Thyroid-Stimulating Hormone
$t_0$	Time of Dosing ( $t=0h$ )
$t_{1/2}$	Apparent Terminal Elimination Half-Life
ULOQ	Upper Limit of Quantification
$V_D$	Apparent Volume of Distribution
$V_D/F$	Apparent Volume of Distribution Affected by the Bioavailability Factor (F)
WBC	White Blood Cell
WHO	World Health Organization
$\lambda_z$	Apparent Terminal Elimination Rate Constant
$\tau$	Dosing Interval

### 3. INTRODUCTION

This Statistical Analysis Plan (SAP) details the statistical methodology to be used in analysing study data and outlines the statistical programming specifications for the Tables, Figures and Listings (TFLs). This document describes the variables and populations, anticipated data transformation and manipulations and other details of the analysis not provided in the Clinical Study Protocol (CSP).

The described analyses are based on the final CSP version 1.0, dated 29NOV2021 [1] and a Protocol Clarification Letter (PCL) dated 25FEB2022. This SAP was finalized prior to data base lock and conduct of the statistical analyses. For each part of the study at the time of the Blind Review Process, a Data Blind Review Process Minute will be prepared and attached to this document (Annex 5).

Additional pharmacokinetic and statistical analysis may be performed to supplement the planned analyses described in this SAP. These supplemental analyses will be identified and presented in the Clinical Study Report (CSR).

### 4. STUDY OBJECTIVES

#### 4.1. Primary

1. To evaluate the plasma and urine pharmacokinetics of givinostat and its metabolites following single and multiple oral doses of givinostat (Part 3).

#### 4.2. Secondary

1. To assess the safety and tolerability of single and multiple doses of givinostat (Part 3).

### 5. INVESTIGATIONAL PLAN

#### 5.1. Study Design

This is a phase I, open-label, 3-part, fixed-sequence, non-randomized study in healthy male and female subjects.

Study parts may be conducted concomitantly.

##### 5.1.1. Rational for Study Design

The rational for study design detailed in the CSP [1].

In Part 3, the pharmacokinetics of givinostat and its metabolites in plasma and urine following single- and multiple-doses of givinostat will be investigated.



## 5.2. Study Plan

### 5.2.1. Part 3 – Pharmacokinetics of Givinostat and its Metabolites Following Single and Multiple Oral Doses of Givinostat

Subjects will be confined in PPD from Day -1 to Day 17.

On Day 1 and Day 13, givinostat 50 mg as oral suspension will be administered as a single dose, in the morning, following an overnight fasting of at least 8 hours and subjects will remain fasted until at least 4 hours post dose. No fluids will be allowed from 1 hour before dosing until 2 hours post dose. Water will be provided ad libitum at all other times.

From Day 5 to Day 12, subjects will receive givinostat 50 mg as oral suspension, twice a day, in the morning and in the evening. On Days 1 and 13, givinostat will be administered with the subjects in a semi-recumbent position and will remain semi-recumbent until at least 4 hours post-dose.

The following assessments will be performed:

- Blood collection for pharmacokinetic analysis on Days 1 to 5 and 9 to 17.
- Urine collection for pharmacokinetic analysis on Days 1 to 5 and 13 to 17 (Baseline urine collection is on day -1).
- Vital signs measurements (BP, PR and RR) on Days 1 and 5 to 13.
- 12-lead ECG on Days 1 and 4 to 13.
- Blood collection for laboratory tests (hematology and biochemistry on Day 9 and hematology on Days 6 and 13).

Subjects will be discharged from PPD on the morning of Day 17 if allowed by the investigator based on their medical condition. The following procedures will be performed:

- Physical examination.
- Collection of body weight.
- Vital signs measurements (BP, PR, RR and body temperature).
- 12-lead ECG.
- Safety laboratory assessments (hematology, biochemistry, coagulation and urinalysis).
- Serum pregnancy test for all female subjects.

Subjects will return to PPD 10 to 14 days after the end of study to undergo additional assessments as required as per protocol.

The total duration of Part 3 for each subject will be up to approximately 8 weeks from Screening to Follow-up visit, divided as follows:

- Screening: up to 21 days
- Treatment Period: Days 1 to 17
- Safety follow-up visit: 12±2 days

## 5.3. Number of Participants and Sample Size Estimation

To ensure enough subjects with evaluable data complete each study part, a total of fifty-four (54) healthy male and female subjects are planned to be enrolled as follows:

- Part 1: twenty-six (26) subjects
- Part 2: twenty (20) subjects

## 5.2. Study Plan

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- Part 1: twenty-six (26) subjects
- Part 2: twenty (20) subjects



- Part 3: eight (8) subjects

Each subject will participate in one study part only.

#### 5.3.1. **Part 3**

A sample size of 8 subjects will be enrolled. The sample size was determined by practical considerations and not based on statistical power calculations.

#### 5.4. **Randomization**

This is a fixed sequence study. Randomization is not applicable.

#### 5.5. **Blinding**

The study will be conducted as open label. Blinding procedures are not applicable.

### 6. **STUDY ASSESSMENTS**

For each study part, a summary of procedures for study assessments are presented in Section 2 of the CSP (Study Flow-Chart and Study Design Diagram) [1].

#### 6.1. **Safety Assessments**

Subjects' safety will be monitored during the study.

Safety assessments will include pre-study medical history, physical examination, vital signs, 12-lead ECG, clinical laboratory tests and adverse event (AE) monitoring. Additional safety measurements may be performed at the discretion of the investigator for reasons related to subject safety.

Medications will be mentioned according to the World Health Organization (WHO) Anatomical Therapeutic Chemical (ATC) classification system.

##### 6.1.1. **Medical History**

Medical history will cover all relevant past or present information related to subject's health at the time of informed consent signature.

Medical history at screening will include past or present relevant cardiovascular, respiratory, renal, genitourinary, gastrointestinal, hepatic, hematological, immunological, endocrine, dermatological, musculoskeletal, neurological, psychiatric, drug and surgical history, or any other diseases or disorders.

Medical history will be referred in accordance with the Medical Dictionary for Regulatory Activities (MedDRA), version 24.1 or higher.

Adverse events related to medical history will be identified.

#### 6.1.2. Physical Examination

Physical examination at screening and end-of-study will include: general appearance; skin; head and neck; thorax and abdomen; pulmonary auscultation; cardiac auscultation; abdomen palpation; limbs; brief neurological examination.

#### 6.1.3. Weight, Height and Body Mass Index

The body height and weight values as well as the body mass index (BMI) will be recorded in the electronic case report form (eCRF) and will be determined at Screening. Only body weight will be determined also at admission and end-of-study.

The subjects' body weight will preferably be measured using the same weighing scale for all subjects and throughout the study. The weighing scale should have a precision of at least 0.5 kg.

#### 6.1.4. Vital Signs

Vital signs will include:

- Systolic blood pressure (SBP).
- Diastolic blood pressure (DBP).
- Pulse rate (PR).
- Respiratory rate (RR).
- Body temperature.

#### 6.1.5. 12 Lead Electrocardiogram

A 12-lead ECG will be performed preferably before the blood collection. The corrected QT interval by Fridericia (QTcF) will be analyzed.

#### 6.1.6. Laboratory Safety Tests

The following laboratory parameters will be tested:

- Hematology:
  - Red blood cell (RBC) count.
  - White blood cell (WBC) count.
  - WBC differential count:
    - Neutrophils.
    - Eosinophils.
    - Basophils.
    - Lymphocytes.
    - Monocytes.
  - Hemoglobin.
  - Mean corpuscular volume (MCV).
  - Mean corpuscular hemoglobin (MCH).
  - Mean corpuscular hemoglobin concentration (MCHC).
  - Coefficient variation of the red cell distribution width (RDW-CV).
  - Hematocrit.
  - Platelet count.
  - Mean platelet volume.



- Coagulation:
  - Prothrombin rate.
  - Prothrombin time.
  - Prothrombin time – international normalized ratio (INR).
  - Activated partial thromboplastin time (aPTT).
- General biochemistry:
  - Total bilirubin.
  - Direct bilirubin.
  - Indirect bilirubin.
  - Alkaline phosphatase (ALP).
  - Amylase.
  - Aspartate aminotransferase (AST).
  - Alanine aminotransferase (ALT).
  - Lactate dehydrogenase (LDH).
  - Cystatin C.
  - C-reactive protein.
  - Gamma-glutamyltransferase (GGT).
  - Creatinine kinase (CK).
  - Total protein.
  - Albumin.
  - Uric acid.
  - Triglycerides.
  - Total cholesterol.
  - Low-density lipoprotein-cholesterol (LDL-C).
  - High-density lipoprotein-cholesterol (HDL-C).
  - Sodium.
  - Potassium.
  - Chloride.
  - Calcium.
  - Magnesium.
  - Glucose.
  - Creatinine.
  - Urea.
  - Thyroid-stimulating hormone (TSH).
  - Estimated creatinine clearance (Cr<sub>CL</sub>).
- Viral Serology:
  - Human Immunodeficiency Virus (HIV):
    - HIV-1 (anti-HIV-1Ab).
    - HIV-2 (anti-HIV-2Ab).
  - Hepatitis B (HBsAg)
  - Hepatitis C (anti-HCVAb).
- Beta-human chorionic gonadotropin (beta-hCG) pregnancy tests:
  - Serum.
  - Urine.
- Urinalysis:
  - pH.

- Specific gravity.
- Protein.
- Hemoglobin.
- Glucose.
- Ketones.
- Bilirubin.
- Nitrites.
- Urobilinogen.
- Microscopy.
- Drugs-of-abuse urine test:
  - Cannabinoids.
  - Opiates.
  - Cocaine.
  - Amphetamines.
  - Benzodiazepines.
- Ethanol urine test.
- Cotinine urine test.
- Follicle-stimulating hormone (FSH) test.
- Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-CoV-2) diagnostic tests.

#### 6.1.7. Adverse Events

The occurrence of clinical AEs will be monitored throughout the study. Clinically significant abnormalities in laboratory safety tests, vital signs and physical examination will be reported as AEs.

Treatment-emergent AEs (TEAEs) are defined as AEs not present prior to first administration of investigational product, or AEs present before first administration of investigational product that worsen after the subject receives the first dose of investigational product. TEAEs that occur after administration of investigational product during the washout of a given period will be assigned to the treatment administered in that period.

The following information will be used for the description of the AEs:

- Reported Term.
- Medical Dictionary for Regulatory Activities (MedDRA) System Organ Class (SOC) coding.
- MedDRA Preferred Term (PT) coding.
- Start date and time.
- End date and time.
- Seriousness.
- Severity:
  - Mild.
  - Moderate.
  - Severe.
- Relationship (causality) for each treatment administered, per part.
  - Reasonably Possible.
  - Not Reasonably Possible.
  - Unknown.



- Action taken:
  - Dose Increased.
  - Dose Not Changed.
  - Dose Rate Reduced.
  - Dose Reduced.
  - Drug Interrupted.
  - Drug Withdrawn.
  - Not Applicable.
  - Unknown.
- Concomitant medication.
- Outcome:
  - Fatal.
  - Not Recovered / Not Resolved.
  - Recovered / Resolved.
  - Recovered / Resolved with Sequelae.
  - Recovering / Resolving.
  - Unknown.
- Most recent study treatment taken.
- Last dosing date.

#### 6.1.8. Previous and Concomitant Medications

A previous medication is any medication for which the end date is prior to first dosing.  
A concomitant medication is any medication ongoing or initiated after first dosing.

For all study parts, the use of any medications including Over-the-Counter (OTC) products (including herbal medicines such as St John's Wort, homeopathic preparations, vitamins, and minerals) is forbidden from 28 days or within 5 half-lives of the medicinal product, whichever is longer, prior to admission up to last sample for pharmacokinetics assessment, except for medications for the treatment of AEs.

Any medication (previous or concomitant) taken within the period of medication restriction must be recorded in the eCRF. If concomitant medication is ongoing at the follow-up visit, no end date will be provided in the eCRF.

For each study part, previous and concomitant medications used by study participants during the study and their judged impact on the pharmacokinetic assessment will be listed in the respective Data Blind Review Meeting/ Process Minute ([Annex 5](#)).

Concomitant medications will be coded according to the WHO ATC classification system.

## 6.2. Pharmacokinetic Assessments

### 6.2.1. Blood Sampling for Pharmacokinetic Assessments

In each study part, while subject is confined at the clinical research unit, blood samples will be taken preferably via an indwelling cannula placed in a vein of an upper limb of the subject. During ambulatory visits, blood samples will be taken by direct venipuncture.

The actual time of all pharmacokinetic blood draws will be recorded and reported for all subjects. The pre-dose blood sample will be collected within 30 minutes before dosing. The post-dose blood samples will be collected within  $\pm 3$  minutes from the scheduled sampling time. Greater deviations will be reported as a protocol deviation and its cause will be recorded.

In case blood sampling for pharmacokinetics and other procedures coincide in time, blood draws will have priority unless other procedures are necessary for assuring subject's safety.

PPD will carry out the determination of plasma levels of givinostat and its metabolites (ITF2374, ITF2375, ITF2440 and ITF2563), midazolam, 1-hydroxymidazolam and total and free dabigatran in accordance with the applicable principles of Good Laboratory Practices (GLP), using a previously validated liquid chromatography with tandem mass spectrometry (LC-MS/MS) analytical method.

The planned lower limit of quantification (LLOQ) and upper limit of quantification (ULOQ) for:

- givinostat are 1 and 100 ng/mL.
- givinostat metabolite ITF2374 are 1 and 100 ng/mL.
- givinostat metabolite ITF2375 are 1 and 100 ng/mL.
- givinostat metabolite ITF2440 are 10 and 500 ng/mL.
- givinostat metabolite ITF2563 are 2 and 125 ng/mL.
- midazolam are 100 and 100000 pg/mL.
- 1-hydroxymidazolam are 100 and 50000 pg/mL.
- total dabigatran are 1 and 400 ng/mL.
- free dabigatran are 1 and 400 ng/mL.

LLOQ represents a value lower than or equal to 1/20 of estimated  $C_{max}$  value. Any adjustment to this range will be documented in the study specific Bioanalytical Report, which will supersede the indicated range.

#### 6.2.1.1. Part 3

A total of forty-eight (48) blood samples will be collected as follows:

- Twenty-two (22) blood samples of 6 mL each will be collected in sodium heparin tubes at pre-dose and at 0.5, 1, 1.5, 2, 2.5, 3, 3.5, 4, 5, 6, 8, 10, 12, 15, 24, 36, 48, 60, 72, 84 and 96 hours after the administration of givinostat, on Days 1 and 13, for the determination of givinostat and its metabolites (ITF2374, ITF2375, ITF2440 and ITF2563) plasma concentrations.
- Four (4) blood samples of 6 mL each will be collected in sodium heparin tubes before the morning dose of givinostat on Days 9, 10, 11 and 12, for the determination of pre-dose ( $C_{trough}$ ) plasma concentration of givinostat and metabolites.

#### 6.2.2. Urine Sampling for Pharmacokinetic Assessments

##### 6.2.2.1. Part 3

Urine will be collected on Day -1 (baseline) and in the following intervals, on Days 1 and 13, for the determination of the amounts of givinostat and its metabolites (ITF2374, ITF2375, ITF2440 and ITF2563) excreted in urine: 0–12, 12–24, 24–36, 36–48, 48–72 and 72–96 post-dose.



The date and actual time of urine collection start and completion will be recorded and reported for all subjects.

PPD will carry out the determination of urine levels of givinostat and its metabolites (ITF2374, ITF2375, ITF2440 and ITF2563) in accordance with the applicable principles of GLP using a previously validated liquid chromatography with tandem mass spectrometry (LC-MS/MS) analytical method.

The planned lower limit of quantification (LLOQ) and upper limit of quantification (ULOQ) for:

- givinostat are 10 and 2500 ng/mL.
- givinostat metabolite ITF2374 are 5 and 2500 ng/mL.
- givinostat metabolite ITF2375 are 4 and 800 ng/mL.
- givinostat metabolite ITF2440 are 150 and 25000 ng/mL.
- givinostat metabolite ITF2563 are 10 and 5000 ng/mL.

### 6.2.3. Plasma Pharmacokinetic Parameters

The following pharmacokinetic parameters will be derived by standard non-compartmental analysis (NCA) methods from the single-dose plasma concentration *versus* time profiles, for:

- Part 3 (Day 1):
  - givinostat.
  - givinostat metabolites (ITF2374, ITF2375, ITF2440 and ITF2563).

Parameter	Description
$C_{max}$	Maximum observed concentration, directly obtained from the observed concentration <i>versus</i> time profile.
$T_{max}$	Time of occurrence of maximum observed concentration.
$AUC_{0-t}^1$	Area under the concentration <i>versus</i> time curve (AUC) from time of dosing ( $t=0h$ ) to the time of the last measurable concentration ( $t_{last}$ ), calculated by the linear-up/log-down trapezoidal method.
$AUC_{0-\infty}$	Total AUC extrapolated to infinity, calculated as $AUC_{0-t} + \frac{C_{last}}{\lambda_z}$ , where $C_{last}$ is the last measurable concentration and $\lambda_z$ is the apparent terminal elimination rate constant.
%AUC <sub>extrap</sub>	Percentage of $AUC_{0-\infty}$ due to extrapolation from the time of the last measurable concentration ( $t_{last}$ ) to infinity, i.e., residual area, calculated as $100 \cdot \frac{AUC_{0-\infty} - AUC_{0-t}}{AUC_{0-\infty}}$ .
$\lambda_z$	Apparent first order elimination rate constant associated with the terminal (log-linear) portion of the concentration <i>versus</i> time curve. The parameter is estimated by linear least square regression analysis using the last three (or more) non-zero concentrations.
$t_{1/2}$	Apparent terminal elimination half-life, calculated as $\frac{\ln(2)}{\lambda_z}$ .

<sup>1</sup> In Part 3, for givinostat and its metabolites the last sampling time corresponds to 96 hours.

The upper and the lower timepoints and the number of timepoints used for  $\lambda_z$  estimation, as well as the goodness of fit for the terminal elimination phase (RSQ), will be reported.  
No values of  $\lambda_z$ ,  $AUC_{0-\infty}$ ,  $\%AUC_{extrap}$  and  $t_{1/2}$  will be reported for cases where  $\lambda_z$  cannot be reliably determined.

The following pharmacokinetic parameters will be derived by standard NCA methods from the multiple-dose plasma concentration *versus* time profiles, for:

- Part 3 (Day 13):
  - givinostat.
  - givinostat metabolites (ITF2374, ITF2375, ITF2440 and ITF2563).

Parameter	Description
$C_{max,ss}$	Maximum observed plasma concentration at steady state, directly obtained from the observed concentration <i>versus</i> time profile.
$T_{max,ss}$	Time of occurrence of maximum observed plasma concentration at steady state.
$AUC_{0-\tau,ss}$	Area under the plasma concentration <i>versus</i> time curve corresponding to dosing interval, i.e., 12 hours, after steady state has been reached. This parameter is calculated by the linear-up/log-down trapezoidal method.
$AUC_{0-t}$	AUC from time of dosing ( $t=0h$ ) to the time of the last measurable concentration ( $t_{last}$ ), calculated by the linear-up/log-down trapezoidal method.
$AUC_{0-\infty}$	Total AUC extrapolated to infinity, calculated as $AUC_{0-t} + \frac{C_{last}}{\lambda_z}$ , where $C_{last}$ is the last measurable concentration and $\lambda_z$ is the apparent terminal elimination rate constant.
$\lambda_z$	Apparent first order elimination rate constant associated with the terminal (log-linear) portion of the concentration <i>versus</i> time curve. The parameter is estimated by linear least square regression analysis using the last three (or more) non-zero concentrations.
$t_{1/2}$	Apparent terminal elimination half-life, calculated as $\frac{\ln(2)}{\lambda_z}$ .
$CL_{ss}/F$	Apparent total body clearance affected by the bioavailability factor (F), calculated as $\frac{Dose}{AUC_{0-\tau,ss}}$ .
$V_D/F$	Apparent volume of distribution affected by F, calculated as $\frac{Dose}{\lambda_z \cdot AUC_{0-\tau,ss}}$ .

The upper and the lower timepoints and the number of timepoints used for  $\lambda_z$  estimation, as well as the RSQ, will be reported.

No values of  $\lambda_z$ ,  $AUC_{0-\infty}$ ,  $\%AUC_{extrap}$  and  $t_{1/2}$  will be reported for cases where  $\lambda_z$  cannot be reliably determined.

For each study part, pharmacokinetic parameters will be estimated from the plasma concentration *versus* time profiles for all subjects in each pharmacokinetic analysis population.

Actual sampling times will be used for the pharmacokinetic analysis. All calculations will be performed using raw data.



In estimating the pharmacokinetic parameters, concentrations below the LLOQ before  $T_{max}$  will be set to zero. After  $T_{max}$ , concentrations below the LLOQ will be considered as missing. However, in case of two or more consecutive concentrations below the LLOQ, the first value will be replaced by  $\frac{1}{2}$  of the LLOQ value and the next values will be considered as missing.

Additional pharmacokinetic parameters may be calculated for the plasma concentration *versus* time profiles, if considered appropriate and justified at the time of the pharmacokinetic analysis.

#### 6.2.4. Urinary Pharmacokinetic Parameters

The following pharmacokinetic parameters will be derived by standard NCA methods from the urine excretion profiles, for the first dose (Day 1, single-dose) and the last dose (Day 13, multiple-dose):

- Part 3:
  - givinostat.
  - givinostat metabolites (ITF2374, ITF2375, ITF2440 and ITF2563).

<i>Parameter</i>	<i>Description</i>
$R_{max}$	Maximum observed urinary excretion rate.
$t_{u_{max}}$	Time of occurrence of maximum urinary excretion rate.
$Amt_{CUM}$	Cumulative amount of drug excreted in urine, calculated as $\sum (Concentration \cdot Volume)$
$AURC_{0-t}$	Area under the urine excretion rate curve from time zero to last measurable observed excretion rate.
$REC\%$	Percentage of drug recovered in urine, calculated for givinostat as $100 \cdot \frac{Amt_{CUM}}{Dose}$ , and for givinostat metabolites as $100 \cdot \frac{AMT_{CUM, Metabolite} / MW_{Metabolite}}{Dose_{Givinostat} / MW_{Givinostat}}$ , where MW represents the molecular weight of the analyte. <i>Note: For givinostat, MW is 421.49 g/mol; for ITF2374, MW is 405.49 g/mol; for ITF2375, MW is 406.47 g/mol; for ITF2440, MW is 257.33 g/mol; and for ITF2563, MW is 178.19 g/mol.</i>
$CL_R/F$	Apparent renal clearance affected by F, calculated as $\frac{Dose}{AURC_{0-t}}$ .

## 7. STUDY ENDPOINTS

### 7.1. Primary

1. Givinostat and its metabolites plasma and urine concentrations and thereof derived pharmacokinetic parameters following single and multiple oral doses of givinostat (Part 3).

### 7.2. Secondary

1. Incidence and severity of AEs; changes in vital signs, physical examination, ECG and clinical laboratory tests following administration of single- and multiple-doses of givinostat (Part 3).

## 8. ANALYSIS POPULATIONS

The analysis populations are defined in accordance with the CSP [1].

For each study part, the subjects to be included in each analysis population will be reported in the respective Data Blind Review Process Minute ([Annex 5](#)).

For each study part, explanation of the reasons for exclusion of subjects from any analysis population will be provided in the respective Data Blind Review Process Minute ([Annex 5](#)) and in the CSR.

The list of subjects who complete each study part will be presented in [Annex 6](#).

### 8.1. Part 3

#### 8.1.1. Part 3 Safety Analysis Population

All subjects who receive at least one dose of the IMP in Part 3 of the study will constitute the Part 3 Safety Analysis Population.

Part 3 safety data analysis will be performed for all subjects in the Part 3 Safety Analysis Population.

#### 8.1.2. Part 3 Pharmacokinetic Analysis Population

##### 8.1.2.1. Part 3 Single-Dose Pharmacokinetic Analysis Population

Part 3 Single-Dose Pharmacokinetic Analysis Population will include all subjects enrolled in Part 3 of the study, who are expected to provide evaluable pharmacokinetic data for the givinostat single-dose administration (Day 1), without deviations affecting pharmacokinetic interpretation.

The following reasons justify the exclusion of the pharmacokinetic data of a subject from the Part 3 single-dose pharmacokinetic analysis population:

- Protocol violation considered to have a potentially relevant effect on the pharmacokinetic results of the study.

*NOTE: These protocol violations will be reported in the Data Blind Review Process Minute ([Annex 5](#)), and their impact will be assessed at the time of pharmacokinetic analysis.*

- Subject experienced vomiting or diarrhea.

*NOTE: Subjects that experienced vomiting or diarrhoea during this study part will be reported in the Data Blind Review Process Minute ([Annex 5](#)), and their exclusion will be assessed at the time of pharmacokinetic analysis.*



- Subject with pre-dose concentration > 5% of the  $C_{max}$  value of the corresponding pharmacokinetic profile.  
*NOTE: This exclusion will be assessed at the time of pharmacokinetic analysis.*
- Subject with lack of any measurable concentrations or only very low plasma concentrations.  
*NOTE: This exclusion will be assessed at the time of pharmacokinetic analysis.*

#### 8.1.2.2. Part 3 Multiple-Dose Pharmacokinetic Analysis Population

Part 3 Multiple-Dose Pharmacokinetic Analysis Population will include all subjects enrolled in Part 3 of the study, who are expected to provide evaluable pharmacokinetic data for the givinostat multiple-dose administration (Day 13), without deviations affecting pharmacokinetic interpretation.

The following reasons justify the exclusion of the pharmacokinetic data of a subject from the Part 3 multiple-dose pharmacokinetic analysis population:

- Protocol violation considered to have a potentially relevant effect on the pharmacokinetic results of the study.  
*NOTE: These protocol violations will be reported in the Data Blind Review Process Minute (Annex 5), and their impact will be assessed at the time of pharmacokinetic analysis.*
- Subject experienced vomiting or diarrhea.  
*NOTE: Subjects that experienced vomiting or diarrhoea during this study part ill be reported in the Data Blind Review Process Minute (Annex 5), and their exclusion will be assessed at the time of pharmacokinetic analysis.*
- Subject with lack of any measurable concentrations or only very low plasma concentrations.  
*NOTE: This exclusion will be assessed at the time of pharmacokinetic analysis.*

## 9. DATA REVIEW / TRANSFORMATION

### 9.1. Data Management

Data handling will be conducted in accordance with the Clinical Data Management section of the CSP and the Data Management Plan developed specifically for this study.

### 9.2. Acceptance of Data

For each study part, TFLs may start being programmed prior to or during the course of the trial. However, the programming of analysis datasets and TFLs will only be concluded and quality-controlled after database soft lock. Only audited data released by the bioanalytical laboratory will be used for programming the final analysis datasets and TFLs.

### 9.3. Data Transformation (CDISC)

Before performing the statistical analysis, all data collected (multiple sources) will be integrated into a common repository, using SAS® version 9.4 or higher.

For standardization and submission purpose, all data will be transformed according to Clinical Data Interchange Standards Consortium (CDISC):



- Study Data Tabulation Model (SDTM) version 1.4 or higher.
- SDTM Implementation Guide (SDTMIG) version 3.2 or higher.
- The following analysis datasets will be generated (Analysis Data Model [ADaM] version 2.1 or higher; ADaM Implementation Guide [ADaMIG], version 1.2 or higher) to support the results and ease the programming activities during the statistical analysis:
  - Subject-Level Analysis Dataset (ADSL).
  - Analysis Data Pharmacokinetic Concentrations (ADPC).
  - Analysis Data Pharmacokinetic Parameters (ADPP).
  - Analysis Data Adverse Events (ADAE).
  - Analysis Data Vital Signs (ADVS).
  - Analysis Data Electrocardiogram Parameters (ADEG).
  - Analysis Data Laboratory Test Results (ADLB).

For the scope of this trial, considering the primary and secondary objectives, six (6) Basic Data Structure (BDS) domains will be generated: ADPC, ADPP, ADAE, ADVS, ADEG and ADLB. These six datasets plus ADSL will be updated for each part. These domains will support the descriptive statistical analyses of the pharmacokinetic concentrations, the pharmacokinetic parameters and safety.

#### ADSL – Subject-Level Analysis Dataset

This analysis domain will contain: general data about the subjects (i.e., age, sex and race), planned and actual allocated treatment analysis, and start and end dates of treatment analysis period. The origin of these data will primarily be the Demographics (DM) and Exposure (EX) SDTM domains. Besides this information, the following variables will be derived:

- *Pharmacokinetic Analysis Set Population Flag 3* – Subjects included in the Part 3 Single-Dose Pharmacokinetic Analysis Population.
- *Pharmacokinetic Analysis Set Population Flag 4* – Subjects included in the Part 3 Multiple-Dose Pharmacokinetic Analysis Population.
- *Completers Population Flag* – Subjects who completed the treatment period.

#### ADPC – Analysis Data Pharmacokinetic Concentrations

This is a BDS dataset that will contain the concentrations for each subject, per treatment analysis, per analyte/metabolite, and per timepoint.

The origin of these data will primarily be the EX and Pharmacokinetic Concentrations (PC) SDTM domains and ADSL ADaM domain.

Besides this information, the following variables will be derived:

- *Pharmacokinetic Analysis Set Population Flag 1.*
- *Drug-Drug Interaction Analysis Set Population Flag 1*
- *Drug-Drug Interaction Analysis Set Population Flag 2*
- *Drug-Drug Interaction Analysis Set Population Flag 3.*
- *Drug-Drug Interaction Analysis Set Population Flag 4.*
- *Drug-Drug Interaction Analysis Set Population Flag 5.*
- *Drug-Drug Interaction Analysis Set Population Flag 6.*
- *Pharmacokinetic Analysis Set Population Flag 2.*
- *Drug-Drug Interaction Analysis Set Population Flag 7.*
- *Pharmacokinetic Analysis Set Population Flag 3.*
- *Pharmacokinetic Analysis Set Population Flag 4.*



#### ADPP – Analysis Data Pharmacokinetic Parameters

This is a BDS dataset that will contain the NCA pharmacokinetic parameters (plasma and urine) for each subject, per treatment analysis and per analyte/metabolite.

Besides the original pharmacokinetic parameter value, the dataset will contain the  $\ln$ -transformed value that is necessary to apply for the specified statistical model(s).

The origin of these data will primarily be the DM and Pharmacokinetic Parameters (PP) SDTM domains.

Besides this information, the following variables will be derived:

- *Pharmacokinetic Analysis Set Population Flag 1.*
- *Drug-Drug Interaction Analysis Set Population Flag 1.*
- *Drug-Drug Interaction Analysis Set Population Flag 2.*
- *Drug-Drug Interaction Analysis Set Population Flag 3.*
- *Drug-Drug Interaction Analysis Set Population Flag 4.*
- *Drug-Drug Interaction Analysis Set Population Flag 5.*
- *Drug-Drug Interaction Analysis Set Population Flag 6.*
- *Pharmacokinetic Analysis Set Population Flag 2.*
- *Drug-Drug Interaction Analysis Set Population Flag 7.*
- *Pharmacokinetic Analysis Set Population Flag 3.*
- *Pharmacokinetic Analysis Set Population Flag 4.*

#### ADAE – Analysis Data Adverse Events Dataset

This is an OCCDS dataset that will contain the adverse events for each subject.

The origin of these data will primarily be the Adverse Events (AE) SDTM domain and ADSL ADaM domain.

#### ADVS – Analysis Data Vital Signs Dataset

This is a BDS dataset that will contain the vital signs results for each subject, per treatment analysis and per timepoint.

The origin of these data will primarily be the Vital Signs (VS) SDTM domain and ADSL ADaM domain.

#### ADEG – Analysis Data Electrocardiogram Parameters Dataset

This is a BDS dataset that will contain the electrocardiogram for each subject, per treatment analysis and per timepoint.

The origin of these data will primarily be the ECG Test Results (EG) SDTM domain and ADSL ADaM domain.

#### ADLB – Analysis Data Laboratory Test Results Dataset

This is a BDS dataset that will contain the safety laboratory test results for each subject and per treatment analysis.

The origin of these data will primarily be the Laboratory Test Results (LB) SDTM domain and ADSL ADaM domain.

## 10. STATISTICAL METHODS



### 10.1. General Considerations

Estimation of the pharmacokinetic parameters and drug-drug interaction comparative bioavailability analysis will be conducted on Phoenix® WinNonlin® version 8.2 or higher (Certara USA Inc, Princeton, NJ). All other statistical analysis will be conducted on SAS® version 9.4 or higher.

Statistical analyses and pharmacokinetic analyses will be performed in accordance with the U.S. FDA guidances [2, 3] and PPD applicable Standard Operating Procedures (SOPs).

Unless specified otherwise, continuous variables will be summarized with the following descriptive statistics: number of observations (n), mean, standard deviation (SD), minimum, median and maximum values. Categorical data will be summarized with frequencies and percentages. For subjects' characteristics and safety analyses, missing data will not be replaced; descriptive statistics and statistical analysis will be performed based on the available data only. All data recorded on discontinued subjects will be listed.

### 10.2. Subjects' Disposition

The number of subjects who completed each study part as well as subjects who withdrew prematurely from the study will be summarized by completion status and reason for withdrawal. Subjects who prematurely withdrew from the study will be listed.

The number of subjects included in each analysis population will be summarized, for each study part.

### 10.3. Protocol Deviations

During the respective Data Blind Review Process, all protocol deviations that occurred during each study part will be assessed in terms of their potential impact on the pharmacokinetic analysis. The result of this assessment will be reported in the respective Data Blind Review Process Minute (Annex 1, Annex 3, and Annex 5).

### 10.4. Plasma Pharmacokinetic Concentrations

For each study part (Part 1, Part 2 and Part 3), for the subjects included in the respective pharmacokinetic analysis population, descriptive statistics [n, geometric mean ( $G_{mean}$ ), arithmetic mean ( $A_{mean}$ ), SD, geometric SD (GSD), coefficient of variation (CV%), geometric CV% (GCV%), two-sided 95% CI of the  $A_{mean}$  and  $G_{mean}$ , median, minimum and maximum] of the plasma concentrations will be presented for each time point, by analyte, and by administration day. Concentrations below the LLOQ will be taken as missing for the calculation of the log-transformed statistics and as zero for the calculation of the remaining summary statistics.

Additionally, for Part 3, for the subjects included in the respective pharmacokinetic analysis population, descriptive statistics of the concentrations of givinostat and its metabolites in plasma will be presented by single and multiple-dose, and by sex. Achievement of steady-state conditions will be determined by visual inspection of the trough plasma concentration-time profile ( $C_{trough}$ ). Analysis of variance of the plasma  $C_{trough}$  may be performed if considered appropriate.



For each study part (Part 1, Part 2 and Part 3), individual (per subject) and  $G_{mean}$  (including 95% CIs) plasma concentration *versus* time profile will be graphically displayed in both linear and semi-logarithmic scales:

- For plotting individual data in linear scale, concentrations below the LLOQ will be substituted by zero.
- For plotting  $G_{mean}$  data in linear scale, concentrations below the LLOQ will be substituted by missing.
- For plotting data in semi-logarithmic scale, concentrations below the LLOQ before  $t_{max}$ , will be substituted by  $\frac{1}{2}$  of the LLOQ value. After  $t_{max}$ , concentrations below the LLOQ will be considered as missing. However, in case of two or more consecutive concentrations below the LLOQ, the first value will be replaced by  $\frac{1}{2}$  of the LLOQ value and the next values will be considered as missing.

Graphic presentation of individual data will be based on actual blood sampling time, except for pre-dose sampling time which will be assumed as  $t_0$ .

Concentration data from non-evaluable periods will be listed and graphically represented separately.

#### 10.5. Plasma Pharmacokinetic Parameters

For each study part, plasma pharmacokinetic parameters will be calculated for the respective pharmacokinetic analysis population.

For each study part, for the subjects included in the pharmacokinetic analysis population, individual plasma pharmacokinetic parameters and descriptive statistics ( $n$ ,  $G_{mean}$ ,  $A_{mean}$ , SD, GSD, CV%, GCV%, two-sided 95% CI of the  $A_{mean}$  and  $G_{mean}$ , median, minimum and maximum), will be presented by investigational product.

If a given pharmacokinetic parameter could not be reliably determined for more than  $\frac{1}{3}$  of the subjects, only the minimum and maximum values will be presented, and the other descriptive statistics will be omitted for that parameter.

Missing pharmacokinetic parameter data will not be imputed.

For Part 3, for the subjects included in the respective pharmacokinetic analysis population, descriptive statistics of the plasma pharmacokinetic parameters of givinostat and its metabolites will be presented by single and multiple-dose, and by sex.

For Part 3, time dependency will be explored by comparing  $AUC_{0-\tau,ss}$  derived from the last dosing day (Day 13), after multiple doses administration, with  $AUC_{0-\infty}$  derived from the first dosing day (Day 1), after a single dose administration. Assessment of linearity will consider whether differences in AUC meet a criterion of  $\pm 25\%$ .

For Part 3, the Accumulation Index for  $C_{max}$  will be derived as the ratio  $\frac{C_{max,ss}}{C_{max}}$  and for AUC it will be derived as the ratio  $\frac{AUC_{0-\tau,ss}}{AUC_{0-t}}$ . Accumulation Index estimates will be summarized using  $A_{mean}$ , SD, SE median, minimum and maximum.

For Part 3, metabolic ratios will be calculated for each metabolite as  $100 \cdot$



$\frac{AUC_{0-\infty, Metabolite}/MW_{Metabolite}}{AUC_{0-\infty, Givinostat}/MW_{Givinostat}}$ , for Day 1 (single-dose), and as  $100 \cdot \frac{AUC_{0-\tau, Metabolite}/MW_{Metabolite}}{AUC_{0-\tau, Givinostat}/MW_{Givinostat}}$ , for Day 13 (multiple-dose).

*Note: MW represents the molecular weight of the analyte. For givinostat, MW is 421.49 g/mol; for ITF2374, MW is 405.49 g/mol; for ITF2375, MW is 406.47 g/mol; for ITF2440, MW is 257.33 g/mol; and for ITF2563, MW is 178.19 g/mol.*

## 10.6. Urinary Data

For Part 3, for the subjects included in the pharmacokinetic analysis population, descriptive statistics (n,  $G_{mean}$ ,  $A_{mean}$ , SD, GSD, CV%, GCV%, two-sided 95% CI of the  $A_{mean}$  and  $G_{mean}$ , median, minimum and maximum) of the urinary excretion profile and urinary cumulative excretion profile will be presented for each collection interval, by analyte, by single (Day 1) and multiple (Day 13) dose, and by sex. Concentrations below the LLOQ will be taken as missing for the calculation of the log-transformed statistics and as zero for the calculation of the remaining summary statistics.

Individual (per subject) and  $G_{mean}$  urinary excretion *versus* time profile and urinary cumulative excretion *versus* time profile will be graphically displayed, in both linear and semi-logarithmic scales:

- For plotting individual data in linear scale, amounts below the LLOQ will be substituted by zero.
- For plotting  $G_{mean}$  data in linear scale, amounts below the LLOQ will be substituted by missing.

Day -1 sampling will be assumed as  $t_0$ .

## 10.7. Urinary Pharmacokinetic Parameters

For Part 3, urinary pharmacokinetic parameters will be calculated for the respective pharmacokinetic analysis population.

For Part 3, for the subjects included in the pharmacokinetic analysis population, individual urinary pharmacokinetic parameters and descriptive statistics (n,  $G_{mean}$ ,  $A_{mean}$ , SD, GSD, CV%, GCV%, two-sided 95% CI of the  $A_{mean}$  and  $G_{mean}$ , median, minimum and maximum), will be presented by analyte, by single and multiple-dose, and by sex.

If a given pharmacokinetic parameter could not be reliably determined for more than  $\frac{1}{3}$  of the subjects, only the minimum and maximum values will be presented, and the other descriptive statistics will be omitted for that parameter.

Missing pharmacokinetic parameter data will not be imputed.

## 11. PRESENTATION OF RESULTS

### 11.1. Statistical Output Specification

Tables will be generated as Rich Text Files (\*.rtf) from SAS®.

Figures and Analysis Outputs will be generated as \*.docx files, exported from Phoenix WinNonlin®



and/or SAS®.

In-text Tables and Figures will preferably be prepared in portrait format. Listings will preferably be prepared in landscape format.

The CSR will be written according to PPD SOPs and templates for reporting bioavailability/bioequivalence trials.

## 11.2. Planned Tables, Figures, and Subject Data Listings

The planned Tables, Figures, and Individual Data Listings for the CSR are listed below.

### 11.2.1. In-text Tables and Figures

For CSR Synopsis:

The following Tables will be produced for direct insertion in the Synopsis of the CSR:

Title
<b>PART 3</b>
Givinostat and Its Main Metabolites (ITF2374 and ITF2375): Summary Statistics of the Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1)
Givinostat Metabolites (ITF2440 and ITF2563): Summary Statistics of the Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1)
Givinostat and Its Main Metabolites (ITF2374 and ITF2375): Summary Statistics of the Plasma Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13)
Givinostat Metabolites (ITF2440 and ITF2563): Summary Statistics of the Plasma Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13)
Givinostat and Its Main Metabolites (ITF2374 and ITF2375): Summary Statistics of the Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1)
Givinostat Metabolites (ITF2440 and ITF2563): Summary Statistics of the Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1)
Givinostat and Its Main Metabolites (ITF2374 and ITF2375): Summary Statistics of the Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13)
Givinostat Metabolites (ITF2440 and ITF2563): Summary Statistics of the Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13)

The following Figures will be produced for direct insertion in the Synopsis of the CSR:

Legend
<b>PART 3</b>
Givinostat: Geometric Mean (95% CI) Plasma Concentration <i>Versus</i> Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat. A – Linear



Legend
Scale; B – Semi-Logarithmic Scale
Givinostat Metabolite (ITF2374): Geometric Mean (95% CI) Plasma Concentration <i>Versus</i> Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat (Day 1). A – Linear Scale; B – Semi-Logarithmic Scale
Givinostat Metabolite (ITF2375): Geometric Mean (95% CI) Plasma Concentration <i>Versus</i> Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat (Day 1). A – Linear Scale; B – Semi-Logarithmic Scale
Givinostat Metabolite (ITF2440): Geometric Mean (95% CI) Plasma Concentration <i>Versus</i> Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat (Day 1). A – Linear Scale; B – Semi-Logarithmic Scale
Givinostat Metabolite (ITF2563): Geometric Mean (95% CI) Plasma Concentration <i>Versus</i> Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat (Day 1). A – Linear Scale; B – Semi-Logarithmic Scale

For CSR Body Text:

The following Tables and Figures will be produced for direct insertion in the body text of the CSR:

In-text Tables	Title
<b>PART 3</b>	
Table 3.A.	Study Flow-Chart – Part 3
Table 3.B.	Identity of Investigational Products – Part 3
Table 3.C.	Summary of Subjects Disposition – Part 3
Table 3.D.	Summary of Demographic Data of the Pharmacokinetic Analysis Populations
Table 3.E.	Givinostat and Metabolites (ITF2374, ITF2375, ITF2440 and ITF2563): Summary Statistics of the Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1)
Table 3.F.	Givinostat and Metabolites (ITF2374, ITF2375, ITF2440 and ITF2563): Summary Statistics of the Plasma Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13)
Table 3.G.	Givinostat and Metabolites (ITF2374, ITF2375, ITF2440 and ITF2563): Summary Statistics of the Plasma Pharmacokinetic Parameters Accumulation Index
Table 3.H.	Givinostat and Metabolites (ITF2374, ITF2375, ITF2440 and ITF2563): Summary Statistics of the Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) <sup>21</sup>
Table 3.I.	Givinostat and Metabolites (ITF2374, ITF2375, ITF2440 and ITF2563): Summary Statistics of the Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) <sup>21</sup>
Table 3.J.	Summary of Treatment-Emergent Adverse Events (TEAEs)

In-text Figures	Legend
<b>PART 3</b>	
Figure 3.A.1.	Givinostat: Geometric Mean (95% CI) Plasma Concentration <i>Versus</i> Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale
Figure 3.A.2.	Givinostat: Geometric Mean (95% CI) Plasma Concentration <i>Versus</i> Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13)



In-text Figures	Legend
	Administration of Givinostat – Semi-Logarithmic Scale
Figure 3.A.3.	Givinostat: Geometric Mean (95% CI) Plasma Concentration <i>Versus</i> Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale
Figure 3.A.4.	Givinostat: Geometric Mean (95% CI) Plasma Concentration <i>Versus</i> Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Semi-Logarithmic Scale
Figure 3.A.5.	Givinostat: Geometric Mean (95% CI) Trough Plasma Concentration <i>Versus</i> Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale
Figure 3.B.1.	Givinostat Metabolite (ITF2374): Geometric Mean (95% CI) Plasma Concentration <i>Versus</i> Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale
Figure 3.B.2.	Givinostat Metabolite (ITF2374): Geometric Mean (95% CI) Plasma Concentration <i>Versus</i> Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Semi-Logarithmic Scale
Figure 3.B.3.	Givinostat Metabolite (ITF2374): Geometric Mean (95% CI) Plasma Concentration <i>Versus</i> Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale
Figure 3.B.4.	Givinostat Metabolite (ITF2374): Geometric Mean (95% CI) Plasma Concentration <i>Versus</i> Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Semi-Logarithmic Scale
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In-text Figures	Legend
	Scale
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In-text Figures	Legend
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#### 11.2.2. Tables, Figures and Graphs Referred to But Not Included in the Text (Section 14 of CSR)

At least the following Tables and Figures should be produced for Section 14 of the CSR.  
If necessary, additional Tables and Figures should be prepared.

The following Tables and Figures will be compiled in a document, to be considered as Section 14 (Tables, Figures and Graph Referred To But Not Included in the Text) of the CSR.

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14.3.2.5.7.	Givinostat Metabolite (ITF2440): Individual Urinary Excretion <i>Versus</i> Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale
14.3.2.5.8.	Givinostat Metabolite (ITF2440): Individual Cumulative Amount Urinary Excretion <i>Versus</i> Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale
14.3.2.5.9.	Givinostat Metabolite (ITF2563): Individual Urinary Excretion <i>Versus</i> Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale
14.3.2.5.10.	Givinostat Metabolite (ITF2563): Individual Cumulative Amount Urinary Excretion <i>Versus</i> Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale



### 11.2.3. List of Subject Data Listings (Section 16.2 of CSR)

The following Data Listings will be produced for Section 16.2. Subject Data Listings of the CSR.

If necessary, additional Data Listings should be prepared.

Listing No.	Title
<b>16.2. SUBJECTS DATA LISTING</b>	
<b>16.2.3. Part 3</b>	
<b>16.2.3.1.</b>	<b>Subject Disposition</b>
16.2.3.1.1.	Subject Disposition
<b>16.2.3.2.</b>	<b>Protocol Deviations</b>
16.2.3.2.1.	Blood Sampling Times Deviations
16.2.3.2.2.	Other Protocol Deviations
<b>16.2.3.3.</b>	<b>Excluded Data from the Pharmacokinetic Analysis</b>
16.2.3.3.1.	Plasma Concentrations
16.2.3.3.1.1.	Individual Data of Plasma Concentrations
16.2.3.3.2.	Plasma Concentration Versus Time Profiles
16.2.3.3.2.1.	Individual Plasma Concentration Versus Time Profiles – Linear Scale
16.2.3.3.2.2.	Individual Plasma Concentration Versus Time Profiles – Semi-Logarithmic Scale
16.2.3.3.3.	Urine Excretion
16.2.3.3.3.1.	Individual Data of Plasma Concentrations
16.2.3.3.4.	Urine Excretion Versus Time Profiles
16.2.3.3.4.1.	Urine Excretion Versus Time Profiles – Linear Scale
<b>16.2.3.4.</b>	<b>Demographic and Other Baseline Data</b>
16.2.3.4.1.	Demographic Data
16.2.3.4.2.	Fertility/Contraception
16.2.3.4.2.1.	Female Fertility/Contraception
16.2.3.4.2.2.	Male Fertility/Contraception
16.2.3.4.3.	Drugs of Abuse, Ethanol and Cotinine
16.2.3.4.4.	Viral Serology at Screening
16.2.3.4.5.	Previous and Concomitant Medication
16.2.3.4.6.	SARS-COV-2 Test
<b>16.2.3.5.</b>	<b>Compliance</b>
16.2.3.5.1.	Investigational Product Administration
<b>16.2.3.6.</b>	<b>Individual Pharmacokinetic Data</b>
16.2.3.6.1.	Givinostat: Individual Pharmacokinetic Data
16.2.3.6.1.1.	Givinostat: Individual Drug Plasma Concentration Data
16.2.3.6.1.2.	Givinostat: Individual Plasma Pharmacokinetic Parameters
16.2.3.6.1.3.	Givinostat: Individual Plasma Pharmacokinetic Profiles
16.2.3.6.1.4.	Givinostat: Individual Urine Excretion Data
16.2.3.6.1.5.	Givinostat: Individual Urine Pharmacokinetic Parameters
16.2.3.6.1.6.	Givinostat: Individual Urine Pharmacokinetic Profiles
16.2.3.6.1.7.	Givinostat: Documentation of Statistical Analysis
16.2.3.6.1.7.1.	Givinostat: Plasma Non-Compartmental Analysis Output
16.2.3.6.1.7.2.	Givinostat: Plasma Non-Compartmental Analysis Plots
16.2.3.6.1.7.3.	Givinostat: Urine Non-Compartmental Analysis Output



Listing No.	Title
16.2.3.6.2.	Givinostat Metabolite (ITF2374): Individual Pharmacokinetic Data
16.2.3.6.2.1.	Givinostat Metabolite (ITF2374): Individual Drug Plasma Concentration Data
16.2.3.6.2.2.	Givinostat Metabolite (ITF2374): Individual Plasma Pharmacokinetic Parameters
16.2.3.6.2.3.	Givinostat Metabolite (ITF2374): Individual Plasma Pharmacokinetic Profiles
16.2.3.6.2.4.	Givinostat Metabolite (ITF2374): Individual Urine Excretion Data
16.2.3.6.2.5.	Givinostat Metabolite (ITF2374): Individual Urine Pharmacokinetic Parameters
16.2.3.6.2.6.	Givinostat Metabolite (ITF2374): Individual Urine Pharmacokinetic Profiles
16.2.3.6.2.7.	Givinostat Metabolite (ITF2374): Documentation of Statistical Analysis
16.2.3.6.2.7.1.	Givinostat Metabolite (ITF2374): Plasma Non-Compartmental Analysis Output
16.2.3.6.2.7.2.	Givinostat Metabolite (ITF2374): Plasma Non-Compartmental Analysis Plots
16.2.3.6.2.7.3.	Givinostat Metabolite (ITF2374): Urine Non-Compartmental Analysis Output
16.2.3.6.3.	Givinostat Metabolite (ITF2375): Individual Pharmacokinetic Data
16.2.3.6.3.1.	Givinostat Metabolite (ITF2375): Individual Drug Plasma Concentration Data
16.2.3.6.3.2.	Givinostat Metabolite (ITF2375): Individual Plasma Pharmacokinetic Parameters
16.2.3.6.3.3.	Givinostat Metabolite (ITF2375): Individual Plasma Pharmacokinetic Profiles
16.2.3.6.3.4.	Givinostat Metabolite (ITF2375): Individual Urine Excretion Data
16.2.3.6.3.5.	Givinostat Metabolite (ITF2375): Individual Urine Pharmacokinetic Parameters
16.2.3.6.3.6.	Givinostat Metabolite (ITF2375): Individual Urine Pharmacokinetic Profiles
16.2.3.6.3.7.	Givinostat Metabolite (ITF2375): Documentation of Statistical Analysis
16.2.3.6.3.7.1.	Givinostat Metabolite (ITF2375): Plasma Non-Compartmental Analysis Output
16.2.3.6.3.7.2.	Givinostat Metabolite (ITF2375): Plasma Non-Compartmental Analysis Plots
16.2.3.6.3.7.3.	Givinostat Metabolite (ITF2375): Urine Non-Compartmental Analysis Output
16.2.3.6.4.	Givinostat Metabolite (ITF2440): Individual Pharmacokinetic Data
16.2.3.6.4.1.	Givinostat Metabolite (ITF2440): Individual Drug Plasma Concentration Data
16.2.3.6.4.2.	Givinostat Metabolite (ITF2440): Individual Plasma Pharmacokinetic Parameters
16.2.3.6.4.3.	Givinostat Metabolite (ITF2440): Individual Plasma Pharmacokinetic Profiles
16.2.3.6.4.4.	Givinostat Metabolite (ITF2440): Individual Urine Excretion Data
16.2.3.6.4.5.	Givinostat Metabolite (ITF2440): Individual Urine Pharmacokinetic Parameters
16.2.3.6.4.6.	Givinostat Metabolite (ITF2440): Individual Urine Pharmacokinetic Profiles
16.2.3.6.4.7.	Givinostat Metabolite (ITF2440): Documentation of Statistical Analysis
16.2.3.6.4.7.1.	Givinostat Metabolite (ITF2440): Plasma Non-Compartmental Analysis



Listing No.	Title
	Output
16.2.3.6.4.7.2.	Givinostat Metabolite (ITF2440): Plasma Non-Compartmental Analysis Plots
16.2.3.6.4.7.3.	Givinostat Metabolite (ITF2440): Urine Non-Compartmental Analysis Output
16.2.3.6.5.	Givinostat Metabolite (ITF2563): Individual Pharmacokinetic Data
16.2.3.6.5.1.	Givinostat Metabolite (ITF2563): Individual Drug Plasma Concentration Data
16.2.3.6.5.2.	Givinostat Metabolite (ITF2563): Individual Plasma Pharmacokinetic Parameters
16.2.3.6.5.3.	Givinostat Metabolite (ITF2563): Individual Plasma Pharmacokinetic Profiles
16.2.3.6.5.4.	Givinostat Metabolite (ITF2563): Individual Urine Excretion Data
16.2.3.6.5.5.	Givinostat Metabolite (ITF2563): Individual Urine Pharmacokinetic Parameters
16.2.3.6.5.6.	Givinostat Metabolite (ITF2563): Individual Urine Pharmacokinetic Profiles
16.2.3.6.5.7.	Givinostat Metabolite (ITF2563): Documentation of Statistical Analysis
16.2.3.6.5.7.1.	Givinostat Metabolite (ITF2563): Plasma Non-Compartmental Analysis Output
16.2.3.6.5.7.2.	Givinostat Metabolite (ITF2563): Plasma Non-Compartmental Analysis Plots
16.2.3.6.5.7.3.	Givinostat Metabolite (ITF2563): Urine Non-Compartmental Analysis Output
<b>16.2.3.7.</b>	<b>Adverse Event Listings (Each Subject)</b>
16.2.3.7.1.	Pre-Treatment Adverse Events
16.2.3.7.2.	Treatment-Emergent Adverse Events
16.2.3.7.3.	Serious Adverse Events (I)
<b>16.2.3.8.</b>	<b>Listings of Laboratory Measurements by Subject</b>
16.2.3.8.1.	Normal Range of Laboratory Values
16.2.3.8.2.	Hematology (I)
16.2.3.8.3.	Hematology (II)
16.2.3.8.4.	Biochemistry (I)
16.2.3.8.5.	Biochemistry (II)
16.2.3.8.6.	Biochemistry (III)
16.2.3.8.7.	Biochemistry (IV)
16.2.3.8.8.	Biochemistry (V)
16.2.3.8.9.	Coagulation
16.2.3.8.10.	Urinalysis
16.2.3.8.11.	Urine Microscopy
16.2.3.8.12.	Pregnancy Test
16.2.3.8.13.	Additional (Not Planned) Laboratory Safety Tests
<b>16.2.3.9.</b>	<b>Vital Signs</b>
<b>16.2.3.10.</b>	<b>12-Lead ECG</b>
<b>16.2.3.11.</b>	<b>Urine Sampling</b>



## 12. REFERENCES

- [1] ITF/2357/55 Clinical Study Protocol, Version 1.0, 29NOV2021
- [2] Food and Drug Administration (FDA), Clinical Drug Interaction Studies - Cytochrome P450 Enzyme- and Transporter-Mediated Drug Interactions Guidance for Industry (2020).



### **13. APPENDICES**

#### **APPENDIX A. Planned Tables, Figures, and Subject Data Listings**

##### **A.1. In-text Tables and Figures – For CSR Synopsis**



A.1.1. Part 3

SYNOPSIS TABLES (1)

**Givinostat and Its Main Metabolites (ITF2374 and ITF2375): Summary Statistics of the Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1)**

Parameter (unit)	Givinostat (n = xx)	ITF2374 (n = xx)	ITF2375 (n = xx)
C <sub>max</sub> (<unitsC>)	xx.xx (xx.x%)	xx.xx (xx.x%)	xx.xx (xx.x%)
T <sub>max</sub> (h)	xx.xx (xx.xx – xx.xx)	xx.xx (xx.xx – xx.xx)	xx.xx (xx.xx – xx.xx)
AUC <sub>0-t</sub> (<unitsA>)	xx.xx (xx.x%)	xx.xx (xx.x%)	xx.xx (xx.x%)
AUC <sub>0-∞</sub> (<unitsA>)	xx.xx (xx.x%)	xx.xx (xx.x%)	xx.xx (xx.x%)
%AUC <sub>extrap</sub> (%)	xx.xx (xx.x%)	xx.xx (xx.x%)	xx.xx (xx.x%)
λ <sub>z</sub> (1/h)	x.xxx (xx.x%)	x.xxx (xx.x%)	x.xxx (xx.x%)
t <sub>1/2</sub> (h)	xx.xx (xx.x%)	xx.xx (xx.x%)	xx.xx (xx.x%)

n – Number of Subjects

Values are geometric mean (G<sub>mean</sub>) with geometric coefficient of variation (GCV%) within parenthesis

T<sub>max</sub> values are median with range between parentheses

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

*Note: This table will present the summary statistics of the pharmacokinetic parameters calculated for Part 3 Pharmacokinetic Analysis Population. In case a pharmacokinetic parameter could not be calculated for one or more subjects in the Part 1 Pharmacokinetic Analysis Population, the pharmacokinetic parameter should be followed by the # symbol and accompanied by a footnote: '# For <Parameter(s)>, n=xx, for <Investigational Product> product <, and n=xx, for <Investigational Product> product'.*



## SYNOPSIS TABLES (2)

### Givinostat Metabolites (ITF2440 and ITF2563): Summary Statistics of the Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1)

Parameter (unit)	ITF2440 (n = xx)	ITF2563 (n = xx)
$C_{max}$ (<unitsC>)	xx.xx (xx.x%)	xx.xx (xx.x%)
$T_{max}$ (h)	xx.xx (xx.xx – xx.xx)	xx.xx (xx.xx – xx.xx)
$AUC_{0-t}$ (<unitsA>)	xx.xx (xx.x%)	xx.xx (xx.x%)
$AUC_{0-\infty}$ (<unitsA>)	xx.xx (xx.x%)	xx.xx (xx.x%)
%AUC <sub>extrap</sub> (%)	xx.xx (xx.x%)	xx.xx (xx.x%)
$\lambda_z$ (1/h)	x.xxx (xx.x%)	x.xxx (xx.x%)
$t_{1/2}$ (h)	xx.xx (xx.x%)	xx.xx (xx.x%)

n – Number of Subjects

Values are geometric mean ( $G_{mean}$ ) with geometric coefficient of variation (GCV%) within parenthesis

$T_{max}$  values are median with range between parentheses

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

Note: This table will present the summary statistics of the pharmacokinetic parameters calculated for Part 3 Pharmacokinetic Analysis Population. In case a pharmacokinetic parameter could not be calculated for one or more subjects in the Part 1 Pharmacokinetic Analysis Population, the pharmacokinetic parameter should be followed by the # symbol and accompanied by a footnote: '# For <Parameter(s)>, n=xx, for <Investigational Product> product <, and n=xx, for <Investigational Product> product>'.

## SYNOPSIS TABLES (3)

### Givinostat and Its Main Metabolites (ITF2374 and ITF2375): Summary Statistics of the Plasma Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13)

Parameter (unit)	Givinostat (n = xx)	ITF2374 (n = xx)	ITF2375 (n = xx)
$C_{max,ss}$ (<unitsC>)	xx.xx (xx.x%)	xx.xx (xx.x%)	xx.xx (xx.x%)
$T_{max,ss}$ (h)	xx.xx (xx.xx – xx.xx)	xx.xx (xx.xx – xx.xx)	xx.xx (xx.xx – xx.xx)
$AUC_{0-t,ss}$ (<unitsA>)	xx.xx (xx.x%)	xx.xx (xx.x%)	xx.xx (xx.x%)
$AUC_{0-t}$ (<unitsA>)	xx.xx (xx.x%)	xx.xx (xx.x%)	xx.xx (xx.x%)
$AUC_{0-\infty}$ (<unitsA>)	xx.xx (xx.x%)	xx.xx (xx.x%)	xx.xx (xx.x%)
$\lambda_z$ (1/h)	x.xxx (xx.x%)	x.xxx (xx.x%)	x.xxx (xx.x%)
$t_{1/2}$ (h)	xx.xx (xx.x%)	xx.xx (xx.x%)	xx.xx (xx.x%)
$CL_{ss}/F$ (L/h)	xx.xx (xx.x%)	xx.xx (xx.x%)	xx.xx (xx.x%)
$V_D/F$ (L)	xx.xx (xx.x%)	xx.xx (xx.x%)	xx.xx (xx.x%)

n – Number of Subjects

Values are geometric mean ( $G_{mean}$ ) with geometric coefficient of variation (GCV%) within parenthesis

$T_{max}$  values are median with range between parentheses

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

*Note: This table will present the summary statistics of the pharmacokinetic parameters calculated for Part 3 Pharmacokinetic Analysis Population. In case a pharmacokinetic parameter could not be calculated for one or more subjects in the Part 1 Pharmacokinetic Analysis Population, the pharmacokinetic parameter should be followed by the # symbol and accompanied by a footnote: '# For <Parameter(s)>, n=xx, for <Investigational Product> product <, and n=xx, for <Investigational Product> product>'.*



## SYNOPSIS TABLES (4)

### Givinostat Metabolites (ITF2440 and ITF2563): Summary Statistics of the Plasma Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13)

Parameter (unit)	ITF2440 (n = xx)	ITF2563 (n = xx)
$C_{max,ss}$ (<unitsC>)	xx.xx (xx.x%)	xx.xx (xx.x%)
$T_{max,ss}$ (h)	xx.xx (xx.xx – xx.xx)	xx.xx (xx.xx – xx.xx)
$AUC_{0-t,ss}$ (<unitsA>)	xx.xx (xx.x%)	xx.xx (xx.x%)
$AUC_{0-t}$ (<unitsA>)	xx.xx (xx.x%)	xx.xx (xx.x%)
$AUC_{0-\infty}$ (<unitsA>)	xx.xx (xx.x%)	xx.xx (xx.x%)
$\lambda_z$ (1/h)	x.xxx (xx.x%)	x.xxx (xx.x%)
$t_{1/2}$ (h)	xx.xx (xx.x%)	xx.xx (xx.x%)
$CL_{ss}/F$ (L/h)	xx.xx (xx.x%)	xx.xx (xx.x%)
$V_D/F$ (L)	xx.xx (xx.x%)	xx.xx (xx.x%)

n – Number of Subjects

Values are geometric mean ( $G_{mean}$ ) with geometric coefficient of variation (GCV%) within parenthesis

$T_{max}$  values are median with range between parentheses

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

*Note: This table will present the summary statistics of the pharmacokinetic parameters calculated for Part 3 Pharmacokinetic Analysis Population. In case a pharmacokinetic parameter could not be calculated for one or more subjects in the Part 1 Pharmacokinetic Analysis Population, the pharmacokinetic parameter should be followed by the # symbol and accompanied by a footnote: '# For <Parameter(s)>, n=xx, for <Investigational Product> product <, and n=xx, for <Investigational Product> product>'.*

## SYNOPSIS TABLES (5)

**Givinostat and Its Main Metabolites (ITF2374 and ITF2375): Summary Statistics of the Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1)**<sup>10</sup>

## SYNOPSIS TABLES (6)

**Givinostat Metabolites (ITF2440 and ITF2563): Summary Statistics of the Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1)**<sup>11</sup>

## SYNOPSIS TABLES (7)

**Givinostat and Its Main Metabolites (ITF2374 and ITF2375): Summary Statistics of the Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13)**<sup>10</sup>

## SYNOPSIS TABLES (8)

**Givinostat Metabolites (ITF2440 and ITF2563): Summary Statistics of the Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13)**<sup>11</sup>



<sup>10</sup> To be generated as follows:

Parameter (unit)	Givinostat (n = xx)	ITF2374 (n = xx)	ITF2375 (n = xx)
R <sub>max,ss</sub> (<units>)	xx.xx (xx.x%)	xx.xx (xx.x%)	xx.xx (xx.x%)
t <sub>u</sub> max (h)	xx.xx (xx.xx – xx.xx)	xx.xx (xx.xx – xx.xx)	xx.xx (xx.xx – xx.xx)
Amt <sub>CUM</sub> (<units>)	xx.xx (xx.x%)	xx.xx (xx.x%)	xx.xx (xx.x%)
AURC <sub>0-t</sub> (<units>)	xx.xx (xx.x%)	xx.xx (xx.x%)	xx.xx (xx.x%)
REC% (%)	xx.xx (xx.x%)	xx.xx (xx.x%)	xx.xx (xx.x%)
CL <sub>R</sub> /F (L/h)	x.xxx (xx.x%)	x.xxx (xx.x%)	x.xxx (xx.x%)

n – Number of Subjects

Values are geometric mean (G<sub>mean</sub>) with geometric coefficient of variation (GCV%) within parenthesis

t<sub>u</sub>max values are median with range between parentheses

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

*Note: This table will present the summary statistics of the pharmacokinetic parameters calculated for Part 3 Pharmacokinetic Analysis Population. In case a pharmacokinetic parameter could not be calculated for one or more subjects in the Part 1 Pharmacokinetic Analysis Population, the pharmacokinetic parameter should be followed by the # symbol and accompanied by a footnote: '# For <Parameter(s)>, n=xx, for <Investigational Product> product <, and n=xx, for <Investigational Product> product'.*

<sup>11</sup> To be generated as follows:

Parameter (unit)	ITF2440 (n = xx)	ITF2563 (n = xx)
R <sub>max,ss</sub> (<units>)	xx.xx (xx.x%)	xx.xx (xx.x%)
t <sub>u</sub> max (h)	xx.xx (xx.xx – xx.xx)	xx.xx (xx.xx – xx.xx)
Amt <sub>CUM</sub> (<units>)	xx.xx (xx.x%)	xx.xx (xx.x%)
AURC <sub>0-t</sub> (<units>)	xx.xx (xx.x%)	xx.xx (xx.x%)
REC% (%)	xx.xx (xx.x%)	xx.xx (xx.x%)
CL <sub>R</sub> /F (L/h)	x.xxx (xx.x%)	x.xxx (xx.x%)

n – Number of Subjects

Values are geometric mean (G<sub>mean</sub>) with geometric coefficient of variation (GCV%) within parenthesis

t<sub>u</sub>max values are median with range between parentheses

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

*Note: This table will present the summary statistics of the pharmacokinetic parameters calculated for Part 3 Pharmacokinetic Analysis Population. In case a pharmacokinetic parameter could not be calculated for one or more subjects in the Part 1 Pharmacokinetic Analysis Population, the pharmacokinetic parameter should be followed by the # symbol and accompanied by a footnote: '# For <Parameter(s)>, n=xx, for <Investigational Product> product <, and n=xx, for <Investigational Product> product'.*

## SYNOPSIS FIGURES (9)

**Givinostat: Geometric Mean (95% CI) Plasma Concentration *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat. A – Linear Scale; B – Semi-Logarithmic Scale**

## SYNOPSIS FIGURES (10)

**Givinostat Metabolite (ITF2374): Geometric Mean (95% CI) Plasma Concentration *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat (Day 1). A – Linear Scale; B – Semi-Logarithmic Scale**

## SYNOPSIS FIGURES (11)

**Givinostat Metabolite (ITF2375): Geometric Mean (95% CI) Plasma Concentration *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat (Day 1). A – Linear Scale; B – Semi-Logarithmic Scale**

## SYNOPSIS FIGURES (12)

**Givinostat Metabolite (ITF2440): Geometric Mean (95% CI) Plasma Concentration *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat (Day 1). A – Linear Scale; B – Semi-Logarithmic Scale**

## SYNOPSIS FIGURES (13)

**Givinostat Metabolite (ITF2563): Geometric Mean (95% CI) Plasma Concentration *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat (Day 1). A – Linear Scale; B – Semi-Logarithmic Scale**



## **A.2. In-text Tables and Figures – For CSR Body Text**

### **A.2.1. Part 3**

#### **Table 1.A. Study Flow-Chart – Part 3**

*Note: To be captured from the Clinical Study Protocol (CSP) and Amendments.*

#### **Table 1.B. Identity of Investigational Products – Part 3**

*Note: To be generated by the Medical Writing (MW).*

Table 1.C. Summary of Subjects Disposition – Part 3

Disposition	Number of Subjects
Admitted to Treatment Period	xx
Discontinued after First Dosing	xx
Reason	
- <Reason>	xx
Part 3 Safety Analysis Population	xx
Part 3 Pharmacokinetic Analysis Population	xx
Completed Part 3 of Study	xx

Program: <SAS Program>  
Execution Date/Time: <ddMMMyyyy HH:MM>



**Table 1.D. Summary of Demographic Data of the Pharmacokinetic Analysis Populations**

Demography	Parameter	Single-Dose	Multiple-Dose
Age (years)	n	XX	XX
	Mean	XX	XX
	SD	XX.X	XX.X
	Median	XX	XX
	Minimum	XX	XX
	Maximum	XX	XX
Sex, n (%)	n	XX	XX
	Male	XX (XX.X%)	XX (XX.X%)
	Female	XX (XX.X%)	XX (XX.X%)
Race, n (%)	n	XX	XX
	<Race>	XX (XX.X%)	XX (XX.X%)
Weight (kg)	n	XX	XX
	Mean	XX.X	XX.X
	SD	XX.XX	XX.XX
	Median	XX.X	XX.X
	Minimum	XX.X	XX.X
	Maximum	XX.X	XX.X
Height (cm)	n	XX	XX
	Mean	XXX	XXX
	SD	XX.X	XX.X
	Median	XXX	XXX
	Minimum	XXX	XXX
	Maximum	XXX	XXX
Body Mass Index (kg/m <sup>2</sup> )	n	XX	XX
	Mean	XX.X	XX.X
	SD	XX.XX	XX.XX
	Median	XX.X	XX.X
	Minimum	XX.X	XX.X
	Maximum	XX.X	XX.X

n – Number of Subjects; SD – Standard Deviation

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

**Table 1.E. Givinostat and Metabolites (ITF2374, ITF2375, ITF2440 and ITF2563): Summary Statistics of the Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1)**

Parameter (unit)	Givinostat (n = xx)	ITF2374 (n = xx)	ITF2375 (n = xx)	ITF2440 (n = xx)	ITF2563 (n = xx)
$C_{max}$ (<unitsC>)	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]
$T_{max}$ (h)	xx.xx (xx.xx – xx.xx)	xx.xx (xx.xx – xx.xx)	xx.xx (xx.xx – xx.xx)	xx.xx (xx.xx – xx.xx)	xx.xx (xx.xx – xx.xx)
$AUC_{0-t}$ (<unitsA>)	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]
$AUC_{0-\infty}$ (<unitsA>)	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]
% $AUC_{extrap}$ (%)	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]
$\lambda_z$ (1/h)	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]
$t_{1/2}$ (h)	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]

n – Number of Subjects

Values are geometric mean ( $G_{mean}$ ) with geometric coefficient of variation (GCV%) within parenthesis and 95% confidence interval of the  $G_{mean}$  within square brackets

$T_{max}$  values are median with range between parentheses

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

*Note: This table will present the summary statistics of the pharmacokinetic parameters calculated for Part 3 Pharmacokinetic Analysis Population. In case a pharmacokinetic parameter could not be calculated for one or more subjects in the Part 1 Pharmacokinetic Analysis Population, the pharmacokinetic parameter should be followed by the # symbol and accompanied by a footnote: '# For <Parameter(s)>, n=xx, for <Investigational Product> product <, and n=xx, for <Investigational Product> product'.*



**Table 1.F. Givinostat and Metabolites (ITF2374, ITF2375, ITF2440 and ITF2563): Summary Statistics of the Plasma Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13)**

Parameter (unit)	Givinostat (n = xx)	ITF2374 (n = xx)	ITF2375 (n = xx)	ITF2440 (n = xx)	ITF2563 (n = xx)
$C_{max,ss}$ (<unitsC>)	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]
$T_{max,ss}$ (h)	xx.xx (xx.xx – xx.xx)	xx.xx (xx.xx – xx.xx)	xx.xx (xx.xx – xx.xx)	xx.xx (xx.xx – xx.xx)	xx.xx (xx.xx – xx.xx)
$AUC_{0-T,ss}$ (<unitsA>)	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]
$AUC_{0-t}$ (<unitsA>)	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]
$AUC_{0-\infty}$ (<unitsA>)	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]
$\lambda_z$ (1/h)	x.xxx (xx.x%) [x.xxx – x.xxx]	x.xxx (xx.x%) [x.xxx – x.xxx]	x.xxx (xx.x%) [x.xxx – x.xxx]	x.xxx (xx.x%) [x.xxx – x.xxx]	x.xxx (xx.x%) [x.xxx – x.xxx]
$t_{1/2}$ (h)	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]
$CL_{ss}/F$ (L/h)	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]
$V_D/F$ (L)	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]

n – Number of Subjects

Values are geometric mean ( $G_{mean}$ ) with geometric coefficient of variation (GCV%) within parenthesis and 95% confidence interval of the  $G_{mean}$  within square brackets

$T_{max,ss}$  values are median with range between parentheses

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

*Note: This table will present the summary statistics of the pharmacokinetic parameters calculated for Part 3 Pharmacokinetic Analysis Population. In case a pharmacokinetic parameter could not be calculated for one or more subjects in the Part 1 Pharmacokinetic Analysis Population, the pharmacokinetic parameter should be followed by the # symbol and accompanied by a footnote: '# For <Parameter(s)>, n=xx, for <Investigational Product> product <, and n=xx, for <Investigational Product> product'.*

**Table 1.G. Givinostat and Metabolites (ITF2374, ITF2375, ITF2440 and ITF2563): Summary Statistics of the Plasma Pharmacokinetic Parameters Accumulation Index**

Parameter (unit)	Givinostat (n = xx)	ITF2374 (n = xx)	ITF2375 (n = xx)	ITF2440 (n = xx)	ITF2563 (n = xx)
C <sub>max</sub>	xx.xx ± xx.xx	xx.xx ± xx.xx	xx.xx ± xx.xx	xx.xx ± xx.xx	xx.xx ± xx.xx
AUC <sub>0-t</sub>	xx.xx ± xx.xx	xx.xx ± xx.xx	xx.xx ± xx.xx	xx.xx ± xx.xx	xx.xx ± xx.xx

n – Number of Subjects

Values are arithmetic mean (A<sub>mean</sub>) ± standard deviation

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

*Note: This table will present the summary statistics of the pharmacokinetic parameters calculated for Part 3 Pharmacokinetic Analysis Population. In case a pharmacokinetic parameter could not be calculated for one or more subjects in the Part 1 Pharmacokinetic Analysis Population, the pharmacokinetic parameter should be followed by the # symbol and accompanied by a footnote: '# For <Parameter(s)>, n=xx, for <Investigational Product> product <, and n=xx, for <Investigational Product> product> '.*



**Table 1.H. Givinostat and Metabolites (ITF2374, ITF2375, ITF2440 and ITF2563): Summary Statistics of the Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1)<sup>21</sup>**

**Table 1.I. Givinostat and Metabolites (ITF2374, ITF2375, ITF2440 and ITF2563): Summary Statistics of the Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13)<sup>21</sup>**

<sup>1</sup> To be generated as follows:

Parameter (unit)	Givinostat (n = xx)	ITF2374 (n = xx)	ITF2375 (n = xx)	ITF2440 (n = xx)	ITF2563 (n = xx)
R <sub>max,ss</sub> (<units>)	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]
t <sub>max</sub> (h)	xx.xx (xx.xx – xx.xx)	xx.xx (xx.xx – xx.xx)	xx.xx (xx.xx – xx.xx)	xx.xx (xx.xx – xx.xx)	xx.xx (xx.xx – xx.xx)
Amt <sub>CUM</sub> (<units>)	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]
AURC <sub>0-t</sub> (<units>)	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]
REC% (%)	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]
CL <sub>R/F</sub> (L/h)	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]	xx.xx (xx.x%) [xx.xx – xx.xx]

n – Number of Subjects

Values are geometric mean (G<sub>mean</sub>) with geometric coefficient of variation (GCV%) within parenthesis and 95% confidence interval of the G<sub>mean</sub> within square brackets

t<sub>max</sub> values are median with range between parentheses

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

*Note: This table will present the summary statistics of the pharmacokinetic parameters calculated for Part 3 Pharmacokinetic Analysis Population. In case a pharmacokinetic parameter could not be calculated for one or more subjects in the Part 1 Pharmacokinetic Analysis Population, the pharmacokinetic parameter should be followed by the # symbol and accompanied by a footnote: '# For <Parameter(s)>, n=xx, for <Investigational Product> product <, and n=xx, for <Investigational Product> product'.*

**Table 1.J. Summary of Treatment-Emergent Adverse Events (TEAEs)**

	<b>Total (N = xx)</b>	
	<b>All TEAEs</b>	<b>Drug-related TEAEs</b>
<b>Number of Subjects {Nr. of TEAEs} (% of Subjects)</b>	xx {xx} (xx%)	xx {xx} (xx%)
<b>MedDRA</b>		
<b>System Organ Class (SOC)</b>		
<b>Preferred Term (PT)</b>		
<TEAEs SOC>	xx {xx} (xx%)	xx {xx} (xx%)
<TEAEs SOC>	xx {xx} (xx%)	xx {xx} (xx%)
<Insert as many rows as deemed necessary>		
<b>Severity</b>		
<b>Mild</b>	xx {xx} (xx%)	xx {xx} (xx%)
<b>Moderate</b>	xx {xx} (xx%)	xx {xx} (xx%)
<b>Severe</b>	xx {xx} (xx%)	xx {xx} (xx%)

MedDRA – Medical Dictionary for Regulatory Activities

Drug-related TEAEs are those which causality was assessed as “Reasonably Possible” by the Investigator.

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>



**Figure 1.A.1. Givinostat: Geometric Mean (95% CI) Plasma Concentration *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

**Figure 1.A.2. Givinostat: Geometric Mean (95% CI) Plasma Concentration *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Semi-Logarithmic Scale**

**Figure 1.A.3. Givinostat: Geometric Mean (95% CI) Plasma Concentration *Versus* Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale**

**Figure 1.A.4. Givinostat: Geometric Mean (95% CI) Plasma Concentration *Versus* Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Semi-Logarithmic Scale**

**Figure 1.A.5. Givinostat: Geometric Mean (95% CI) Trough Plasma Concentration *Versus* Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale**

**Figure 1.B.1. Givinostat Metabolite (ITF2374): Geometric Mean (95% CI) Plasma Concentration *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

**Figure 1.B.2. Givinostat Metabolite (ITF2374): Geometric Mean (95% CI) Plasma Concentration *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Semi-Logarithmic Scale**

**Figure 1.B.3. Givinostat Metabolite (ITF2374): Geometric Mean (95% CI) Plasma Concentration *Versus* Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale**

**Figure 1.B.4. Givinostat Metabolite (ITF2374): Geometric Mean (95% CI) Plasma Concentration *Versus* Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Semi-Logarithmic Scale**

**Figure 1.B.5. Givinostat Metabolite (ITF2374): Geometric Mean (95% CI) Trough Plasma Concentration *Versus* Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale**

**Figure 1.C.1. Givinostat Metabolite (ITF2375): Geometric Mean (95% CI) Plasma Concentration Versus Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

**Figure 1.C.2. Givinostat Metabolite (ITF2375): Geometric Mean (95% CI) Plasma Concentration Versus Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Semi-Logarithmic Scale**

**Figure 1.C.3. Givinostat Metabolite (ITF2375): Geometric Mean (95% CI) Plasma Concentration Versus Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale**

**Figure 1.C.4. Givinostat Metabolite (ITF2375): Geometric Mean (95% CI) Plasma Concentration Versus Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Semi-Logarithmic Scale**

**Figure 1.C.5. Givinostat Metabolite (ITF2375): Geometric Mean (95% CI) Trough Plasma Concentration Versus Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale**

**Figure 1.D.1. Givinostat Metabolite (ITF2440): Geometric Mean (95% CI) Plasma Concentration Versus Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

**Figure 1.D.2. Givinostat Metabolite (ITF2440): Geometric Mean (95% CI) Plasma Concentration Versus Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Semi-Logarithmic Scale**

**Figure 1.D.3. Givinostat Metabolite (ITF2440): Geometric Mean (95% CI) Plasma Concentration Versus Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale**

**Figure 1.D.4. Givinostat Metabolite (ITF2440): Geometric Mean (95% CI) Plasma Concentration Versus Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Semi-Logarithmic Scale**

**Figure 1.D.5. Givinostat Metabolite (ITF2440): Geometric Mean (95% CI) Trough Plasma Concentration Versus Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale**



**Figure 1.E.1. Givinostat Metabolite (ITF2563): Geometric Mean (95% CI) Plasma Concentration Versus Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

**Figure 1.E.2. Givinostat Metabolite (ITF2563): Geometric Mean (95% CI) Plasma Concentration Versus Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Semi-Logarithmic Scale**

**Figure 1.E.3. Givinostat Metabolite (ITF2563): Geometric Mean (95% CI) Plasma Concentration Versus Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale**

**Figure 1.E.4. Givinostat Metabolite (ITF2563): Geometric Mean (95% CI) Plasma Concentration Versus Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Semi-Logarithmic Scale**

**Figure 1.E.5. Givinostat Metabolite (ITF2563): Geometric Mean (95% CI) Trough Plasma Concentration Versus Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale**

**Figure 1.F.1. Givinostat: Geometric Mean (95% CI) of Urinary Excretion *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

**Figure 1.F.2. Givinostat: Geometric Mean (95% CI) of Cumulative Amount Urinary Excretion *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

**Figure 1.G.1. Givinostat Metabolite (ITF2374): Geometric Mean (95% CI) of Urinary Excretion *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

**Figure 1.G.2. Givinostat Metabolite (ITF2374): Geometric Mean (95% CI) of Cumulative Amount Urinary Excretion *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

**Figure 1.H.1. Givinostat Metabolite (ITF2375): Geometric Mean (95% CI) of Urinary Excretion *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

**Figure 1.H.2. Givinostat Metabolite (ITF2375): Geometric Mean (95% CI) of Cumulative Amount Urinary Excretion *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

**Figure 1.I.1. Givinostat Metabolite (ITF2440): Geometric Mean (95% CI) of Urinary Excretion *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

**Figure 1.I.2. Givinostat Metabolite (ITF2440): Geometric Mean (95% CI) of Cumulative Amount Urinary Excretion *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

**Figure 1.J.1. Givinostat Metabolite (ITF2563): Geometric Mean (95% CI) of Urinary Excretion *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

**Figure 1.J.2. Givinostat Metabolite (ITF2563): Geometric Mean (95% CI) of Cumulative Amount Urinary Excretion *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**



### A.3. Tables, Figures and Graphs Referred to But Not Included in the Text (Section 14 of CSR)

#### 14.1.1. SAFETY DATA –

##### A.3.1. Part 3

#### 14.2. PART 3

##### 14.2.1. DEMOGRAPHIC DATA – PART 3

###### 14.2.1.1. Demographic Data – Part 3 Safety Analysis Population <sup>2</sup>

###### 14.2.1.2. Demographic Data – Part 3 Single-Dose Pharmacokinetic Analysis Population <sup>2</sup>

###### 14.2.1.3. Demographic Data – Part 3 Multiple-Dose Pharmacokinetic Analysis Population <sup>2</sup>

<sup>2</sup> To be generated as follows:

Subject No.	Sex	Race	Age (years)	Height (cm)	Weight (kg)	BMI (kg/m <sup>2</sup> )
<Subject ID>	<Male / Female>	<Race>	XX	XXX	XX.X	XX.X
<Insert as many rows as deemed necessary>						
		n	XX	XX	XX	XX
		Mean	XX	XXX	XX.X	XX.X
		SD	XX.X	XX.X	XX.XX	XX.XX
		Median	XX	XXX	XX.X	XX.X
		Minimum	XX	XXX	XX.X	XX.X
		Maximum	XX	XXX	XX.X	XX.X

BMI – Body Mass Index; n – Number of Subjects; SD – Standard Deviation

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

###### 14.2.1.4. Subjects Prematurely Discontinued After First Dosing

Subject No.	Time of Discontinuation	Reason for Discontinuation
<Subject ID>	<Day # >	<Reason>
<Insert as many rows as deemed necessary>		

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

## 14.2.2. PHARMACOKINETIC DATA – PART 3

### 14.2.2.1. Plasma Pharmacokinetic Data

#### 14.2.2.1.1. Deviations from Blood Sampling Schedule

##### 14.2.2.1.1.1. Deviations from Blood Sampling Schedule

Subject No.	Day of Dose	Analyte	Planned Time (h:min)	Deviation (h:min)	Reason for Deviation
<Subject>	<Day #>	<Analyte>	<Planned blood sampling time>	<Deviation time>	<Reason for deviation (e.g. Difficulty with blood sampling / Subject unavailable at scheduled time)>

<Insert as many rows as deemed necessary>

Program: <SAS Program>  
Execution Date/Time: <ddMMMyyyy HH:MM>

##### 14.2.2.1.1.2. Missing Blood Samples

Subject No.	Day of Dose	Analyte	Planned Time (h:min)	Reason for Missing Blood Sample
<Subject ID>	<Day #>	<Analyte>	<Planned blood sampling time>	<Reason for missing blood sample>

<Insert as many rows as deemed necessary>

Program: <SAS Program>  
Execution Date/Time: <ddMMMyyyy HH:MM>



#### 14.2.2.1.1.3. Other Important Protocol Deviations

Subject No.	Day of Dose	Analyte	Category	Description
<Subject ID>	<Day #>	<Analyte>	<Deviation category>	<Description of deviation>
<Insert as many rows as deemed necessary>				

Program: <SAS Program>  
Execution Date/Time: <ddMMMyyyy HH:MM>

#### **14.2.2.1.2. Plasma Concentrations**

**14.2.2.1.2.1. Givinostat: Individual Data and Descriptive Statistics of Plasma Concentrations Following Single-Dose Administration of Givinostat (Day 1) <sup>3</sup>**

**14.2.2.1.2.2. Givinostat: Individual Data and Descriptive Statistics of Plasma Concentrations Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>3</sup>**

**14.2.2.1.2.3. Givinostat: Individual Data and Descriptive Statistics of Plasma Concentrations Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>3</sup>**

**14.2.2.1.2.4. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Plasma Concentrations Following Single-Dose Administration of Givinostat (Day 1) <sup>3</sup>**

**14.2.2.1.2.5. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Plasma Concentrations Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>3</sup>**

**14.2.2.1.2.6. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Plasma Concentrations Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>3</sup>**

**14.2.2.1.2.7. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Plasma Concentrations Following Single-Dose Administration of Givinostat (Day 1) <sup>3</sup>**

**14.2.2.1.2.8. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Plasma Concentrations Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>3</sup>**

**14.2.2.1.2.9. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Plasma Concentrations Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>3</sup>**

**14.2.2.1.2.10. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Plasma Concentrations Following Single-Dose Administration of Givinostat (Day 1) <sup>3</sup>**

**14.2.2.1.2.11. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Plasma Concentrations Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>3</sup>**

**14.2.2.1.2.12. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Plasma Concentrations Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>3</sup>**

**14.2.2.1.2.13. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Plasma Concentrations Following Single-Dose Administration of Givinostat (Day 1) <sup>3</sup>**

**14.2.2.1.2.14. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Plasma Concentrations Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>3</sup>**

**14.2.2.1.2.15. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Plasma Concentrations Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>3</sup>**



<sup>3</sup> To be generated as follows:

Subject No.	Pre-dose	Time Post-Dose (h)									
		XX	XX	XX	XX	XX	XX	XX	XX	XX	XX
<Subject ID>	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
<Insert as many rows as deemed necessary>											
n	XX	XX	XX	XX	XX	XX	XX	XX	XX	XX	XX
G <sub>mean</sub> (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
A <sub>mean</sub> (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
GSD (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
SD (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
GCV%	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X
CV%	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X
G <sub>mean</sub> 95% CI Lower (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
G <sub>mean</sub> 95% CI Upper (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
A <sub>mean</sub> 95% CI Lower (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
A <sub>mean</sub> 95% CI Upper (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
Median (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
Minimum (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
Maximum (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX

n – Number of Subjects; G<sub>mean</sub> – Geometric Mean; A<sub>mean</sub> – Arithmetic Mean; GSD – Geometric Standard Deviation; SD – Standard Deviation; GCV% – Geometric Coefficient of Variation; CV% – Coefficient of Variation; G<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Geometric Mean; G<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Geometric Mean; A<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Arithmetic Mean; A<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Arithmetic Mean.

BLQ – Below the Limit of Quantification (LLOQ = <LLOQ> <units>) of the assay (taken as missing for the calculation of log-transformed statistics and as zero for the calculation of the remaining summary statistics)

ND – Not Done

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>



**14.2.2.1.2.16. Givinostat: Individual Data and Descriptive Statistics of Plasma Concentrations Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) <sup>4</sup>**

**14.2.2.1.2.17. Givinostat: Individual Data and Descriptive Statistics of Plasma Concentrations Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Male Subjects <sup>4</sup>**

**14.2.2.1.2.18. Givinostat: Individual Data and Descriptive Statistics of Plasma Concentrations Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Female Subjects <sup>4</sup>**

**14.2.2.1.2.19. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Plasma Concentrations Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) <sup>4</sup>**

**14.2.2.1.2.20. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Plasma Concentrations Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Male Subjects <sup>4</sup>**

**14.2.2.1.2.21. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Plasma Concentrations Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Female Subjects <sup>4</sup>**

**14.2.2.1.2.22. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Plasma Concentrations Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) <sup>4</sup>**

**14.2.2.1.2.23. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Plasma Concentrations Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Male Subjects <sup>4</sup>**

**14.2.2.1.2.24. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Plasma Concentrations Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Female Subjects <sup>4</sup>**

**14.2.2.1.2.25. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Plasma Concentrations Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) <sup>4</sup>**

**14.2.2.1.2.26. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Plasma Concentrations Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Male Subjects <sup>4</sup>**

**14.2.2.1.2.27. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Plasma Concentrations Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Female Subjects <sup>4</sup>**

**14.2.2.1.2.28. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Plasma Concentrations Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) <sup>4</sup>**

**14.2.2.1.2.29. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Plasma**



**Concentrations Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Male Subjects<sup>4</sup>**

**14.2.2.1.2.30. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Plasma Concentrations Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Female Subjects<sup>4</sup>**

<sup>4</sup> To be generated as follows:

Subject No.	Pre-Dose					Time Post-Day 13 Dose (h)					
	Day 9	Day 10	Day 11	Day 12	Day 13	1.11	1.11	1.11	1.11	1.11	1.11
<Subject ID>	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
<Insert as many rows as deemed necessary>											
n	XX	XX	XX	XX	XX	XX	XX	XX	XX	XX	XX
G <sub>mean</sub> (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
A <sub>mean</sub> (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
GSD (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
SD (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
GCV%	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X
CV%	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X
G <sub>mean</sub> 95% CI Lower (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
G <sub>mean</sub> 95% CI Upper (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
A <sub>mean</sub> 95% CI Lower (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
A <sub>mean</sub> 95% CI Upper (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
Median (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
Minimum (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
Maximum (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX

n – Number of Subjects; G<sub>mean</sub> – Geometric Mean; A<sub>mean</sub> – Arithmetic Mean; GSD – Geometric Standard Deviation; SD – Standard Deviation; GCV% – Geometric Coefficient of Variation; CV% – Coefficient of Variation; G<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Geometric Mean; G<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Geometric Mean; A<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Arithmetic Mean; A<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Arithmetic Mean.

BLQ – Below the Limit of Quantification (LLOQ = <LLOQ> <units>) of the assay (taken as missing for the calculation of log-transformed statistics and as zero for the calculation of the remaining summary statistics)

ND – Not Done

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>



#### **14.2.2.1.3. Plasma Pharmacokinetic Parameters**

**14.2.2.1.3.1. Givinostat: Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) <sup>3</sup>**

**14.2.2.1.3.2. Givinostat: Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>3</sup>**

**14.2.2.1.3.3. Givinostat: Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>3</sup>**

**14.2.2.1.3.4. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) <sup>3</sup>**

**14.2.2.1.3.5. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>3</sup>**

**14.2.2.1.3.6. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>3</sup>**

**14.2.2.1.3.7. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) <sup>3</sup>**

**14.2.2.1.3.8. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>3</sup>**

**14.2.2.1.3.9. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>3</sup>**

**14.2.2.1.3.10. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) <sup>3</sup>**

**14.2.2.1.3.11. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>3</sup>**

**14.2.2.1.3.12. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>3</sup>**

**14.2.2.1.3.13. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) <sup>3</sup>**

**14.2.2.1.3.14. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>3</sup>**

**14.2.2.1.3.15. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>4</sup>**

<sup>4</sup> To be generated as follows:

Subject No.	C <sub>max</sub> (<unitsC>)	T <sub>max</sub> (h)	AUC <sub>0-t</sub> (<unitsA>)	AUC <sub>0-∞</sub> (<unitsA>)	%AUC <sub>extrap</sub> (%)	λ <sub>z</sub> (1/h)	t <sub>1/2</sub> (h)
<Subject ID>	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
<Insert as many rows as deemed necessary>							
n	XX	XX	XX	XX	XX	XX	XX
G <sub>mean</sub>	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
A <sub>mean</sub>	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
GSD	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
SD	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
GCV%	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X
CV%	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X
G <sub>mean</sub> 95% CI Lower	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
G <sub>mean</sub> 95% CI Upper	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
A <sub>mean</sub> 95% CI Lower	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
A <sub>mean</sub> 95% CI Upper	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
Median	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
Minimum	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
Maximum	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX

n – Number of Subjects; G<sub>mean</sub> – Geometric Mean; A<sub>mean</sub> – Arithmetic Mean; GSD – Geometric Standard Deviation; SD – Standard Deviation; GCV% – Geometric Coefficient of Variation; CV% – Coefficient of Variation; G<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Geometric Mean; G<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Geometric Mean; A<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Arithmetic Mean; A<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Arithmetic Mean.

NC – Not Calculated

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>



**14.2.2.1.3.16. Plasma Metabolic Ratios for Givinostat Metabolites (ITF2374, ITF2375, ITF2440 and ITF2563) Following Single-Dose Administration of Givinostat (Day 1)**

Subject No.	ITF2374	ITF2375	ITF2440	ITF2563
<Subject ID>	XX.XX	XX.XX	XX.XX	XX.XX
<Insert as many rows as deemed necessary>				
n	XX	XX	XX	XX
G <sub>mean</sub>	XX.XX	XX.XX	XX.XX	XX.XX
A <sub>mean</sub>	XX.XX	XX.XX	XX.XX	XX.XX
GSD	XX.XX	XX.XX	XX.XX	XX.XX
SD	XX.XX	XX.XX	XX.XX	XX.XX
GCV%	XX.X	XX.X	XX.X	XX.X
CV%	XX.X	XX.X	XX.X	XX.X
G <sub>mean</sub> 95% CI Lower	XX.XX	XX.XX	XX.XX	XX.XX
G <sub>mean</sub> 95% CI Upper	XX.XX	XX.XX	XX.XX	XX.XX
A <sub>mean</sub> 95% CI Lower	XX.XX	XX.XX	XX.XX	XX.XX
A <sub>mean</sub> 95% CI Upper	XX.XX	XX.XX	XX.XX	XX.XX
Median	XX.XX	XX.XX	XX.XX	XX.XX
Minimum	XX.XX	XX.XX	XX.XX	XX.XX
Maximum	XX.XX	XX.XX	XX.XX	XX.XX

n – Number of Subjects; G<sub>mean</sub> – Geometric Mean; A<sub>mean</sub> – Arithmetic Mean; GSD – Geometric Standard Deviation; SD – Standard Deviation; GCV% – Geometric Coefficient of Variation; CV% – Coefficient of Variation; G<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Geometric Mean; G<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Geometric Mean; A<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Arithmetic Mean; A<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Arithmetic Mean.

NC – Not Calculated

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

**14.2.2.1.3.17. Givinostat: Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) <sup>4</sup>**

**14.2.2.1.3.18. Givinostat: Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) – Male Subjects <sup>4</sup>**

**14.2.2.1.3.19. Givinostat: Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) – Female Subjects <sup>4</sup>**

**14.2.2.1.3.20. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) <sup>4</sup>**

**14.2.2.1.3.21. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) – Male Subjects <sup>4</sup>**

**14.2.2.1.3.22. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) – Female Subjects <sup>4</sup>**

**14.2.2.1.3.23. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) <sup>4</sup>**

**14.2.2.1.3.24. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) – Male Subjects <sup>4</sup>**

**14.2.2.1.3.25. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) – Female Subjects <sup>4</sup>**

**14.2.2.1.3.26. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) <sup>4</sup>**

**14.2.2.1.3.27. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) – Male Subjects <sup>4</sup>**

**14.2.2.1.3.28. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) – Female Subjects <sup>4</sup>**

**14.2.2.1.3.29. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) <sup>4</sup>**

**14.2.2.1.3.30. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Plasma**



## Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) – Male Subjects <sup>4</sup>

### 14.2.2.1.3.31. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Plasma Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) – Female Subjects <sup>4</sup>

<sup>4</sup> To be generated as follows:

Subject No.	C <sub>max,ss</sub> ( <b>&lt;unitsC&gt;</b> )	T <sub>max,ss</sub> (h)	AUC <sub>0-τ,ss</sub> ( <b>&lt;unitsA&gt;</b> )	AUC <sub>0-t</sub> ( <b>&lt;unitsA&gt;</b> )	AUC <sub>0-∞</sub> ( <b>&lt;unitsA&gt;</b> )	λ <sub>z</sub> (1/h)	t <sub>1/2</sub> (h)	CL <sub>ss</sub> /F (L/h)	V <sub>D</sub> /F (L)
<Subject ID>	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX	XX.XX	XX.XX
<Insert as many rows as deemed necessary>									
n	XX	XX	XX	XX	XX	XX	XX	XX	XX
G <sub>mean</sub>	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX	XX.XX	XX.XX
A <sub>mean</sub>	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX	XX.XX	XX.XX
GSD	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX	XX.XX	XX.XX
SD	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX	XX.XX	XX.XX
GCV%	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X
CV%	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X
G <sub>mean</sub> 95% CI Lower	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX	XX.XX	XX.XX
G <sub>mean</sub> 95% CI Upper	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX	XX.XX	XX.XX
A <sub>mean</sub> 95% CI Lower	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX	XX.XX	XX.XX
A <sub>mean</sub> 95% CI Upper	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX	XX.XX	XX.XX
Median	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX	XX.XX	XX.XX
Minimum	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX	XX.XX	XX.XX
Maximum	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX	XX.XX	XX.XX

n – Number of Subjects; G<sub>mean</sub> – Geometric Mean; A<sub>mean</sub> – Arithmetic Mean; GSD – Geometric Standard Deviation; SD – Standard Deviation; GCV% – Geometric Coefficient of Variation; CV% – Coefficient of Variation; G<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Geometric Mean; G<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Geometric Mean; A<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Arithmetic Mean; A<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Arithmetic Mean.

NC – Not Calculated

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

**14.2.2.1.3.32. Givinostat: Individual Data and Descriptive Statistics of Accumulation Index <sup>4</sup>**

**14.2.2.1.3.33. Givinostat: Individual Data and Descriptive Statistics of Accumulation Index – Male Subjects <sup>4</sup>**

**14.2.2.1.3.34. Givinostat: Individual Data and Descriptive Statistics of Accumulation Index – Female Subjects <sup>4</sup>**

**14.2.2.1.3.35. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Accumulation Index <sup>4</sup>**

**14.2.2.1.3.36. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Accumulation Index – Male Subjects <sup>4</sup>**

**14.2.2.1.3.37. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Accumulation Index – Female Subjects <sup>4</sup>**

**14.2.2.1.3.38. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Accumulation Index <sup>4</sup>**

**14.2.2.1.3.39. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Accumulation Index – Male Subjects <sup>4</sup>**

**14.2.2.1.3.40. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Accumulation Index – Female Subjects <sup>4</sup>**

**14.2.2.1.3.41. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Accumulation Index <sup>4</sup>**

**14.2.2.1.3.42. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Accumulation Index – Male Subjects <sup>4</sup>**

**14.2.2.1.3.43. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Accumulation Index – Female Subjects <sup>4</sup>**

**14.2.2.1.3.44. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Accumulation Index <sup>4</sup>**

**14.2.2.1.3.45. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Accumulation Index – Male Subjects <sup>4</sup>**

**14.2.2.1.3.46. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Accumulation Index – Female Subjects <sup>4</sup>**



<sup>4</sup> To be generated as follows:

Subject No.	C <sub>max</sub>	AUC
<Subject ID>	xx.xx	xx.xx
<Insert as many rows as deemed necessary>		
n	xx	xx
A <sub>mean</sub>	xx.xx	xx.xx
SD	xx.xx	xx.xx
SE	xx.x	xx.x
Median	xx.xx	xx.xx
Minimum	xx.xx	xx.xx
Maximum	xx.xx	xx.xx

n – Number of Subjects; A<sub>mean</sub> – Arithmetic Mean; SD – Standard Deviation; SE – Standard Error.

NC – Not Calculated

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

**14.2.2.1.3.47. Plasma Metabolic Ratios for Givinostat Metabolites (ITF2374, ITF2375, ITF2440 and ITF2563) Following Multiple-Dose Administration of Givinostat (Day 13)**

Subject No.	ITF2374	ITF2375	ITF2440	ITF2563
<Subject ID>	XX.XX	XX.XX	XX.XX	XX.XX
<Insert as many rows as deemed necessary>				
n	XX	XX	XX	XX
G <sub>mean</sub>	XX.XX	XX.XX	XX.XX	XX.XX
A <sub>mean</sub>	XX.XX	XX.XX	XX.XX	XX.XX
GSD	XX.XX	XX.XX	XX.XX	XX.XX
SD	XX.XX	XX.XX	XX.XX	XX.XX
GCV%	XX.X	XX.X	XX.X	XX.X
CV%	XX.X	XX.X	XX.X	XX.X
G <sub>mean</sub> 95% CI Lower	XX.XX	XX.XX	XX.XX	XX.XX
G <sub>mean</sub> 95% CI Upper	XX.XX	XX.XX	XX.XX	XX.XX
A <sub>mean</sub> 95% CI Lower	XX.XX	XX.XX	XX.XX	XX.XX
A <sub>mean</sub> 95% CI Upper	XX.XX	XX.XX	XX.XX	XX.XX
Median	XX.XX	XX.XX	XX.XX	XX.XX
Minimum	XX.XX	XX.XX	XX.XX	XX.XX
Maximum	XX.XX	XX.XX	XX.XX	XX.XX

n – Number of Subjects; G<sub>mean</sub> – Geometric Mean; A<sub>mean</sub> – Arithmetic Mean; GSD – Geometric Standard Deviation; SD – Standard Deviation; GCV% – Geometric Coefficient of Variation; CV% – Coefficient of Variation; G<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Geometric Mean; G<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Geometric Mean; A<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Arithmetic Mean; A<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Arithmetic Mean.

NC – Not Calculated

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>



#### 14.2.2.1.4. Log-Linear Regression Parameters for $\lambda_z$ Estimation

##### 14.2.2.1.4.1. Givinostat: Log-Linear Regression Parameters for $\lambda_z$ Estimation Following Single-Dose Administration of Givinostat (Day 1) <sup>6</sup>

##### 14.2.2.1.4.2. Givinostat Metabolite (ITF2374): Log-Linear Regression Parameters for $\lambda_z$ Estimation Following Single-Dose Administration of Givinostat (Day 1) <sup>6</sup>

##### 14.2.2.1.4.3. Givinostat Metabolite (ITF2375): Log-Linear Regression Parameters for $\lambda_z$ Estimation Following Single-Dose Administration of Givinostat (Day 1) <sup>6</sup>

##### 14.2.2.1.4.4. Givinostat Metabolite (ITF2440): Log-Linear Regression Parameters for $\lambda_z$ Estimation Following Single-Dose Administration of Givinostat (Day 1) <sup>6</sup>

##### 14.2.2.1.4.5. Givinostat Metabolite (ITF2563): Log-Linear Regression Parameters for $\lambda_z$ Estimation Following Single-Dose Administration of Givinostat (Day 1) <sup>6</sup>

##### 14.2.2.1.4.6. Givinostat: Log-Linear Regression Parameters for $\lambda_z$ Estimation Following Multiple-Dose Administration of Givinostat (Day 13) <sup>6</sup>

##### 14.2.2.1.4.7. Givinostat Metabolite (ITF2374): Log-Linear Regression Parameters for $\lambda_z$ Estimation Following Multiple-Dose Administration of Givinostat (Day 13) <sup>6</sup>

##### 14.2.2.1.4.8. Givinostat Metabolite (ITF2375): Log-Linear Regression Parameters for $\lambda_z$ Estimation Following Multiple-Dose Administration of Givinostat (Day 13) <sup>6</sup>

##### 14.2.2.1.4.9. Givinostat Metabolite (ITF2440): Log-Linear Regression Parameters for $\lambda_z$ Estimation Following Multiple-Dose Administration of Givinostat (Day 13) <sup>6</sup>

##### 14.2.2.1.4.10. Givinostat Metabolite (ITF2563): Log-Linear Regression Parameters for $\lambda_z$ Estimation Following Multiple-Dose Administration of Givinostat (Day 13) <sup>6</sup>

<sup>6</sup> To be generated as follows:

Subject No.	Lower TLIN (h)	Upper TLIN (h)	No. of Time Points	RSQ
<Subject ID>	xx.xx	xx.xx	xx	x.xx
<Insert as many rows as deemed necessary>				

Lower TLIN – Lower Time Point Used in Regression Analysis; Upper TLIN – Upper Time Point Used in Regression Analysis;

RSQ – Goodness of Fit for the Terminal Elimination Phase

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

#### 14.2.2.2. Urinary Pharmacokinetic Data

##### 14.2.2.2.1. Deviations from Urine Sampling Schedule

###### 14.2.2.2.1.1. Missing Urine Samples

Subject No.	Day of Dose	Planned Start Time (h:min)	Planned End Time (h:min)	Reason for Missing Urine Sample
<Subject ID>	<Day #>	<Planned blood sampling time>	<Planned blood sampling time>	<Reason for missing blood sample>
<Insert as many rows as deemed necessary>				
Program: <SAS Program>				
Execution Date/Time: <ddMMMyyyy HH:MM>				



#### **14.2.2.2.2. Urinary Excretion Profile**

##### **14.2.2.2.2.1. Givinostat: Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) <sup>3</sup>**

##### **14.2.2.2.2.2. Givinostat: Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>3</sup>**

##### **14.2.2.2.2.3. Givinostat: Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>3</sup>**

##### **14.2.2.2.2.4. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) <sup>3</sup>**

##### **14.2.2.2.2.5. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>3</sup>**

##### **14.2.2.2.2.6. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>3</sup>**

##### **14.2.2.2.2.7. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) <sup>3</sup>**

##### **14.2.2.2.2.8. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>3</sup>**

##### **14.2.2.2.2.9. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>3</sup>**

##### **14.2.2.2.2.10. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) <sup>3</sup>**

##### **14.2.2.2.2.11. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>3</sup>**

##### **14.2.2.2.2.12. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>3</sup>**

##### **14.2.2.2.2.13. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) <sup>3</sup>**

##### **14.2.2.2.2.14. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>3</sup>**

##### **14.2.2.2.2.15. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>3</sup>**



**14.2.2.2.16. Givinostat: Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) <sup>3</sup>**

**14.2.2.2.17. Givinostat: Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>3</sup>**

**14.2.2.2.18. Givinostat: Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>3</sup>**

**14.2.2.2.19. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) <sup>3</sup>**

**14.2.2.2.20. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>3</sup>**

**14.2.2.2.21. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>3</sup>**

**14.2.2.2.22. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) <sup>3</sup>**

**14.2.2.2.23. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>3</sup>**

**14.2.2.2.24. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>3</sup>**

**14.2.2.2.25. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) <sup>3</sup>**

**14.2.2.2.26. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>3</sup>**

**14.2.2.2.27. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>3</sup>**



**14.2.2.2.28. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics Cumulative Amount Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) <sup>3</sup>**

**14.2.2.2.29. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>3</sup>**

**14.2.2.2.30. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>3</sup>**

<sup>4</sup> To be generated as follows:

Subject No.	Pre-Dose (Day -1)	Post-Day 1 Dose					
		0h – 12h	12h – 24h	24h – 36h	36h – 48h	48h – 72h	72h – 96h
<Subject ID>	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
<Insert as many rows as deemed necessary>							
n	XX	XX	XX	XX	XX	XX	XX
G <sub>mean</sub> (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
A <sub>mean</sub> (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
GSD (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
SD (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
GCV%	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X
CV%	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X
G <sub>mean</sub> 95% CI Lower (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
G <sub>mean</sub> 95% CI Upper (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
A <sub>mean</sub> 95% CI Lower (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
A <sub>mean</sub> 95% CI Upper (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
Median (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
Minimum (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
Maximum (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX

n – Number of Subjects; G<sub>mean</sub> – Geometric Mean; A<sub>mean</sub> – Arithmetic Mean; GSD – Geometric Standard Deviation; SD – Standard Deviation; GCV% – Geometric Coefficient of Variation; CV% – Coefficient of Variation; G<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Geometric Mean; G<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Geometric Mean; A<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Arithmetic Mean; A<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Arithmetic Mean.

ND – Not Done

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>



**14.2.2.2.31. Givinostat: Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) <sup>3</sup>**

**14.2.2.2.32. Givinostat: Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) – Male Subjects <sup>3</sup>**

**14.2.2.2.33. Givinostat: Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) – Female Subjects <sup>3</sup>**

**14.2.2.2.34. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) <sup>3</sup>**

**14.2.2.2.35. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) – Male Subjects <sup>3</sup>**

**14.2.2.2.36. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) – Female Subjects <sup>3</sup>**

**14.2.2.2.37. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) <sup>3</sup>**

**14.2.2.2.38. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) – Male Subjects <sup>3</sup>**

**14.2.2.2.39. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) – Female Subjects <sup>3</sup>**

**14.2.2.2.40. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) <sup>3</sup>**

**14.2.2.2.41. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) – Male Subjects <sup>3</sup>**

**14.2.2.2.42. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) – Female Subjects <sup>3</sup>**

**14.2.2.2.43. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) <sup>3</sup>**

**14.2.2.2.44. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) – Male Subjects <sup>3</sup>**

**14.2.2.2.45. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) – Female Subjects <sup>3</sup>**



**14.2.2.2.2.46. Givinostat: Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) <sup>3</sup>**

**14.2.2.2.2.47. Givinostat: Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) – Male Subjects <sup>3</sup>**

**14.2.2.2.2.48. Givinostat: Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) – Female Subjects <sup>3</sup>**

**14.2.2.2.2.49. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) <sup>3</sup>**

**14.2.2.2.2.50. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) – Male Subjects <sup>3</sup>**

**14.2.2.2.2.51. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) – Female Subjects <sup>3</sup>**

**14.2.2.2.2.52. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) <sup>3</sup>**

**14.2.2.2.2.53. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) – Male Subjects <sup>3</sup>**

**14.2.2.2.2.54. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) – Female Subjects <sup>3</sup>**

**14.2.2.2.2.55. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) <sup>3</sup>**

**14.2.2.2.2.56. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) – Male Subjects <sup>3</sup>**

**14.2.2.2.2.57. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) – Female Subjects <sup>3</sup>**



**14.2.2.2.58. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics Cumulative Amount Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13)<sup>3</sup>**

**14.2.2.2.59. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) – Male Subjects<sup>3</sup>**

**14.2.2.2.60. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Cumulative Amount Urinary Excretion Profile Following Multiple-Dose Administration of Givinostat (Day 13) – Female Subjects<sup>3</sup>**

<sup>4</sup> To be generated as follows:

Subject No.	Post-Day 13 Dose					
	0h – 12h	12h – 24h	24h – 36h	36h – 48h	48h – 72h	72h – 96h
<Subject ID>	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
<Insert as many rows as deemed necessary>						
n	XX	XX	XX	XX	XX	XX
G <sub>mean</sub> (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
A <sub>mean</sub> (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
GSD (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
SD (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
GCV%	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X
CV%	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X
G <sub>mean</sub> 95% CI Lower (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
G <sub>mean</sub> 95% CI Upper (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
A <sub>mean</sub> 95% CI Lower (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
A <sub>mean</sub> 95% CI Upper (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
Median (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
Minimum (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX
Maximum (<units>)	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX	XX.XX

n – Number of Subjects; G<sub>mean</sub> – Geometric Mean; A<sub>mean</sub> – Arithmetic Mean; GSD – Geometric Standard Deviation; SD – Standard Deviation; GCV% – Geometric Coefficient of Variation; CV% – Coefficient of Variation; G<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Geometric Mean; G<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Geometric Mean; A<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Arithmetic Mean; A<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Arithmetic Mean.

ND – Not Done

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>



#### **14.2.2.2.3. Urinary Pharmacokinetic Parameters**

**14.2.2.2.3.1. Givinostat: Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) <sup>4</sup>**

**14.2.2.2.3.2. Givinostat: Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>4</sup>**

**14.2.2.2.3.3. Givinostat: Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>4</sup>**

**14.2.2.2.3.4. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) <sup>4</sup>**

**14.2.2.2.3.5. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>4</sup>**

**14.2.2.2.3.6. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>4</sup>**

**14.2.2.2.3.7. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) <sup>4</sup>**

**14.2.2.2.3.8. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>4</sup>**

**14.2.2.2.3.9. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>4</sup>**

**14.2.2.2.3.10. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) <sup>4</sup>**

**14.2.2.2.3.11. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>4</sup>**

**14.2.2.2.3.12. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>4</sup>**

**14.2.2.2.3.13. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) <sup>4</sup>**

**14.2.2.2.3.14. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects <sup>4</sup>**

**14.2.2.2.3.15. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects <sup>4</sup>**

<sup>4</sup> To be generated as follows:

Subject No.	R <sub>max</sub> ( <i>&lt;units&gt;</i> )	t <sub>u,max</sub> (h)	Amt <sub>CUM</sub> ( <i>&lt;units&gt;</i> )	AURC <sub>0-t</sub> ( <i>&lt;units&gt;</i> )	REC % (%)	CL <sub>R/F</sub> (L/h)
<Subject ID>	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX
<Insert as many rows as deemed necessary>						
n	XX	XX	XX	XX	XX	XX
G <sub>mean</sub>	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX
A <sub>mean</sub>	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX
GSD	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX
SD	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX
GCV%	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X
CV%	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X
G <sub>mean</sub> 95% CI Lower	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX
G <sub>mean</sub> 95% CI Upper	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX
A <sub>mean</sub> 95% CI Lower	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX
A <sub>mean</sub> 95% CI Upper	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX
Median	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX
Minimum	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX
Maximum	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX

n – Number of Subjects; G<sub>mean</sub> – Geometric Mean; A<sub>mean</sub> – Arithmetic Mean; GSD – Geometric Standard Deviation; SD – Standard Deviation; GCV% – Geometric Coefficient of Variation; CV% – Coefficient of Variation; G<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Geometric Mean; G<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Geometric Mean; A<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Arithmetic Mean; A<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Arithmetic Mean.

NC – Not Calculated

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>



**14.2.2.2.3.16. Percentage of Dose Recovered in Urine for Givinostat and Givinostat Metabolites (ITF2374, ITF2375, ITF2440 and ITF2563) Following Single-Dose Administration of Givinostat (Day 1)**

Subject No.	Givinostat	ITF2374	ITF2375	ITF2440	ITF2563	Total
<Subject ID>	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
<Insert as many rows as deemed necessary>						
n	XX	XX	XX	XX	XX	XX
G <sub>mean</sub>	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
A <sub>mean</sub>	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
GSD	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
SD	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
GCV%	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X
CV%	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X
G <sub>mean</sub> 95% CI Lower	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
G <sub>mean</sub> 95% CI Upper	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
A <sub>mean</sub> 95% CI Lower	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
A <sub>mean</sub> 95% CI Upper	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
Median	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
Minimum	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
Maximum	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX

n – Number of Subjects; G<sub>mean</sub> – Geometric Mean; A<sub>mean</sub> – Arithmetic Mean; GSD – Geometric Standard Deviation; SD – Standard Deviation; GCV% – Geometric Coefficient of Variation; CV% – Coefficient of Variation; G<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Geometric Mean; G<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Geometric Mean; A<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Arithmetic Mean; A<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Arithmetic Mean.

Total corresponds to the sum of percentage of dose recovered in urine for all analytes

NC – Not Calculated

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

**14.2.2.2.3.17. Givinostat: Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) <sup>4</sup>**

**14.2.2.2.3.18. Givinostat: Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) – Male Subjects <sup>4</sup>**

**14.2.2.2.3.19. Givinostat: Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) – Female Subjects <sup>4</sup>**

**14.2.2.2.3.20. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) <sup>4</sup>**

**14.2.2.2.3.21. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) – Male Subjects <sup>4</sup>**

**14.2.2.2.3.22. Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) – Female Subjects <sup>4</sup>**

**14.2.2.2.3.23. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) <sup>4</sup>**

**14.2.2.2.3.24. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) – Male Subjects <sup>4</sup>**

**14.2.2.2.3.25. Givinostat Metabolite (ITF2375): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) – Female Subjects <sup>4</sup>**

**14.2.2.2.3.26. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) <sup>4</sup>**

**14.2.2.2.3.27. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) – Male Subjects <sup>4</sup>**

**14.2.2.2.3.28. Givinostat Metabolite (ITF2440): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) – Female Subjects <sup>4</sup>**



**14.2.2.2.3.29. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) <sup>4</sup>**

**14.2.2.2.3.30. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) – Male Subjects <sup>4</sup>**

**14.2.2.2.3.31. Givinostat Metabolite (ITF2563): Individual Data and Descriptive Statistics of Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13) – Female Subjects <sup>4</sup>**

<sup>4</sup> To be generated as follows:

Subject No.	R <sub>max</sub> ( <i>&lt;units&gt;</i> )	t <sub>u,max</sub> (h)	Amt <sub>CUM</sub> ( <i>&lt;units&gt;</i> )	AURC <sub>0-t</sub> ( <i>&lt;units&gt;</i> )	REC % (%)	CL <sub>R/F</sub> (L/h)
<Subject ID>	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX
<Insert as many rows as deemed necessary>						
n	XX	XX	XX	XX	XX	XX
G <sub>mean</sub>	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX
A <sub>mean</sub>	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX
GSD	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX
SD	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX
GCV%	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X
CV%	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X
G <sub>mean</sub> 95% CI Lower	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX
G <sub>mean</sub> 95% CI Upper	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX
A <sub>mean</sub> 95% CI Lower	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX
A <sub>mean</sub> 95% CI Upper	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX
Median	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX
Minimum	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX
Maximum	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	X.XXX

n – Number of Subjects; G<sub>mean</sub> – Geometric Mean; A<sub>mean</sub> – Arithmetic Mean; GSD – Geometric Standard Deviation; SD – Standard Deviation; GCV% – Geometric Coefficient of Variation; CV% – Coefficient of Variation; G<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Geometric Mean; G<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Geometric Mean; A<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Arithmetic Mean; A<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Arithmetic Mean.

NC – Not Calculated

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

**14.2.2.2.3.32. Percentage of Dose Recovered in Urine for Givinostat and Givinostat Metabolites (ITF2374, ITF2375, ITF2440 and ITF2563) Following Multiple-Dose Administration of Givinostat (Day 13)**

Subject No.	Givinostat	ITF2374	ITF2375	ITF2440	ITF2563	Total
<Subject ID>	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
<Insert as many rows as deemed necessary>						
n	XX	XX	XX	XX	XX	XX
G <sub>mean</sub>	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
A <sub>mean</sub>	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
GSD	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
SD	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
GCV%	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X
CV%	XX.X	XX.X	XX.X	XX.X	XX.X	XX.X
G <sub>mean</sub> 95% CI Lower	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
G <sub>mean</sub> 95% CI Upper	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
A <sub>mean</sub> 95% CI Lower	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
A <sub>mean</sub> 95% CI Upper	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
Median	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
Minimum	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
Maximum	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX

n – Number of Subjects; G<sub>mean</sub> – Geometric Mean; A<sub>mean</sub> – Arithmetic Mean; GSD – Geometric Standard Deviation; SD – Standard Deviation; GCV% – Geometric Coefficient of Variation; CV% – Coefficient of Variation; G<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Geometric Mean; G<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Geometric Mean; A<sub>mean</sub> 95% CI Lower – Lower Limit of the 95% Confidence Interval for the Arithmetic Mean; A<sub>mean</sub> 95% CI Upper – Upper Limit of the 95% Confidence Interval for the Arithmetic Mean.

Total corresponds to the sum of percentage of dose recovered in urine for all analytes

NC – Not Calculated

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>



#### **14.2.2.3. Plasma Concentration *Versus* Time Profiles of All Subjects**

##### **14.2.2.3.1. Givinostat: Plasma Concentration *Versus* Time Profiles of All Subjects Following Single-Dose Administration of Givinostat (Day 1) – Linear Scale**

##### **14.2.2.3.2. Givinostat: Plasma Concentration *Versus* Time Profiles of All Subjects Following Multiple-Dose Administration of Givinostat (Day 13) – Linear Scale**

##### **14.2.2.3.3. Givinostat Metabolite (ITF2374): Plasma Concentration *Versus* Time Profiles of All Subjects Following Single-Dose Administration of Givinostat (Day 1) – Linear Scale**

##### **14.2.2.3.4. Givinostat Metabolite (ITF2374): Plasma Concentration *Versus* Time Profiles of All Subjects Following Multiple-Dose Administration of Givinostat (Day 13) – Linear Scale**

##### **14.2.2.3.5. Givinostat Metabolite (ITF2375): Plasma Concentration *Versus* Time Profiles of All Subjects Following Single-Dose Administration of Givinostat (Day 1) – Linear Scale**

##### **14.2.2.3.6. Givinostat Metabolite (ITF2375): Plasma Concentration *Versus* Time Profiles of All Subjects Following Multiple-Dose Administration of Givinostat (Day 13) – Linear Scale**

##### **14.2.2.3.7. Givinostat Metabolite (ITF2440): Plasma Concentration *Versus* Time Profiles of All Subjects Following Single-Dose Administration of Givinostat (Day 1) – Linear Scale**

##### **14.2.2.3.8. Givinostat Metabolite (ITF2440): Plasma Concentration *Versus* Time Profiles of All Subjects Following Multiple-Dose Administration of Givinostat (Day 13) – Linear Scale**

##### **14.2.2.3.9. Givinostat Metabolite (ITF2563): Plasma Concentration *Versus* Time Profiles of All Subjects Following Single-Dose Administration of Givinostat (Day 1) – Linear Scale**

##### **14.2.2.3.10. Givinostat Metabolite (ITF2563): Plasma Concentration *Versus* Time Profiles of All Subjects Following Multiple-Dose Administration of Givinostat (Day 13) – Linear Scale**



#### **14.2.2.4. Individual Plasma Concentration *Versus* Time Profiles**

**14.2.2.4.1. Givinostat: Individual Plasma Concentration *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

**14.2.2.4.2. Givinostat: Individual Plasma Concentration *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Semi-Logarithmic Scale**

**14.2.2.4.3. Givinostat: Individual Plasma Concentration *Versus* Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale**

**14.2.2.4.4. Givinostat: Individual Plasma Concentration *Versus* Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Semi-Logarithmic Scale**

**14.2.2.4.5. Givinostat: Individual Trough Plasma Concentration *Versus* Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale**

**14.2.2.4.6. Givinostat Metabolite (ITF2374): Individual Plasma Concentration *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

**14.2.2.4.7. Givinostat Metabolite (ITF2374): Individual Plasma Concentration *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Semi-Logarithmic Scale**

**14.2.2.4.8. Givinostat Metabolite (ITF2374): Individual Plasma Concentration *Versus* Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale**

**14.2.2.4.9. Givinostat Metabolite (ITF2374): Individual Plasma Concentration *Versus* Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Semi-Logarithmic Scale**

**14.2.2.4.10. Givinostat Metabolite (ITF2374): Individual Trough Plasma Concentration *Versus* Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale**

**14.2.2.4.11. Givinostat Metabolite (ITF2375): Individual Plasma Concentration *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

**14.2.2.4.12. Givinostat Metabolite (ITF2375): Individual Plasma Concentration *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Semi-Logarithmic Scale**

**14.2.2.4.13. Givinostat Metabolite (ITF2375): Individual Plasma Concentration *Versus* Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale**

**14.2.2.4.14. Givinostat Metabolite (ITF2375): Individual Plasma Concentration *Versus* Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Semi-Logarithmic Scale**

**14.2.2.4.15. Givinostat Metabolite (ITF2375): Individual Trough Plasma Concentration *Versus* Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale**



**14.2.2.4.16. Givinostat Metabolite (ITF2440): Individual Plasma Concentration *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

**14.2.2.4.17. Givinostat Metabolite (ITF2440): Individual Plasma Concentration *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Semi-Logarithmic Scale**

**14.2.2.4.18. Givinostat Metabolite (ITF2440): Individual Plasma Concentration *Versus* Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale**

**14.2.2.4.19. Givinostat Metabolite (ITF2440): Individual Plasma Concentration *Versus* Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Semi-Logarithmic Scale**

**14.2.2.4.20. Givinostat Metabolite (ITF2440): Individual Trough Plasma Concentration *Versus* Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale**

**14.2.2.4.21. Givinostat Metabolite (ITF2563): Individual Plasma Concentration *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

**14.2.2.4.22. Givinostat Metabolite (ITF2563): Individual Plasma Concentration *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Semi-Logarithmic Scale**

**14.2.2.4.23. Givinostat Metabolite (ITF2563): Individual Plasma Concentration *Versus* Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale**

**14.2.2.4.24. Givinostat Metabolite (ITF2563): Individual Plasma Concentration *Versus* Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Semi-Logarithmic Scale**

**14.2.2.4.25. Givinostat Metabolite (ITF2563): Individual Trough Plasma Concentration *Versus* Time Profile Following Multiple-Dose Administration of Givinostat (Day 9 to Day 13) – Linear Scale**

#### **14.2.2.5. Individual Urinary Excretion *Versus* Time Profiles**

##### **14.2.2.5.1. Givinostat: Individual Urinary Excretion *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

##### **14.2.2.5.2. Givinostat: Individual Cumulative Amount Urinary Excretion *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

##### **14.2.2.5.3. Givinostat Metabolite (ITF2374): Individual Urinary Excretion *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

##### **14.2.2.5.4. Givinostat Metabolite (ITF2374): Individual Cumulative Amount Urinary Excretion *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

##### **14.2.2.5.5. Givinostat Metabolite (ITF2375): Individual Urinary Excretion *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

##### **14.2.2.5.6. Givinostat Metabolite (ITF2375): Individual Cumulative Amount Urinary Excretion *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

##### **14.2.2.5.7. Givinostat Metabolite (ITF2440): Individual Urinary Excretion *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

##### **14.2.2.5.8. Givinostat Metabolite (ITF2440): Individual Cumulative Amount Urinary Excretion *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

##### **14.2.2.5.9. Givinostat Metabolite (ITF2563): Individual Urinary Excretion *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**

##### **14.2.2.5.10. Givinostat Metabolite (ITF2563): Individual Cumulative Amount Urinary Excretion *Versus* Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale**



### 14.2.3. SAFETY DATA – PART 3

#### 14.2.3.1. Adverse Events

##### 14.2.3.1.1. Pre-Treatment Adverse Events

Subject No.	SOC MedDRA PT (Reported Term)	SAE? (Yes/No)	Adverse Event Date and Time		Maximal Severity	Causality	Medication Required?	Outcome
			Start	End				
<Subject ID>	<SOC> <TEAE in MedDRA PT> (<TEAE reported term>)	<Yes / No>	DDMMYYYY- hh:mm	DDMMYYYY- hh mm	<Maximal Severity of the AE>	<Causality of the AE>	<Yes/No>	<Outcome of the AE>
<Insert as many rows as deemed necessary>								

SOC – System Organ Class; MedDRA – Medical Dictionary for Regulatory Activities; PT – Preferred Term; SAE – Serious Adverse Event

NK – Not Known

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

#### 14.2.3.1.2. Treatment-Emergent Adverse Events (I)

Subject No.	Day of Dose	SOC MedDRA PT (Reported Term)	SAE? (Yes/No)	Last Treatment Date and Time	Adverse Event Date and Time	
					Start	Start
<Subject ID>	<Day #>	<SOC> <TEAE in MedDRA PT> (<TEAE reported term>)	<Yes / No>	<DDMMYYYY hh mm>	<DDMMYYYY hh:mm>	<DDMMYYYY hh:mm>

<Insert as many rows as deemed necessary>

SOC – System Organ Class; MedDRA – Medical Dictionary for Regulatory Activities; PT – Preferred Term; SAE – Serious Adverse Event

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

#### 14.2.3.1.3. Treatment-Emergent Adverse Events (II)

Subject No.	Day of Dose	SOC MedDRA PT (Reported Term)	Maximal Severity	Causality	Medication Required?	Outcome
<Subject ID>	<Day #>	<SOC> <TEAE in MedDRA PT> (<TEAE reported term>)	<Maximal Severity of the TEAE>	<Causality of the TEAE>	<Yes/No>	<Outcome of the TEAE>

<Insert as many rows as deemed necessary>

SOC – System Organ Class; MedDRA – Medical Dictionary for Regulatory Activities; PT – Preferred Term; SAE – Serious Adverse Event

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>



#### **14.2.3.2. Vital Signs**

##### **14.2.3.2.1. Descriptive Statistics of Vital Signs Parameters <sup>9</sup>**

##### **14.2.3.2.2. Descriptive Statistics of Change from Baseline of Vital Signs Parameters <sup>9</sup>**

##### **14.2.3.2.3. Shifts in Vital Signs Parameters <sup>10</sup>**

#### **14.2.3.3. 12-Lead ECG**

##### **14.2.3.3.1. Descriptive Statistics of 12-Lead ECG Parameters <sup>9</sup>**

##### **14.2.3.3.2. Descriptive Statistics of Change from Baseline of 12-Lead ECG Parameters <sup>9</sup>**

##### **14.2.3.3.3. Shifts in 12-Lead ECG Parameters <sup>10</sup>**

#### **14.2.3.4. Hematology**

##### **14.2.3.4.1. Descriptive Statistics of Hematology Parameters <sup>9</sup>**

##### **14.2.3.4.2. Descriptive Statistics of Change from Baseline of Hematology Parameters <sup>9</sup>**

##### **14.2.3.4.3. Shifts in Hematology Parameters <sup>10</sup>**

#### **14.2.3.5. Biochemistry**

##### **14.2.3.5.1. Descriptive Statistics of Biochemistry Parameters <sup>9</sup>**

##### **14.2.3.5.2. Descriptive Statistics of Change from Baseline of Biochemistry Parameters <sup>9</sup>**

##### **14.2.3.5.3. Shifts in Biochemistry Parameters <sup>10</sup>**

#### **14.2.3.6. Coagulation**

##### **14.2.3.6.1. Descriptive Statistics of Coagulation Parameters <sup>9</sup>**

##### **14.2.3.6.2. Descriptive Statistics of Change from Baseline of Coagulation Parameters <sup>9</sup>**

##### **14.2.3.6.3. Shifts in Coagulation Parameters <sup>10</sup>**

#### **14.2.3.7. Urinalysis**

##### **14.2.3.7.1. Descriptive Statistics of Urinalysis Parameters <sup>9</sup>**

##### **14.2.3.7.2. Descriptive Statistics of Change from Baseline of Urinalysis Parameters <sup>9</sup>**

### 14.2.3.7.3. Shifts in Urinalysis Parameters <sup>10</sup>

<sup>9</sup> To be generated as follows:

Parameter	Study Day	Protocol Time	n	Mean	SD	SE	Median	Minimum	Maximum
<Parameter (unit)><Day>		<Protocol Time>	xx	x.xxx	x.xxx	x.xxx	x.xxx	x.xxx	x.xxx
		<Protocol Time>							
		<Protocol Time>							
		<Protocol Time>							
<Insert as many rows as deemed necessary>									

n – number of subjects; SD – Standard Deviation; SE – Standard Error

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

<sup>10</sup> To be generated as follows:

Parameter	Study Day	Protocol Time	Classification		Nr. of Subjects
			Baseline	Post-Baseline Assessment	
<Parameter (unit)> <Day>		<Protocol Time>	<Low/ Normal/ High>	<Low/ Normal/ High>	xx
		<Protocol Time>	<Low/ Normal/ High>	<Low/ Normal/ High>	xx
<Insert as many rows as deemed necessary>					

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>



#### 14.2.3.8. Body Weight

##### 14.2.3.8.1. Descriptive Statistics of Body Weight <sup>9</sup>

##### 14.2.3.8.2. Descriptive Statistics of Body Weight – Male Subjects <sup>9</sup>

##### 14.2.3.8.3. Descriptive Statistics of Body Weight – Female Subjects <sup>9</sup>

##### 14.2.3.8.4. Descriptive Statistics of Change from Baseline of Body Weight <sup>9</sup>

##### 14.2.3.8.5. Descriptive Statistics of Change from Baseline of Body Weight – Male Subjects <sup>9</sup>

##### 14.2.3.8.6. Descriptive Statistics of Change from Baseline of Body Weight – Female Subjects <sup>9</sup>

<sup>9</sup> To be generated as follows:

Visit	n	Mean	SD	SE	Median	Minimum	Maximum
<Day>	xx	x.xxx	x.xxx	x.xxx	x.xxx	x.xxx	x.xxx
<Insert as many rows as deemed necessary>							

n – number of subjects; SD – Standard Deviation; SE – Standard Error

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

#### A.4. List of Subject Data Listings (Section 16.2 of CSR)

##### A.4.1. Part 3

##### 16.2.1. PART 3

##### 16.2.1.1. Subject Disposition

##### 16.2.1.1.1. Subject Disposition

Screening No.	Date ICF Signed	Subject Eligible?	Subject Given an Unique Subject No.?	Unique Subject No.	Inclusion Criteria Not Met	Exclusion Criteria Met	Other Reason for Non-Eligibility	Reason for Non-Randomization
<Screening number>	<DDMMYYYY>	<Yes/No>	<Yes/No>	<Unique Subject No>	<List all inclusion criteria that were not met>	<List all exclusion criteria that were met>	<List other reasons for non-eligibility>	<List reason for non-randomization>
<Insert as many rows as deemed necessary>								

ICF – Informed Consent Form

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>



## 16.2.1.2. Protocol Deviations

### 16.2.1.2.1. Blood Sampling Times Deviations

Subject No.	Sampling Timepoint	Sampling Date and Time		Deviation Reason
		Scheduled/Target	Actual	
<Subject ID>	<Scheduled Time or unscheduled>	<DDMMYYYY hh:mm>	<DDMMYYYY hh:mm>	<Reason for deviation>

<Insert as many rows as deemed necessary>

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

### 16.2.1.2.2. Other Protocol Deviations

Subject No.	Protocol Phase	Category	Description	Classification
<Subject ID>	<Protocol Phase #>	<Deviation Category>	<Description of the deviation>	<Important / Not important>

<Insert as many rows as deemed necessary>

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

### 16.2.1.3. Excluded Data from the Pharmacokinetic Analysis

#### 16.2.1.3.1. Plasma Concentrations

##### 16.2.1.3.1.1. Individual Data of Plasma Concentrations

Subject No.	Day of Dosing	Analyte	Pre-dose	Time Post-Dose (h)									
				I.XX	I.XX	I.XX	I.XX	I.XX	I.XX	I.XX	I.XX	I.XX	I.XX
<Subject ID>	<Day #>	<Analyte>	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX	XX.XX
<Insert as many rows as deemed necessary>													
BLQ – Below the Limit of Quantification (LLOQ = <LLOQ> <unitsC>) of the assay													
ND – Not Done													
Program: <SAS Program>													
Execution Date/Time: <ddMMMyyyy HH:MM>													

#### 16.2.1.3.2. Plasma Concentration *Versus* Time Profiles

##### 16.2.1.3.2.1. Individual Plasma Concentration *Versus* Time Profiles – Linear Scale

##### 16.2.1.3.2.2. Individual Plasma Concentration *Versus* Time Profiles – Semi-Logarithmic Scale

*Note: These Figures will be extracted from Phoenix® WinNonlin® 8.2 or higher. A figure will be produced for each subject with data excluded from the pharmacokinetic analysis.*



### 16.2.1.3.3. Urine Excretion

#### 16.2.1.3.3.1. Individual Data of Plasma Concentrations

Subject No.	Day of Dosing	Analyte	Pre-dose	Time Post-Dose (h)					
				0h – 12h	12h – 24h	24h – 36h	36h – 48h	48h – 72h	72h – 96h
<Subject ID>	<Day #>	<Analyte>	xx.xx	xx.xx	xx.xx	xx.xx	xx.xx	xx.xx	xx.xx

<Insert as many rows as deemed necessary>

BLQ – Below the Limit of Quantification (LLOQ = <LLOQ> <unitsC>) of the assay

ND – Not Done

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

### 16.2.1.3.4. Urine Excretion *Versus* Time Profiles

#### 16.2.1.3.4.1. Urine Excretion *Versus* Time Profiles – Linear Scale

*Note: These Figures will be extracted from Phoenix® WinNonlin® 8.2 or higher. A figure will be produced for each subject with data excluded from the pharmacokinetic analysis.*

#### 16.2.1.4. Demographic and Other Baseline Data

##### 16.2.1.4.1. Demographic Data

Subject No.	Date ICF Signed	Date of Birth	Sex	Race	Age (years)	Height (cm)	Weight (kg)	BMI (kg/m <sup>2</sup> )
<Subject ID>	<DDMMYYYY>	<MMYYYY>	<M/F>	<Race>	xx	xxx	xx.x	xx.x

<Insert as many rows as deemed necessary>

ICF – Informed Consent Form; BMI – Body Mass Index; M – Male; F – Female

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>



#### 16.2.1.4.2. Fertility/Contraception

##### 16.2.1.4.2.1. Female Fertility/Contraception

Subject No.	Childbearing Potential?	If Yes, Birth Control Method	Breast Feeding?	Comments
<Subject ID>	<Yes/No>	<Birth Control Method>	<Yes/No>	<Comments>
<Insert as many rows as deemed necessary>				

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

##### 16.2.1.4.2.2. Male Fertility/Contraception

Subject No.	Sexually Active?	Agrees to Use Condom?	Female partner agrees to use a highly effective method of contraception?	Agree not to donate sperm from first dose administration until at least 90 days after the last study drug administration?	Comments
<Subject ID>	<Yes/No>	<Yes/No>	<Yes/No>	<Yes/No>	<Comments>
<Insert as many rows as deemed necessary>					

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

#### 16.2.1.4.3. Drugs of Abuse, Ethanol and Cotinine

Subject No.	Date and Time	Amphetamines	Benzodiazepines	Cocaine	Cannabinoids	Opiates	Ethanol	Cotinine
<Subject ID>	<DDMMYYYY hh:mm>	<Positive / Negative>	<Positive / Negative>	<Positive / Negative>	<Positive / Negative>	<Positive / Negative>	<Positive / Negative>	<Positive / Negative>
<Insert as many rows as deemed necessary>								

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>



#### 16.2.1.4.4. Viral Serology at Screening

Subject No.	Actual Date and Time	Protocol Phase	HIV-1 & HIV-2	Hepatitis B	Hepatitis C
<Subject ID>	<DDMMYYYYYY hh:mm>	<Protocol Phase>	<Qualitative Value>	<Qualitative Value>	<Qualitative Value>
<Insert as many rows as deemed necessary>					

HIV – Human Immunodeficiency Virus

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

#### 16.2.1.4.5. Previous and Concomitant Medication

Subject No.	Drug Name [ATC Code]	Indication	Pharmaceutical Form	Dose (units)	Frequency	Route	Date and Time	
							Start	End
<Subject ID>	<Generic name [XXXXXXXX]>	<Therapeutic >indication	<Pharmaceutical Form>	<Dose (units)>	<Frequency>	<Route>	<DDMMYYYY>	<DDMMYYYY or Ongoing>
<Insert as many rows as deemed necessary>								

ATC – Anatomical Therapeutic Chemical

NK – Not Known

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>



#### 16.2.1.4.6. SARS-COV-2 Test

Subject No.	Actual Date	Protocol Phase	Result	If Unscheduled, Reason
<Subject ID>	<DDMMYYYY>	<Protocol Phase>	<Not detectable, Detectable, Inconclusive >	<Reason>
<Insert as many rows as deemed necessary>				
Program: <SAS Program>				
Execution Date/Time: <ddMMMyyyy HH:MM>				

*Note: The "If Unscheduled, Reason" column will only be generated if there is an Unscheduled Reason for any of the evaluations.*

#### 16.2.1.5. Compliance

##### 16.2.1.5.1. Investigational Product Administration

Subject No.	Date	Actual Clock Time	Dose (units)	Investigational Product Formulation	Route of Administration	Hands and Mouth Check?	150 mL of water taken?	Comments
<Subject ID>	<DDMMYY YY/N/A>	<hh mm/ N/A>	<Dose (units)>	<Formulation>	<Route>	<Yes/No/ N/A>	<Yes/No/ N/A>	<Comments>

<Insert as many rows as deemed necessary>

N/A – Not Applicable

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>



#### **16.2.1.6. Individual Pharmacokinetic Data**

##### **16.2.1.6.1. Givinostat: Individual Pharmacokinetic Data**

###### **16.2.1.6.1.1. Givinostat: Individual Drug Plasma Concentration Data**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.1.2.1. for single-dose data and 14.3.2.1.2.16. for multiple-dose data'*

###### **16.2.1.6.1.2. Givinostat: Individual Plasma Pharmacokinetic Parameters**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.1.3.1. for single-dose data and 14.3.2.1.3.16. for multiple-dose data'*

###### **16.2.1.6.1.3. Givinostat: Individual Plasma Pharmacokinetic Profiles**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.4.1. to 14.3.2.4.2.'*

###### **16.2.1.6.1.4. Givinostat: Individual Urine Excretion Data**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.2.1. for single-dose data and 14.3.2.2.16. for multiple-dose data'*

###### **16.2.1.6.1.5. Givinostat: Individual Urine Pharmacokinetic Parameters**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.2.3.1. for single-dose data and 14.3.2.2.3.16. for multiple-dose data'*

###### **16.2.1.6.1.6. Givinostat: Individual Urine Pharmacokinetic Profiles**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.5.1. to 14.3.2.5.2.'*

###### **16.2.1.6.1.7. Givinostat: Documentation of Statistical Analysis**

###### **16.2.1.6.1.7.1. Givinostat: Plasma Non-Compartmental Analysis Output**

###### **16.2.1.6.1.7.2. Givinostat: Plasma Non-Compartmental Analysis Plots**

###### **16.2.1.6.1.7.3. Givinostat: Urine Non-Compartmental Analysis Output**

*Note: The documentation of statistical analysis will be extracted Phoenix® WinNonlin® 8.2 or higher.*

#### **16.2.1.6.2. Givinostat Metabolite (ITF2374): Individual Pharmacokinetic Data**

##### **16.2.1.6.2.1. Givinostat Metabolite (ITF2374): Individual Drug Plasma Concentration Data**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.1.2.4. for single-dose data and 14.3.2.1.2.19. for multiple-dose data'*

##### **16.2.1.6.2.2. Givinostat Metabolite (ITF2374): Individual Plasma Pharmacokinetic Parameters**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.1.3.4. for single-dose data and 14.3.2.1.3.19. for multiple-dose data'*

##### **16.2.1.6.2.3. Givinostat Metabolite (ITF2374): Individual Plasma Pharmacokinetic Profiles**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.4.3. to 14.3.2.4.4.'*

##### **16.2.1.6.2.4. Givinostat Metabolite (ITF2374): Individual Urine Excretion Data**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.2.2.4. for single-dose data and 14.3.2.2.2.19. for multiple-dose data'*

##### **16.2.1.6.2.5. Givinostat Metabolite (ITF2374): Individual Urine Pharmacokinetic Parameters**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.2.3.4. for single-dose data and 14.3.2.2.3.19. for multiple-dose data'*

##### **16.2.1.6.2.6. Givinostat Metabolite (ITF2374): Individual Urine Pharmacokinetic Profiles**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.5.3. to 14.3.2.5.4.'*

#### **16.2.1.6.2.7. Givinostat Metabolite (ITF2374): Documentation of Statistical Analysis**

##### **16.2.1.6.2.7.1. Givinostat Metabolite (ITF2374): Plasma Non-Compartmental Analysis Output**

##### **16.2.1.6.2.7.2. Givinostat Metabolite (ITF2374): Plasma Non-Compartmental Analysis Plots**

##### **16.2.1.6.2.7.3. Givinostat Metabolite (ITF2374): Urine Non-Compartmental Analysis Output**

*Note: The documentation of statistical analysis will be extracted Phoenix<sup>®</sup> WinNonlin<sup>®</sup> 8.2 or higher.*



#### **16.2.1.6.3. Givinostat Metabolite (ITF2375): Individual Pharmacokinetic Data**

##### **16.2.1.6.3.1. Givinostat Metabolite (ITF2375): Individual Drug Plasma Concentration Data**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.1.2.7. for single-dose data and 14.3.2.1.2.22. for multiple-dose data'*

##### **16.2.1.6.3.2. Givinostat Metabolite (ITF2375): Individual Plasma Pharmacokinetic Parameters**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.1.3.7. for single-dose data and 14.3.2.1.3.22. for multiple-dose data'*

##### **16.2.1.6.3.3. Givinostat Metabolite (ITF2375): Individual Plasma Pharmacokinetic Profiles**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.4.5. to 14.3.2.4.6.'*

##### **16.2.1.6.3.4. Givinostat Metabolite (ITF2375): Individual Urine Excretion Data**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.2.7. for single-dose data and 14.3.2.2.22. for multiple-dose data'*

##### **16.2.1.6.3.5. Givinostat Metabolite (ITF2375): Individual Urine Pharmacokinetic Parameters**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.2.3.7. for single-dose data and 14.3.2.2.3.22. for multiple-dose data'*

##### **16.2.1.6.3.6. Givinostat Metabolite (ITF2375): Individual Urine Pharmacokinetic Profiles**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.5.5. to 14.3.2.5.6.'*

##### **16.2.1.6.3.7. Givinostat Metabolite (ITF2375): Documentation of Statistical Analysis**

###### **16.2.1.6.3.7.1. Givinostat Metabolite (ITF2375): Plasma Non-Compartmental Analysis Output**

###### **16.2.1.6.3.7.2. Givinostat Metabolite (ITF2375): Plasma Non-Compartmental Analysis Plots**

###### **16.2.1.6.3.7.3. Givinostat Metabolite (ITF2375): Urine Non-Compartmental Analysis Output**

*Note: The documentation of statistical analysis will be extracted Phoenix<sup>®</sup> WinNonlin<sup>®</sup> 8.2 or higher.*

#### **16.2.1.6.4. Givinostat Metabolite (ITF2440): Individual Pharmacokinetic Data**

##### **16.2.1.6.4.1. Givinostat Metabolite (ITF2440): Individual Drug Plasma Concentration Data**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.1.2.14. for single-dose data and 14.3.2.1.2.28. for multiple-dose data'*

##### **16.2.1.6.4.2. Givinostat Metabolite (ITF2440): Individual Plasma Pharmacokinetic Parameters**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.1.3.14. for single-dose data and 14.3.2.1.3.28. for multiple-dose data'*

##### **16.2.1.6.4.3. Givinostat Metabolite (ITF2440): Individual Plasma Pharmacokinetic Profiles**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.4.9. to 14.3.2.4.10.'*

##### **16.2.1.6.4.4. Givinostat Metabolite (ITF2440): Individual Urine Excretion Data**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.2.2.14. for single-dose data and 14.3.2.2.2.28. for multiple-dose data'*

##### **16.2.1.6.4.5. Givinostat Metabolite (ITF2440): Individual Urine Pharmacokinetic Parameters**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.2.3.14. for single-dose data and 14.3.2.2.3.28. for multiple-dose data'*

##### **16.2.1.6.4.6. Givinostat Metabolite (ITF2440): Individual Urine Pharmacokinetic Profiles**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.5.9. to 14.3.2.5.10.'*

##### **16.2.1.6.4.7. Givinostat Metabolite (ITF2440): Documentation of Statistical Analysis**

###### **16.2.1.6.4.7.1. Givinostat Metabolite (ITF2440): Plasma Non-Compartmental Analysis Output**

###### **16.2.1.6.4.7.2. Givinostat Metabolite (ITF2440): Plasma Non-Compartmental Analysis Plots**

###### **16.2.1.6.4.7.3. Givinostat Metabolite (ITF2440): Urine Non-Compartmental Analysis Output**

*Note: The documentation of statistical analysis will be extracted Phoenix<sup>®</sup> WinNonlin<sup>®</sup> 8.2 or higher.*



#### **16.2.1.6.5. Givinostat Metabolite (ITF2563): Individual Pharmacokinetic Data**

##### **16.2.1.6.5.1. Givinostat Metabolite (ITF2563): Individual Drug Plasma Concentration Data**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.1.3.7. for single-dose data and 14.3.2.1.2.28. for multiple-dose data'*

##### **16.2.1.6.5.2. Givinostat Metabolite (ITF2563): Individual Plasma Pharmacokinetic Parameters**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.1.3.7. for single-dose data and 14.3.2.1.3.28. for multiple-dose data'*

##### **16.2.1.6.5.3. Givinostat Metabolite (ITF2563): Individual Plasma Pharmacokinetic Profiles**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.4.9. to 14.3.2.4.10.'*

##### **16.2.1.6.5.4. Givinostat Metabolite (ITF2563): Individual Urine Excretion Data**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.2.7. for single-dose data and 14.3.2.2.28. for multiple-dose data'*

##### **16.2.1.6.5.5. Givinostat Metabolite (ITF2563): Individual Urine Pharmacokinetic Parameters**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.2.3.5. for single-dose data and 14.3.2.2.3.28. for multiple-dose data'*

##### **16.2.1.6.5.6. Givinostat Metabolite (ITF2563): Individual Urine Pharmacokinetic Profiles**

*Note: The following sentence should be inserted in this section: 'No further information is provided; refer to sections 14.3.2.5.9. to 14.3.2.5.10.'*

##### **16.2.1.6.5.7. Givinostat Metabolite (ITF2563): Documentation of Statistical Analysis**

###### **16.2.1.6.5.7.1. Givinostat Metabolite (ITF2563): Plasma Non-Compartmental Analysis Output**

###### **16.2.1.6.5.7.2. Givinostat Metabolite (ITF2563): Plasma Non-Compartmental Analysis Plots**

###### **16.2.1.6.5.7.3. Givinostat Metabolite (ITF2563): Urine Non-Compartmental Analysis Output**

*Note: The documentation of statistical analysis will be extracted Phoenix<sup>®</sup> WinNonlin<sup>®</sup> 8.2 or higher.*

## 16.2.1.7. Adverse Event Listings (Each Subject)

### 16.2.1.7.1. Pre-Treatment Adverse Events <sup>7</sup>

### 16.2.1.7.2. Treatment-Emergent Adverse Events <sup>7</sup>

<sup>7</sup> To be generated as follows:

Subject No.	Day of Dose	Treatment	Reported Term	Serious?	Maximal Severity	Causality	Action Taken	Medication Required?	Outcome	Adverse Event Start Date and Time	Adverse Event End Date and Time	Comments
<Subject ID>	<Day #>	<Investigational Product>	<AE Reported Term>	<Yes/No>	<Mild, Moderate or Severe>	<Not Reasonably Possible or Reasonably Possible>	<Dose Increased, Dose Not Changed, Dose Rate Reduced, Dose Reduced, Drug Interrupted, Drug Withdrawn, Not Applicable or Unknown>	<Yes or No>	<Fatal, Not Recovered / Not Resolved, Recovered / Resolved, Recovered / Resolved with Sequelae, Recovering / Resolving or Unknown>	<DDMMYY hh:mm>	<DDMMYY hh:mm or Ongoing>	<Comments>

<Insert as many rows as deemed necessary>

NK – Not Known

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

Note: The "Investigational Product" column will only be generated for Table 16.2.3.7.2. If an Adverse Event is related to Medical History, that information will be reported in 'Comments'. The "Comments" column will only be generated if there are reported comments for the adverse events.



#### 16.2.1.7.3. Serious Adverse Events (I)

Subject No.	Reported Term	Reason for Seriousness	Details
<Subject ID>	<AE Reported Term>	<Reason>	<SAE details>
<Insert as many rows as deemed necessary>			
Program: <SAS Program>			
Execution Date/Time: <ddMMMyyyy HH:MM>			

## 16.2.1.8. Listings of Laboratory Measurements by Subject

### 16.2.1.8.1. Normal Range of Laboratory Values

					Normal Range			
					Female		Male	
Category	Analyte	Age	Units	LLN	ULN	LLN	ULN	Qualitative
<Category>	<Laboratory Analyte>	<Age interval>	<Units>	<Lower Limit>	<Upper Limit>	<Lower Limit>	<Upper Limit>	<Qualitative Values>
<Insert as many rows as deemed necessary>								

LLN – Lower Limit Normal; ULN – Upper Limit Normal

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>



#### 16.2.1.8.2. Hematology (I)

Subject No.	Actual Date and Time	Protocol Phase	Hemoglobin (g/dL)	RBC (10 <sup>12</sup> /L)	Hematocrit (%)	MCV (fL)	MCH (pg)	MCHC (g/dL)	RDW-CV (%)	Total WBC (10 <sup>9</sup> /L)	If Unscheduled, Reason
<Subject ID>	<DDMMYY YYY hh mm>	<Protocol Phase>	xx	xx	xx	xx	xx	xx	xx	xx	<Reason>

<Insert as many rows as deemed necessary>

RBC – Red Blood Cell; MCV – Mean Corpuscular Volume; MCH – Mean Corpuscular Hemoglobin; MCHC – Mean Corpuscular Hemoglobin Concentration; RDW-CV – Coefficient Variation of the Red Cell Distribution Width; WBC – White Blood Cell

L – Low; H – High; CR – Clinically Relevant; NCR – Not Clinically Relevant; RU – Relevance Unknown

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

*Note: The “If Unscheduled, Reason” column will only be generated if there is an Unscheduled Reason for one of the evaluated parameters.*

### 16.2.1.8.3. Hematology (II)

Subject No.	Actual Date and Time	Protocol Phase	Neutrophils (10 <sup>9</sup> /L)	Eosinophils (10 <sup>9</sup> /L)	Basophils (10 <sup>9</sup> /L)	Lymphocytes (10 <sup>9</sup> /L)	Monocytes (10 <sup>9</sup> /L)	Platelets (10 <sup>9</sup> /L)	Mean Platelet Volume (fL)	If Unscheduled, Reason
<Subject ID>	<DDMMYY YYY hh:mm>	<Protocol Phase>	xx	xx	xx	xx	xx	xx	xx	<Reason>
<Insert as many rows as deemed necessary>										

L – Low; H – High; CR – Clinically Relevant; NCR – Not Clinically Relevant; RU – Relevance Unknown

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

*Note: The “If Unscheduled, Reason” column will only be generated if there is an Unscheduled Reason for one of the evaluated parameters.*



#### 16.2.1.8.4. Biochemistry (I)

Subject No.	Actual Date and Time	Protocol Phase	AST (IU/L)	ALT (IU/L)	GGT (IU/L)	LDH (IU/L)	ALP (IU/L)	CK (IU/L)	If Unscheduled, Reason
<Subject ID>	<DDMMYY YY hh mm>	<Protocol Phase>	xx	xxx	xx	xx	xx	xx	<Reason>
<Insert as many rows as deemed necessary>									

L – Low; H – High; CR – Clinically Relevant; NCR – Not Clinically Relevant; RU – Relevance Unknown

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

*Note: The “If Unscheduled, Reason” column will only be generated if there is an Unscheduled Reason for one of the evaluated parameters.*

#### 16.2.1.8.5. Biochemistry (II)

Subject No.	Actual Date and Time	Protocol Phase	Creatinine (mg/dL)	eGFR	Urea (mg/dL)	Sodium (mmol/L)	Potassium (mmol/L)	Calcium (mg/dL)	If Unscheduled, Reason
<Subject ID>	<DDMMYY YY hh mm>	<Protocol Phase>	xx	xxx	xx	xx	xx	xx	<Reason>
<Insert as many rows as deemed necessary>									

L – Low; H – High; CR – Clinically Relevant; NCR – Not Clinically Relevant; RU – Relevance Unknown

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

*Note: The “If Unscheduled, Reason” column will only be generated if there is an Unscheduled Reason for one of the evaluated parameters.*



#### 16.2.1.8.6. Biochemistry (III)

Subject No.	Actual Date and Time	Protocol Phase	Magnesium (mg/dL)	Glucose (mg/dL)	Total Cholesterol (mg/dL)	Triglycerides (mg/dL)	Albumin (g/dL)	Total Protein (g/dL)	If Unscheduled, Reason
<Subject ID>	<DDMMYY YY hh mm>	<Protocol Phase>	xx	xxx	xx	xx	xx	xx	<Reason>
<Insert as many rows as deemed necessary>									

L – Low; H – High; CR – Clinically Relevant; NCR – Not Clinically Relevant; RU – Relevance Unknown

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

*Note: The "If Unscheduled, Reason" column will only be generated if there is an Unscheduled Reason for one of the evaluated parameters.*

#### 16.2.1.8.7. Biochemistry (IV)

Subject No.	Actual Date and Time	Protocol Phase	Uric Acid (mg/dL)	Total Bilirubin (mg/dL)	Direct Bilirubin (mg/dL)	Indirect Bilirubin (mg/dL)	TSH (mIU/L)	Chloride (mmol/L)	If Unscheduled, Reason
<Subject ID>	<DDMMYY YY hh mm>	<Protocol Phase>	XX	XXX	XX	XX	XX	XX	<Reason>
<Insert as many rows as deemed necessary>									

L – Low; H – High; CR – Clinically Relevant; NCR – Not Clinically Relevant; RU – Relevance Unknown

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

*Note: The "If Unscheduled, Reason" column will only be generated if there is an Unscheduled Reason for one of the evaluated parameters.*



#### 16.2.1.8.8. Biochemistry (V)

Subject No.	Actual Date and Time	Protocol Phase	Amylase (g/dL)	LDL Cholesterol (mg/dL)	HDL Cholesterol (mg/dL)	FSH (mg/dL)	C-Reactive Protein	Cystain C	If Unscheduled, Reason
<Subject ID>	<DDMMYY YY hh mm>	<Protocol Phase>	XX	XXX	XX	XX	XX	XX	<Reason>
<Insert as many rows as deemed necessary>									

L – Low; H – High; CR – Clinically Relevant; NCR – Not Clinically Relevant; RU – Relevance Unknown

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

*Note: The "If Unscheduled, Reason" column will only be generated if there is an Unscheduled Reason for one of the evaluated parameters.*

#### 16.2.1.8.9. Coagulation

Subject No.	Actual Date and Time	Protocol Phase	Prothrombin Rate (%)	Prothrombin Time (sec)	INR	aPTT (sec)	If Unscheduled, Reason
<Subject ID>	<DDMMYYYY> hh:mm	<Protocol Phase>	xx	xx.x	x.xx	xx.x	<Reason>

<Insert as many rows as deemed necessary>

INR – International Normalized Ratio; aPTT – Activated Partial Thromboplastin Time  
L – Low; H – High; CR – Clinically Relevant; NCR – Not Clinically Relevant; RU – Relevance Unknown  
Program: <SAS Program>  
Execution Date/Time: <ddMMMyyyy HH:MM>

*Note: The “If Unscheduled, Reason” column will only be generated if there is an Unscheduled Reason for one of the evaluated parameters.*



#### 16.2.1.8.10. Urinalysis

Subject No.	Actual Date and Time	Protocol Phase	pH	Specific Gravity	Protein	Hemoglobin	Glucose	Ketones	Bilirubin	Nitrites	Urobilinogen	If Unscheduled, Reason
<Subject ID>	<DDMM MYYY hh:mm>	<Protocol Phase>	xx.x	x.xxx	<Qualitative Value>	<Qualitative Value>	<Qualitative Value>	<Qualitative Value>	<Qualitative Value>	<Qualitative Value>	<Qualitative Value>	<Reason>

<Insert as many rows as deemed necessary>

L – Low; H – High; NCR – Not Clinically Relevant; CR – Clinically Relevant; RU - Relevance Unknown

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

*Note: The "If Unscheduled, Reason" column will only be generated if there is an Unscheduled Reason for one of the evaluated parameters.*

#### 16.2.1.8.11. Urine Microscopy

Subject No.	Actual Date and Time	Protocol Phase	Squamous Epithelial Cells (/uL)	Erythrocytes (/uL)	Leukocytes (/uL)	Observations	If Unscheduled, Reason
<Subject ID>	<DDMMYYYY hh:mm>	<Protocol Phase>	<Qualitative Value>	xx	xx	<Observations>	<Reason>
<Insert as many rows as deemed necessary>							

L – Low; H – High; CR – Clinically Relevant; NCR – Not Clinically Relevant; RU – Relevance Unknown

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

*Note: The "If Unscheduled, Reason" column will only be generated if there is an Unscheduled Reason for one of the evaluated parameters.*



#### 16.2.1.8.12. Pregnancy Test

Subject No.	Actual Date and Time	Protocol Phase	Matrix	Result	If Unscheduled, Reason
<Subject ID>	<DDMMYYYY hh:mm>	<Protocol Phase>	<Serum or Urine>	<Positive or Negative>	<Reason>
<Insert as many rows as deemed necessary>					
Program: <SAS Program>					
Execution Date/Time: <ddMMMyyyy HH:MM>					

*Note: The "If Unscheduled, Reason" column will only be generated if there is an Unscheduled Reason for one of the evaluations.*

16.2.1.8.13. Additional (Not Planned) Laboratory Safety Tests

Subject No.	Actual Date and Time	Protocol Phase	Investigational Product	Parameter	Results	Reason
<Subject ID>	<DDMMYYYY hh mm>	<Protocol Phase>	<Investigational Product>	<Parameter>	<Results>	<Reason>
<Insert as many rows as deemed necessary>						

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>



### 16.2.1.9. Vital Signs

Subject No.	Scheduled Time	Actual Date and Time	Protocol Phase	Investigational Product	SBP (mmHg)	DBP (mmHg)	Pulse Rate (beats/min)	Body Temperature (°C)	Respiratory Rate (beats/min)	If Unscheduled, Reason
<Subject ID>	<Scheduled Time or Unscheduled>	<DDMMYY YY hh:mm>	<Protocol Phase>	<Investigational Product>	xx	xx	xx	xx	xx	<Reason>

<Insert as many rows as deemed necessary>

SBP – Systolic Blood Pressure; DBP – Diastolic Blood Pressure  
L – Low; H – High; CR – Clinically Relevant; NCR – Not Clinically Relevant  
Program: <SAS Program>  
Execution Date/Time: <ddMMYY HH:MM>

*Note: The “If Unscheduled, Reason” column will only be generated if there is an Unscheduled Reason for one of the evaluated parameters.*

#### 16.2.1.10. 12-Lead ECG

Subject No.	Scheduled Time	Actual Date and Time	Protocol Phase	Investigational Product	QTcF (msec)	Result	If Abnormal, Reason	If Abnormal, Clinically Significant?	If Unscheduled, Reason
<Subject ID>	<Scheduled Time or Unscheduled>	<DDMMYY YY hh:mm>	<Protocol Phase>	<Investigational Product>	<xxx>	<Abnormal/ Normal>	<Reason for being considered abnormal>	<Yes / No>	<Reason>

<Insert as many rows as deemed necessary>

Program: <SAS Program>

Execution Date/Time: <ddMMMyyyy HH:MM>

*Note: The "If Unscheduled, Reason" column will only be generated if there is an Unscheduled Reason for any of the evaluations.*



#### 16.2.1.11. Urine Sampling

Subject No.	Day of Dose	Protocol Time	Collection Time		Urine Volume
			Start	End	
<Subject ID>	<Day #>	<Protocol Time>	<DDMMYYYY hh:mm>	<DDMMYYYY hh:mm>	<Abnormal/ Normal>
<Insert as many rows as deemed necessary>					
Program: <SAS Program>					
Execution Date/Time: <ddMMMyyyy HH:MM>					

#### 14. ANNEXES

[Annex 5: Data Blind Review Process Minute – Part 3](#)

[Annex 6: List of Subjects Dosed – Part 3](#)



- Part 3: eight (8) subjects

Each subject will participate in one study part only.

#### 5.3.1. **Part 3**

A sample size of 8 subjects will be enrolled. The sample size was determined by practical considerations and not based on statistical power calculations.

#### 5.4. **Randomization**

This is a fixed sequence study. Randomization is not applicable.

#### 5.5. **Blinding**

The study will be conducted as open label. Blinding procedures are not applicable.

### 6. **STUDY ASSESSMENTS**

For each study part, a summary of procedures for study assessments are presented in Section 2 of the CSP (Study Flow-Chart and Study Design Diagram) [1].

#### 6.1. **Safety Assessments**

Subjects' safety will be monitored during the study.

Safety assessments will include pre-study medical history, physical examination, vital signs, 12-lead ECG, clinical laboratory tests and adverse event (AE) monitoring. Additional safety measurements may be performed at the discretion of the investigator for reasons related to subject safety.

Medications will be mentioned according to the World Health Organization (WHO) Anatomical Therapeutic Chemical (ATC) classification system.

##### 6.1.1. **Medical History**

Medical history will cover all relevant past or present information related to subject's health at the time of informed consent signature.

Medical history at screening will include past or present relevant cardiovascular, respiratory, renal, genitourinary, gastrointestinal, hepatic, hematological, immunological, endocrine, dermatological, musculoskeletal, neurological, psychiatric, drug and surgical history, or any other diseases or disorders.

Medical history will be referred in accordance with the Medical Dictionary for Regulatory Activities (MedDRA), version 24.1 or higher.

Adverse events related to medical history will be identified.

#### 6.1.2. **Physical Examination**

Physical examination at screening and end-of-study will include: general appearance; skin; head and neck; thorax and abdomen; pulmonary auscultation; cardiac auscultation; abdomen palpation; limbs; brief neurological examination.

#### 6.1.3. **Weight, Height and Body Mass Index**

The body height and weight values as well as the body mass index (BMI) will be recorded in the electronic case report form (eCRF) and will be determined at Screening. Only body weight will be determined also at admission and end-of-study.

The subjects' body weight will preferably be measured using the same weighing scale for all subjects and throughout the study. The weighing scale should have a precision of at least 0.5 kg.

#### 6.1.4. **Vital Signs**

Vital signs will include:

- Systolic blood pressure (SBP).
- Diastolic blood pressure (DBP).
- Pulse rate (PR).
- Respiratory rate (RR).
- Body temperature.

#### 6.1.5. **12 Lead Electrocardiogram**

A 12-lead ECG will be performed preferably before the blood collection. The corrected QT interval by Fridericia (QTcF) will be analyzed.

#### 6.1.6. **Laboratory Safety Tests**

The following laboratory parameters will be tested:

- Hematology:
  - Red blood cell (RBC) count.
  - White blood cell (WBC) count.
  - WBC differential count:
    - Neutrophils.
    - Eosinophils.
    - Basophils.
    - Lymphocytes.
    - Monocytes.
  - Hemoglobin.
  - Mean corpuscular volume (MCV).
  - Mean corpuscular hemoglobin (MCH).
  - Mean corpuscular hemoglobin concentration (MCHC).
  - Coefficient variation of the red cell distribution width (RDW-CV).
  - Hematocrit.
  - Platelet count.
  - Mean platelet volume.



- Specific gravity.
- Protein.
- Hemoglobin.
- Glucose.
- Ketones.
- Bilirubin.
- Nitrites.
- Urobilinogen.
- Microscopy.
- Drugs-of-abuse urine test:
  - Cannabinoids.
  - Opiates.
  - Cocaine.
  - Amphetamines.
  - Benzodiazepines.
- Ethanol urine test.
- Cotinine urine test.
- Follicle-stimulating hormone (FSH) test.
- Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-CoV-2) diagnostic tests.

#### 6.1.7. Adverse Events

The occurrence of clinical AEs will be monitored throughout the study.

Clinically significant abnormalities in laboratory safety tests, vital signs and physical examination will be reported as AEs.

Treatment-emergent AEs (TEAEs) are defined as AEs not present prior to first administration of investigational product, or AEs present before first administration of investigational product that worsen after the subject receives the first dose of investigational product. TEAEs that occur after administration of investigational product during the washout of a given period will be assigned to the treatment administered in that period.

The following information will be used for the description of the AEs:

- Reported Term.
- Medical Dictionary for Regulatory Activities (MedDRA) System Organ Class (SOC) coding.
- MedDRA Preferred Term (PT) coding.
- Start date and time.
- End date and time.
- Seriousness.
- Severity:
  - Mild.
  - Moderate.
  - Severe.
- Relationship (causality) for each treatment administered, per part.
  - Reasonably Possible.
  - Not Reasonably Possible.
  - Unknown.

- Action taken:
  - Dose Increased.
  - Dose Not Changed.
  - Dose Rate Reduced.
  - Dose Reduced.
  - Drug Interrupted.
  - Drug Withdrawn.
  - Not Applicable.
  - Unknown.
- Concomitant medication.
- Outcome:
  - Fatal.
  - Not Recovered / Not Resolved.
  - Recovered / Resolved.
  - Recovered / Resolved with Sequelae.
  - Recovering / Resolving.
  - Unknown.
- Most recent study treatment taken.
- Last dosing date.

#### 6.1.8. Previous and Concomitant Medications

A previous medication is any medication for which the end date is prior to first dosing.  
A concomitant medication is any medication ongoing or initiated after first dosing.

For all study parts, the use of any medications including Over-the-Counter (OTC) products (including herbal medicines such as St John's Wort, homeopathic preparations, vitamins, and minerals) is forbidden from 28 days or within 5 half-lives of the medicinal product, whichever is longer, prior to admission up to last sample for pharmacokinetics assessment, except for medications for the treatment of AEs.

Any medication (previous or concomitant) taken within the period of medication restriction must be recorded in the eCRF. If concomitant medication is ongoing at the follow-up visit, no end date will be provided in the eCRF.

For each study part, previous and concomitant medications used by study participants during the study and their judged impact on the pharmacokinetic assessment will be listed in the respective Data Blind Review Meeting/ Process Minute ([Annex 5](#)).

Concomitant medications will be coded according to the WHO ATC classification system.

## 6.2. Pharmacokinetic Assessments

### 6.2.1. Blood Sampling for Pharmacokinetic Assessments

In each study part, while subject is confined at the clinical research unit, blood samples will be taken preferably via an indwelling cannula placed in a vein of an upper limb of the subject. During ambulatory visits, blood samples will be taken by direct venipuncture.



The actual time of all pharmacokinetic blood draws will be recorded and reported for all subjects. The pre-dose blood sample will be collected within 30 minutes before dosing. The post-dose blood samples will be collected within  $\pm 3$  minutes from the scheduled sampling time. Greater deviations will be reported as a protocol deviation and its cause will be recorded.

In case blood sampling for pharmacokinetics and other procedures coincide in time, blood draws will have priority unless other procedures are necessary for assuring subject's safety.

PPD will carry out the determination of plasma levels of givinostat and its metabolites (ITF2374, ITF2375, ITF2440 and ITF2563), midazolam, 1-hydroxymidazolam and total and free dabigatran in accordance with the applicable principles of Good Laboratory Practices (GLP), using a previously validated liquid chromatography with tandem mass spectrometry (LC-MS/MS) analytical method.

The planned lower limit of quantification (LLOQ) and upper limit of quantification (ULOQ) for:

- givinostat are 1 and 100 ng/mL.
- givinostat metabolite ITF2374 are 1 and 100 ng/mL.
- givinostat metabolite ITF2375 are 1 and 100 ng/mL.
- givinostat metabolite ITF2440 are 10 and 500 ng/mL.
- givinostat metabolite ITF2563 are 2 and 125 ng/mL.
- midazolam are 100 and 100000 pg/mL.
- 1-hydroxymidazolam are 100 and 50000 pg/mL.
- total dabigatran are 1 and 400 ng/mL.
- free dabigatran are 1 and 400 ng/mL.

LLOQ represents a value lower than or equal to 1/20 of estimated  $C_{max}$  value. Any adjustment to this range will be documented in the study specific Bioanalytical Report, which will supersede the indicated range.

#### 6.2.1.1. Part 3

A total of forty-eight (48) blood samples will be collected as follows:

- Twenty-two (22) blood samples of 6 mL each will be collected in sodium heparin tubes at pre-dose and at 0.5, 1, 1.5, 2, 2.5, 3, 3.5, 4, 5, 6, 8, 10, 12, 15, 24, 36, 48, 60, 72, 84 and 96 hours after the administration of givinostat, on Days 1 and 13, for the determination of givinostat and its metabolites (ITF2374, ITF2375, ITF2440 and ITF2563) plasma concentrations.
- Four (4) blood samples of 6 mL each will be collected in sodium heparin tubes before the morning dose of givinostat on Days 9, 10, 11 and 12, for the determination of pre-dose ( $C_{trough}$ ) plasma concentration of givinostat and metabolites.

#### 6.2.2. Urine Sampling for Pharmacokinetic Assessments

##### 6.2.2.1. Part 3

Urine will be collected on Day -1 (baseline) and in the following intervals, on Days 1 and 13, for the determination of the amounts of givinostat and its metabolites (ITF2374, ITF2375, ITF2440 and ITF2563) excreted in urine: 0–12, 12–24, 24–36, 36–48, 48–72 and 72–96 post-dose.



The date and actual time of urine collection start and completion will be recorded and reported for all subjects.

PPD will carry out the determination of urine levels of givinostat and its metabolites (ITF2374, ITF2375, ITF2440 and ITF2563) in accordance with the applicable principles of GLP using a previously validated liquid chromatography with tandem mass spectrometry (LC-MS/MS) analytical method.

The planned lower limit of quantification (LLOQ) and upper limit of quantification (ULOQ) for:

- givinostat are 10 and 2500 ng/mL.
- givinostat metabolite ITF2374 are 5 and 2500 ng/mL.
- givinostat metabolite ITF2375 are 4 and 800 ng/mL.
- givinostat metabolite ITF2440 are 150 and 25000 ng/mL.
- givinostat metabolite ITF2563 are 10 and 5000 ng/mL.

### 6.2.3. Plasma Pharmacokinetic Parameters

The following pharmacokinetic parameters will be derived by standard non-compartmental analysis (NCA) methods from the single-dose plasma concentration *versus* time profiles, for:

- Part 3 (Day 1):
  - givinostat.
  - givinostat metabolites (ITF2374, ITF2375, ITF2440 and ITF2563).

Parameter	Description
$C_{max}$	Maximum observed concentration, directly obtained from the observed concentration <i>versus</i> time profile.
$T_{max}$	Time of occurrence of maximum observed concentration.
$AUC_{0-t}^1$	Area under the concentration <i>versus</i> time curve (AUC) from time of dosing ( $t=0h$ ) to the time of the last measurable concentration ( $t_{last}$ ), calculated by the linear-up/log-down trapezoidal method.
$AUC_{0-\infty}$	Total AUC extrapolated to infinity, calculated as $AUC_{0-t} + \frac{C_{last}}{\lambda_z}$ , where $C_{last}$ is the last measurable concentration and $\lambda_z$ is the apparent terminal elimination rate constant.
%AUC <sub>extrap</sub>	Percentage of $AUC_{0-\infty}$ due to extrapolation from the time of the last measurable concentration ( $t_{last}$ ) to infinity, i.e., residual area, calculated as $100 \cdot \frac{AUC_{0-\infty} - AUC_{0-t}}{AUC_{0-\infty}}$ .
$\lambda_z$	Apparent first order elimination rate constant associated with the terminal (log-linear) portion of the concentration <i>versus</i> time curve. The parameter is estimated by linear least square regression analysis using the last three (or more) non-zero concentrations.
$t_{1/2}$	Apparent terminal elimination half-life, calculated as $\frac{\ln(2)}{\lambda_z}$ .

<sup>1</sup> In Part 3, for givinostat and its metabolites the last sampling time corresponds to 96 hours.



In estimating the pharmacokinetic parameters, concentrations below the LLOQ before  $T_{max}$  will be set to zero. After  $T_{max}$ , concentrations below the LLOQ will be considered as missing. However, in case of two or more consecutive concentrations below the LLOQ, the first value will be replaced by  $\frac{1}{2}$  of the LLOQ value and the next values will be considered as missing.

Additional pharmacokinetic parameters may be calculated for the plasma concentration *versus* time profiles, if considered appropriate and justified at the time of the pharmacokinetic analysis.

#### 6.2.4. Urinary Pharmacokinetic Parameters

The following pharmacokinetic parameters will be derived by standard NCA methods from the urine excretion profiles, for the first dose (Day 1, single-dose) and the last dose (Day 13, multiple-dose):

- Part 3:
  - givinostat.
  - givinostat metabolites (ITF2374, ITF2375, ITF2440 and ITF2563).

<i>Parameter</i>	<i>Description</i>
$R_{max}$	Maximum observed urinary excretion rate.
$t_{u_{max}}$	Time of occurrence of maximum urinary excretion rate.
$Amt_{CUM}$	Cumulative amount of drug excreted in urine, calculated as $\sum (Concentration \cdot Volume)$
$AURC_{0-t}$	Area under the urine excretion rate curve from time zero to last measurable observed excretion rate.
$REC\%$	Percentage of drug recovered in urine, calculated for givinostat as $100 \cdot \frac{Amt_{CUM}}{Dose}$ , and for givinostat metabolites as $100 \cdot \frac{Amt_{CUM, Metabolite} / MW_{Metabolite}}{Dose_{Givinostat} / MW_{Givinostat}}$ , where MW represents the molecular weight of the analyte. <i>Note: For givinostat, MW is 421.49 g/mol; for ITF2374, MW is 405.49 g/mol; for ITF2375, MW is 406.47 g/mol; for ITF2440, MW is 257.33 g/mol; and for ITF2563, MW is 178.19 g/mol.</i>
$CL_R/F$	Apparent renal clearance affected by F, calculated as $\frac{Dose}{AURC_{0-t}}$ .

## 7. STUDY ENDPOINTS

### 7.1. Primary

1. Givinostat and its metabolites plasma and urine concentrations and thereof derived pharmacokinetic parameters following single and multiple oral doses of givinostat (Part 3).

### 7.2. Secondary

1. Incidence and severity of AEs; changes in vital signs, physical examination, ECG and clinical laboratory tests following administration of single- and multiple-doses of givinostat (Part 3).

## 8. ANALYSIS POPULATIONS

The analysis populations are defined in accordance with the CSP [1].

For each study part, the subjects to be included in each analysis population will be reported in the respective Data Blind Review Process Minute ([Annex 5](#)).

For each study part, explanation of the reasons for exclusion of subjects from any analysis population will be provided in the respective Data Blind Review Process Minute ([Annex 5](#)) and in the CSR.

The list of subjects who complete each study part will be presented in [Annex 6](#).

### 8.1. Part 3

#### 8.1.1. Part 3 Safety Analysis Population

All subjects who receive at least one dose of the IMP in Part 3 of the study will constitute the Part 3 Safety Analysis Population.

Part 3 safety data analysis will be performed for all subjects in the Part 3 Safety Analysis Population.

#### 8.1.2. Part 3 Pharmacokinetic Analysis Population

##### 8.1.2.1. Part 3 Single-Dose Pharmacokinetic Analysis Population

Part 3 Single-Dose Pharmacokinetic Analysis Population will include all subjects enrolled in Part 3 of the study, who are expected to provide evaluable pharmacokinetic data for the givinostat single-dose administration (Day 1), without deviations affecting pharmacokinetic interpretation.

The following reasons justify the exclusion of the pharmacokinetic data of a subject from the Part 3 single-dose pharmacokinetic analysis population:

- Protocol violation considered to have a potentially relevant effect on the pharmacokinetic results of the study.

*NOTE: These protocol violations will be reported in the Data Blind Review Process Minute ([Annex 5](#)), and their impact will be assessed at the time of pharmacokinetic analysis.*

- Subject experienced vomiting or diarrhea.

*NOTE: Subjects that experienced vomiting or diarrhoea during this study part will be reported in the Data Blind Review Process Minute ([Annex 5](#)), and their exclusion will be assessed at the time of pharmacokinetic analysis.*



and/or SAS®.

In-text Tables and Figures will preferably be prepared in portrait format. Listings will preferably be prepared in landscape format.

The CSR will be written according to PPD SOPs and templates for reporting bioavailability/bioequivalence trials.

## 11.2. Planned Tables, Figures, and Subject Data Listings

The planned Tables, Figures, and Individual Data Listings for the CSR are listed below.

### 11.2.1. In-text Tables and Figures

For CSR Synopsis:

The following Tables will be produced for direct insertion in the Synopsis of the CSR:

Title
<b>PART 3</b>
Givinostat and Its Main Metabolites (ITF2374 and ITF2375): Summary Statistics of the Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1)
Givinostat Metabolites (ITF2440 and ITF2563): Summary Statistics of the Plasma Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1)
Givinostat and Its Main Metabolites (ITF2374 and ITF2375): Summary Statistics of the Plasma Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13)
Givinostat Metabolites (ITF2440 and ITF2563): Summary Statistics of the Plasma Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13)
Givinostat and Its Main Metabolites (ITF2374 and ITF2375): Summary Statistics of the Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1)
Givinostat Metabolites (ITF2440 and ITF2563): Summary Statistics of the Urinary Pharmacokinetic Parameters Following Single-Dose Administration of Givinostat (Day 1)
Givinostat and Its Main Metabolites (ITF2374 and ITF2375): Summary Statistics of the Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13)
Givinostat Metabolites (ITF2440 and ITF2563): Summary Statistics of the Urinary Pharmacokinetic Parameters Following Multiple-Dose Administration of Givinostat (Day 13)

The following Figures will be produced for direct insertion in the Synopsis of the CSR:

Legend
<b>PART 3</b>
Givinostat: Geometric Mean (95% CI) Plasma Concentration <i>Versus</i> Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat. A – Linear



In-text Figures	Legend
	Amount Urinary Excretion <i>Versus</i> Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale
Figure 3.J.1.	Givinostat Metabolite (ITF2563): Geometric Mean (95% CI) of Urinary Excretion <i>Versus</i> Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale
Figure 3.J.2.	Givinostat Metabolite (ITF2563): Geometric Mean (95% CI) of Cumulative Amount Urinary Excretion <i>Versus</i> Time Profile Following Single-Dose (Day 1) and Multiple-Dose (Day 13) Administration of Givinostat – Linear Scale

#### 11.2.2. Tables, Figures and Graphs Referred to But Not Included in the Text (Section 14 of CSR)

At least the following Tables and Figures should be produced for Section 14 of the CSR.  
If necessary, additional Tables and Figures should be prepared.

The following Tables and Figures will be compiled in a document, to be considered as Section 14 (Tables, Figures and Graph Referred To But Not Included in the Text) of the CSR.

End-of-Text Tables	Title
14.3. PART 3	
14.3.1. DEMOGRAPHIC DATA – PART 3	
14.3.1.1.	Demographic Data – Part 3 Safety Analysis Population
14.3.1.2.	Demographic Data – Part 3 Single-Dose Pharmacokinetic Analysis Population
14.3.1.3.	Demographic Data – Part 3 Multiple-Dose Pharmacokinetic Analysis Population
14.3.1.4.	Subjects Prematurely Discontinued After First Dosing
14.3.2. PHARMACOKINETIC DATA – PART 3	
14.3.2.1.	Plasma Pharmacokinetic Data
14.3.2.1.1.	Deviations from Blood Sampling Schedule
14.3.2.1.1.1.	Deviations from Blood Sampling Schedule
14.3.2.1.1.2.	Missing Blood Samples
14.3.2.1.1.3.	Other Important Protocol Deviations
14.3.2.1.2.	Plasma Concentrations
14.3.2.1.2.1.	Givinostat: Individual Data and Descriptive Statistics of Plasma Concentrations Following Single-Dose Administration of Givinostat (Day 1)
14.3.2.1.2.2.	Givinostat: Individual Data and Descriptive Statistics of Plasma Concentrations Following Single-Dose Administration of Givinostat (Day 1) – Male Subjects
14.3.2.1.2.3.	Givinostat: Individual Data and Descriptive Statistics of Plasma Concentrations Following Single-Dose Administration of Givinostat (Day 1) – Female Subjects
14.3.2.1.2.4.	Givinostat Metabolite (ITF2374): Individual Data and Descriptive Statistics of Plasma Concentrations Following Single-Dose Administration of Givinostat (Day 1)



### 11.2.3. List of Subject Data Listings (Section 16.2 of CSR)

The following Data Listings will be produced for Section 16.2. Subject Data Listings of the CSR.

If necessary, additional Data Listings should be prepared.

Listing No.	Title
<b>16.2. SUBJECTS DATA LISTING</b>	
<b>16.2.3. Part 3</b>	
<b>16.2.3.1.</b>	<b>Subject Disposition</b>
16.2.3.1.1.	Subject Disposition
<b>16.2.3.2.</b>	<b>Protocol Deviations</b>
16.2.3.2.1.	Blood Sampling Times Deviations
16.2.3.2.2.	Other Protocol Deviations
<b>16.2.3.3.</b>	<b>Excluded Data from the Pharmacokinetic Analysis</b>
16.2.3.3.1.	Plasma Concentrations
16.2.3.3.1.1.	Individual Data of Plasma Concentrations
16.2.3.3.2.	Plasma Concentration Versus Time Profiles
16.2.3.3.2.1.	Individual Plasma Concentration Versus Time Profiles – Linear Scale
16.2.3.3.2.2.	Individual Plasma Concentration Versus Time Profiles – Semi-Logarithmic Scale
16.2.3.3.3.	Urine Excretion
16.2.3.3.3.1.	Individual Data of Plasma Concentrations
16.2.3.3.4.	Urine Excretion Versus Time Profiles
16.2.3.3.4.1.	Urine Excretion Versus Time Profiles – Linear Scale
<b>16.2.3.4.</b>	<b>Demographic and Other Baseline Data</b>
16.2.3.4.1.	Demographic Data
16.2.3.4.2.	Fertility/Contraception
16.2.3.4.2.1.	Female Fertility/Contraception
16.2.3.4.2.2.	Male Fertility/Contraception
16.2.3.4.3.	Drugs of Abuse, Ethanol and Cotinine
16.2.3.4.4.	Viral Serology at Screening
16.2.3.4.5.	Previous and Concomitant Medication
16.2.3.4.6.	SARS-COV-2 Test
<b>16.2.3.5.</b>	<b>Compliance</b>
16.2.3.5.1.	Investigational Product Administration
<b>16.2.3.6.</b>	<b>Individual Pharmacokinetic Data</b>
16.2.3.6.1.	Givinostat: Individual Pharmacokinetic Data
16.2.3.6.1.1.	Givinostat: Individual Drug Plasma Concentration Data
16.2.3.6.1.2.	Givinostat: Individual Plasma Pharmacokinetic Parameters
16.2.3.6.1.3.	Givinostat: Individual Plasma Pharmacokinetic Profiles
16.2.3.6.1.4.	Givinostat: Individual Urine Excretion Data
16.2.3.6.1.5.	Givinostat: Individual Urine Pharmacokinetic Parameters
16.2.3.6.1.6.	Givinostat: Individual Urine Pharmacokinetic Profiles
16.2.3.6.1.7.	Givinostat: Documentation of Statistical Analysis
16.2.3.6.1.7.1.	Givinostat: Plasma Non-Compartmental Analysis Output
16.2.3.6.1.7.2.	Givinostat: Plasma Non-Compartmental Analysis Plots
16.2.3.6.1.7.3.	Givinostat: Urine Non-Compartmental Analysis Output