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ICH HARMONISED GUIDELINE

DRUG INTERACTION STUDIES
M12

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ICH Consensus Guideline

Table of Contents

Table of Contents	i
1. INTRODUCTION.....	1
1.1 Objective	1
1.2 Background	1
1.3 Scope.....	1
1.4 General Principles	2
2. IN VITRO EVALUATION.....	4
2.1 Evaluation of Metabolism-Mediated Interactions	4
2.1.1 Drug as a Substrate of Metabolizing Enzymes.....	4
2.1.2 Drug as an Inhibitor of CYP Enzymes	5
2.1.2.1 Reversible Inhibition	5
2.1.2.2 Time-Dependent Inhibition.....	6
2.1.3 Drug as an Inhibitor of UGTs	6
2.1.4 Drug as an Inducer of CYP Enzymes	7
2.1.4.1 Basic ‘mRNA Fold-Change’ Method.....	8
2.1.4.2 Correlation Methods	8
2.1.4.3 Basic Kinetic Model	9
2.1.4.4 Additional Considerations Related to Induction	9
2.2 Evaluation of Transporter-Mediated Interactions	10
2.2.1 Drug as a Substrate of Transporters.....	10
2.2.1.1 Data Analysis and Interpretation	11
2.2.2 Drug as an Inhibitor of Transporters.....	11
2.2.3 Drug as an Inducer of Transporters.....	12
2.3 DDI Potential of Metabolites	12
2.3.1 Metabolite as a Substrate	13
2.3.2 Metabolite as an Inhibitor	13
2.3.3 Metabolite as an Inducer.....	14
3. CLINICAL EVALUATION	14
3.1 Types of Clinical DDI Studies (Terminology)	14

3.1.1 Stand-alone and Nested DDI Studies	15
3.1.2 DDI Studies with Index Precipitants and Index Substrates.....	15
3.1.3 DDI Studies with Expected Concomitant Drugs	16
3.1.4 Cocktail Approach.....	16
3.1.5 Biomarker Approach	16
3.2 Study Planning and Considerations for Clinical DDI Studies	16
3.2.1 Study Design.....	16
3.2.1.1 Study Population and Number of Subjects	16
3.2.1.2 Dose.....	17
3.2.1.3 Single or Multiple Doses	17
3.2.1.4 Route of Administration and Formulations.....	18
3.2.1.5 Parallel Versus Crossover Studies	18
3.2.1.6 Timing of Drug Administration	19
3.2.1.7 Co-Medications and Other Extrinsic Factors Affecting DDIs.....	19
3.2.1.8 Sample and Data Collection.....	20
3.2.2 Specific Considerations for Nested DDI Studies.....	20
3.2.3 Considerations for CYP-Mediated Interactions.....	21
3.2.3.1 The Investigational Drug as a Substrate for CYP Enzymes.....	21
3.2.3.2 The Investigational Drug as an Inhibitor or an Inducer of CYP Enzymes	22
3.2.4 Considerations for Evaluation of UGT-Mediated Interactions.....	23
3.2.4.1 Investigational Drug as a Substrate of UGTs	23
3.2.4.2 Investigational Drug as an Inhibitor of UGTs	24
3.2.4.3 Investigational Drug as an Inducer of UGTs.....	24
3.2.5 Considerations for Evaluation of Transporter-Mediated Interactions	24
3.2.5.1 Investigational Drug as a Substrate of Transporters.....	24
3.2.5.2 Investigational Drug as an Inhibitor of Transporters.....	26
3.2.5.3 Investigational Drug as an Inducer of Transporters.....	26
3.2.6 Cocktail Studies-Considerations for CYP or Transporter Cocktail Studies	27
3.2.7 Considerations for Biomarker Approach	27
3.2.7.1 Investigational Drug as an Inhibitor of Hepatic OATP1B.....	27
4. OTHER TOPICS	28
4.1 Pharmacogenetics	28
4.2 Therapeutic Protein DDIs	29
4.2.1 Proinflammatory Cytokine-Related Mechanism	29
4.2.2 Antibody-Drug Conjugates.....	30
5. REPORTING AND INTERPRETING CLINICAL DDI STUDY RESULTS.....	30

5.1 Pharmacokinetic Data Analysis	31
5.1.1 Non-Compartmental Analysis (NCA)	31
5.1.2 Population PK Analysis.....	31
5.2 Reporting DDI Results.....	31
5.3 Interpreting DDI Study Results	32
5.3.1 Determination of No-Effect Boundaries.....	32
5.3.2 Investigational Drug as a Precipitant of DDIs: Classification System.....	32
5.3.3 Extrapolating Study Results	33
5.3.3.1 Extrapolating Complex Scenarios.....	34
6. RISK ASSESSMENT AND MANAGEMENT	34
7. APPENDICES	35
7.1 Glossary	35
7.2 Protein Binding	38
7.3 In Vitro Evaluation of Metabolism-Based DDIs	39
7.3.1 In Vitro Systems.....	39
7.3.2 Investigational Drug as an Enzyme Substrate: Reaction Phenotyping.....	40
7.3.2.1 Metabolic Pathway Identification	41
7.3.2.2 Metabolic Enzyme Identification.....	41
7.3.3 Investigational Drug as an Enzyme Inhibitor	42
7.3.4 Investigational Drug as an Enzyme Inducer	43
7.4 In Vitro Evaluation of Transporter-Based DDIs.....	45
7.4.1 In Vitro Systems.....	45
7.4.2 Investigational Drug as a Transporter Substrate.....	47
7.4.3 Investigational Drug as a Transporter Inhibitor.....	48
7.5 Predictive Modeling.....	48
7.5.1 Using Mechanistic Static Models for DDI Predictions.....	49
7.5.1.1 Evaluation of an Investigational Drug as a DDI Precipitant of CYP-Mediated DDIs	49
7.5.1.2 Evaluation of Investigational Drug as an Object of CYP-Mediated DDIs	49
7.5.1.3 Evaluation of The Potential for Transporter-Mediated DDIs.....	51
7.5.2 Using PBPK Models to Predict Enzyme or Transporter-Based DDIs.....	51
7.5.2.1 Potential Applications of PBPK to the Evaluation of CYP-Mediated DDIs.....	51
7.5.2.1.1 Modeling Considerations - PBPK for Evaluation of CYP Interactions for Drugs as Substrates	52
7.5.2.1.2 Modeling Considerations - PBPK for Evaluation of CYP Interactions for Drugs as Precipitants.....	52

7.5.2.2 Potential Applications of PBPK to the Evaluation of Transporter-Mediated DDIs .	53
7.5.2.2.1 Modeling Considerations - Drug as a Transporter Substrate	54
7.5.2.2.2 Modeling Considerations - Drug as a Transporter Inhibitor	54
7.6 List of Drugs that can be used in In Vitro Studies	54
7.6.1 CYP Enzymes.....	54
7.6.1.1 CYP Enzyme Substrates for In Vitro Studies.....	54
7.6.1.2 CYP Enzymes Inhibitors/inducers for In Vitro Studies	55
7.6.2 UGTs.....	56
7.6.2.1 UGT Substrates for In Vitro Studies.....	56
7.6.2.2 UGT Inhibitors for In Vitro Studies.....	56
7.6.3 Transporters	57
7. 7 List of Drugs that can be used in Clinical Studies	57
7.7.1 CYPs Enzymes	57
7.7.1.1 CYP Enzyme Substrates for Clinical Studies.....	57
7.7.1.2 CYP Enzyme Inhibitors for Clinical Studies	59
7.7.1.3 CYP Enzyme Inducers for Clinical Studies	59
7.7.2 UGTs.....	60
7.7.3 Transporters	61
7.7.3.1 Transporter Substrates for Clinical Studies.....	61
7.7.3.2 Transporter Inhibitors for Clinical Studies.....	63
8. REFERENCES.....	64

1. INTRODUCTION

1.1 Objective

This guideline provides recommendations to promote a consistent approach in designing, conducting, and interpreting enzyme- or transporter-mediated *in vitro* and clinical drug-drug interaction (DDI) studies during the development of a therapeutic product. A consistent approach will reduce uncertainty for the pharmaceutical industry to meet the requirements of multiple regulatory agencies and lead to more efficient utilization of resources. In addition, this approach will lead to the effective and safe treatment for patients who take multiple medications.

1.2 Background

In clinical practice, patients are often prescribed more than one drug, which can result in a DDI. Some patients, in particular fragile older patients or patients with serious or multiple health issues, can be prescribed a large number of different drugs (i.e., polypharmacy). The occurrence of DDIs is a common clinical problem that can increase the risk of adverse effects, sometimes leading to hospital admissions. Alternatively, some DDIs can reduce or enhance efficacy of the treatment. Hence, it is important to consider an investigational drug's potential to interact with other drugs.

Regional guidelines for investigations of DDIs have been available for decades and have undergone several updates as scientific progress has been made. In general, the proposed approach to the evaluation of interaction potential of investigational drugs has been similar between regions, but despite harmonization initiatives, some differences have remained. This ICH guideline aims to harmonize recommendations for *in vitro* and clinical evaluation of DDIs.

This guideline provides general recommendations on how to evaluate the DDI potential of an investigational drug. It is recognized that the DDI evaluation is generally tailored based on the specific drug, intended patient population, and therapeutic context. Alternative approaches may be acceptable if properly justified. The focus of the guideline is the development of new drugs, but if new scientific information regarding the potential for DDIs is obtained after drug approval, additional DDI evaluation should be considered.

1.3 Scope

The scope of the guideline is limited to pharmacokinetic interactions, with a focus on metabolic enzyme- and transporter-mediated interactions. These aspects in general apply to the development of small chemical molecules. DDI evaluation of biologics is covered briefly, with focus on monoclonal antibodies and antibody-drug conjugates. Recommendations are provided on how to investigate interactions mediated by inhibition or induction of enzymes or transporters, both *in vitro* and *in vivo* (the terms 'clinical' and '*in vivo*' are used interchangeably in this document), and on how to translate the results to appropriate treatment recommendations. The guideline also includes recommendations on how to address metabolite-mediated interactions. The use of model-based data evaluations and DDI predictions are also covered.

The development and emergence of other modalities, such as oligonucleotides, small interfering ribose nucleic acids, and peptides is acknowledged. However, these modalities are out of scope for this guideline. Where applicable, regional guidelines should be considered.

Other types of pharmacokinetic interactions, such as impact on absorption (e.g., gastric pH change, gastric motility change, formation of chelation or complexation), food effects, or protein binding displacement, are not part of this document and may be covered by regional guidelines. Similarly, DDIs that are a result of pharmacodynamic interactions are beyond the scope of this guideline.

1.4 General Principles

The potential for an investigational drug to cause DDIs should be investigated in a stepwise manner during drug development. The potential for an investigational drug to cause pharmacokinetic interactions both as an object (effect of other drugs on the investigational drug) and as a precipitant (effect of the investigational drug on concomitant drugs) should be evaluated. All aspects mentioned below are further expanded and discussed later in the document. Note that historically, some regions have used the term “victim” instead of “object” and the term “perpetrator” instead of “precipitant.” Because objects of DDIs are substrates of enzymes and/or transporters, the term “substrate” in this guideline refers to drugs that may be objects of DDIs.

Evaluating the potential of an investigational drug as an object of a metabolic enzyme- or transporter-mediated DDI involves identification of the principal routes of the drug’s elimination. For drugs whose major elimination routes are not urinary excretion of unchanged parent drug or that are not biologics eliminated through unspecific catabolism, the keystone of the identification of principal elimination routes is a well-performed clinical mass balance study. In some instances, (e.g., if a large part of the dose is found as unchanged parent drug in feces), an absolute bioavailability study can also be a useful complement to aid interpretation of principal elimination. Using data from the mass balance study, the quantitative contributions of the different elimination pathways should be estimated based on the amount of dose excreted as primary and secondary metabolites along specific routes. For quantitatively important elimination pathways, *in vitro* and clinical studies should be used to identify the main enzymes and/or transporter proteins involved in these pathways. The ability to predict interactions affecting the investigational drug is dependent on the identification of these proteins.

Evaluating the DDI potential of an investigational drug as a precipitant involves characterizing the effect of the drug on enzymes and transporters. This evaluation often starts with *in vitro* experiments to elucidate potential DDI mechanisms. Identification of DDI risks should then be followed by clinical DDI studies based on mechanistic knowledge, and the results should be translated to appropriate clinical management recommendations for drugs as a precipitant of DDIs.

The results of DDI evaluations inform the protocols for clinical studies in patients regarding the use of concomitant drugs. Information about the interaction potential should be gained as early in drug development as practically possible to ensure safety and avoid unnecessary restrictions of concomitant medications and/or exclusion of patients who require the concomitant medications in clinical studies. The timing of the different non-clinical and clinical studies is dependent on the context and type of product; some general recommendations are given below. Predictive modeling (see Section 7.5) can also assist evaluation of the DDI potential.

- *In vitro* data on the investigational drug as a substrate of metabolic enzymes is generally recommended to be obtained before starting the clinical phase in patients to evaluate metabolic stability and identify the potential main metabolic pathway(s) and enzyme(s) that metabolize the investigational drug (reaction phenotyping studies). If *in vitro* studies suggest the possibility of a clinically significant interaction with inhibitors or inducers of a metabolic enzyme, additional actions (e.g., clinical DDI studies) may be needed prior to studies in patients. Until appropriate additional actions are taken, a conservative strategy, such as excluding patients on certain concomitant drugs that are inhibitors and/or inducers, may be needed.
- ADME (absorption, distribution, metabolism and excretion) properties determine whether *in vitro* data of the investigational drug as a substrate for transport proteins should be collected. If a drug has limited absorption or is expected to undergo significant active hepatic uptake, biliary excretion, or active renal secretion as unchanged drug, the relevant transporter(s) should be identified *in vitro* before initiating clinical studies in patients to avoid restrictions on concomitant medications in protocols.
- *In vitro* data on the effects of the investigational drug as a precipitant on the major cytochrome P450 (CYP) enzymes and transporters is generally recommended to be available before initiating larger studies in patients.
- The results of the mass balance study are generally recommended to be available before starting phase 3. Based on results of the mass balance study and *in vitro* studies, clinical studies with strong index enzyme inhibitors and inducers should be considered to confirm and quantify the main metabolism pathways and define the risk for clinically significant DDIs.
- The pharmacokinetic DDI potential of metabolites with significant plasma exposure or pharmacological activity (see Section 2.3) are recommended to be considered similarly as for the parent drug, but these investigations can generally be completed later in development when more knowledge about the exposure and activity of metabolites is available.

2. IN VITRO EVALUATION

2.1 Evaluation of Metabolism-Mediated Interactions

In vitro studies are important first steps to identify risks for a drug to be an object or precipitant of DDIs related to inhibition or induction of drug metabolizing enzymes.

2.1.1 Drug as a Substrate of Metabolizing Enzymes

Typically, an *in vitro* screening to identify the main enzymes responsible for the metabolism of a new drug is performed early in drug development. If oxidative metabolism is important, the identification of catalyzing enzymes usually starts by determining whether the investigational drug is an *in vitro* substrate for the most common CYP enzymes involved in drug metabolism: CYP1A2, CYP2B6, CYP2C8, CYP2C9, CYP2C19, CYP2D6, and CYP3A (CYP3A4 and CYP3A5) using *in vitro* phenotyping experiments. If the drug is not found to undergo metabolism by these major CYP enzymes, other enzymes can be investigated. These additional enzymes can include, but are not limited to:

- Other CYP enzymes, including CYP2A6, CYP2E1, CYP2J2, and CYP4F2, and other phase 1 enzymes, including alcohol/aldehyde dehydrogenase (ADH/ALDH), aldehyde oxidase (AO), carboxylesterase (CES), flavin monooxygenase (FMO), monoamine oxidase (MAO), and xanthine oxidase (XO).
- Phase 2 enzymes: The most frequently evaluated phase 2 enzymes, Uridine 5'-diphosphoglucuronosyltransferases (UGTs), are responsible for glucuronide conjugation of drugs and metabolites. Potential UGTs to be investigated *in vitro* include: UGT1A1, 1A3, 1A4, 1A6, 1A9, 1A10, 2B4, 2B7, 2B10, 2B15, and 2B17.
- Other phase 2 enzymes, including glutathione S-transferases (GSTs), N-acetyltransferases (NATs), sulfotransferases (SULTs).

Details on the experimental setup for *in vitro* studies to identify enzymes catalyzing the main elimination pathways are given in Sections 7.3.1 and 7.3.2.

In vitro phenotyping, metabolic profiling, and mass balance studies are generally used to identify and quantify the various elimination pathways of a drug. An enzyme that is estimated to contribute to $\geq 25\%$ of total elimination generally needs additional clinical characterization to quantify the risk of interaction with the investigational drug as a substrate. The characterization is often done by performing clinical DDI studies using a strong index inhibitor of the enzyme when available (Refer to Section 3.2.3.1). For some enzymes, pharmacogenetic studies may substitute for clinical DDI studies with a strong index inhibitor (refer to Section 4.1). A clinical study with a strong inducer is also generally needed to fully characterize the risk of DDI, since inducers can up-regulate expression of multiple enzymes and some transporters (except CYP2D6, which is generally considered not inducible by drugs) (Refer to Section 3.2.3.1).

2.1.2 Drug as an Inhibitor of CYP Enzymes

An investigational drug's potential to inhibit CYP1A2, CYP2B6, CYP2C8, CYP2C9, CYP2C19, CYP2D6, and CYP3A in both a reversible manner and time-dependent manner (TDI) should be evaluated.

2.1.2.1 Reversible Inhibition

In the reversible inhibition experiments, a K_i (inhibition constant) is usually determined experimentally or estimated based on half-maximal inhibitory concentration (IC_{50}) using unbound concentrations (refer to Section 7.3.3). If the initial experiments testing a sufficiently high concentration of the investigational drug already indicate that the $K_{i,u}$ will be markedly higher than the cutoffs given (see below), the risk for clinical inhibition can normally be excluded without further data.

The risk for reversible enzyme inhibition can be excluded based on *in vitro* data ("basic method") if:

$$K_{i,u} > 50 \times C_{max,u} \text{ (i.e., } \frac{C_{max,u}}{K_{i,u}} < 0.02)$$

$K_{i,u}$ is the unbound inhibition constant.

$C_{max,u}$ is the unbound C_{max} at the highest recommended dose at steady state.

For the purposes of calculating the unbound measure of concentration, the measured $f_{u,p}$ (fraction unbound in plasma) can be used for all drugs, including highly protein bound drugs (i.e., >99% protein binding), if the accuracy and precision of the protein binding measurement has been demonstrated. Such a demonstration should include validation data of the protein binding assay including bioanalytical method and appropriate positive controls (i.e., drugs with high binding to relevant plasma proteins). If the reliability of $f_{u,p}$ measurements <1% cannot be demonstrated, a default value of 1% unbound fraction in plasma (i.e., $f_{u,p} = 0.01$) should be used (see Section 7.2 for details). This consideration for $f_{u,p}$ applies in other contexts where basic method, mechanistic static, and dynamic models (often referred as physiologically based pharmacokinetic (PBPK) modeling) can be used to interpret the *in vitro* results of enzyme and transporter inhibition/induction experiments.

For orally administered drugs that are inhibitors of CYP3A, the risk of intestinal CYP3A inhibition can be excluded if:

$$K_{i,u} > 0.1 \times \frac{\text{maximum clinical dose}}{250 \text{ mL}} \text{ (i.e., } \frac{\text{Dose}/250 \text{ mL}}{K_{i,u}} < 10)$$

If risk for clinical inhibition cannot be excluded using this basic method, mechanistic static and/or PBPK models can be used to interpret the *in vitro* experiment results (refer to Section 7.5). If *in vitro* data and modeling do not exclude the risk for clinically relevant inhibition, a clinical DDI study with a sensitive index substrate should be conducted.

If a clinical study using a substrate for an enzyme that was inhibited *in vitro* by an investigational drug with a low $K_{i,u}$ shows lack of inhibition, then the risk for clinical inhibition can be excluded for other enzymes having a larger $K_{i,u}$. Such an inference should be made only for the enzymes that are expressed at the same site and for which the inhibition potencies are determined in the same HLM/hepatocyte batch (rank order approach) (1). Of note, an orally administered drug can inhibit intestinal metabolic enzymes (e.g., CYP3A) in addition to hepatic enzymes. In such situations, the risk for inhibition of CYP3A in the gastrointestinal (GI) tract should be considered even if systemic inhibition of CYP3A can be excluded using the rank order approach based on a negative clinical study on another CYP enzyme. In the presence of inhibitory metabolites of an investigational drug, their contribution should also be considered when using rank order approach to determine if clinical studies should be conducted.

2.1.2.2 Time-Dependent Inhibition

If an *in vitro* assay (described in Section 7.3.3) indicates an increased enzyme inhibition potential with drug pre-incubation, the inhibitory kinetic parameters (K_I and k_{inact}) will be determined and the following equation can be used as the basic method to evaluate the risk for TDI (2). The risk for clinical inhibition can be excluded based on *in vitro* data if:

$$\frac{(k_{obs} + k_{deg})}{k_{deg}} < 1.25$$

$$\text{where } k_{obs} = \frac{(k_{inact} \times 5 \times C_{max,u})}{(K_{I,u} + 5 \times C_{max,u})}$$

k_{obs} is the apparent first-order inactivation rate constant of the affected enzyme.

k_{deg} is the apparent first-order degradation rate constant of the affected enzyme (refer to Table 6).

$K_{I,u}$ is the unbound inhibitor concentration causing half-maximal inactivation.

k_{inact} is the maximal inactivation rate constant.

$C_{max,u}$ is the maximal unbound plasma concentration of the inhibitor drug at steady state. The $f_{u,p}$ should be set to 1% if the reliability of $f_{u,p}$ measurements <1% cannot be demonstrated (also refer to Section 2.1.2.1).

Note: $C_{max,u}$ and $K_{I,u}$ should be expressed in the same unit (e.g., in a molar concentration unit).

If the above ratio is ≥ 1.25 , mechanistic static and/or PBPK models can be used to interpret the *in vitro* experiment results (refer to Section 7.5). If *in vitro* data and modeling do not exclude the risk for clinical inhibition, a clinical DDI study with a sensitive index substrate should be conducted. The rank order approach, mentioned above for reversible inhibitors, does not apply to TDIs.

2.1.3 Drug as an Inhibitor of UGTs

It is recognized that a drug which is not a substrate of an enzyme can still be an inhibitor. However, considering the generally limited magnitude of UGT inhibition-mediated DDIs (3), a routine evaluation of investigational drugs to inhibit UGTs may not be warranted. If direct glucuronidation is the major elimination pathway of an investigational drug, it is recommended to study *in vitro* whether the drug can inhibit UGTs. Potential UGTs to be studied include UGT1A1, UGT1A4,

UGT1A9, UGT2B7, and UGT2B15. The evaluation is usually performed using recombinant UGTs or human liver microsome (HLM) with relatively selective substrates (refer to Table 8, Section 7.6.2.1 for an illustrative list of substrates) (4). When an investigational drug will be commonly administered with a drug that is mainly metabolized by direct glucuronidation, it is recommended to evaluate *in vitro* the potential inhibitory effect of the investigational drug on the UGT isoform(s) responsible for the elimination of the other drug.

Due to limited availability of data from clinical DDI studies that evaluate inhibition of UGT isoenzymes, cutoffs for determining DDI risk using basic models like those for CYP enzymes have not been established. This is an area of ongoing research and, in the interim, sponsors can consider the same criterion that is applied to CYPs (i.e., $C_{\max,u}/K_{i,u} < 0.02$) or propose an alternative with justification.

2.1.4 Drug as an Inducer of CYP Enzymes

An investigational drug's potential to induce enzymes via activation of nuclear receptors pregnane X receptor (PXR), constitutive androstane receptor (CAR), aryl hydrocarbon receptor (AhR), and if relevant other drug regulation pathways, should be evaluated. For technical advice on the experiments, refer to Section 7.3.4.

To assess the DDI liability of a drug as an inducer, studies should be performed in human hepatocytes from at least 3 individual donors, and the extent of enzyme induction should be measured at the mRNA level. The enzymes CYP3A4, CYP2B6 and CYP1A2 should always be included as markers of induction mediated via PXR/CAR (CYP3A4, CYP2B6) and AhR (CYP1A2). Enzymes CYP3A4 and CYP2C8, CYP2C9, and CYP2C19 are induced via activation of the PXR. CYP2C8, CYP2C9, and CYP2C19 are generally less inducible compared to CYP3A4. Therefore, the potential of the investigational drug to induce CYP2C8, CYP2C9, and CYP2C19 should be evaluated *in vitro* and/or *in vivo* when it has been shown to be an inducer of CYP3A4 in a clinical study. A negative result for induction in a clinical study with a sensitive CYP3A4 substrate can be used to rule out an investigational drug's potential to induce CYP2C8, CYP2C9, and CYP2C19 enzymes provided that the potential of CYP3A4 inhibition by the drug and its major metabolite(s) can be excluded. When *in vitro* induction of CYP2C19 is evaluated, the mRNA responses to inducers are often limited, and thus the activity should be measured using a probe substrate to evaluate the CYP2C19 induction potential of the investigational drug (See Section 7.3.4).

As described below, there are several methods that can be used to interpret mRNA data from *in vitro* induction experiments and to assess the *in vivo* potential of a drug to induce enzymes. It is recommended to first use the basic qualitative method (mRNA fold-change). If the basic method indicates induction potential, the evaluation can continue using more quantitative approaches (e.g., correlation methods), provided it is possible to study a wide range of concentrations of the

investigational drug to determine induction parameters (e.g., $E_{\max}$ and EC_{50}). For the more quantitative approaches, one qualified batch of hepatocytes is sufficient. The basic method only uses *in vitro* data from the investigational drug, whereas correlation methods compare the induction response of the drug to that of multiple established clinical inducers of the enzyme of interest.

In addition, mechanistic static or PBPK models can potentially be used (refer to Section 7.5). If a risk for induction cannot be excluded based on *in vitro* data and modeling, clinical studies with sensitive substrates of the enzymes of interest should be conducted.

2.1.4.1 Basic ‘mRNA Fold-Change’ Method

The induction results should be evaluated separately for each donor. The levels of mRNA should be compared to the control (vehicle) incubations, and a fold-change over the vehicle control should be calculated. *In vivo* induction potential cannot be excluded if the drug in hepatocytes from at least one donor meets both of the following criteria, and further evaluation of the induction potential should be conducted:

- increases mRNA expression of a CYP enzyme in a concentration-dependent manner; and
- the fold-change of CYP mRNA expression is ≥ 2 -fold at concentrations $\leq 50 \times C_{\max,u}$.

When the response of the positive control is <6 -fold increase in mRNA, the induction potential cannot be ruled out for an investigational drug that increases CYP enzyme mRNA less than 2-fold of the vehicle control but more than 20% of the response of the positive control. Further evaluation is recommended when there is an inconclusive finding (see Section 7.3.4 and ICH M12 Questions and Answers Document).

To calculate the percent of the response to the positive control, the following equation should be used:

$$\% \text{ of positive control} = \frac{(\text{mRNA fold increase of test drug treated cells} - 1)}{(\text{mRNA fold increase of positive control} - 1)} \times 100$$

2.1.4.2 Correlation Methods

Correlation methods compare the induction effect of the investigational drug to that of established *in vivo* inducers of the enzyme of interest (5). The magnitude of an *in vivo* induction effect (e.g., area under the curve (AUC) ratio of sensitive substrate in the presence and absence of inducers) of an investigational drug is predicted based on a calibration curve of relative induction scores (RIS, see equation below) or $C_{\max,u}/EC_{50u}$ versus the *in vivo* induction effect for a set of known inducers of the same enzyme (also refer to Section 7.3.4). If the predicted AUC ratio > 0.8 , the analysis can be used to exclude the risk for *in vivo* induction.

$$RIS = \frac{E_{max} \times C_{max,u}}{EC_{50,u} + C_{max,u}}$$

$EC_{50,u}$ is the unbound concentration causing half the maximal effect.

E_{max} is the maximum induction effect.

$C_{max,u}$ is the unbound maximum plasma concentration of a drug at steady state. The $f_{u,p}$ should be set to 1% if the reliability of $f_{u,p}$ measurements <1% cannot be demonstrated (also refer to Section 2.1.2.1).

Sometimes, E_{max} or $EC_{50,u}$ cannot be estimated due to an incomplete *in vitro* induction profile (e.g., limited by solubility or cytotoxicity of tested drug). An alternative correlation approach can be used if the method is validated (6).

2.1.4.3 Basic Kinetic Model

Mechanistic models have been proposed to predict the sum of different interaction processes (reversible inhibition, TDI, induction) systemically as well as in the GI tract. This approach is further discussed in Section 7.5.

A limited version of this approach is described as below. If $R > 0.8$, the analysis can be used to exclude the risk for *in vivo* induction.

$$R = \frac{1}{1 + d \times \frac{(E_{max} \times 10 \times C_{max,u})}{(EC_{50,u} + 10 \times C_{max,u})}}$$

R is predicted AUC ratio of a sensitive enzyme substrate with and without an inducer

$C_{max,u}$ is the unbound maximum plasma concentration in plasma. The $f_{u,p}$ should be set to 1% if the reliability of is 0.01, $f_{u,p}$ measurements <1% cannot be demonstrated (also refer to Section 2.1.2.1).

d is the scaling factor (7). If the scaling factor has not been determined in a calibrated hepatocyte batch (see Section 7.1.4), $d=1$ should be used.

If the above methods indicate that the investigational drug has the potential to induce metabolizing enzymes (using specific cutoff values mentioned above or developed by individual laboratories for these methods), the enzyme induction potential of the investigational drug should be further investigated by conducting a clinical DDI study with a sensitive index substrate or using mechanistic models (refer to Section 7.5).

2.1.4.4 Additional Considerations Related to Induction

Currently, *in vitro* methods to evaluate UGT induction are not well established. If an investigational drug has been observed to be an inducer of CYP enzymes via activation of nuclear receptors such as PXR or CAR, it is likely that UGTs regulated through these receptors will be induced. Refer to Section 3.2.4.3 which describes conducting clinical DDI studies mediated by induction of UGT for more considerations.

In vitro induction studies can also detect enzyme down-regulation. However, research in this area is presently very limited, and the mechanisms behind these effects are unclear. If concentration-dependent down-regulation (<50% of the mRNA control response) is observed *in vitro* and is not attributable to cytotoxicity, additional *in vitro* or clinical studies can be considered to understand the potential clinical consequences.

2.2 Evaluation of Transporter-Mediated Interactions

2.2.1 Drug as a Substrate of Transporters

P-glycoprotein (P-gp) and breast cancer resistance protein (BCRP) are efflux transporters expressed in the GI tract and can affect the oral bioavailability of drugs. Thus, the possibility of being a substrate of P-gp and/or BCRP is often evaluated *in vitro* for investigational drugs given orally. Because P-gp and BCRP are also expressed in the liver (P-gp, BCRP) and kidneys (P-gp), an *in vitro* study should be considered for a drug if biliary excretion or active renal secretion is a major elimination pathway of the drug. In addition, evaluating P-gp- and BCRP-mediated transport may be useful in the assessment of the drug's brain penetration.

Organic anion transporting polypeptide (OATP)1B1 and OATP1B3 are important hepatic uptake transporters. Examination of whether an investigational drug is a substrate for OATP1B1 and 1B3 should be considered if hepatic metabolism or biliary excretion accounts for $\geq 25\%$ of elimination of a drug or if the pharmacological target of a drug is in the liver.

Renal uptake transporters (Organic anion transporter (OAT)1, OAT3, and Organic cation transporter (OCT)2) and renal efflux transporters (Multidrug and toxin extrusion protein (MATE)1 and MATE2-K) are often involved in active renal secretion of drugs. *In vitro* studies to evaluate a drug as a substrate of these transporters should be considered if the drug clearance by renal active secretion is $\geq 25\%$ of its systemic clearance. Assuming there is no reabsorption (e.g., passive reabsorption is equal to passive secretion, and there is no active reabsorption), active secretion clearance can be calculated as $(CL_r - (f_{u,p} \times GFR))$, where GFR is glomerular filtration rate and CL_r is renal clearance. If pharmacokinetic data following intravenous administration are not available, systemic clearance may be computed by multiplying apparent total clearance by estimated bioavailability.

In addition to the above-mentioned transporters, *in vitro* evaluation of a drug as substrate of additional transporters may be considered on a case-by-case basis as new information emerges and our understanding evolves. For example, multidrug resistance-associated protein 2 (MRP2) is also an efflux transporter in similar locations as P-gp and BCRP; OATP2B1 is an uptake transporter present in the liver and intestine and is involved in absorption of certain drugs; and OCT1 is a hepatic transporter mediating the uptake of some drugs into the liver. The decision to evaluate additional transporters can take into consideration the site of action, passive permeability, and knowledge about absorption and elimination pathways of a drug.

2.2.1.1 Data Analysis and Interpretation

When examining the possibility that an investigational drug is a substrate of transporters, *in vitro* studies should be performed using experimental systems with the transporter activity confirmed using probe substrates and inhibitors (refer to Tables 10 and 11, Section 7.6.3 for some examples). Further details about considerations when performing *in vitro* studies are described in Sections 7.4.1 and 7.4.2.

For uptake studies, if there is significant uptake of a tested drug in transporter-expressed cells relative to the vehicle control-transfected cells (e.g., ≥ 2 -fold of controls), and the uptake in transporter-expressed cells can be inhibited by more than 50% by a known inhibitor of the transporter, the tested drug can be considered a substrate of the transporter examined.

For bidirectional transport studies, if there is significant directional transport of a tested drug in transporter-expressed cells relative to un-transfected or parental cells (e.g., net efflux ratio ≥ 2) or Caco-2 cells (e.g., efflux ratio ≥ 2), and the efflux ratio can be inhibited by more than 50% by a known inhibitor of the transporter, the tested drug can be considered as a substrate of the transporter examined.

A cutoff other than 2 or a specific relative ratio to positive controls can be used if prior experience with the cell system used justifies these alternative methods. Sponsors can also propose criteria for vesicle assays based on prior experience and internal data.

If *in vitro* studies indicate that a drug is a substrate of a transporter, clinical studies should be considered. Refer to Section 3.2.5.1 for more details.

2.2.2 Drug as an Inhibitor of Transporters

Studies should be conducted to evaluate whether an investigational drug is an inhibitor of P-gp, BCRP, OATP1B1, OATP1B3, OAT1, OAT3, OCT2, MATE1 and MATE2-K. Sponsors can consider evaluating the inhibition potential of a drug on other transporters as necessary. The decision to evaluate additional transporters can take into consideration if commonly coadministered drugs are substrates for these transporters. *In vitro* studies should be performed using an experimental system whose transport activity is confirmed using probe substrates and inhibitors (see Section 7.6.3 for more details). Considerations about how *in vitro* studies should be conducted are described in Sections 7.4.1 and 7.4.3.

The risk for transporter inhibition by an investigational drug in humans can be excluded based on *in vitro* data using the basic methods and the criteria outlined in Table 1. The contribution of drug metabolites to transporter inhibition should also be considered (see Section 2.3.2).

Table 1: Recommended ratio and cutoff value for drug as inhibitor of transporters

P-gp or BCRP	$IC_{50,u}^* > 0.1 \times (\text{Dose}/250 \text{ mL})$ (i.e., $(\text{Dose}/250 \text{ mL})/IC_{50,u} < 10$) for orally administered drugs
OATP1B1 or OATP1B3	$IC_{50,u} > 10 \times C_{\max, \text{inlet},u}^{\#}$ (i.e., $C_{\max, \text{inlet},u}/IC_{50,u} < 0.1$)
OAT1, OAT3, OCT2	$IC_{50,u} > 10 \times C_{\max,u}$ (i.e., $C_{\max,u}/IC_{50,u} < 0.1$)
MATE1/MATE2-K	$IC_{50,u} > 50 \times C_{\max,u}$ (i.e., $C_{\max,u}/IC_{50,u} < 0.02$)

$C_{\max,u}$ is unbound maximal plasma concentration of an inhibitor at steady state after therapeutic dose.

* The $K_{i,u}$ of an inhibitor approaches $IC_{50,u}$ when substrate concentration is much less than K_m assuming competitive inhibition (8).

$C_{\max, \text{inlet},u}$ is estimated unbound maximum plasma concentration of an inhibitor at liver inlet. $C_{\max, \text{inlet},u} = f_{u,p} \times (C_{\max} + (Fa \times Fg \times ka \times \text{Dose})/Qh/R_B)$ (36). If unknown, $Fa = 1$, $Fg = 1$ and $k = 0.1/\text{min}$ can be used as a worst-case estimate. The $f_{u,p}$ should be set to 1% if the reliability of $f_{u,p}$ measurements < 1% cannot be demonstrated (also refer to Section 2.1.2.1).

The recommended ratio and cutoff value for P-gp or BCRP is for orally administered drugs. If the investigational drug is administered parenterally or if it is a metabolite formed post-absorption that inhibits P-gp or BCRP, $IC_{50,u} > 50 \times C_{\max,u}$ (i.e., $C_{\max,u}/IC_{50,u} < 0.02$) can be used.

The cutoff values in Table 1 were determined based on *in vitro*-to-*in vivo* extrapolation analyses mainly using IC_{50} and are based on limited published data. Other cutoff values can be proposed if justified based on *in vitro*-to-*in vivo* extrapolation and a calibration of the specific *in vitro* systems with known inhibitors and non-inhibitors of these transporter systems.

If the above analysis indicates that a drug inhibits a transporter, a clinical study should be considered based on whether the likely concomitant medications used in the indicated patient populations are known substrates of the inhibited transporter and the safety profiles of those substrates. Alternatively, the inhibition potential of a drug can be evaluated using mechanistic static models, PBPK modeling, or endogenous biomarkers. These approaches should be supported by submission of evidence supporting validity of the methods.

2.2.3 Drug as an Inducer of Transporters

Currently, *in vitro* methods to evaluate transporter induction are not well established. If an investigational drug has been observed to be an inducer of CYP enzymes via activation of nuclear receptors such as PXR or CAR, it is likely that transporters regulated through these receptors will be induced, such as P-gp. Refer to Section 3.2.5, which describes conducting clinical DDI studies mediated by transporters, for more considerations.

2.3 DDI Potential of Metabolites

Assessing the DDI liability of an investigational drug's metabolites often starts with *in vitro* experiments and generally uses the same strategies as those for parent drugs. The need to evaluate

the DDI potential of metabolites is based on pharmacologic activity or plasma exposure, as described below.

2.3.1 Metabolite as a Substrate

The risk of DDIs through altered formation or elimination of a metabolite should be investigated if available non-clinical or clinical data indicate that a change in metabolite exposure can result in clinically meaningful alteration of efficacy or safety of a drug (“target” as well as “off-target” effects). The enzymes responsible for formation and elimination of a metabolite should be identified *in vitro* if the metabolite contributes to an *in vivo* target effect to a similar or greater extent than the parent drug. The contribution of the metabolite to efficacy should be estimated by taking into account unbound metabolite and parent drug exposures (e.g., AUC expressed in molar units) in humans, pharmacological potency (e.g., receptor binding affinity, enzyme inhibitory potency), and if available, data related to target tissue distribution. Identification of the enzyme(s) responsible for formation and elimination of the active metabolite should be studied in the same manner as the identification of the enzymes involved in the elimination of the parent drug. If the plasma protein binding of the metabolite is high, the measured $f_{u,p}$ can be used if the accuracy and precision of protein binding measurement has been demonstrated (see Section 7.2 for details). Similarly, if a metabolite is suspected to cause substantial adverse effects based on available nonclinical or clinical information, the enzymes involved in the formation and elimination of that metabolite should be identified. Similar to metabolic phenotyping for parent drugs, the characterization of enzymes involved in metabolite formation and metabolism should also start with major CYP enzymes and can examine other enzymes when appropriate.

The general principles described above can also be applied to characterization of a metabolite as a substrate of major transporters, with consideration of the relevance of transporter-mediated distribution or elimination in the disposition of a metabolite.

Whether a sponsor should conduct a clinical DDI study with an inhibitor or inducer of an enzyme or a transporter depends on the estimated fraction of formation or elimination of a metabolite mediated by an enzyme or transporter, how much the metabolite contributes to the clinical effect, the exposure-response relationship of the metabolite if known, and likely concomitant medications that affect the enzyme or transporter.

2.3.2 Metabolite as an Inhibitor

If *in vitro* assessments suggest that the parent drug alone does not inhibit major CYP enzymes/transporters or is not expected to inhibit enzymes/transporters clinically, DDI liability due to metabolites as inhibitors can still exist. As a pragmatic rule, it is recommended to investigate the CYP enzyme and transporter inhibitory potential of metabolites that have total $AUC_{\text{metabolite}} \geq 25\%$ of AUC_{parent} and also account for at least 10% of drug-related material in circulation (i.e., considered as a major metabolite, often determined based on radioactivity data).

If *in vitro* assessments suggest that the parent drug inhibits major CYP enzymes and transporters and clinical DDI studies are planned, *in vitro* assessments of metabolites as inhibitor of those enzymes or transporters may not be needed. The inhibition potential of metabolites would be implicitly reflected in the clinical DDI study along with the parent drug, unless clinically relevant exposures of the metabolite cannot be adequately represented in the clinical DDI study (i.e., the study duration does not allow the metabolite to accumulate). It is noted that *in vitro* assessments of metabolites can become useful in interpreting the results of DDI studies.

Based on the results of *in vitro* DDI assessments of a metabolite, the determination of whether to conduct a clinical DDI study follows the same approaches as those for the parent drug. Usually, metabolites are irrelevant for the evaluation of intestinal CYP or transporter inhibition except when formed substantially in the gut or intestinal cells. If basic methods suggest that the metabolite(s) could have *in vivo* DDI liability, and a mechanistic static or PBPK model is then used to evaluate the DDI risk of a drug, metabolite(s) should be incorporated in those models.

2.3.3 Metabolite as an Inducer

While metabolites can induce CYP enzymes, the *in vitro* evaluation of the parent drug as a potential inducer could also reflect induction by metabolites because metabolites can be generated during incubation of the parent drug with hepatocytes. However, when the drug is a prodrug or when a metabolite is mainly formed extra-hepatically, *in vitro* evaluation of a metabolite's induction potential on CYP enzymes is recommended if the metabolite is a major metabolite and has $AUC_{\text{metabolite}}/AUC_{\text{parent}} \geq 25\%$ and accounts for at least 10% of drug-related material in circulation. Based on the results of *in vitro* assessments of the metabolite, the determination of whether to conduct a clinical DDI study follows the same approach as for the parent drug.

3. CLINICAL EVALUATION

3.1 Types of Clinical DDI Studies (Terminology)

There are different study types that can be conducted to determine the presence or absence of a clinical DDI and the magnitude of the DDI if one exists. The study types described in this section are not mutually exclusive. The specific goal of a study should be considered when determining the type of study to conduct.

Regulatory decision-making generally relies upon prospective studies specifically designed to evaluate the potential for DDIs. Retrospective evaluation of drug concentrations from studies not designed to evaluate DDIs rarely includes sufficient accuracy and precision to provide an adequate assessment. DDIs identified or ruled out using a retrospective analysis may need to be confirmed using a prospective evaluation.

In some situations, predictive modeling approaches (mechanistic static or PBPK) can be used to translate *in vitro* results to the clinical setting, without a clinical DDI study. The scenarios and best practice considerations are described in Section 7.5.

3.1.1 Stand-alone and Nested DDI Studies

A stand-alone DDI study is a clinical study with the primary objective of determining the presence or absence of a clinical DDI and the magnitude of the DDI. Alternatively, a nested DDI study evaluates DDIs as part of larger studies in patients (e.g., Phase 2/3) for which DDI evaluation is not the primary objective. The nested DDI evaluation is prospectively planned and appropriately designed (refer to Section 3.2.2 for more details).

3.1.2 DDI Studies with Index Precipitants and Index Substrates

Precipitants (inhibitors or inducers) and objects (substrates) with well-understood and predictable pharmacokinetic and DDI properties with regard to level of inhibition, induction, or metabolic pathway are known as “index drugs.” The most common purpose of studies conducted with these drugs is to estimate the greatest magnitude of interaction for the studied pathway. For drugs that are evaluated as objects of a DDI, the greatest magnitude of interaction generally results from concomitant administration of a strong index inhibitor or inducer of the drug’s metabolic pathway(s). For drugs evaluated as precipitants of DDIs, the greatest magnitude of interaction generally results from concomitant administration of the drug with a sensitive index substrate.

A distinctive feature of index studies is that the results usually can be extrapolated to other drug combinations. Thus, after conducting a study with an index inhibitor, one can assume that other inhibitors of equal strength for that metabolic pathway will generally have a similar DDI effect size. Additionally, if one concludes that the change in drug exposure following a concomitant strong index inhibitor is not clinically relevant, the same can be concluded for all other inhibitors for that particular metabolic pathway without additional studies. Results from DDI studies with index precipitants or substrates are also used to help design DDI studies with commonly used concomitant medications in the investigational drug’s target population.

A list of index drugs (either as substrates, inhibitors, or inducers of CYPs) is presented in Section 7.7.1.

Index substrates or precipitants have not been identified for transporters and several metabolic pathways (e.g., CYP2B6, UGTs). The lack of index substrates or precipitants is mainly due to selectivity issues. However, information similar to that provided by studies with index precipitants or substrates (i.e., the likelihood of a DDI due to a specific pathway) is often important for these pathways. Although index substrates and precipitants have not been identified, Sections 7.7.2 and 7.7.3 list examples of drugs that can be useful for DDI studies because they provide informative results and explain the limitations of the drugs. However, extrapolating the results of these studies

can be more difficult and complex than extrapolating results from studies with index drugs.

3.1.3 DDI Studies with Expected Concomitant Drugs

In addition to DDI studies with index drugs, as described above, it can be informative to conduct studies that investigate DDIs between the investigated drug and drugs likely to be administered to the target population. These studies can also be considered when a drug is used as an add-on to other therapies or as part of a fixed dose combination. When choosing drugs to evaluate in these studies, sponsors should consider the mechanistic understanding of the potential for DDIs (based on *in vitro* studies and clinical studies with index drugs) and the relative frequency of co-administration. Another situation where DDI evaluation is often based on the likelihood of co-administration is for transporter-mediated pathways and several metabolic pathways (UGTs; CYP2B6) because of a general lack of index substrates or precipitants for these pathways.

DDI studies with expected concomitant drugs can provide information about specific dose adjustments for the investigational drug and/or concomitant drug or inform the need for monitoring for adverse effects or reduced efficacy. However, although these studies are often informative to patients and medical professionals, the results may be difficult to extrapolate to other drugs.

3.1.4 Cocktail Approach

A cocktail study includes the simultaneous administration of substrates of multiple enzymes and/or transporters to study subjects. A cocktail approach can simultaneously evaluate a drug's inhibition or induction potential for multiple enzymes and transporters if the study is properly designed and conducted (refer to Section 3.2.6 for additional details).

3.1.5 Biomarker Approach

An emerging approach in the evaluation of DDI risk assessment is the deployment of endogenous biomarkers that are substrates for drug metabolism and transport. This approach is enabled by the measurement of endogenous biomarkers in plasma and/or urine before and after administration of the investigational drug. The biomarker-based approach may provide an early indication of DDI potential of a drug as a precipitant via specific pathway by monitoring endogenous biomarkers in clinical studies (refer to Section 3.2.7 for additional details).

3.2 Study Planning and Considerations for Clinical DDI Studies

The objective of most DDI studies is to determine the ratio of a measure of substrate drug exposure (e.g., AUC ratio) in the presence and absence of a precipitant drug. The following considerations are important when designing prospective clinical DDI studies to unambiguously determine this ratio.

3.2.1 Study Design

3.2.1.1 Study Population and Number of Subjects

Most clinical DDI studies can be conducted using healthy subjects, under the assumption that

findings in healthy subjects translate to findings in the intended patient population. However, safety considerations can prevent the use of healthy subjects in studies of certain drugs. For some drugs, use of the intended patient population in DDI studies can allow for evaluation of pharmacodynamic endpoints that cannot be studied in healthy subjects, in addition to pharmacokinetic endpoints.

The number of subjects included in a DDI study should be sufficient to provide a reliable estimate of the magnitude and variability of the interaction (See ICH M12 Questions and Answers Document).

3.2.1.2 Dose

For studies intended to identify the interaction of greatest magnitude, the doses of the precipitant drug used in DDI studies should maximize the possibility of identifying a DDI. Thus, the maximum dose and the shortest dosing interval of the precipitant under the recommended clinical use should generally be evaluated.

If the substrate drug has dose-proportional pharmacokinetics, sponsors can study any dose in the range where exposure to the drug increases in a dose-proportional manner. If the substrate drug does not have dose-proportional pharmacokinetics, the therapeutic dose most likely to demonstrate the largest magnitude DDI should be used. When there are safety concerns, lower doses of the substrate drug, including doses lower than therapeutic doses, can be used.

For studies with anticipated concomitant drugs when a clinically significant DDI is anticipated based on prior *in vitro* or clinical DDI information, it can be informative to build a dose adjustment of the substrate drug into the study to allow identification of doses that can be administered together in clinical practice. In such a scenario, a clinically relevant dose of the precipitant should be used.

3.2.1.3 Single or Multiple Doses

The precipitant drug is often administered in a multiple-dose regimen in DDI studies. However, sponsors can evaluate single-dose administration of a precipitant if the interaction potential is only relevant during absorption (e.g., inhibition of intestinal P-gp or BCRP).

In addition, DDI studies can evaluate single-dose administration of a precipitant if the exposure of precipitant following a single dose is representative of exposure at steady-state and if the precipitant is not a potential inducer or time-dependent inhibitor. The single dose can be the therapeutic dose or a higher dose, depending on accumulation. Safety of a higher dose would need to be understood before conducting the study. When studied with a substrate with a long half-life, it may be necessary to administer a precipitant multiple times to cover the full time-course of the substrate exposure. The duration of the treatment with the precipitant should be long enough to

cover at least 90% of the plasma concentration-time curve of the substrate, considering that its half-life may be longer in the presence of an inhibitor drug. However, if the substrate has a very long terminal half-life that does not allow dosing with the precipitant to cover the full plasma concentration-time curve, population pharmacokinetic analysis or PBPK analysis can be used to bridge the clinical DDI study results to the maximum effect on the exposure of the substrate.

If a metabolite of the precipitant has a long half-life or has demonstrated time-dependent inhibition of the enzyme being evaluated in the DDI study, the duration of the treatment with the parent drug should be sufficient for steady-state of enzyme inhibition due to parent drug and metabolite to be reached.

Inducers should be administered as multiple doses to ensure the maximal induction of a specific pathway. The duration should consider the time to reach steady-state of the inducer, the turnover time of the affected enzyme or transporter, and the half-life of substrate drug. Typical pre-treatment durations range from 7 to 14 days.

When there are multiple mechanisms of interactions for a specific precipitant, single-dose administration can be appropriate in certain situations (e.g., evaluation of rifampin as an inhibitor of OATP1B1), while multiple-dose administration can be appropriate in other situations (e.g., evaluation of rifampin as a CYP3A inducer).

If the substrate demonstrates time-dependent pharmacokinetics (clearance changes over time), multiple-dose administration of the substrate and a precipitant should be evaluated. If the substrate does not demonstrate time-dependent pharmacokinetics, the substrate can be administered as single doses, and the observed magnitude increase in exposure can be extrapolated to steady-state conditions.

3.2.1.4 Route of Administration and Formulations

The route of administration of the investigational drug evaluated in DDI studies should generally be the one planned for routine clinical use. When multiple routes of administration are developed for clinical use, the route of drug administration for DDI studies should be selected based on the expected mechanisms of the DDIs and the similarity of the concentration-time profiles for the parent drug and metabolites after different routes of administration.

Formulation-related differences in DDIs are also reported to occur (9,10). The possibility of formulation differences in interaction potential should be considered when extrapolating interaction results between formulations. In general, DDI potential can be extrapolated between formulations by comparing their rate and extent of absorption.

3.2.1.5 Parallel Versus Crossover Studies

Crossover studies (one-sequence or randomized) are preferred over parallel study designs to reduce variability. Duration of the washout period should be based on the pharmacokinetics of the substrate and the precipitant, the anticipated impact on the substrate's half-life, and the duration necessary for enzyme activity to return to baseline or for potential pharmacodynamic effects to return to pre-treatment levels (if pharmacodynamic effects are also assessed). In some situations, additional periods can be informative. The situations include evaluation of the time it takes for enzyme activity to return to normal following removal of an inducer or time-dependent inhibitor, evaluation of two drugs that may affect each other (each drug alone and in combination), or evaluation of the effects of acute and chronic treatment of a drug.

Parallel, two-arm studies can be appropriate when a crossover study design is not feasible, such as when one of the drugs (or major active metabolite, if applicable) has a long half-life. Typically, parallel-design studies call for larger sample sizes than crossover studies and subjects should be matched for intrinsic factors likely to affect pharmacokinetics.

3.2.1.6 Timing of Drug Administration

In most DDI studies, the precipitant and substrate drugs can be administered at the same time. However, the timing of administration of the precipitant is critical if it is both an inhibitor and an inducer. In such scenarios, the timing of administration of the precipitant and the pharmacokinetic sampling times should consider the objective of the DDI study. For an index study, to ensure the maximum induction effect is identified, staggered administration of the precipitant and substrate is recommended to prevent inhibition from masking induction. The anticipated clinical dosing scenario should be used if the objective of the study is evaluation of concomitant medications.

If a large part of an interaction occurs during absorption or first pass, staggered dosing schedules can be studied (clinical study or PBPK) to understand whether such a method is a viable mitigation strategy for the DDI.

When evaluating the interaction between drugs that require different food conditions for optimal absorption, the timing of drug administration should be adjusted to maximize the potential to detect an interaction (i.e., index studies) and/or to reflect the clinically relevant conditions (i.e., DDI study with potential concomitant drugs).

3.2.1.7 Co-Medications and Other Extrinsic Factors Affecting DDIs

To reduce variability in the magnitude of DDIs, use of the following should be excluded to the extent possible during DDI studies: other medications, dietary/nutritional supplements, herbal supplements, tobacco, smoking, alcohol, foods, and fruit juices that may affect the expression or function of enzymes and transporters. The exclusion should begin for a sufficient time before subjects enter the study and continue for the duration of the study.

3.2.1.8 Sample and Data Collection

Sampling times for pharmacokinetic assessment should be sufficient to characterize the AUC_{0-inf} (for single-dose studies) or the AUC_{0-tau} (for multiple-dose studies) and C_{max} of the substrate drug administered alone and under conditions of the anticipated interaction. Data on additional pharmacokinetic parameters should be collected based on the pharmacokinetic or pharmacological relevance for the proposed indication (e.g., the minimum concentration (C_{min}), partial AUC). The sampling times for single-dose studies should be planned so that the mean difference between the AUC_{0-t} and the AUC_{0-inf} is less than 20%, accounting for potential longer elimination half-life due to the DDI. Samples collected should contain the moieties needed to interpret study results; in most cases, the moiety needed to interpret results will be the parent drug. Metabolite concentrations should be determined if they provide information about the effect of a DDI on safety or efficacy or if the data inform the mechanism of the drug interaction. For example, when a clinical DDI study evaluates drugs that may interact via multiple pathways, measuring metabolites may help determine the enzymes and/or transporters responsible for the interaction. In addition to collecting samples for pharmacokinetic assessment of the object drug, optional collection of a few sparse samples for the inhibitor or inducer can help ensure the plasma concentrations of the inhibitor or inducer are within the expected range. In addition, urine samples can be collected for understanding DDIs involving renal transporters,

When *in vitro* data provide a plausible DDI mechanism that cannot be evaluated with systemic drug exposure, collection and analysis of pharmacodynamic data can be informative. One possible scenario where this could occur is when transporter inhibition alters access of the drug to specific organs or tissues. In such scenarios, clinical consequences such as altered efficacy or increased toxicity resulting from altered tissue distribution of a substrate drug, can be measured as pharmacodynamic endpoints, and *in vitro* evidence of a drug's interaction potential can support data interpretation.

3.2.2 Specific Considerations for Nested DDI Studies

Nested DDI studies are clinical DDI investigations that are part of other studies (e.g., phase 2/3) in which the assessment of DDI is not the primary objective. However, these studies are designed prospectively to investigate DDIs as an exploratory or secondary objective. Nested DDI studies are usually used to evaluate the drug as an object of interactions with concomitant drugs and sometimes can also be used to assess the drug as a precipitant. The results of such analyses can be informative, and sometimes conclusive, when the clinical studies are adequately designed to detect significant changes in drug exposure due to DDIs. An advantage of nested DDI studies is the fact that they are conducted in a patient population and may more closely represent the anticipated clinical setting. However, nested DDI studies can also be challenging because they call for careful attention to study design and data collection. In some cases, PBPK modeling can assist the design of nested DDI studies (refer to Section 7.5.2).

A nested DDI study can evaluate the effect of concomitant drugs that are used for the full duration of the clinical trial or those that are added in response to the patient's condition during the trial. Concomitant drugs to be evaluated should be prespecified. The drugs are typically selected because there is a mechanistic reason to anticipate an interaction. Relevance in the patient population is also a consideration. The study design can specify individual drugs or a grouping, based on mechanism (e.g., strong CYP3A inhibitors). However, if a grouping is evaluated it is important to consider the potential for differences in the effect of different drugs in the group and the effect of the potential variability on data analysis and translation of the findings (11).

Simulations can be used to determine the appropriate number of pharmacokinetic samples and to assist in the selection of sampling times. A power analysis can also be performed to estimate the minimum effect size that is likely to be detected with acceptable precision in a study using a given number of patients on a concomitant drug.

Collection of the following data is critical to ensure interpretable results: timing of drug administration (investigational drug and concomitant drug), drug dose, timing relative to food (when relevant), other concomitant drugs, and pharmacokinetic sampling date and time (actual, not scheduled). It is also important to document the start date and discontinuation date of the concomitant drug relative to when an interaction will be observed, particularly when the concomitant drug is an inducer or time-dependent inhibitor.

A nested DDI study is typically evaluated using population pharmacokinetic analysis, which should be performed according to well-established scientific practice using a model that is robust and fit-for-purpose. The sample collection plan for population pharmacokinetic analysis for the DDI assessment should be established prior to conducting the study. In general, the standard analysis approach is a binary evaluation that includes the concomitant drug as a categorical covariate. Sponsors should consider whether their selected analysis methods will provide the desired level of precision in DDI evaluation. Regardless of analysis method, all assumptions should be stated.

In some instances, unplanned analyses of potential DDIs in phase 2/3 trials are conducted to explain clinical study results, such as safety or efficacy issues in a group of patients, or to screen for potential DDIs not anticipated at the time the trials were designed. If the data collected meet the criteria described in this section, except for preplanned DDI assessment, it can be possible to draw conclusions about the presence or absence of an interaction. In situations where the data do not permit an accurate assessment of a DDI, a further assessment of the DDI potential should be conducted.

3.2.3 Considerations for CYP-Mediated Interactions

3.2.3.1 The Investigational Drug as a Substrate for CYP Enzymes

When evaluating the investigational drug as a substrate, the first clinical DDI studies should, in general, determine the effects of a strong index inhibitor and a strong index inducer on the investigational drug. Moderate index inhibitors or inducers can be used if strong index inhibitors or inducers are not available for a particular enzyme. Some of these inhibitors and inducers can also affect other metabolism and/or transporter pathways; thus, when selecting index inhibitors and inducers for prospective DDI studies, all metabolic and transport pathways of the investigational drug should be considered. Studies with other strong inhibitors and inducers of CYP enzymes can also be appropriate, considering the criteria listed in Section 7.7.1. If the investigational drug is a substrate for multiple enzymes and/or transporters, measuring metabolites can, in some cases, help interpret study results and interaction mechanisms.

If a DDI study with a strong index inhibitor or inducer indicates no DDI is present, additional clinical studies with other inhibitors or inducers of the same enzyme are not needed. However, because a negative DDI study may reveal that the enzyme proposed to be the major metabolizing enzyme based on *in vitro* data is not contributing to the elimination of the drug, this may instead indicate that further investigations of alternative pathways should be considered.

If a DDI study with strong index inhibitors or inducers indicates that there is a clinically relevant interaction, evaluating the impact of moderate inhibitors or inducers can be useful to gain a full understanding of the investigational drug's DDI potential. The evaluated moderate inhibitors and inducers may be anticipated concomitant medications in the intended patient population. The effect of the additional inhibitors and inducers can be evaluated in a clinical interaction study, or in some cases, modeling approaches can provide additional information (refer to Section 7.5). If it is anticipated that co-administration with strong inducers or inhibitors should be avoided, a DDI study with a moderate inducer or inhibitor may be preferable as the initial study.

If the investigational drug is subject to significant metabolism by a genetically polymorphic enzyme for which a well-defined poor metabolizer phenotype exists that results in non-functional enzyme activity, a comparison of the pharmacokinetic parameters of the drug in individuals with the poor metabolizer phenotype versus those with a normal metabolizer phenotype can substitute for an interaction study for that particular pathway (refer to Section 4.1).

3.2.3.2 The Investigational Drug as an Inhibitor or an Inducer of CYP Enzymes

When studying an investigational drug as a potential inhibitor or inducer of a CYP enzyme, the sensitive index substrate selected for the initial clinical studies should be sensitive to changes in activity or amount of the CYP enzyme being evaluated (refer to Section 7.7.1). Because some substrates are not specific for one CYP enzyme and sometimes are also substrates of transporters, the most appropriate substrate should be selected considering the inhibitor/inducer characteristics of the investigational drug, based on available *in vitro* and clinical data. Other CYP enzyme

substrates can also be appropriate. If the substrate drug is metabolized by more than one enzyme, measuring metabolites sometimes can help with interpretation of study results.

If the initial study with a sensitive index substrate is negative, studies with less sensitive substrates of the enzyme are not needed. If an initial study determines that an investigational drug either inhibits or induces the metabolism of sensitive index substrates, further evaluations using other substrates (e.g., relevant co-medications) can be useful. The magnitude of the effect of the investigational drug on the sensitive index substrate and the potential for concomitant use with other drugs that are substrates of the same enzyme should be considered.

If the investigational drug is both an inducer and an inhibitor of an enzyme, the net effect of the drug on enzyme function may be time dependent. The timing of pharmacokinetic endpoints should permit an understanding of the changes in effects over time, when relevant. To achieve this understanding, the pharmacokinetics of the substrate drug should be evaluated at early and late time points during the administration of the investigational drug in the test period. The observed effect of reversible inhibition may be more pronounced in the beginning of the treatment and the induction may be most pronounced after ending the treatment.

3.2.4 Considerations for Evaluation of UGT-Mediated Interactions

3.2.4.1 Investigational Drug as a Substrate of UGTs

Based on limited literature evidence, the magnitude of DDIs mediated through inhibition of UGTs (reflected by AUC ratio of a substrate in the presence of an inhibitor compared to no inhibitor) is typically less than the magnitude observed with CYP inhibition (3). For an investigational drug that is mainly eliminated by direct glucuronidation, clinical DDI studies with UGT inhibitors should be conducted on a case-by-case basis, considering the safety profile of the drug and the likelihood of its concomitant use with inhibitors of that UGT isoform (refer to Table 16, Section 7.7.2 for some examples of UGT inhibitors). Some UGT substrates are also substrates of other enzymes or transporters, and the interaction with a UGT inhibitor may involve other mechanisms when the UGT inhibitor also affects those enzymes or transporters. Thus, it may be valuable to also measure the glucuronide conjugate concentrations in addition to the UGT substrate itself. The change of glucuronide metabolite relative to the parent drug may provide insight into the underlying mechanism of interaction. In addition, some glucuronide metabolites are active or reactive and may significantly contribute to efficacy or safety of a drug. In such cases, the concentrations of glucuronide conjugates should be measured in addition to parent drug concentrations.

Genetic variation in certain UGT enzymes (for example, UGT1A1, UGT2B7, UGT2B10, UGT2B15, and UGT2B17) has been reported to contribute to variation in the pharmacokinetics of drugs metabolized by UGTs. In certain cases, comparative pharmacokinetic data in subjects with

various UGT genotypes can be used to identify the importance of the UGT pathway(s) in the elimination of a drug and to estimate the extent of DDI with inhibitors of UGT.

In addition, UGTs can be induced, for example, by certain PXR agonists (e.g., moderate or strong CYP3A inducers). The impact of inducers on an investigational drug that is mainly metabolized by UGTs should also be considered and evaluated depending on the likelihood of its concomitant use with UGT inducers and the dose/exposure-efficacy relationship of the investigational drug.

3.2.4.2 Investigational Drug as an Inhibitor of UGTs

As indicated in Section 2.1.3, considering the generally limited magnitude of UGT inhibition-mediated DDIs, a routine evaluation of UGT inhibition by an investigational drug may not be warranted. Following the *in vitro* assessment described in Section 2.1.3, a decision on whether to perform a clinical DDI study to evaluate the effect of a drug as a UGT inhibitor should also take into consideration the likelihood of the drug's concomitant use with known substrates of the UGT isoform (refer to Table 15, Section 7.7.2 for examples) and the safety profiles of those substrates.

3.2.4.3 Investigational Drug as an Inducer of UGTs

There is limited understanding about gene expression of UGTs. However, limited clinical DDI studies indicate certain UGTs may be induced by agonists of PXR and/or CAR, which also regulate CYP3A4 expression. UGTs are less inducible than CYP3A4. Thus, for a drug found to induce CYP3A4 *in vitro* and further evaluated with a clinical DDI study, the effect of the drug on CYP3A4 substrates may inform its potential induction effect on UGTs. In such a situation, if a drug reduces the AUC of a sensitive substrate of CYP3A by $\geq 50\%$, a clinical DDI study should be considered taking into account the following: the magnitude of exposure change of the CYP3A substrate, the likelihood of concomitant use of the investigational drug with UGT substrates, whether there are other enzymes/transporters involved in the pharmacokinetics of UGT substrates which can also be regulated by PXR/CAR agonists, and the dose or exposure-efficacy relationship of those UGT substrates. It is noted that some CYP3A4 inducers have their induction effect overridden by their inhibition effect on CYP3A. Thus, while those drugs inhibit CYP3A4 in clinical studies, they may exhibit induction effects on UGTs.

3.2.5 Considerations for Evaluation of Transporter-Mediated Interactions

3.2.5.1 Investigational Drug as a Substrate of Transporters

If *in vitro* studies indicate that the investigational drug is a transporter substrate, sponsors should determine whether to conduct clinical DDI studies based on the drug's passive permeability, route of administration, absorption and elimination, putative site of action, safety profile, dose or exposure-response (efficacy and safety) relationship, and likely concomitant drugs that are known inhibitors or inducers of the transporters. The information in Table 2 helps guide when a clinical DDI study should be considered for investigational drugs that are transporter substrates *in vitro*:

Table 2: Considerations for clinical evaluation of drug as substrate of transporters

Transporters	When a clinical DDI study should be considered
P-gp and BCRP	When intestinal absorption is limited, or biliary excretion/active renal secretion is a major elimination pathway.
OATP1B1 and OATP1B3	When hepatic (metabolic/biliary) elimination is a significant clearance pathway ($\geq 25\%$) for the investigational drug or the action site of the drug is in liver, and the drug's properties support the importance of active uptake of the drug into the liver.
OAT1 and OAT3, OCT2, MATE1, and MATE2-K	When the investigational drug undergoes significant active renal secretion (i.e., accounting for $\geq 25\%$ of systemic clearance)

When evaluating an investigational drug as a substrate in transporter-mediated DDIs, the selected precipitant drug should be a known inhibitor of the transporter under investigation. Because of a general lack of index precipitants for transporter-mediated pathways, the choice of transporter precipitant is generally based on the likelihood of concomitant use (e.g., to obtain clinically relevant DDI information that can inform labeling regarding the management of a DDI). Some examples are provided in Section 7.7.3.2, Table 19.

DDI studies with transporter inhibitors can be used to understand the underlying mechanisms of DDIs or to determine the anticipated largest magnitude DDI. If *in vitro* studies indicate a drug is a substrate of multiple transporters, a clinical study can be conducted with a broad inhibitor of multiple transporters to determine the anticipated largest magnitude DDI. For example, cyclosporine, which inhibits intestinal P-gp and BCRP and hepatic OATPs, can be used as the inhibitor in a DDI study. Negative results from this kind of study may rule out the need to further evaluate the drug as a substrate for any of the individual transporters. If the study result is positive, additional studies with more selective inhibitors of specific transporter pathways can be conducted to determine the impact of inhibition of each transporter on the disposition of the substrate drug. The same paradigm can apply to an investigational drug that is a substrate for both transporters and metabolic enzymes (e.g., CYP3A and P-gp).

If the goal of the study is to determine the role of a specific pathway in the pharmacokinetics of a substrate drug and resulting DDIs due to that pathway, then a more selective inhibitor should be used. Use of these inhibitors in clinical studies can provide a mechanistic understanding of transporter-mediated DDIs. Some transporters, including OATP1B1 and BCRP, are encoded by genetically polymorphic genes (SLCO1B1 and ABCG2, respectively) for which phenotypes with reduced functionality exist. Similar to drugs that are substrates of CYPs encoded by polymorphic genes, the relative contribution of a specific transporter to the disposition of the investigational drug can be evaluated in subjects with different transporter genotypes, if a non-functional phenotype exists (refer to Section 4.1).

Examples of transporter inhibitors are listed in Section 7.7.3.2. Many of them not only inhibit the specified transporters but also can inhibit other transporters and/or CYP enzymes. Thus, extrapolation of results from transporter inhibition studies to other drugs can be challenging. Interpretation of the study results should consider the knowledge of transport and metabolic pathways for the investigational drug.

3.2.5.2 Investigational Drug as an Inhibitor of Transporters

If *in vitro* studies indicate that the investigational drug is a transporter inhibitor, the determination of whether to conduct a clinical DDI study should be based on likely concomitant drugs and safety considerations. When studying the investigational drug's potential to act as an inhibitor drug for a transporter, a substrate drug whose pharmacokinetic profile is markedly altered by coadministration of known inhibitors of that transporter and is also a likely concomitant drug is preferred. Some examples of transporter substrates that can be used in DDI studies are listed in Section 7.7.3.1. Because many drugs are substrates of multiple transporters and/or enzymes, the observed clinical interactions can be a result of the modulation of multiple pathways if the investigational drug is also an inhibitor or inducer for those pathways. Extrapolation of results from these studies to other drugs can thus be challenging. The choice of substrates can be determined by the therapeutic area of the investigational drug and the likely concomitant drugs that are known substrates of the transporters.

In some cases, an alteration in drug transport may not be fully reflected by changes in plasma concentrations of parent drug alone. Therefore, measurement of metabolite or pharmacodynamic markers to reflect altered distribution to the organs expressing the transporter may be considered to help interpret the potential for an interaction.

Recent literature reports indicate potential utility of endogenous substrates for some drug transporters (See Section 3.2.7.1). Evaluating the change in exposure of the endogenous substrate when the investigational drug is administered may provide information regarding the drug's potential as a transporter inhibitor.

3.2.5.3 Investigational Drug as an Inducer of Transporters

Since P-gp is co-regulated with CYP3A, for example by agonists of PXR and/or CAR, but is less inducible than CYP3A (12,13), if an investigational drug reduces the AUC of a sensitive substrate of CYP3A by 50% or more (i.e., being a moderate or strong inducer), a further clinical study to evaluate potential induction effect of the drug on P-gp substrates should be considered. This case-by-case consideration should take into account the following factors: the magnitude of CYP3A substrate AUC change by the investigational drug, the likelihood of concomitant use of the drug with P-gp substrates, whether there are other enzymes/transporters involved in the pharmacokinetics of P-gp substrates which can also be regulated by PXR and/or CAR agonists, and the dose or exposure-efficacy relationship of P-gp substrates. It is noted that some CYP3A4

inducers have their induction effect overridden by their inhibition effect on CYP3A. Thus, while those drugs inhibit CYP3A4 in clinical studies, they may exhibit induction effects on P-gp. Sponsors should also consider whether to conduct clinical DDI studies to evaluate the potential effect of a drug on other transporters regulated through the same pathways as CYP3A.

3.2.6 Cocktail Studies-Considerations for CYP or Transporter Cocktail Studies

A cocktail approach can simultaneously evaluate a drug's inhibition or induction potential for multiple CYPs and transporters if the study is properly designed. Results from a well-conducted cocktail study that includes all elements of a prospective DDI study can be interpreted the same way as results from any other well-conducted DDI study (See 3.2.1.1. through 3.2.1.8). The criteria for the selection of cocktail drugs are: (a) the substrates are specific for individual CYP enzymes or transporters; and (b) there are no interactions among the substrates. If these criteria are not met, the lack of specificity or the interaction among substrates should be understood and incorporated into the interpretation of the study results. It should be noted that findings obtained with a microdose of a substrate cannot always be extrapolated to a therapeutic dose of that substrate.

3.2.7 Considerations for Biomarker Approach

An alternative approach to assess an investigational drug's potential as a precipitant is evaluating the change in exposure of a well-characterized endogenous substrate. Sufficient analytical validation should be conducted to ensure an adequate level of quality and consistency to allow for reliable interpretation of the results. Reported examples of biomarkers include, but are not limited to, plasma coproporphyrin I (hepatic OATP1B1/3), plasma and urine N¹-methylnicotinamide and N¹-methyladenosine (renal OCT2, MATE1, MATE2K), plasma pyridoxic acid (renal OAT1/3), and plasma 4β-hydroxycholesterol/cholesterol ratio and urine 6β-hydroxycortisol/cortisol ratio for CYP3A (14,15,16,17). It should be noted that not all endogenous biomarkers are validated and characterized in terms of their performance characteristics such as sensitivity, selectivity, specificity, dynamic range, correlation with pharmacokinetic parameters of the probe drugs, and variability (due to factors such as diet, age, exercise, diurnal variation, and disease state) (18,19). Availability of such data can enable sponsor-regulator dialogue regarding the prioritization, need for, and design of endogenous biomarker-based DDI evaluation.

3.2.7.1 Investigational Drug as an Inhibitor of Hepatic OATP1B

As an example, recent literature reports support the utility of plasma coproporphyrin I (CPI) for evaluation of hepatic OATP1B inhibition potential. Monitoring of plasma CPI can be incorporated in early human healthy volunteer pharmacokinetic studies, such as phase 1 single or multiple dose escalation studies. Plasma CPI measured prior to the administration of the investigational drug (pre-dose) represents baseline concentration and baseline AUC_{0-t} (baseline AUC_{0-t} = baseline CPI x t). Serial samples for CPI following the administration of the investigational drug will allow the characterization of C_{max} and AUC of CPI. The metrics of interest are the ratio of CPI C_{max} and

AUC_{0-t} after administration of an investigational drug to a baseline assessment. If the ratio is less than 1.25, this indicates a low likelihood of a clinical DDI via OATP1B inhibition (20).

4. OTHER TOPICS

4.1 Pharmacogenetics

Pharmacogenetic variations in genes encoding drug metabolizing enzymes or drug transporters can affect the pharmacokinetics of a drug, increase interindividual variability in drug exposure, affect safety or efficacy, and alter the magnitude of DDIs. Important pharmacogenes include those that encode phase 1 (e.g., CYP2C9, CYP2C19, CYP2D6) and phase 2 (e.g., NAT2, UGT1A1) drug metabolizing enzymes as well as genes that encode drug transporters (e.g., BCRP, OATP1B1). Polymorphisms in metabolizing enzymes can lead to increased, normal, decreased, or absent enzyme activity resulting in ultra-rapid (UM), normal or extensive (NM or EM, hereafter referred to as NM), intermediate (IM), and poor (PM) metabolizers, respectively. Polymorphisms in drug transporters can increase or decrease transport of a drug across membranes. These drug metabolizing enzyme and transporter polymorphisms can affect the systemic or tissue concentrations of a drug and/or its metabolite(s).

The scope of this section is limited to the evaluation of the impact of pharmacogenetics on DDIs and on DDI evaluation. While the considerations described below use metabolizing enzymes as examples, the concept can also be applicable to transporters with polymorphisms.

If an investigational drug is a substrate/inhibitor for a polymorphic enzyme and a DDI study with an index inhibitor/substrate is conducted to evaluate pharmacokinetic changes, it is recommended to prospectively characterize the subjects' genotypes. Exclusion of PMs is recommended, to allow characterization of the greatest magnitude of interaction. If PMs are not excluded, the DDI effect should be evaluated separately in subjects with different phenotypes (e.g., PM, IM, and NM), as relevant.

If an investigational drug is subject to metabolism by an enzyme with a well-defined PM phenotype (for example, CYP2D6, CYP2C19), exposure in PM is expected to be similar to the effect of a strong inhibitor of that pathway. A comparison of the pharmacokinetic parameters of the drug in individuals with the PM phenotype with those with a NM phenotype can substitute for a DDI study of that pathway with a strong inhibitor. Similarly, the exposures in subjects with a polymorphic PM phenotype could be estimated using the results of a clinical DDI study with a strong inhibitor. If there is a significant difference in exposure between individuals with the PM and NM phenotypes, further studies to evaluate the DDI potential with moderate inhibitors or inducers of the specific enzyme should be considered.

When an enzyme encoded by a polymorphic gene is one of two major elimination routes of an investigational drug, the interaction effects of inhibiting the other enzymes is expected to vary in

different phenotypes of the polymorphic enzyme. In a DDI study evaluating the impact of inhibitors of the other enzyme, prospective genotyping and enrichment of subjects with absent or decreased function of the polymorphic gene besides NM subjects can help assess the interaction effects in the various phenotypes. Because the DDI magnitude may become large in PMs or IMs of the polymorphic enzyme when combined with an inhibitor of a parallel pathway, depending on the safety profile of the drug, different doses should be considered in those subjects. PBPK modeling can be useful to supplement such studies or to extrapolate the interaction effects in different genotypes (refer to Section 7.5.2).

A retrospective pharmacogenetic analysis can help elucidate reasons for a high variability in a DDI study. When study enrollment is not based on the genotype of a polymorphic metabolizing enzyme or transporter, a retrospective analysis of the metabolizing enzyme or transporter of interest can help to characterize differences in the magnitude of the DDI across genotype groups and explain why some subjects have unanticipated increases or decreases in drug concentrations.

Guidance on DNA sample collection for prospective and retrospective pharmacogenetic analysis can be found elsewhere. The frequency of certain pharmacogenetic variations can vary across populations. Therefore, when performing pharmacogenetic analysis, an individual's race/ethnicity should be considered. In addition, regional regulations on sampling and analyzing human derived materials need to be followed.

4.2 Therapeutic Protein DDIs

In general, the risk of pharmacokinetic DDIs is lower for therapeutic proteins compared to small molecules. The *in vitro* assays that are applicable for small molecules are generally not applicable to proteins.

When evaluating the potential for a DDI between therapeutic proteins and small molecules or between multiple therapeutic proteins, the mechanisms of a potential DDI should be considered, taking into account the pharmacology and clearance of the therapeutic proteins as well as any co-administered medications in the patient population.

DDI risk for emerging modalities such as oligonucleotides, small interfering ribonucleic acids, modified ribose nucleic acids, and peptides are out of scope for the M12 guideline.

4.2.1 Proinflammatory Cytokine-Related Mechanism

Certain therapeutic proteins may exert an indirect effect on expression of CYP enzymes and thus affect the pharmacokinetics of small molecules. Therapeutic proteins that are proinflammatory cytokines (e.g., peginterferon) or that can increase cytokine levels can down-regulate the expression of CYP enzymes, thereby decreasing the metabolism of drugs that are CYP substrates and increasing their exposure levels. The increase in cytokine levels as a result of drug treatment

can be transient or persistent; sponsors should consider this increase when determining whether to conduct a DDI study as well as the design of that study.

Conversely, therapeutic proteins that reduce the elevated cytokine levels (e.g., inhibitors of tumor necrosis factor) can relieve the CYP down-regulation from an inflammatory environment (e.g., rheumatoid arthritis), thereby increasing CYP expression and activity and reducing exposure for CYP substrates.

If the investigational drug is a cytokine or a cytokine modifier, sponsors should consider whether to perform a clinical DDI study to evaluate the effects of the investigational therapeutic protein on sensitive substrates for CYP enzymes. Known drug effects on metabolism in disease states with similar or higher inflammatory burden, differences in exposure levels of sensitive CYP substrates in healthy subjects versus patients in the indicated population, and the magnitude of the drug effect on cytokine levels should be considered when determining whether to conduct a clinical study. In some cases, a DDI study in the relevant indicated population should be conducted to further inform instructions for use of the drug. Important design aspects include the disease type and severity in the included patients and the dose and treatment time of the precipitant drug.

4.2.2 Antibody-Drug Conjugates

For antibody-drug conjugates (ADCs), the small molecule drug component conjugated to the antibody component can be released in unconjugated form. Therefore, the DDI potential of both the antibody and the small molecule drug component should be considered. In general, for the small molecule component, the potential to inhibit or induce enzymes and transporters should be addressed in a similar manner as described for small molecules in this guideline. In many cases, however, the systemic concentration of small molecule drug component might be too low to act as a precipitant clinically.

It is important to understand the formation, distribution, and elimination kinetics of the small molecule and to assess the systemic exposure of the small molecule drug component of the ADC. It might be necessary to evaluate the small molecule component (administered as an ADC) as a substrate drug, in particular if increased levels of free drug may be associated with safety concerns. Understanding the exposure-response relationship of the components of the ADC is important in determining whether to conduct DDI studies and their significance.

5. REPORTING AND INTERPRETING CLINICAL DDI STUDY RESULTS

A DDI study report should include and justify the study design and data analysis method based on what is known about the mechanism of the DDI and the pharmacokinetic properties of the precipitant and substrate drugs. Data analysis of pharmacokinetic parameters (and pharmacodynamic parameters, when relevant) should include all subjects enrolled in the study who have evaluable pharmacokinetic (and/or pharmacodynamic) data. If a subject is dropped from

the study or has incomplete plasma concentration sampling during a treatment period, the possibility that the observation is due to an interaction should be considered. When indicated, the interaction effect should be presented with and without the individuals proposed for exclusion and a short description of each dropout should be provided.

5.1 Pharmacokinetic Data Analysis

5.1.1 Non-Compartmental Analysis (NCA)

The following exposure measures should be determined for each subject: AUC_{0-inf} , AUC_{0-t} , the percent extrapolated from AUC_{0-t} to AUC_{0-inf} , C_{max} , and time to C_{max} (T_{max}). For multiple-dose studies, C_{max} , C_{min} , AUC_{TAU} at steady-state should also be reported. Additional parameters can help to interpret the pharmacokinetic results: clearance (CL or CL/F), half-life, and volume of distribution. Parameters for metabolites, when measured, should also be presented. NCA can be used to evaluate stand-alone DDI studies conducted to evaluate the investigational drug as an object or precipitant.

5.1.2 Population PK Analysis

Pharmacokinetic data collected in nested DDI studies should typically be evaluated using population pharmacokinetic methods. DDIs should be evaluated using all plausible structural elements of the pharmacokinetic model (e.g., clearance (CL or CL/F), relative bioavailability, rate of absorption). Population pharmacokinetic analyses should derive pharmacokinetic parameters appropriate for the study design and pharmacokinetic properties of the drug, such as AUC and C_{max} . For multiple-dose studies, C_{max} , C_{min} and AUC_{0-TAU} at steady-state should be reported.

5.2 Reporting DDI Results

Typical pharmacokinetic endpoints for DDI studies should include changes in drug exposure parameters for the substrate drug, such as AUC, C_{max} , and, when applicable, C_{min} (see section 5.1.1). Pharmacokinetic results of DDI studies should be reported as the geometric mean ratio of the observed pharmacokinetic exposure measures with and without the precipitant drug and the associated 90 percent confidence interval. Measures of the observed variability of the interaction, such as the range of AUC or C_{max} ratios for individuals in a cross-over study, should be reported.

There are multiple methods of displaying the data, and sponsors are encouraged to select the most appropriate methods based on the data and the situation. Data can be displayed graphically, for example, by using forest plots. A comparison of the individual pharmacokinetic parameters with and without concomitant medication can also be presented graphically (e.g., as spaghetti plots or individual ratio plots). The proportion of individuals for whom exposure extends beyond the no-effect boundary (see section 5.3.1) can also be presented.

If pharmacodynamic endpoints are also assessed in the DDI study, the results should be reported and summarized.

5.3 Interpreting DDI Study Results

5.3.1 Determination of No-Effect Boundaries

The results of a DDI study should be interpreted based on the no-effect boundaries for the substrate drug. No effect-boundaries represent the interval within which a change in systemic exposure measure is considered not significant enough to warrant clinical action (e.g., avoiding coadministration, dose or schedule adjustment, or additional therapeutic monitoring).

It is preferable for no-effect boundaries to be developed based on exposure-response relationships derived from clinical trial results, as well as other relevant information for the substrate drug (e.g., safety data and the maximum-tolerated dose). A good understanding of exposure-response relationships for desirable and undesirable drug effects, as well as knowledge of the variability of exposures in the indicated population, facilitates data interpretation. In the absence of a defined exposure-response relationship, the totality of evidence should be considered when determining the clinical impact of a DDI. Sometimes, a 90% confidence interval of 80%-125% is proposed as a default no-effect boundary. This is in general an acceptable approach but is considered overly conservative for most drugs, where small changes in exposure are unlikely to be of clinical consequence.

In general, the point estimate for the ratio between the exposure of the substrate with and without the precipitant can be used to describe the magnitude of the interaction and to determine whether interventions such as dose adjustments should be considered. Sponsors should also consider the variability of the interaction. The number of subjects included in the study should be sufficient to provide a reliable estimate of the magnitude and variability of the interaction (See ICH M12 Questions and Answers Document).

5.3.2 Investigational Drug as a Precipitant of DDIs: Classification System

The classification system assists in the extrapolation of DDI study results to drugs that have not been evaluated in a clinical DDI study.

If an investigational drug is a CYP inhibitor, it can be classified as a strong, moderate, or weak inhibitor based on its effect on a sensitive index CYP substrate. The convention is to categorize CYP inhibition in the following way:

- A strong inhibitor increases the AUC of a sensitive index CYP substrate ≥ 5 -fold.
- A moderate inhibitor increases the AUC of a sensitive index CYP substrate ≥ 2 - to < 5 -fold.
- A weak inhibitor increases the AUC of a sensitive index CYP substrate ≥ 1.25 - to < 2 -fold.

If an investigational drug is a CYP inducer, it can be classified as a strong, moderate, or weak inducer based on its effect on a sensitive index CYP substrate. The convention is to categorize CYP induction in the following way:

- A strong inducer decreases the AUC of a sensitive index CYP substrate by $\geq 80\%$.
- A moderate inducer decreases the AUC of a sensitive index CYP substrate by ≥ 50 to $< 80\%$.
- A weak inducer decreases the AUC of a sensitive index CYP substrate by ≥ 20 to $< 50\%$.

These categories generally describe the effect of the investigational drug when given at the highest clinical dose and the shortest dosing interval within its therapeutic dose range/dosing regimen. It is noted that the effects of some inhibitors or inducers are dose dependent.

Although CYP inhibitor and inducer classifications are typically based on DDI studies with sensitive index substrates, if the metabolic properties of a different sensitive substrate are well understood, it can be possible to classify the investigational drug based on a study with the alternative substrate.

Currently, there are no classification systems for transporters or non-CYP enzymes. The interacting mechanisms may involve other transporters and/or enzymes, making it challenging to classify inhibitors using the same criteria as those for CYP enzymes. In addition, the magnitude of DDIs mediated by many of the transporters or non-CYP enzymes (e.g., UGTs) has a more limited range.

5.3.3 Extrapolating Study Results

Clinical evaluation of all possible combinations of drugs is not feasible. When possible, results from DDI studies should be extrapolated to other drugs and clinical situations. Results from DDI studies with index drugs generally represent the largest magnitude interaction by a specific mechanism and can be used to predict the magnitude of other interactions by the same mechanism. The classification system for CYP inhibitors and inducers assists the extrapolation. For example, if there is no effect on the exposure of an investigational drug when co-administered with a strong CYP3A index inhibitor, then one can generally assume that there is no effect when other strong, moderate, or weak CYP3A4 inhibitors are co-administered with the investigational drug. If administration of a strong CYP2D6 index inhibitor results in a significant increase in exposure of the investigational drug, these results can be directly extrapolated to other strong CYP2D6 inhibitors. In some cases, extrapolation of positive findings to moderate and weak inhibitors can be possible using mechanistic modeling (See Section 7.5).

Because of the lack of specific transporter substrates and inhibitors and the possible interplay with metabolism, it is generally challenging to extrapolate results from DDI studies evaluating transporter-mediated DDIs or transporter-metabolism interactions from one drug to other drugs. However, if the ADME properties of the investigational drug and potential concomitant drugs are well understood, it is possible to extrapolate to transporter-mediated interactions with other concomitant drugs.

5.3.3.1 Extrapolating Complex Scenarios

Most DDI studies evaluate the interaction between two drugs and consider the effect on single transporters or enzymes. However, DDIs for a specific drug may result from a combination of mechanisms, patients may receive more than two potentially interacting drugs, and the magnitude of an interaction may be different in different populations. Some of the resulting “complex DDI scenarios” are listed below:

- Concurrent inhibition of an enzyme and a transporter by a drug or drugs;
- Concurrent inhibition of multiple transporters by a drug or drugs;
- Concurrent inhibition and induction of a drug’s metabolic pathways, involving one or more enzymes;
- Increased inhibition of drug elimination by use of inhibitors of more than one enzyme that metabolizes the drug;
- Inhibition of an enzyme other than the genetic polymorphic enzyme in poor metabolizers taking a substrate that is metabolized by both enzymes;
- Effect of enzyme/transporter inhibitors in subjects with varying degrees of impairment of drug eliminating organs (e.g., liver or kidney);
- The two drugs affect one another’s pharmacokinetics (both act as precipitant and object).

When there are multiple factors that affect the absorption and disposition of an investigational drug as well as multiple mechanisms of DDIs, sponsors should consider evaluating the effect of the combination of mechanisms and/or individual factors on drug exposure to facilitate risk assessment and provide recommendations. The complex scenarios can be evaluated by integrating knowledge from the relevant *in vitro* and clinical studies as well as endogenous biomarker data. Predictive modeling can sometimes be used to determine whether a clinical study would be informative or to inform the design of clinical studies.

6. RISK ASSESSMENT AND MANAGEMENT

Risk assessment should inform the use of DDI management strategies (i.e., DDI prevention and risk minimization strategies). A DDI is clinically relevant if concomitant use of the drugs leads to safety, effectiveness, or tolerability concerns greater than those present when the drugs are administered alone.

In general, DDI management strategies should result in drug concentrations of the substrate drug falling within the no-effect boundaries. The risk assessment and development of risk minimization strategies should consider the following factors:

- The exposure-response relationships for safety and efficacy;

- The variability of the observed DDI data, if available;
- The expected duration of concomitant drug use (e.g., acute, short-term, or chronic use of one or both drugs);
- The anticipated timing of the introduction of the concomitant medication;
- The mechanism of the DDI (e.g., reversible or time-dependent inhibition, induction, combined inhibition and induction);
- The availability of monitoring parameters (e.g., therapeutic drug monitoring, laboratory tests);
- The ability to interrupt the investigational drug or concomitant interacting medication and the availability of other therapeutic options for either drug;
- The clinical importance of the relevant adverse outcome relative to the clinical benefit of the drugs;

After considering the items above, DDI management strategies can include the following: (Note that there may be regional regulatory differences in how recommendations are worded.)

- Contraindicating or avoiding concomitant use;
- Temporarily discontinuing one of the interacting drugs;
- Modifying the dosing regimen of one of the drugs;
- Staggering drug administration (e.g., administer the investigational drug at a different time than a concomitant drug);
- Implementing specific monitoring strategies (e.g., therapeutic drug monitoring, laboratory testing);
- Replacing one of the interacting drugs with a drug not expected to interact.

7. APPENDICES

7.1 Glossary

ADC: antibody-drug conjugate

ADME: absorption, distribution, metabolism, and/or excretion

AUC: area under the concentration vs. time curve

AUC_{0-inf}: AUC extrapolated to infinity

AUC_{0-t}: AUC from time zero to time of last quantifiable observation (t)

AUC_{0-tau}: AUC for one dosing interval following multiple doses, generally at steady state

AUCR: ratio of AUCs of the substrate (object) drug in the presence and absence of a precipitant drug

BCRP: breast cancer resistance protein

CAR: constitutive androstane receptor

C_{max}: maximum concentration of the drug after dosing

C_{max,u}: unbound C_{max}

C_{max,inlet,u}: estimated unbound C_{max} of an inhibitor at liver inlet.

C_{min}: minimum concentration during one dosing interval at steady state

CYP: cytochrome P450

DDI: drug-drug interaction

EC₅₀: concentration causing half maximal effect

E_{max}: maximum induction effect

fm: fraction of systemic clearance of the substrate mediated by the CYP enzyme that is subject to inhibition/induction.

f_{u,p}: unbound fraction in plasma

HLM: human liver microsome

IC₅₀: half-maximal inhibitory concentration

IC_{50,u}: unbound IC₅₀

Index precipitant: A drug recommended for use in a stand-alone clinical DDI study because it has a well-established potency and selectivity profile that causes a defined degree of inhibition or induction of a given elimination pathway when administered with a sensitive and specific substrate of that pathway.

Index substrate: A drug recommended for use in a stand-alone clinical DDI study as substrate because it has a well-established sensitivity and specificity profile that demonstrates a defined degree of change in exposures when administered with a strong inhibitor or inducer for that specific elimination pathway.

Investigational drug: A medicinal product or a drug under development that is investigated as to

its potential to act as an affecting drug or an affected drug

k_{deg} : apparent first-order degradation rate constant of the affected enzyme

K_i : inhibition constant causing half-maximal inactivation

$K_{i,u}$: unbound inhibition constant causing half-maximal inactivation

k_{inact} : maximal inactivation rate constant

K_m : Michaelis-Menton constant

K_{obs} : apparent first-order inactivation rate constant of the affected enzyme

MATE: multidrug and toxin extrusion

MRP: multidrug resistance-associated protein

NADPH: nicotinamide adenine dinucleotide phosphate (reduced form)

Nested DDI study: A DDI investigation that is part of a clinical study (e.g., phase 2/3) in which the assessment of DDI is not the primary objective.

No-effect boundaries: Interval within which a change in a systemic exposure measure is considered not significant enough to warrant clinical action (e.g., dose or schedule adjustment, additional therapeutic monitoring, avoid use)

OAT: organic anion transporter

OATP: organic anion transporting polypeptide

Object: A substrate of enzyme or transporter

OCT: organic cation transporter

pAUC: partial AUC. Area under the concentration vs. time curve between two specific time points

PBPK: physiologically-based pharmacokinetic

P-gp: P-glycoprotein

Precipitant: A drug that can induce or inhibit an enzyme or a transporter.

Probe substrate: A drug used in *in vitro* studies that measure individuals enzyme inhibition or induction properties of an investigational drug. The probe substrate should be selective, or the formation of a specific metabolite should be selective for the evaluated enzyme.

PXR: pregnane X receptor

Stand-alone DDI studies: A clinical DDI study with the primary objective of determining the presence or absence of a clinical DDI and the magnitude of the DDI.

Studies with index precipitants and index substrates: Clinical DDI studies conducted with index precipitants or index substrates that aim to investigate the greatest magnitude of interaction with the investigational drug for the studied pathway and which results usually can be extrapolated to other drug combinations.

TDI: time-dependent inhibition

T_{max}: time to C_{max}

t_{1/2}: elimination half-life

UGT: uridine diphosphate (UDP)-glucuronosyl transferase

7.2 Protein Binding

A key parameter of interest in predicting the risk of DDIs for an investigational drug as a precipitant is plasma protein binding. Historically, considering uncertainties in protein binding measurements for highly bound drugs, regulatory agencies have recommended the $f_{u,p}$ (fraction unbound in plasma) be set at 0.01 (i.e., 1%). This conservative approach was adopted to protect against false negative DDI predictions. Recent advances have resulted in the development of methodologies that allow accurate and precise measurement of protein binding for highly protein bound drugs. When choosing a protein binding assay, it is important to ensure the suitability of a given assay methodology for the investigational drug. The next step is to demonstrate the protein binding assay is precise and accurate. Such a demonstration should include validation data of the protein binding assay qualified with appropriate positive controls (i.e., range of compounds with high binding to relevant plasma proteins). The bioanalytical methods used in these experiments should have appropriate precision and accuracy at the required sensitivity range (i.e., Calibration standards and Quality Controls at 15%; Lower Limit of Quantification at 20%). Protein binding assays that have demonstrated these features can be used to determine the protein binding of an investigational drug (21). When novel and emerging protein binding assay methodologies are being established, the assay method should be qualified against an orthogonal method that is previously established and accepted for a range of highly bound compounds (22).

Of note, the protein binding experiments for the investigational drug should always include a positive control for which the assay was previously qualified to ensure in-study performance. The positive control should be demonstrated to be within 3-fold of the mean $f_{u,p}$ value that was previously reported with the qualification of the protein binding assay method.

7.3 *In Vitro* Evaluation of Metabolism-Based DDIs

7.3.1 *In Vitro* Systems

Various hepatic *in vitro* systems can be used to evaluate the risk for enzyme-mediated interactions for an investigational drug, including:

- Subcellular human liver tissue fractions such as microsomal systems (human liver microsomes (HLM); containing CYP450 and UGT enzymes), supernatants after 9000 g centrifugation of liver homogenate (S9; containing microsomal as well as cytosolic enzymes such as sulfotransferases, glutathione transferases, aldehyde dehydrogenase, aldehyde oxidase and alcohol dehydrogenase), and cytosol (adding co-factors as appropriate). For HLM, a pool of at least 10 donors is suggested.
- Recombinant, such as human CYP and UGT enzymes. These systems usually express only one single enzyme.
- Human liver tissues, including freshly prepared or cryopreserved hepatocytes that preserve enzyme architecture and contain the full complement of phase 1 and 2 drug metabolizing enzymes. For phenotyping and inhibition experiments, hepatocytes pooled from at least 5-10 donors is suggested, whereas for induction experiments usually hepatocytes from at least 3 individual donors should be used.

The *in vitro* systems used should be robust and reproducible.

Microsomal protein concentrations should be minimised, and standardised assay conditions (e.g., buffer strength, and pH) should be used. An incubation time and an enzyme amount that result in linear formation of the metabolite (at an initial rate of the metabolite formation) is recommended.

For phenotyping experiments, the system should be characterized with *in vitro* probe substrates to prove the activity of each enzyme. In general, a probe substrate should be selective (e.g., predominantly metabolized by a single enzyme), or a specific metabolite of a probe substrate is primarily formed by a single enzyme. A list of examples of probe substrates with their marker reactions can be found in Table 4, Section 7.6.1.1 For studies of time-dependent inhibition or induction, appropriate inhibitors or inducers should be included as positive controls (refer to Section 7.6.1 for more details).

For enzyme inhibition studies, if the investigational drug is metabolized by the enzymes present in the incubation, the probe substrate should, if possible, have a markedly faster metabolism rate than the investigational drug to minimize the influence of investigational drug metabolism (decreasing concentrations) on the estimation of inhibitory parameters.

Robust analytical methods should be used to quantify an investigational drug and its relevant metabolite(s) in phenotyping experiments as well as probe substrates and/or their relevant metabolites in inhibition and induction experiments (when enzyme activities are measured). Good laboratory practice (GLP) standard is not required, but a full description of the analytical methods employed should be provided.

It is recognized that obtaining high drug concentrations in the *in vitro* studies of enzyme inhibition or induction may not be possible in some circumstances due to poor aqueous solubility or cell toxicity. If limited by solubility, co-solvents can be used to reach the highest concentration possible. Any organic solvents should be used at low concentrations (<1% volume/volume and preferably < 0.5%) because some solvents can inhibit or activate enzymes. The experiment should include a solvent (vehicle) control, and when appropriate (e.g., when uncommon solvents are used), also a no-solvent control to evaluate potential effect of solvent on enzyme reaction. There is at present much uncertainty regarding how to interpret *in vitro* inhibition and induction data when sufficiently high concentrations cannot be tested; absence of a clinical DDI can be further justified by using other methods to calculate inhibition and induction parameters (6) and by considering pharmacokinetic characteristics of the compound, such as solubility limited absorption, dose- and time dependency, etc. Otherwise, clinical studies are recommended to test the DDI potential of these compounds.

Limited drug stability or non-specific binding in the incubations (e.g., with apparatus, microsomes or hepatocytes) can also create experimental challenges in *in vitro* studies of enzyme inhibition or induction. Actual unbound concentrations of the drug in the *in vitro* system (e.g., incubation medium) should in general be used for extrapolating *in vitro* results to clinical scenarios. Non-specific binding can be measured experimentally (e.g., using equilibrium dialysis) or predicted using *in silico* methods. For highly lipophilic drugs, it is preferred to experimentally determine non-specific binding (23).

For induction experiments, sponsors are encouraged to measure concentrations of the parent drug in the medium on the last day of incubation with hepatocytes and non-specific binding should also be considered. When measured concentrations in medium are <80% of concentrations due to non-metabolism/transporter related confounders, sponsors should discuss the potential impact of the discrepancy on data interpretation with regulatory authorities (24,25).

7.3.2 Investigational Drug as an Enzyme Substrate: Reaction Phenotyping

Drug metabolizing enzyme identification studies, often referred to as reaction phenotyping studies, identify the specific enzymes contributing to the main elimination pathways of a drug. Along with other information (e.g., clinical pharmacokinetics, mass-balance study, pharmacogenetic data or available DDI data), *in vitro* phenotyping data are often used to identify and quantify elimination pathways of an investigational drug.

Although the main focus of this guideline is on hepatic CYP involved metabolism, in order to identify the metabolic pathways for the individual investigational drug, non-CYP enzyme-based metabolism and metabolism occurring in extra-hepatic tissues should also be considered for certain drugs.

7.3.2.1 Metabolic Pathway Identification

Metabolic pathway identification experiments should be performed early in drug development to identify the number and structures of metabolites formed when a drug is metabolized and to investigate whether the metabolic pathways are parallel or sequential. These experiments use HLM, intact human liver systems (e.g., hepatocytes), or recombinant enzyme systems. Data obtained from metabolic pathway identification experiments help to determine whether and how to conduct a reaction phenotyping study.

7.3.2.2 Metabolic Enzyme Identification

Reaction phenotyping can be done in HLM or hepatocytes using selective enzyme inhibitors and/or in human recombinant enzymes. When using individual human recombinant enzymes, the difference in the amount and enzyme activity of CYPs between the recombinant CYP enzyme systems and the human liver should be considered. Whenever possible, all experiments should be conducted with drug concentrations relevant to the clinical setting, and under initial rate conditions (e.g., linearity of metabolite production rates with respect to time and enzyme concentrations).

The contribution of individual enzymes to the overall metabolism of an investigational drug can be examined by measurement of parent drug depletion or measurement of metabolite formation. For the latter method, all the major metabolites should have been identified and quantified in metabolite formation experiments. The use of a radiolabelled drug substrate is advantageous because samples can be analysed using liquid chromatography coupled with a radioactivity detector and a mass spectrometer to identify and quantify drug-related species. Evaluation of individual isomers of racemic drugs is recommended when it is important to understand the different disposition characteristics of each isomer (e.g., when two isomers have different pharmacological activities).

Some chemical inhibitors are not specific for an individual CYP enzyme. The selectivity and potency of inhibitors should be verified in the same experimental conditions using probe substrates for each CYP enzyme (see Section 7.6.1.1 for more details). If specific antibodies are used instead of inhibitors, the inhibitory effect of an antibody to a CYP enzyme should be tested at sufficiently low and high concentrations to establish a titration curve and ensure the maximal inhibition of a particular pathway (ideally resulting in >80% inhibition). The effect of an antibody should be verified using probe substrates of each CYP isoform in the same experimental conditions.

For UGT enzymes, *in vitro* studies are most commonly performed with HLM or recombinant UGT enzymes as the enzyme source. When HLM are used as enzyme source, either addition of alamethicin or sonication is necessary for activating HLM and add BSA to prevent inhibition by long-chain fatty acid (4). Determination of the contribution of each UGT isoform to the overall elimination is sometimes challenging due to lack of selective inhibitors, variability of results depending on experiment conditions, and instability of glucuronide metabolite in biologic matrices from a mass balance study.

7.3.3 Investigational Drug as an Enzyme Inhibitor

The potential of an investigational drug to inhibit CYP enzymes is usually investigated using selective probe substrates to determine the type of inhibition (e.g., reversible inhibition or time-dependent inhibition (TDI)) and measure of inhibition potency (e.g., $K_{i,u}$ for reversible inhibition, and $K_{i,u}$ and k_{inact} for TDI). The *in vitro* systems used for these studies include pooled HLM, microsomes obtained from recombinant CYP-expression systems, or pooled human hepatocytes.

For reversible inhibition, experiments with a high concentration of test drug can be performed first to study its inhibition potential on a particular enzyme (e.g., $50 \times C_{max,u}$ or $0.1 \times \text{Dose}/250 \text{ mL}$, refer to Section 2.1.2.1). If clinical interaction cannot be excluded at the high concentration, lower drug concentrations should be tested to estimate the drug's $IC_{50,u}$ or $K_{i,u}$ value; it is recommended to examine at least four different concentrations of the investigational drug. To determine the $K_{i,u}$ for inhibition, varying concentrations of both the inhibitor and substrate, to cover ranges above and below the substrate's K_m , should be tested. For competitive inhibition or uncompetitive inhibition, $IC_{50,u}/2$ can be used as an estimate for $K_{i,u}$ if the substrate concentration in the incubation is the same as its K_m value (26). If the substrate concentration is much less than the K_m value, then the $IC_{50,u}$ value will approximate the $K_{i,u}$ value for a competitive inhibitor. More accurate estimation of the $K_{i,u}$ value can be derived from the $IC_{50,u}$ value using the Cheng-Prusoff equation (8). For non-competitive inhibition, $K_{i,u}$ value is equal to $IC_{50,u}$ regardless of substrate concentration used (27). Thus, $IC_{50,u}/2$ can still be used as a conservative estimate.

Evaluation of *in vitro* TDI potential of an investigational drug consists of two steps. The first step is a screening step to identify potential for TDI of CYP enzymes, and the second step measures the inhibition potency. There are various assays to identify TDI of CYP enzymes. For example, TDI can be detected by assessing a difference in IC_{50} curves generated with and without a pre-incubation with nicotinamide adenine dinucleotide phosphate (NADPH) (i.e., IC_{50} shift), decreases in enzyme activity (measurement of the pseudo first-order rate constant, k_{obs}), or percent activity loss with the inactivator over time. In the IC_{50} shift assay, pooled HLM should typically be pre-incubated for 30 min with the investigational drug with or without NADPH. A left shift of the IC_{50} curve (e.g., ≥ 1.5 - or 2-fold) from the samples pre-incubated with NADPH compared to those without, suggests a potential for enzyme inactivation by the investigational drug. A known TDI should be included as positive control, which should provide a response that is within an

established historical range to demonstrate sensitivity of the experimental system, otherwise the assay should be repeated. Dilution and non-dilution methods can be used for the IC₅₀ shift assay. Dilution assays may be more sensitive for screening purposes because inactivation of the enzymes occurs at higher concentrations of the investigational drugs whereas non-dilution method may be useful in instances of poor solubility of the drug or metabolically labile compounds.

To rule out a TDI, the decreases in CYP enzyme activity with time can also be evaluated at a single concentration of the investigational drug (e.g., k_{obs} or percent activity loss). A reduction in CYP enzyme activity greater than a pre-defined threshold for the assay (e.g., of >20% reduction in activity or a k_{obs} value of $>0.01 \text{ min}^{-1}$) can be used to define a positive result.

When a drug is identified as a TDI with initial assessment as described above, definitive *in vitro* studies should be performed to obtain TDI parameters (i.e., k_{inact} and $K_{I,u}$) in pooled HLM for DDI predictions. Human hepatocytes and rhCYP can also be considered for TDI assessment.

7.3.4 Investigational Drug as an Enzyme Inducer

The potential of an investigational drug to act as an inducer of CYP enzymes is normally investigated in plateable, cryopreserved or freshly isolated, human hepatocytes. Alternative *in vitro* systems such as immortalized hepatic cell lines and cell receptor assays can be used, but the results from these studies are generally considered supportive rather than definitive in nature. If an alternative *in vitro* system is used as the main method, the sponsor should provide a justification supporting the appropriateness of the *in vitro* system as well as data interpretation.

It is recommended to measure the extent of enzyme induction at the mRNA level. Enzyme activity can also be measured, but measuring only the enzyme activity is usually not recommended, as the induction could be masked when the investigational drug is also an inhibitor. However, when *in vitro* induction of CYP2C19 is evaluated, enzyme activity should be measured, since its mRNA change is often limited even in response to positive control (28).

Regardless of which *in vitro* system and endpoint are chosen, the system should be qualified to show that all major CYP enzymes are functional and inducible with positive controls. The response of positive controls (measured as mRNA fold change) is normally at least a 6-fold increase for CYP1A2, CYP2B6, and CYP3A4, which is considered indicative of satisfactory sensitivity of hepatocyte lots (refer to Section 2.1.4.1) (29). *In vitro* induction of CYP2C8, CYP2C9, and CYP2C19 can also be measured, but their mRNA change is often limited even in response to the positive control (e.g., rifampicin). Therefore, a 6-fold increase of the positive control for these enzymes is not mandatory as a guarantee of sensitivity. Conclusive interpretation of the results is often challenging because of the limited fold of increase in mRNA of these enzymes to positive controls.

Duration of incubation of an investigational drug is usually 48-72 hours to allow complete induction to occur. Justification should be provided for shorter incubation times (28). Incubations normally include a daily addition of the investigational drug, and the medium containing the drug should be changed regularly. More frequent addition of a drug can be considered if its stability is low. The optimal time course for incubation should allow detection of enzyme induction without causing cell toxicity. If cytotoxicity occurs, reduced incubation durations can be used if adequate sensitivity of the assay can be demonstrated.

Quality and viability of the cultured hepatocytes should be verified and documented by cell morphology and biochemistry tests. A suitable viability assessment is normally performed before and at the end of the incubation period to certify that cell toxicity is not influencing the induction response. If toxicity/loss of viability is observed, influence on the study results should be discussed in the study report, and clinical studies may be considered.

If hepatocytes from a donor: (a) do not respond satisfactorily to the positive induction controls, (b) demonstrate viability <80% before addition of the investigational drug, or (c) demonstrate viability <70% at the end of the incubation compared to solvent (vehicle) control, the cells should be replaced by hepatocytes from a new donor.

To rule out that the investigational drug is an *in vitro* inducer, an induction study including 3 donor hepatocytes should be run with at least 3 replicates of 3-5 different concentrations of the test investigational drug, encompassing $50 \times C_{\max,u}$. The basic mRNA fold-change method can be used to evaluate clinical induction potential based upon the criteria described earlier (refer to Section 2.1.4.1).

When there is an induction signal, 5-8 concentrations are recommended for an accurate determination of $E_{\max}$ and $EC_{50,u}$. The sponsor can further use the correlation method or mechanistic static models to predict the magnitude of a clinical induction effect of an investigational drug. These methods utilize full concentration-response curves for induction, to estimate $E_{\max}$ and $EC_{50,u}$ of the investigational drug. In addition, to use these methods, a batch of hepatocytes should be 'calibrated' (5). For the correlation method, a large set of inducers ($n \geq 8$) covering the full clinical induction potency range and including at least 2 weak inducers, are recommended for calibration. $E_{\max}$ and $EC_{50,u}$ are determined for all inducers and a correlation is established between a certain matrix (incorporating $E_{\max}$ and/or $EC_{50,u}$ and clinical concentrations of inducers) and clinical change in the AUC of a sensitive substrate of a specific enzyme (e.g., midazolam for CYP3A) for each inducer. For the mechanistic method, an empirical calibration factor, 'd' factor to enable *in vitro*-to-*in vivo* induction scaling, should be determined for a hepatocyte batch. The 'd' factor can be estimated by correlating the predicted and observed induction effects (i.e., AUC ratio of a sensitive substrate of a particular enzyme) of a set of known

inducers and performing a linear regression to identify a 'd' value that can minimize the prediction error (7). If the 'd' factor is not estimated, it should be set as a default value of 1.

For the correlation or mechanistic static methods, sponsors can use only one hepatocyte donor. The calibration can be established once for that batch of hepatocytes rather than multiple times for each experiment with investigational drugs. When performing the *in vitro* study evaluating the induction potential of an investigational drug, criteria for acceptable assay variability should be established. To confirm performance and sensitivity of the experimental system, at least 2 of the inducers (weak and strong) of the calibration set should be included as controls. The responses of the controls should fall within the defined assay variability in order to utilize the calibration set of that hepatocyte batch. If this method is used, both the calibration data set/calibration report and the data on the investigational drug should be submitted.

7.4 *In Vitro* Evaluation of Transporter-Based DDIs

7.4.1 *In Vitro* Systems

Various *in vitro* transporter assays can be used to evaluate the risk for transporter-mediated interactions of an investigational drug. Selecting the *in vitro* model can depend on the purpose of the study and the questions to be addressed. Available *in vitro* systems include:

- Membrane vesicles

In vitro systems using inside-out membrane vesicles from cells transfected with a transporter can be used to evaluate whether an investigational drug is a substrate or inhibitor of efflux transporters such as P-gp or BCRP but may fail to identify highly permeable drugs or highly non-specific binding drugs as substrates.

P-gp and BCRP assays using membrane vesicles should directly measure the adenosine triphosphate (ATP)-dependent, transporter-mediated uptake of drugs. For studies evaluating a drug as substrate of transporters, control groups should be included, for example, AMP-treatment or non-transfected vesicles.

- Bi-directional transport assays with cell-based systems

Bidirectional assays can be used to evaluate whether an investigational drug is a substrate or inhibitor of efflux transporters such as P-gp or BCRP.

The permeability of the substrate should be investigated in both directions and evaluated under conditions for which the transport rate is linear. The apparent permeability (P_{app}) of the drug in both the AP→BL (absorption: apical to basolateral) and BL→AP (efflux: basolateral to apical) directions can be calculated, as well as an efflux ratio (ER) of BL→AP to AP→BL.

$$ER = \frac{P_{app,BL-AP}}{P_{app,AP-BL}}$$

When using transfected cell lines, efflux ratios of the transfected cell line should be compared with appropriate control conditions to account for endogenous transporter activity and non-specific binding. One approach is to compare the efflux ratios from transfected cell line to the parental or empty vector-transfected cell line.

$$Net\ ER = \frac{ER_{transfected}}{ER_{parental}}$$

The integrity of monolayer membrane should be measured before and after experiments by examining whether transepithelial/transendothelial electrical resistance (TEER) values or permeability of paracellular markers fall within the pre-defined acceptance range.

- Uptake assays with cell-based systems

Uptake assays can be used to evaluate whether an investigational drug is a substrate or inhibitor of solute carrier (SLC) transporters such as OCTs, OATs, OATPs and MATEs, but can also be used to investigate efflux transporters.

When transfected cell lines are used to evaluate whether a drug is a substrate of a transporter, the drug uptake in the transfected cell line should be compared to the parental or empty vector-transfected cell line, or a comparison of the uptake with or without an inhibitor of the transporter should be performed. When assessing a drug as an inhibitor of a transporter, evaluation of the uptake of a known probe substrate using transporter-transfected cell lines alone can be sufficient. Besides transfected cell lines, human hepatocytes or hepatic cell lines in suspension or plated can be used.

The model system and experimental conditions should be validated, including culture and transport assay conditions. Transport studies should be performed under linear transport rate conditions (probe substrate concentration used is usually below its K_m for the transporter). Appropriate positive controls should be included in the test study to ensure the validity of the study's results. The assays should be optimized to ensure consistent transporter function (e.g., uptake, efflux) with control experiments (e.g., positive and negative controls for substrates/inhibitors (refer to Tables 10 and 11, Section 7.6.3 for some examples), non-transfected control cells). The following conditions should be considered whenever applicable: the source of the membrane vesicles or cells, cell culture conditions (e.g., cell passage number, seeding density, monolayer age), probe substrate/inhibitor concentrations, incubation time, buffer/pH conditions, sampling interval, and methods for estimating parameters such as the IC_{50} , K_i , and K_m . In addition, adding serum or plasma proteins to the media can also affect transport activity.

Laboratory acceptance criteria for study results should be established (e.g., monolayer integrity, passive permeability, efflux or uptake of probe substrates, K_m for a probe substrate, IC_{50} for a probe inhibitor). The K_m value of a probe substrate or the IC_{50} value of a probe inhibitor should be comparable to literature-reported values.

The substrate should be readily measured with no interference from the assay matrix.

Any organic solvents should only be used at low concentrations (< 1% volume/volume and preferably < 0.5%) because some solvents can affect cell integrity or transporter function. The experiment should include a solvent (vehicle) control, and when appropriate (e.g., uncommon solvents are used), also a no-solvent control to evaluate potential effect of solvent on transporter activity.

Several factors may cause actual drug concentrations in the *in vitro* assays to deviate from nominal concentrations, including poor aqueous solubility, non-specific binding, and instability. Sponsors are encouraged to measure drug concentrations in the medium. Correction for binding or stability or solubility issues should be considered when interpreting the data.

7.4.2 Investigational Drug as a Transporter Substrate

The concentration range of an investigational drug should be relevant to the site of transport and should be based on the expected clinical concentration range. For transporters expressed in multiple organs (e.g., P-gp, BCRP), the sponsors should provide justification for the choice of concentrations, taking into consideration the sites where the transporter is likely to play a role for drug disposition. When a range of drug concentrations is relevant, it is important to ensure that low concentrations are included, as high concentrations may saturate transporters that are still active at lower drug concentrations.

If the *in vitro* system expresses multiple transporters (e.g., Caco-2 cells, hepatocytes), the sponsor should conduct additional experiments to confirm the findings with two or more known potent inhibitors, including the ones that are relatively specific for individual transporters.

If active transport is concluded, the passive permeability in the absence of transporters is one of the factors that could be taken into account to estimate the clinical importance of the transporter. For intestinal transporters, the role of these transporters may be limited if the permeability in the absence of transporters is high ($\geq$ the permeability constant of the highly permeable control drug). In this case, the effect of active drug transport may be negligible as compared to the passive, concentration-gradient driven absorption of the drug. To estimate the permeability of a drug in the absence of transporters, for bi-directional assays (e.g., Caco-2 cells) the permeability constant can, for example, be determined at concentrations high enough to completely saturate the transporters

(assessed as an ER ratio of 0.5 – 2). If this approach is used, it should be established that the cell monolayer is unaffected. Alternatively, the permeability of a drug can be measured in the presence of a broad inhibitor of transporters. The investigation should include a well validated, high, and low permeable reference substance (for example, metoprolol and mannitol; refer to the ICH M9 guideline for more details).

7.4.3 Investigational Drug as a Transporter Inhibitor

Normally the investigation of transporter inhibition starts with testing a high concentration of the test drug, for example, $10 \times C_{\max,u}$ for OAT1/3 and OCT2, $50 \times C_{\max,u}$ for MATEs, $10 \times$ liver inlet $C_{\max,u}$ for OATP1B1/3, and $0.1 \times$ the highest therapeutic dose/250 mL for orally administered P-gp or BCRP inhibitors. The drug concentration should, however, not exceed the drug's solubility limits or cause deleterious effects (e.g., cytotoxicity) in the cells. There is at present much uncertainty regarding how to extrapolate *in vitro* results to *in vivo* when sufficiently high concentrations cannot be tested. Thus, the general recommendation is to test the DDI potential of these compounds *in vivo*, unless *in vitro* results are sufficiently justified.

If the test drug demonstrates inhibitory activity at the recommended cutoff concentration, the sponsor should test additional concentrations to estimate IC_{50} or K_i values. The sponsor should evaluate at least four concentrations of the investigational drug with the probe substrate. The sponsor can then compare IC_{50} or K_i values to clinical plasma or estimated intestinal concentrations of a drug to predict the potential for DDIs.

For some transporters (e.g., OATP1B1 and OATP1B3) and experimental systems, it can be relevant to determine IC_{50} or K_i following pre-incubation with the investigational drug, since some inhibitors demonstrated more inhibition potency after pre-incubation (30,31,32). This is an area of emerging information, and sponsors are encouraged to follow current literature for information on transporters of interest and relevant experimental protocols.

7.5 Predictive Modeling

This section describes how mechanistic modeling approaches can be used to: (1) characterize the potential for DDIs, (2) indicate whether a dedicated clinical DDI study should be conducted, and (3) support clinical recommendations in the absence of a clinical DDI study. The modeling approaches discussed are static mechanistic models and dynamic mechanistic models (also known as PBPK models).

Various mathematical and mechanistic modeling approaches in conjunction with findings from *in vitro* and early clinical studies can help translate *in vitro* observations into predictions of potential clinical DDIs.

Section 2 of this guideline describes the evaluation of *in vitro* metabolism and transporter studies to determine whether further evaluation of a drug as an object or precipitant of CYP enzyme- or

transporter-mediated interactions should be conducted. If those assessments indicate further evaluations should be conducted, they may be conducted using mechanistic static models or PBPK models, as described in 7.5.1 and 7.5.2. For some drug development programs, multiple approaches for assessing DDI risk may be feasible.

Depending on the results of the mechanistic static or PBPK modeling, follow-up clinical DDI studies could be needed.

The use of appropriate *in vitro* experimental conditions is critical to any model used for a quantitative prediction.

7.5.1 Using Mechanistic Static Models for DDI Predictions

A mechanistic static model incorporates detailed drug disposition and drug interaction mechanisms for both interacting and substrate drugs (2, 33, 34). The model includes the effect of reversible and time dependent enzyme inhibition, as well as enzyme induction. Thus, the model can estimate the effect of several interaction processes. The overall effect of the precipitant drug on the substrate drug is represented as AUCR (ratio of the AUC of the substrate drug in the presence and absence of the precipitant drug) and is given by the equation below. Input parameters should be sufficiently supported by data and/or scientific literature.

7.5.1.1 Evaluation of an Investigational Drug as a DDI Precipitant of CYP-Mediated DDIs

For a drug that is both an inhibitor and an inducer of an enzyme, in addition to the combination of inhibition and induction, a drug's inhibition potential alone (A and B only, assuming C is equal to 1 in the equation below), and induction potential alone (C only, assuming A and B are equal to 1 in the equation below) should be conducted. Concurrent prediction can lead to a false negative prediction if the inhibition potential is over-predicted, thus masking the induction effect (35). If the induction potential is over-predicted, it will mask the inhibition effect.

7.5.1.2 Evaluation of Investigational Drug as an Object of CYP-Mediated DDIs

In principle, mechanistic static models can be used to predict DDI effects with a less potent precipitants after the model has been confirmed with index precipitants.

Equation to calculate AUCR of the substrate drugs (AUC plus investigational drug/AUC minus investigational drug)

$$AUCR = \left(\frac{1}{[A_g \times B_g \times C_g] \times (1 - F_g) + F_g} \right) \times \left(\frac{1}{[A_h \times B_h \times C_h] \times f_m + (1 - f_m)} \right)$$

The equation assumes that the drug has negligible extrahepatic clearance.

A is the effect of reversible inhibitions.

B is the effect of TDI.

C is the effect of induction.

F_g is the fraction available after intestinal metabolism.

f_m is the fraction of hepatic clearance of the substrate mediated by the CYP enzyme that is subject to inhibition/induction.

Subscripts ‘h’ denote liver.

Subscripts ‘g’ denote gut.

Table 3: Equations to calculate AUCR of the substrate drug for reversible and time-dependent inhibition

	Gut	Liver
Reversible inhibition	$A_g = \frac{1}{1 + \frac{[I]_g}{K_i}}$	$A_h = \frac{1}{1 + \frac{[I]_h}{K_i}}$
Time-dependent inhibition	$B_g = \frac{k_{deg,g}}{k_{deg,g} + \frac{[I]_g \times k_{inact}}{[I]_g + K_I}}$	$B_h = \frac{k_{deg,h}}{k_{deg,h} + \frac{[I]_h \times k_{inact}}{[I]_h + K_I}}$
Induction	$C_g = 1 + \frac{d \times E_{max} \times [I]_g}{[I]_g + EC_{50}}$	$C_h = 1 + \frac{d \times E_{max} \times [I]_h}{[I]_h + EC_{50}}$

Each value can be estimated with the following equations:

$$[I]_h = f_{u,p} \times (C_{max} + (Fa \times Fg \times ka \times Dose) / Qh / R_B) \quad (36).$$

$$[I]_g = Fa \times ka \times Dose / Q_{en} \quad (35).$$

$f_{u,p}$ is the unbound fraction in plasma. The $f_{u,p}$ should be set to 1% if the reliability of $f_{u,p}$ measurements <1% cannot be demonstrated (also refer to Section 2.1.2.1). Since the potential impact of $f_{u,p}$ on the prediction of DDI is high, sensitivity analyses for $f_{u,p}$ should be provided for highly protein bound drugs.

C_{max} is the maximal total (free and bound) inhibitor concentration in the plasma at steady state.

Fa is the fraction absorbed after oral administration; a value of 1 should be used when the data are not available.

Fg is the fraction available after intestinal metabolism; a value of 1 should be used when the data are not available.

ka is the first order absorption rate constant in vivo; a value of 0.1 min⁻¹ (36) can be used when the data are not available.

Q_{en} is the blood flow through enterocytes (e.g., 18 L/hr/70 kg) (37).

Qh is the hepatic blood flow (e.g., 97 L/hr/70 kg) (38).

R_B is the blood-to-plasma concentration ratio.

d is a scaling factor determined in a calibrated hepatocyte batch based on positive control inducers (7, 33). If not determined, it is assumed to be 1. A different value can be used if supported by prior experience with the system used.

Reports of modeling exercises and results should provide support for input parameters based on data and/or scientific literature.

If the model estimates AUCR between 0.80 to 1.25, the risk of a clinically relevant interaction is low, and additional evaluations of the drug as a precipitant for the studied enzyme are not needed. If AUCR is outside 0.80 to 1.25, further evaluation should be conducted to quantify the effect. Alternatively, sponsors should provide sufficient justification(s) if no further assessments are planned.

Mechanistic static models are currently used to determine whether the potential for a DDI can be ruled out. This use, along with the current equations used for drug concentrations in the gut and liver (above), can be overly conservative and thus result in false positive results. However, when more relevant drug concentrations in gut and liver are considered, mechanistic static models can provide quantitative estimates of DDIs (2, 34).

For quantitative estimation of an interaction, reports of results should include justifications for both system- and drug-dependent parameters and sensitivity analyses to cover for uncertainties in model parameters, when relevant.

7.5.1.3 Evaluation of The Potential for Transporter-Mediated DDIs

Although there are fewer examples, the mechanistic static models can be used to evaluate transporter-mediated DDIs, when the fraction of substrate drug transported at tissues relevant to the transporter of interest (f_T) is derived from clinical data. For evaluating a drug as an inhibitor, the f_T of a substrate should be confirmed with clinical DDI studies with other inhibitors. The potential applications and considerations listed for PBPK modeling (refer Section 7.5.2.2) are generally also relevant for mechanistic static modeling.

7.5.2 Using PBPK Models to Predict Enzyme or Transporter-Based DDIs

PBPK models can assist in the evaluation of the DDI potential of an investigational drug and/or a metabolite as an object or precipitant of enzyme or transporter-mediated interactions. When PBPK modeling is used to support drug development and regulatory decisions, it is important to justify any model assumptions, the physiological and biochemical plausibility of the model, variability, and uncertainty measures. PBPK analysis reports should include a description of the context of use for the model, model structure and development plan, the sources and justifications for both system- and drug-dependent parameters, and an adequate sensitivity analysis plan. When using predefined models (structural and error), the software version and any deviations from predefined models should be described.

In general, broad recommendations for PBPK model verification, validation and the reporting of the results are beyond the scope of this guideline (refer to regional guidelines where available). Instead, this guideline describes the utility of PBPK modeling for the evaluation of DDIs, with the understanding that models should be demonstrated as fit-for-purpose. Specific best practice considerations for use of PBPK modeling for the evaluation of DDIs are also described below.

7.5.2.1 Potential Applications of PBPK to the Evaluation of CYP-Mediated DDIs

Related to evaluation of CYP-mediated DDIs, PBPK models can help select key DDI studies for a development program and inform the study design for clinical DDI studies. They can also be used to explain pharmacokinetic observations, such as observed pharmacokinetic differences that are due to genetic polymorphism.

When evaluating a drug as a potential object of CYP-mediated DDIs, PBPK models can be used to predict DDI effects with a less potent precipitant after the model has been confirmed with index precipitants. They can also predict clinically relevant DDI scenarios, such as the effect following multiple dose administration of the object drug if only single dose administration is evaluated in a clinical DDI study.

When evaluating a drug as a potential precipitant of CYP-mediated DDIs, PBPK models can be used to support the lack of clinical DDI potential and to predict DDI effects under different dosing regimens after the model has been confirmed with a sensitive index substrate.

7.5.2.1.1 Modeling Considerations - PBPK for Evaluation of CYP Interactions for Drugs as Substrates

Sponsors should consider the following when using PBPK modeling to predict the DDI potential of the investigational drug (including clinically relevant metabolite(s)) as a CYP enzyme substrate:

- The base PBPK model of the investigational drug should describe the available clinical pharmacokinetic data using different dosing regimens (e.g., a dose proportionality study, repeated dosing) and dosing routes (e.g., intravenous or oral).
- The major metabolic and other elimination pathways should be quantitatively assigned in all relevant tissues in the investigational drug's model according to available *in vitro* and clinical data.
- The uncertainty of the PBPK model parameters should be assessed using sensitivity analysis.
- The index precipitant models should describe the available clinical pharmacokinetic data using different dosing regimens (e.g., a dose proportionality study) and, as appropriate, different dosing routes (e.g., intravenous or oral).
- The acceptability of index precipitant models should be independently confirmed with regard to their modulating effect on the pharmacokinetics of sensitive enzyme substrates in humans.
- If complex metabolic and transport mechanisms are expected, the substrate and precipitant models should include the relevant disposition and interaction mechanisms and should be identifiable and deemed fit-for-purpose.

7.5.2.1.2 Modeling Considerations - PBPK for Evaluation of CYP Interactions for Drugs as Precipitants

Sponsors should consider the following when using PBPK modeling to predict the drug interaction potential of an investigational drug (including clinically relevant metabolite(s)) as a CYP enzyme precipitant:

- The base PBPK model of the investigational precipitant (and its metabolites, when relevant) should describe the available clinical pharmacokinetic data using different dosing regimens (e.g., a dose proportionality study, repeated dosing) and, as appropriate, dosing routes (e.g., intravenous or oral).
- The DDI parameters should be assigned in the precipitant's model according to available *in vitro* and clinical data such as clinical DDI study(ies).
- For precipitants that exhibit both inhibition and induction, the inhibition and induction mechanisms should be separately considered, in addition to the combination of inhibition and induction, to ensure a conservative prediction of clinical enzyme inhibition or induction. In most cases, the clinically relevant effect of interest is the combined effect.
- The sensitive index substrate models should describe the available clinical pharmacokinetic data using different dosing regimens (e.g., a dose proportionality study) and as appropriate, different dosing routes (e.g., intravenous or oral).
- The sensitive index substrate models should be independently confirmed with regard to the effect of a strong index precipitant-mediated altered enzyme activity on its pharmacokinetics in humans.
- The simulation should include the highest clinical dose and shortest dosing interval of the investigational precipitant. The pharmacokinetics and modulating effect of the highest dose should be confirmed before use in the simulation.
- Sensitivity analyses should be conducted for parameters exhibiting high levels of uncertainty. For example, since the potential impact of $f_{u,p}$ on the prediction of DDI is high, sensitivity analyses for $f_{u,p}$ is expected for highly protein bound drugs.

7.5.2.2 Potential Applications of PBPK to the Evaluation of Transporter-Mediated DDIs

Related to evaluation of transporter-mediated DDIs, PBPK models can be used to support the initial study design for clinical DDI studies when a DDI liability is identified.

When evaluating a drug as a potential object of transporter-mediated DDIs, PBPK models can be used to explain pharmacokinetic observations, such as pharmacokinetic differences that are due to genetic polymorphism (e.g., OATP1B1). PBPK models can also be used to explore involvement of specific transporters in a drug's ADME and DDI liability.

When evaluating a drug as a potential inhibitor of transporter-mediated DDIs, PBPK models may support negative DDI prediction when the drug is an *in vitro* inhibitor for a basolateral uptake

transporter (e.g., OAT1/3). They can also be used to evaluate the effect of an investigational drug on the pharmacokinetics of a transporter substrate with a well characterized pathway.

7.5.2.2.1 Modeling Considerations - Drug as a Transporter Substrate

In general, quantitatively confirming the model regarding the involvement of the specific transporter in the relevant organ(s) is challenging. Comprehensive model exploration and/or clinical studies should be conducted for quantitative model confirmation.

7.5.2.2.2 Modeling Considerations - Drug as a Transporter Inhibitor

In general, when using PBPK models to evaluate a drug as a transporter inhibitor, the substrate model should be confirmed for the relevant transporter(s). Further, the analysis report should include a sensitivity analysis addressing uncertainties related to the precipitant's inhibition constant and the concentration that is relevant for the DDI.

7.6 List of Drugs that can be used in In Vitro Studies

7.6.1 CYP Enzymes

7.6.1.1 CYP Enzyme Substrates for In Vitro Studies

Probe substrates are used to measure inhibitor/inducer properties of a drug candidate on individual CYP enzymes (see Table 4 for examples of substrates; the list is not exhaustive, and sponsors can use other substrates/metabolites with appropriate justification). The substrates should be selective, or the formation of a specific metabolite is selective for a CYP enzyme. Concentration of the substrate should be at or below its K_m for the reaction.

Table 4: Examples of probe substrates for CYP enzymes (in vitro studies)

CYP Enzyme	Probe substrate	Marker reaction
CYP1A2	Phenacetin 7-Ethoxyresorufin	Phenacetin O-deethylation 7-Ethoxyresorufin-O-deethylation
CYP2B6	Bupropion Efavirenz	Bupropion hydroxylation Efavirenz hydroxylation
CYP2C8	Paclitaxel Amodiaquine	Paclitaxel 6 α -hydroxylation Amodiaquine N-deethylation
CYP2C9	S-warfarin Diclofenac	S-warfarin 7-hydroxylation Diclofenac 4'-hydroxylation
CYP2C19	S-Mephenytoin	S-Mephenytoin 4'-hydroxylation
CYP2D6	Bufuralol Dextromethorphan	Bufuralol 1'-hydroxylation Dextromethorphan O-demethylation
CYP3A (recommend using two structurally different substrates)	Midazolam Testosterone	Midazolam 1'-hydroxylation Testosterone 6 β -hydroxylation

7.6.1.2 CYP Enzymes Inhibitors/inducers for In Vitro Studies

The enzyme inhibitors and inducers are used to phenotype individual CYP enzymes involved in the drug candidate metabolism *in vitro*. In general, the inhibitors/inducers should be selective at the concentration used. The following tables are provided to help sponsors design *in vitro* studies and to evaluate the interaction potential (Tables 5-7). These tables are not exhaustive, and sponsors can use other inhibitors/inducers with appropriate justification.

Table 5: Examples of inhibitors for CYP enzymes (*in vitro* studies)

CYP Enzyme	Inhibitor
CYP1A2	α -Naphthoflavone, Furafylline*
CYP2B6	Clopidogrel*, Ticlopidine*, Thiotepa*
CYP2C8	Gemfibrozil glucuronide*, Montelukast, Phenelzine*
CYP2C9	Sulfaphenazole, Tienilic acid*
CYP2C19	Loratadine, Ticlopidine*
CYP2D6	Paroxetine*, Quinidine
CYP3A	Azamulin*, Itraconazole, Ketoconazole, Troleandomycin*

* Designated as time dependent inhibitor. When used, those inhibitors should be pre-incubated with the experimental system.

Table 6: Representative values of the turnover rate constant (K_{deg}) and half-life ($t_{1/2}$) of major CYP enzymes to aid in the assessment of time-dependent inhibition

Enzymes (hepatic)	$t_{1/2}$ (hr)	K_{deg} (/min)
CYP1A2	38	0.00030
CYP2B6	32	0.00036
CYP2C8	22	0.00053
CYP2C9	104	0.00011
CYP2C19	26	0.00044
CYP2D6	51	0.00023
CYP3A4	36	0.00032
CYP3A4 (intestinal)	24	0.00048

Table 7: Examples of inducers for CYP enzymes (*In Vitro* Studies)

CYP Enzyme	Inducer
CYP1A2	Omeprazole
CYP2B6	Phenobarbital
CYP2C8	Rifampicin
CYP2C9	Rifampicin
CYP2C19	Rifampicin
CYP3A4	Rifampicin

7.6.2 UGTs

7.6.2.1 UGT Substrates for In Vitro Studies

The list provided in Table 8 is not exhaustive, and sponsors can use other substrates with appropriate justification.

Table 8: Examples of substrates for UGTs (In Vitro Studies)

UGT enzyme	Substrate
UGT1A1	β -Estradiol, PF-06409577
UGT1A3	Telmisartan
UGT1A4	Trifluoperazine, 1'-Hydroxymidazolam
UGT1A6	Deferiprone, 5-Hydroxytryptophol, Serotonin
UGT1A9	Mycophenolic acid, Propofol
UGT2B7	Morphine, Zidovudine
UGT2B10	Cotinine, RO5263397
UGT2B15	S-Oxazepam
UGT2B17	Testosterone

7.6.2.2 UGT Inhibitors for In Vitro Studies

There is a lack of relatively selective inhibitors for UGT1A3, UGT1A6, UGT2B7, and UGT2B15. In the absence of selective inhibitors, a combination of methods including use of recombinant UGT isoform, HLM expressing polymorphic variants of UGT isoform (where appropriate), the relative activity factor (RAF) or relative expression factor (REF) approach, and activity correlation approach can be employed. Comparative studies with multiple inhibitors can also help assess the involvement of a particular isoform. When using individual recombinant enzyme preparations, the difference in the amount and enzyme activity of UGTs between the recombinant enzyme systems and the human liver should be considered.

The list provided in Table 9 is not exhaustive, and sponsors can use other inhibitors with appropriate justification.

Table 9: Examples of inhibitors for UGTs (In Vitro Studies)

UGT enzyme	Inhibitor
UGT1A1	Nilotinib, Regorafenib
UGT1A3	-
UGT1A4	Hecogenin
UGT1A6	-
UGT1A9	Magnolol, Niflumic acid
UGT2B7	16 α - and 16 β -Phenyllongifolol*, fluconazole**
UGT2B10	Desloratadine
UGT2B15	-
UGT2B17	Imatinib

*16 α - and 16 β -Phenyllongifolol also inhibit UGT2B4. Their effects on UGT2B10 remains unknown.

** Fluconazole also inhibits UGT2B10 and UGT2B17.

7.6.3 Transporters

Some substrates are not specific for an individual transporter. When an experimental system expressing multiple transporters is used, a more specific substrate is preferred. The following tables provide examples of transporter substrate and inhibitors for *in vitro* studies (Tables 10 and 11).

Table 10: Examples of substrates for transporters (*In Vitro* Studies)

Transporter	Substrate
P-gp	Digoxin, <i>N</i> -methyl-quinidine (NMQ), Quinidine, Vinblastine
BCRP	Estrone-3-sulfate, 2-amino-1-methyl-6-phenylimidazo[4,5-b]pyridine (PhIP), Prazosin, Rosuvastatin, Sulfasalazine
OATP1B1, OATP1B3	Cholecystokinin octapeptide (CCK-8, selective for OATP1B3), Estradiol-17 β -glucuronide, Pitavastatin, Pravastatin, Rosuvastatin
OAT1	Adefovir, Cidofovir, <i>p</i> -aminohippurate (PAH), Tenofovir
OAT3	Benzylpenicillin, Estrone-3-sulfate, Methotrexate
MATE1, MATE2-K	Creatinine, Metformin, 1-methyl-4-phenylpyridinium (MPP+), Tetraethylammonium (TEA)
OCT2	Creatinine, Metformin, Tetraethylammonium (TEA)

Table 11: Examples of inhibitors for transporters (*In Vitro* Studies)

Transporter	Inhibitor
P-gp	GF120918 (dual P-gp/BCRP inhibitor), Verapamil, Valspodar (PSC833), Zosuquidar (LY335979)
BCRP	Fumitremorgin C, GF120918 (dual P-gp/BCRP inhibitor), Ko143, Novobiocin
OATP1B1, OATP1B3	Bromosulphophthalein (BSP), Cyclosporine, Rifampin, Rifamycin SV
OAT1, OAT3	Benzylpenicillin*, Probenecid
MATE1, MATE2-K	Cimetidine, Pyrimethamine, Quinidine
OCT2	Cimetidine, Clonidine, Pyrimethamine, Verapamil

* Relatively selective inhibitor for OAT3.

7.7 List of Drugs that can be used in Clinical Studies

7.7.1 CYPs Enzymes

7.7.1.1 CYP Enzyme Substrates for Clinical Studies

Ideally, drugs selections should be based on sensitivity, specificity, safety profiles, and reported clinical DDI studies with inhibitors, as well as an absence of studies that indicate the drug does not meet the criteria described below:

- Index substrates predictably exhibit exposure increase due to inhibition of a given metabolic pathway and results are available from prospective clinical DDI studies. These drugs can be safely administered with potential inhibitors, sometimes with a dose reduction.
- Sensitive index substrates are index drugs that demonstrate an increase in AUC of ≥ 5 -fold with strong index inhibitors of a given metabolic pathway in clinical DDI studies.
- Moderately sensitive substrates are drug that demonstrate an increase in AUC of ≥ 2 - to < 5 -fold with strong index inhibitors of a given metabolic pathway in clinical DDI studies.

Sponsors are encouraged to consider the unique characteristics of each drug when designing DDI studies. For example, a drug could be a substrate for multiple CYPs or a CYP plus a transporter. In such a case, the selection of an index drug for a study should take into consideration the knowledge about the potential precipitant (enzymes and/or transporters which it could inhibit). The drugs listed in Table 12 below have been identified as appropriate index substrates for clinical DDI studies. Other drugs can be proposed, considering the criteria above.

Table 12: Examples of index substrates for CYP enzymes (Clinical studies)

CYP Enzyme	Sensitive index substrate (unless otherwise noted)	Comments
CYP1A2	Caffeine	
CYP2B6	Bupropion	Bupropion is metabolized by CYP2B6 and non-CYP enzymes. Thus, by itself is not a sensitive substrate. Hydroxybupropion should also be measured since it is primarily formed by CYP2B6. Hydroxybupropion concentration changes should be considered when determining clinical significance, since it is the major active moiety.
CYP2C8	Repaglinide	Also metabolized by CYP3A though to a lesser extent. Transported by OATP1B1.
CYP2C9	S-warfarin, Flurbiprofen, Tolbutamide	Moderately sensitive substrates
CYP2C19	Omeprazole	Also metabolized by CYP3A though to a lesser extent. Measurement of metabolite concentrations should be considered when there are multiple interacting mechanisms involved.
CYP2D6	Desipramine, Dextromethorphan, Nebivolol	Dextromethorphan is also metabolized by CYP3A to a lesser extent.
CYP3A	Midazolam, Triazolam	

7.7.1.2 CYP Enzyme Inhibitors for Clinical Studies

Index inhibitors predictably inhibit metabolism via a given pathway, and results are available from prospective clinical DDI studies. Strong and moderate inhibitors are drugs that increase the AUC of sensitive index substrates of a given metabolic pathway ≥ 5 -fold and ≥ 2 - to < 5 -fold, respectively.

Ideally, index inhibitors should be selected based on potency and selectivity of inhibition, safety profiles, availability of reported clinical DDI studies with different clinical substrates, as well as an absence of studies that indicate the drug does not meet the criteria described above.

Sponsors are encouraged to consider the unique characteristics of each drug when designing DDI studies. For example, a drug could inhibit multiple CYPs or a CYP plus a transporter. Sponsors should select an index inhibitor for a study based on knowledge about the potential CYPs and transporters involved with the substrate's disposition.

The drugs listed in Table 13 below have been identified as appropriate index inhibitors for clinical DDI studies. Other drugs can be proposed, considering the criteria described above.

Table 13: Examples of index inhibitors for CYP enzymes (Clinical Studies)

CYP Enzyme	Strong index inhibitors	Comments
CYP1A2	Fluvoxamine	Also, strong inhibitor of CYP2C19; weak inhibitor of CYP2C9, CYP2D6, and CYP3A.
CYP2B6	<i>No strong index inhibitors are available for CYP2B6</i>	Ticlopidine can be used as a CYP2B6 inhibitor. It decreases hydroxybupropion formation by more than 80%. Ticlopidine is also a strong inhibitor of CYP2C19.
CYP2C8	Gemfibrozil	Also inhibits OATP1B1 and OAT3.
CYP2C9	Fluconazole (moderate inhibitor)	Also strong inhibitor of CYP2C19; moderate inhibitor CYP3A.
CYP2C19	Fluvoxamine Fluconazole	Fluvoxamine: Also, strong inhibitor of CYP1A2; weak inhibitor of CYP2C9, CYP2D6, and CYP3A. Fluconazole: Also, moderate inhibitor of CYP2C9 and CYP3A.
CYP2D6	Fluoxetine Paroxetine	Fluoxetine: Also, strong inhibitor of CYP 2C19.
CYP3A	Clarithromycin Itraconazole	Clarithromycin and itraconazole both inhibit P-gp. Itraconazole also inhibits BCRP.

7.7.1.3 CYP Enzyme Inducers for Clinical Studies

Inducers in Table 14 below were selected based on potency of induction, safety profiles, and availability of clinical DDI studies with different clinical substrates. Due to the mechanisms of induction, inducers usually regulate the expression of multiple enzymes and transporters.

Strong and moderate inducers decrease the AUC of sensitive index substrates of a given metabolic pathway by $\geq 80\%$ and $\geq 50\%$ to $< 80\%$, respectively.

Table 14: Examples of inducers for CYP enzymes (Clinical Studies) - the list is not exhaustive and other inducers can be used

CYP Enzyme	Strong inducers	Moderate inducers
CYP1A2*		Phenytoin, Rifampin, Cigarette smoking
CYP2B6	Carbamazepine	Rifampin, Efavirenz
CYP2C8		Rifampin
CYP2C9		Rifampin
CYP2C19	Rifampin	
CYP3A	Carbamazepine, Phenytoin, Rifampin	Efavirenz

*CYP1A2: Phenytoin, rifampin, and cigarette smoking are weak-to-moderate inducers based on limited number of clinical DDI studies conducted with caffeine, tizanidine, and theophylline.

7.7.2 UGTs

UGT substrates and inhibitors/inducers that are useful for clinical DDI studies are listed below (Tables 15-17). UGT substrates, inhibitors, and inducers are less established (see Section 3.2.4). Other substrates/inhibitors/inducers can be used with appropriate justifications.

Table 15: Examples of substrates for UGTs (Clinical Studies)

UGT enzyme	Substrates	Comments
UGT1A1	Bictegravir Cabotegravir Dolutegravir SN-38	Also metabolized by CYP3A Also glucuronidated by UGT1A9 Also metabolized by CYP3A and glucuronidated by UGT1A3 and UGT1A9 Active metabolite of irinotecan. Also glucuronidated by UGT1A9
UGT1A4	Lamotrigine Pexidartinib	Also glucuronidated by UGT2B7 Also metabolized by CYP3A
UGT1A9	Dapagliflozin Ertugliflozin Sotagliflozin	Also metabolized by CYPs Also glucuronidated by UGT2B4 and UGT2B7 Also metabolized by CYP3A
UGT2B7	Indomethacin S-Naproxen Zidovudine	Indomethacin and naproxen form acyl glucuronides that are unstable. Since there is possibility that the acyl glucuronides are transported by OAT1/3, alternative options may be considered if co-administered with an OAT1/3 inhibitor Also glucuronidated by UGT2B4

UGT2B15	Lorazepam Oxazepam	S-lorazepam and S-oxazepam are glucuronidated by UGT2B15. R-enantiomers are also glucuronidated by other UGT2B isoforms and UGT1A9
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Table 16: Examples of inhibitors for UGTs (Clinical Studies)

UGT enzyme	Inhibitors	Comments
UGT1A1	Atazanavir	Also, an inhibitor of CYP3A
UGT1A4	Probenecid	Also, an inhibitor of OAT1 and OAT3
UGT1A9	Mefenamic Acid	Also, an inhibitor of UGT2B7
UGT2B7	Probenecid	Also, an inhibitor of OAT1 and OAT3
UGT2B15	Probenecid	Also, an inhibitor of OAT1 and OAT3

Table 17: Examples of inducers for UGT (Clinical Studies)

UGT enzyme	Inducers
UGT1A1	Carbamazepine, Efavirenz, Phenobarbital, Rifampin, St. John's wort, Tipranavir combined with ritonavir
UGT1A4	Carbamazepine, Lopinavir combined with ritonavir, Phenobarbital, Phenytoin, Rifampin
UGT1A9	Rifampin
UGT2B7	Rifampin
UGT2B15	Rifampin, Phenytoin

7.7.3 Transporters

7.7.3.1 Transporter Substrates for Clinical Studies

Transporter substrates that are useful for clinical DDI studies are listed in Table 18 below. Many of them are substrates of multiple transporters and/or enzymes. Thus, the extrapolation of results from these studies to other drugs can be challenging and as indicated earlier (See Section 3.2.5), index substrates are not available for transporters. Interpretation of the study results should take into consideration the knowledge of the transporter inhibition properties for the investigational drug as well as its effect on metabolic enzymes. It is most useful to select a transporter substrate that is likely to be administered in the intended patient population for the investigational drug.

The listed substrates exhibit markedly altered pharmacokinetic profiles following co-administration of known inhibitors of the transporter, meeting the criteria below. In addition, they are generally safe for use in clinical DDI studies.

Criteria

The criteria below were used to select recommended transporter substrates for use in DDI studies to characterize a drug's transporter inhibition properties. Results from studies conducted with clinically relevant doses were used for selection of drugs. When possible, drugs most relevant for global drug development programs were selected.

- P-gp: (1) AUC fold-increase ≥ 2 with itraconazole, quinidine, or verapamil co-administration, (2) *in vitro* transport by P-gp expression systems, and (3) not extensively metabolized *in vivo*.
- BCRP: (1) AUC fold-increase ≥ 2 with pharmacogenetic alteration of ABCG2 (421C>A) and (2) *in vitro* transport by BCRP expression systems.
- OATP1B1/OATP1B3: (1) AUC fold-increase ≥ 2 with rifampin (single dose) or cyclosporine co-administration, or pharmacogenetic alteration of SLCO1B1 (521T>C) and (2) *in vitro* transport by OATP1B1 or OATP1B3 expression systems.
- OAT1/OAT3: (1) AUC fold-increase ≥ 2 with probenecid co-administration, (2) fraction excreted into urine as an unchanged drug ≥ 0.5 , and (3) *in vitro* transport by OAT1 and/or OAT3 expression systems.
- OCT2/MATEs: (1) AUC fold-increase ≥ 2 with dolutegravir or pyrimethamine; (2) fraction excreted into urine as an unchanged drug ≥ 0.5 , and (3) *in vitro* transport by OCT2 and/or MATEs expression system.

Note: The list is not exhaustive, and sponsors can use substrates that are not listed in the table if the drug's transport properties are well understood and similar to the criteria above.

Table 18: Examples of substrates for transporters (Clinical Studies)

Transporter	Substrates	Comments*
P-gp	Dabigatran etexilate Digoxin*** Fexofenadine	Dabigatran etexilate** – only affected by intestinal P-gp. Fexofenadine – also substrate for OATP1B1, 1B3 and 2B1.
BCRP	Rosuvastatin Sulfasalazine	Rosuvastatin – also a substrate for OATP1B1, 1B3, 2B1, and OAT3. Sulfasalazine – only affected by intestinal BCRP.
OATP1B1, OATP1B3	Atorvastatin Bosentan Pitavastatin Pravastatin Rosuvastatin Simvastatin acid	Atorvastatin – also a substrate of BCRP, P-gp, and CYP3A. Pravastatin – also a substrate of MRP2 and OAT3. Rosuvastatin – also a substrate for BCRP, OAT3, and OATP2B1. Simvastatin – also a substrate of CYP3A.
OAT1 OAT3	Adefovir Baricitinib Cefaclor Furosemide Oseltamivir carboxylate Penicillin G	Adefovir**** – Higher contribution of OAT1 than OAT3. Baricitinib, cefaclor and Penicillin G – Higher contribution of OAT3 than OAT1. Furosemide – dual substrate of OAT1/OAT3 is also a substrate of BCRP, OATP2B1, and UGT.

MATE1, MATE2-K, OCT2	Metformin	
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**Due to the evolving nature of the understanding, some of the drugs listed in the table could be substrates of other transporters that are not listed here.*

*** Dabigatran etexilate is a pro-drug and converted by carboxylesterase (CES) to dabigatran which is the measured moiety (dabigatran is not a substrate of P-gp). Thus, for correct interpretation of clinical DDI results, preassessment of the inhibitory effects of an investigational drug on CES activity should be considered.*

**** For P-gp, renal inhibition can be determined using renal clearance of digoxin.*

***** Adefovir dipivoxil is a pro-drug and converted by carboxylesterase (CES) to adefovir, which is a substrate of OAT1 and OAT3. Adefovir is the measured moiety in the DDI study. Thus, for correct interpretation of clinical DDI results, preassessment of the inhibitory effect of an investigational drug on CES activity should be considered. Adefovir dipivoxil is a substrate of P-gp.*

7.7.3.2 Transporter Inhibitors for Clinical Studies

Transporter inhibitors that are useful for clinical DDI studies are listed in Table 19 below. Many of them not only inhibit the specified transporters but also inhibit some other transporters and/or CYP enzymes. Thus, extrapolation of results from these studies to other drugs can be challenging as indicated earlier (See Section 3.2.5), index inhibitors are not available for transporters. Interpretation of the study results should take into consideration the knowledge of transport and metabolic/elimination pathways for the investigational drug. It is most useful to select a transporter inhibitor that is likely to be administered in the intended patient population for the investigational drug.

The listed inhibitors lead to markedly altered pharmacokinetic profiles of known substrates of the transporter following co-administration, meeting the criteria below. In addition, they are generally safe for use in clinical DDI studies.

Criteria

The criteria below were used to select recommended transporter inhibitors for use in DDI studies to characterize a drug's properties as a transporter substrate. Results from studies conducted with clinically relevant doses were used for selection of drugs. When possible, drugs most relevant for global drug development programs were selected.

- P-gp: (1) AUC fold-increase of digoxin, dabigatran, or fexofenadine ≥ 2 with co-administration and (2) *in vitro* inhibitor.
- BCRP: (1) AUC fold-increase of rosuvastatin ≥ 2 or close to 2-fold with co-administration and (2) *in vitro* inhibitor.
- OATP1B1/OATP1B3: (1) AUC fold-increase ≥ 2 for at least one of the clinical substrates with co-administration and (2) *in vitro* inhibitor.

- OAT1/OAT3: (1) AUC fold-increase ≥ 2 for at least one of the clinical substrates with co-administration and (2) *in vitro* inhibitor.
- OCT2/MATE: (1) AUC fold-increase of metformin ≥ 2 with co-administration and (2) *in vitro* inhibitor.

Note: The list is not exhaustive, and sponsors can use inhibitors that are not listed in the table if the drug's transporter inhibition properties are well understood and similar to the criteria above.

Table 19: Examples of inhibitors for transporters (Clinical Studies)

Transporter	Inhibitor	Comments
P-gp	Itraconazole Quinidine Verapamil	Itraconazole – also inhibits BCRP and CYP3A Verapamil – also inhibits CYP3A
BCRP	Cyclosporine Darolutamide Fostamatinib	Cyclosporine – also inhibits CYP3A, MRP2, OATP1B1, OATP1B3, and P-gp. Fostamatinib – also inhibits P-gp
OATP1B1, OATP1B3	Rifampin (single dose) Cyclosporine	Rifampin – also inhibits P-gp Cyclosporine – also inhibits CYP3A, MRP2, P-gp and BCRP
OAT1, OAT3	Probenecid	Probenecid – also inhibits OATP1B1.
MATE1, MATE2-K, OCT2	Dolutegravir Pyrimethamine	Dolutegravir – in general a more potent inhibitor for OCT2 than for MATEs Pyrimethamine – a relatively specific inhibitor of MATEs.

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