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Physiologically Based Pharmacokinetic Modeling for Quantitative Prediction of Exposure to a Human Disproportionate Metabolite of the Selective Na_v1.7 Inhibitor DS-1971a, a Mixed Substrate of Cytochrome P450 and Aldehyde Oxidase, Using Chimeric Mice With Humanized Liver^S

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ABSTRACT

In a previous study on the human mass balance of DS-1971a, a selective Na_v1.7 inhibitor, its CYP2C8-dependent metabolite M1 was identified as a human disproportionate metabolite. The present study assessed the usefulness of pharmacokinetic evaluation in chimeric mice grafted with human hepatocytes (PXB-mice) and physiologically based pharmacokinetic (PBPK) simulation of M1. After oral administration of radiolabeled DS-1971a, the most abundant metabolite in the plasma, urine, and feces of PXB-mice was M1, while those of control SCID mice were aldehyde oxidase-related metabolites including M4, suggesting a drastic difference in the metabolism between these mouse strains. From a qualitative perspective, the metabolite profile observed in PXB-mice was remarkably similar to that in humans, but the quantitative evaluation indicated that the area under the plasma concentration-time curve (AUC) ratio of M1 to DS-1971a (M1/P ratio) was approximately only half of that in humans. A PXB-mouse-derived PBPK model was then constructed to achieve a more accurate prediction, giving an M1/P ratio (1.3) closer to that in humans (1.6) than the observed value in PXB-mice (0.69). In addition, simulated maximum plasma concentration and AUC values of M1 (3429 ng/ml and 17,116 ng·h/ml, respectively) were similar to those in humans (3180 ng/ml and 18,400 ng·h/ml, respectively). These results suggest that PBPK modeling incorporating pharmacokinetic parameters obtained with PXB-mice is useful for quantitatively predicting exposure to human disproportionate metabolites.

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SIGNIFICANCE STATEMENT

The quantitative prediction of human disproportionate metabolites remains challenging. This paper reports on a successful case study on the practical estimation of exposure (C_{max} and AUC) to DS-1971a and its CYP2C8-dependent, human disproportionate metabolite M1, by PBPK simulation utilizing pharmacokinetic parameters obtained from PXB-mice and in vitro kinetics in human liver fractions. This work adds to the growing knowledge regarding metabolite exposure estimation by static and dynamic models.

Introduction

Human pharmacokinetics (PK) of parent compounds has been successfully predicted by in vitro to in vivo extrapolation and allometric

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methods (Obach et al., 1997; Tang et al., 2007), but the estimation of metabolite PK is still challenging. In fact, there are many cases where human disproportionate metabolites have not been quantitatively characterized until clinical studies.

A few papers indicated that exposure levels of metabolites of cytochrome P450 (P450) substrates such as midazolam, imipramine, (R)-4-((4-(((4-(((4-(((tetrahydrofuran-3-yl)oxy)benzo[d]isoxazol-3-yl)oxy)methyl)piperidin-1-yl)methyl)tetrahydro-2H-pyran-4-ol (also known as TBPT), and losartan were successfully estimated (Nguyen et al., 2016; Obach et al., 2018; Callegari et al., 2020), while even fewer papers described successful attempts for substrates where non-P450 enzymes were involved. Two approaches can predict metabolite

ABBREVIATIONS: AO, aldehyde oxidase; AUC, area under the plasma concentration-time curve; BW, body weight; CL, clearance; CL_{int}, intrinsic clearance; C_{max} , maximum plasma concentration; CL_{uint} , unbound intrinsic clearance; DMA, dimethylacetamide; eq., equation; f_{u} , unbound fraction; HLM, human liver microsome; HPLC, high-performance liquid chromatography; LC-MS/MS, liquid chromatography-tandem mass spectrometry; ka, absorption constant; kt, transit constant; M/P ratio, metabolite to parent area under the plasma concentration-time curve ratio; P450, cytochrome P450; PBPK, physiologically based pharmacokinetic; PK, pharmacokinetics; Rb, blood to plasma concentration ratio; T_{max}, time to reach the maximum plasma concentration; Vd, distribution volume.

exposure: static and dynamic models. Although static models enable a point-in-time calculation of metabolite-to-parent area under the plasma concentration-time curve (AUC) ratios (M/P ratios), dynamic models provide us with not only M/P ratios but also important PK parameters including maximum plasma concentration ($C_{\rm max}$) and the AUC values. Dynamic models include the physiologically based pharmacokinetic (PBPK) models in which various kinds of animal and human physiologic parameters, such as blood flow and organ volumes, are involved. Since substantial information about physiology and complex systems is required for the construction of a full PBPK model, researchers frequently use commercially available software, including Simcyp (Nguyen et al., 2016; Obach et al., 2018) and Gastroplus (Yamada et al., 2020). Meanwhile, customized PBPK models (Gill et al., 2013; Ito et al., 2020) are also developed on a fit-for-purpose basis.

PXB-mouse are chimeric mice prepared by the injection of human hepatocytes into the spleen of immunodeficient mice. More than 70% of liver cells in PXB-mice are replaced with human hepatocytes, and various kinds of human metabolizing enzymes and transporters are expressed in the liver of PXB-mice (Okumura et al., 2007; Hasegawa et al., 2012; Tateno et al., 2013). PXB-mice can be used for estimating human PK parameters by the allometric scaling method (Sanoh et al., 2015), and in vivo metabolite profiles in PXB-mice qualitatively reflect those in humans (Kato et al., 2020). Additionally, urinary and biliary excretion of metabolites in PXB-mice reflects that in humans (Sanoh et al., 2012; Kato et al., 2020). Meanwhile, caution should be paid to the influence of the remaining mouse hepatocytes on experimental results, and the accuracy of human metabolite prediction would depend on a case-by-case basis, as illustrated by the detection of metabolites produced by remaining mouse hepatocytes (Kamimura et al., 2010).

DS-1971a is a selective voltage-gated sodium channel (Na_v)1.7 inhibitor under a clinical trial for neuropathic pain (Shinozuka et al., 2020). Nonclinical evaluation indicated that more than 90% of radiolabeled DS-1971a was orally absorbed in male ICR mice and monkeys, and various kinds of oxidized metabolites with species differences in abundance were observed (Asano et al., 2021). Male ICR mice had the highest exposure to the aldehyde oxidase (AO)-dependent metabolite, M4, while the P450-produced metabolite, M2, was the predominant metabolite in male monkeys (Fig. 1). Thus, DS-1971a was identified as a mixed AO and P450 substrate. In vitro incubation study with human liver S9 in the presence of NADPH indicated that M1, a regioisomer of M2, was the potential major metabolite in humans.

A human mass balance study of radiolabeled DS-1971a in the fed condition demonstrated that more than 70% of the dose was absorbed, and the predominant metabolite was M1 in human plasma, urine, and feces (Asano et al., 2022). The exposure level of M1 in human plasma was 1.5-fold higher than that of the parent compound (DS-1971a), and M1 exposure in animals used for safety assessments (mice and monkeys) was much smaller than that in humans, so M1 was found to be a human disproportionate metabolite. In addition, incubation studies with human liver microsomes (HLMs) pretreated with P450-specific inhibitors and with recombinant P450 have indicated that CYP2C8 plays a key role in M1 formation (Asano et al., 2022).

Although previous studies on M1 formation in animals and humans suggested that M1 was a human disproportionate metabolite from a retrospective view, a critical question remains: How can the human exposure level of M1 as a human disproportionate metabolite be estimated before clinical trials? To answer this question, we performed PK evaluation in PXB-mice, which are likely the only animal species possessing human-like metabolic activity toward DS-1971a. Further, a PBPK model was constructed to predict the exposure levels of DS-1971a and M1. We here demonstrate the usefulness of the PBPK model derived

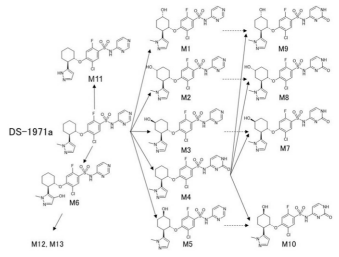


Fig. 1. The proposed metabolic pathway of DS-1971a from nonclinical studies. The map was prepared based on our previous report (Asano et al., 2021). Solid and dashed arrows indicate the metabolic pathways with high and low possibility, respectively.

from PXB-mice for estimating exposure of a human disproportionate metabolite

Materials and Methods

Chemicals and Reagents

DS-1971a (5-chloro-2-fluoro-4- $\{[(1S,2R)-2-(1-methyl-1H-pyrazol-5-yl)cyclo$ hexyl]oxy}-N-(pyrimidin-4-yl)benzenesulfonamide), a stable-isotope-labeled racemic mixture of DS-1971a (5-chloro-2-fluoro-4-{[(1RS,2SR)-2-[1-(13C,2H3)methyl-1*H*-pyrazol-5-yl]cyclohexyl]oxy}-*N*-(pyrimidin-4-yl)benzenesulfonamide), M1 (5-chloro-2-fluoro-4- $\{[(1S,2R,5S)-5-hydroxy-2-(1-methyl-1H-pyrazol-5-hydroxy-2-(1-methyl-1H-py$ yl)cyclohexyl]oxy}-N-(pyrimidin-4-yl)benzenesulfonamide), M2 (5-chloro-2-fluoro-4-{[(1S,2R,4R)-4-hydroxy-2-(1-methyl-1*H*-pyrazol-5-yl)cyclohexyl]oxy}-N-(pyrimidin-4-yl)benzenesulfonamide), M3 (5-chloro-2-fluoro-4-{[(1S,2R,4S)-4hydroxy-2-(1-methyl-1*H*-pyrazol-5-yl)cyclohexyl]oxy}-*N*-(pyrimidin-4-yl)benzenesulfonamide), M4 (5-chloro-2-fluoro-4-{[(1S,2R)-2-(1-methyl-1H-pyrazol-5-yl)cyclohexyl]oxy}-N-(2-oxo-1,2-dihydropyrimidin-4-yl)benzenesulfonamide), a racemic mixture of M5 (5-chloro-2-fluoro-4-{[(1RS,2SR,5SR)-5-hydroxy-2-(1-methyl-1*H*-pyrazol-5-yl)cyclohexyl]oxy}-*N*-(pyrimidin-4-yl)benzenesulfonamide), M6 (5-chloro-2-fluoro-4-{[(1S,2R)-2-(4-hydroxy-1-methyl-1H-pyrazol-5yl)cyclohexyl]oxy}-N-(pyrimidin-4-yl)benzenesulfonamide), a racemic mixture of M7 (5-chloro-2-fluoro-4-{[(1RS,2SR,4RS)-4-hydroxy-2-(1-methyl-1H-pyrazol-5yl)cyclohexyl]oxy}-N-(2-oxo-1,2-dihydropyrimidin-4-yl)benzenesulfonamide), a racemic mixture of M8 (5-chloro-2-fluoro-4-{[(1RS,2SR,4SR)-4-hydroxy-2-(1methyl-1*H*-pyrazol-5-yl)cyclohexyl]oxy}-*N*-(2-oxo-1,2-dihydropyrimidin-4-yl)benzenesulfonamide), a racemic mixture of M9 (5-chloro-2-fluoro-4-{[(1RS, 2SR,5RS)-5-hydroxy-2-(1-methyl-1*H*-pyrazol-5-yl)cyclohexyl]oxy}-*N*-(2-oxo-1, 2-dihydropyrimidin-4-yl)benzenesulfonamide), a racemic mixture of M10 (5-chloro-2fluoro-4-{[(1RS,2SR,5SR)-5-hydroxy-2-(1-methyl-1H-pyrazol-5-yl)cyclohexyl]oxy}-N-(2-oxo-1,2-dihydropyrimidin-4-yl)benzenesulfonamide), M11 (5-chloro-2-fluoro-4- $\{[(1S,2R)-2-(1H-pyrazol-5-yl)cyclohexyl]oxy\}-N-(pyrimidin-4-yl)benzenesulfo$ namide), M12 (5-chloro-2-fluoro-4-({(1S,2S)-2-[(4R)-4-hydroxy-1-methyl-5oxo-4,5-dihydro-1*H*-pyrazol-4-yl]cyclohexyl}oxy)-*N*-(pyrimidin-4-yl)benzenesulfonamide), and M13 (5-chloro-2-fluoro-4-({(1S,2S)-2-[(4S)-4-hydroxy-1methyl-5-oxo-4,5-dihydro-1*H*-pyrazol-4-yl]cyclohexyl}oxy)-*N*-(pyrimidin-4-yl)benzenesulfonamide) were synthesized at Daiichi Sankyo Co., Ltd. (Tokyo, Japan). [3H]DS-1971a (999 GBq/mmol, 37 MBq/ml in ethanol; radiochemical purity, 99%) (Fig. 1C) was purchased from Sekisui Medical Co., Ltd. (Tokyo, Japan). Pooled liver microsome and S9 from humans was purchased from XenoTech (Kansas City, KS). Potassium phosphate buffer (KPB; pH 7.4), and NADPHregenerating system solutions A and B were purchased from Corning (Woburn, MA). Fresh human blood and plasma were provided by Daiichi Sankyo Co., Ltd. (Tokyo, Japan). Dimethylacetamide (DMA) was obtained from Nacalai Tesque (Kyoto, Japan). Saline was obtained from Otsuka Pharmaceutical Factory, Inc. (Tokushima, Japan). PXB cells whose replacement index was around 90% were purchased from PhoenixBio (Hiroshima, Japan).

Animals

Animal experiments were approved by the Institutional Animal Care and Use Committee and performed in accordance with the animal welfare guidelines of Daiichi Sankyo Co., Ltd., which is accredited by AAALAC International. Male CB-17 SCID mice (10 weeks old) were obtained from Charles River Laboratories Japan, Inc. (Yokohama, Japan). Male PXB-mice (12–18 weeks old) whose replacement index was around 90% were purchased from PhoenixBio (Hiroshima, Japan).

PK Study of DS-1971a and M1 in SCID Mice and PXB-Mice

DS-1971a dissolved in vehicle (10% DMA and 90% saline) was intravenously and orally administered to fasted SCID mice and PXB-mice (N = 3) at a dose of 10 mg/5 ml/kg. M1 dissolved in the same vehicle was intravenously injected into fasted PXB-mice (N = 3) at a dose of 1 mg/5 ml/kg. Plasma samples were collected 0.083 and/or 0.25, 0.5, 1, 2, 4, 7, and 24 hours later. Extraction and liquid chromatography-tandem mass spectrometry (LC-MS/MS) measurement were performed as previously described (Asano et al., 2021). PK parameters [i.e., total plasma clearance and plasma distribution volume in a steady state (Vd_{ss}) after intravenous administration, C_{max} , time to reach the maximum plasma concentration (T_{max}) , apparent oral plasma clearance after oral administration, and AUC from the time of dosing to the time of the last observation after both intravenous and oral administration] were calculated based on a noncompartmental analysis technique using WinNonlin 6.3 (Certara, Princeton, NJ). Oral bioavailability of DS-1971a was calculated by dividing the AUC of DS-1971a after oral administration by that after intravenous administration and multiplying times 100. M1/P ratio was obtained by dividing the AUC of M1 by the AUC of DS-1971a after oral dosing.

Calculation of Dose-Normalized AUC of M1

Dose-normalized AUC in various species were calculated as follows.

Normalized AUC = observed AUC ×
$$\frac{\text{oral dose in humans } \div \text{ human BW}}{\text{oral dose in various species}}$$
 (1)

Here, oral dose in humans is 400 mg and human body weight (BW) represents BW of 70 kg for humans (Davies and Morris, 1993). Oral dose (3-10 mg/kg) and observed AUC in ICR mice, monkeys, rats, and dogs are cited from Asano et al. (2021, 2022), and those in SCID mice and PXB-mice are described in this paper.

Mass Balance Study of Radiolabeled DS-1971a in SCID Mice and PXB-Mice

[3 H]DS-1971a dissolved in vehicle (10% DMA and 90% saline) was orally administered to fasted SCID mice and PXB-mice (N=3) at a dose of 10 mg/5 ml/kg. Urine and feces were collected at 0 to 72 hours post-administration. Fifty microliters of urine or homogenized feces were mixed with 2 ml of tissue solubilizer Solvable (PerkinElmer, Waltham, MA) and 10 ml of the scintillator Hionic-fluor, and radioactivity was measured using a Tri-Carb 3110TR liquid scintillation counter (PerkinElmer). The excretion ratio of radioactivity in urine and feces (percentage of dose) was calculated by dividing the excreted radioactivity by the administered radioactivity.

Metabolite Detection in Plasma, Liver, Urine, and Feces of SCID Mice and PXB-Mice and Calculation of Urinary, Biliary, and Metabolic Clearance in PXB-Mice

After oral administration of [3 H]DS-1971a dissolved in vehicle (10% DMA and 90% saline) to fasted SCID mice and PXB-mice (N=3) at a dose of 10 mg/5 ml/kg, plasma and liver samples were collected from mice 1 hour later. In addition, urinary and fecal samples collected in the mass balance study described earlier were used for metabolite detection. Extraction and radio-HPLC analysis of collected plasma, liver, urine, and feces were performed as previously described (Asano et al., 2021). Based on the radiochromatograms of urinary and fecal extracts, the composition ratio and amounts of DS-1971a and M1 excreted into urine and feces were calculated. Throughout the manuscript, second subscripts "p" and "m" of abbreviations represent parent and metabolite, respectively. The hepatic biliary plasma clearance (CLbp and

CLbm), renal plasma clearance (CLrp and CLrm), and hepatic metabolic plasma clearance (CLmp and CLmm) of DS-1971a and M1 in PXB-mice were calculated as follows:

CLbp in PXB-mice (mL/min/kg) = Apparent oral plasma clearance of DS-1971a

$$\times \frac{\text{Amount of DS-1971a excreted in feces}}{\text{Dose}} \quad (2)$$

CLbm in PXB-mice (mL/min/kg) = Apparent oral plasma clearance of M1

$$\times \frac{\text{Amount of M1 excreted in feces}}{\text{Dose}}$$
 (3)

$$CLrp in PXB-mice (mL/min/kg) = \frac{Amount of DS-1971a excreted in urine}{Plasma AUC of DS-1971a}$$
(4)

$$CLrm in PXB-mice (mL/min/kg) = \frac{Amount of M1 excreted in urine}{Plasma AUC of M1}$$
 (5)

CLmp in PXB-mice (mL/min/kg)

=
$$Total \ plasma \ clearance \ of \ DS-1971a \ (CLp, total) - CLbp - CLrp$$
 (6)

$$CL$$
mm in PXB - mice (mL/min/kg)

=
$$Total \ plasma \ clearance \ of \ M1 \ (CLm, total) - CLbm - CLrm$$
 (7)

Note that biliary plasma clearance (*CL*bp and *CL*bm) were calculated based on previous reports (Yang et al., 2009) with some modifications, assuming that the fecal recovery of DS-1971a and M1 was equal to the biliary excretion because no further metabolism of either compound was confirmed when they were incubated for 24 hours with fecal homogenates from PXB-mice (data not shown), and both fraction absorbed (*F*a) and intestinal availability (*F*g) of DS-1971a in PXB-mice were calculated to be 1.0 (described in the Appendix). Total plasma clearance (*CL*p, total and *CL*m, total), which were determined in intravenous PK study of DS-1971a and M1, were assumed to be the sum of biliary plasma clearance (*CL*bp and *CL*bm), renal plasma clearance (*CL*rp and *CL*rm), and hepatic metabolic plasma clearance (*CL*mp and *CL*mm).

Human Clearance and Vd_{ss} Estimation by Single Species Allometric Scaling of PK Parameters in PXB-Mice

Based on a previous report (Sanoh et al., 2015), the plasma clearance (CLbp, CLbm, CLrm, CLmp, and CLmm) and plasma volume of distribution in a steady state (Vd_{ss}) of DS-1971a and M1 in humans were estimated using the following equations:

Human plasma
$$CL$$
s = PXB-mouse plasma CL s

$$\times (\text{human BW/PXB-mouse BW})^{0.814}$$
 (8)

Human plasma $Vd_{ss} = PXB$ -mouse plasma Vd_{ss}

$$\times (\text{human BW/PXB-mouse BW})^{0.9}$$
 (9)

Here, BW represents BW of 70 kg for humans (Davies and Morris, 1993) and 0.020 kg for PXB-mice (the measured value of the purchased PXB-mice).

Measurement of Concentration-Based Rb Values of DS-1971a and M1

DS-1971a or M1 (1 μ M) was incubated in human blood at 37°C for 5 minutes. The final concentration of organic solvent was less than 1.0%. Incubated human blood samples were centrifuged at 21,600 g for 3 minutes to separate supernatant plasma fractions. DS-1971a and M1 concentrations in the separated supernatant plasma samples from the incubated blood (referred to as "supernatant sample") were quantified by LC-MS/MS. Concentration-based blood to plasma ratios of DS-1971a (Rbp) and M1 (Rbm) were calculated by dividing the tested concentration (1 μ M) by that in "supernatant sample," assuming no degradation during incubation.

Calculation of Hepatic Availability of DS-1971a (Fh,p) and M1 (Fh,m) in PXB-Mice and Humans

Hepatic availability of DS-1971a and M1 (Fh,p and Fh,m) was calculated based on the following equations:

$$Fh, p = 1 - \frac{CLmp + CLbp}{Rbn \times Oh}$$
 (10)

$$Fh, p = 1 - \frac{CLmp + CLbp}{Rbp \times Qh}$$

$$Fh, m = 1 - \frac{CLmm + CLbm}{Rbm \times Qh}$$
(10)

where Qh represents hepatic blood flow of 90 ml/min/kg for PXB-mice and 21 ml/min/kg for humans (Davies and Morris, 1993) and hepatic plasma clearance was assumed to be the total of biliary (CLbp and CLbm) and metabolic plasma clearance (CLmp and CLmm).

Measurement of Liver KP value in PXB-Mice

PXB-mouse plasma and liver samples collected for the metabolite detection experiment were used for measurement of liver KP values. Based on the radioactive concentration of the plasma and liver homogenates measured using a Tri-Carb 3110TR liquid scintillation counter (PerkinElmer, Waltham, MA) and radiochromatograms obtained in the metabolite detection experiment, liverplasma partition coefficients of DS-1971a (Kphp) and M1 (Kphm) were calculated by dividing the liver concentrations of DS-1971a and M1 by their plasma concentrations, respectively.

Incubation of DS-1971a With HLM

DS-1971a (0.1, 0.3, 1, 3, 10, and 30 μ M) was incubated with HLM (0.5 mg protein/ml in 100 mM KPB, pH 7.4) in the presence of the NADPH-regenerating system at 37°C for 10 minutes in triplicate, and the concentrations of DS-1971a and M1 were determined by LC-MS/MS. The final concentration of organic solvent was less than 1.0%. The depletion rate for DS-1971a and formation rate for M1 were calculated by dividing the measured concentrations of both compounds by the protein concentration and incubation time using Microsoft Excel 2010. Based on the obtained depletion or formation rate for DS-1971a and M1, $K_{\rm m}$ and $V_{\rm max}$ were determined by a simple $E_{\rm max}$ model (model 101) using WinNonlin 6.3 based on the following equation:

$$E = \frac{E_{\text{max}} \times \mathbf{C}}{\mathbf{C} + EC_{50}} \tag{12}$$

where E, C, EC_{50} , and E_{max} represent metabolic reaction velocity, DS-1971a concentration, K_{m} , and V_{max} as per the Michaelis-Menten equation, respectively. Intrinsic clearance (CLint) for DS-1971a depletion and M1 formation was calculated by dividing the V_{max} values by the corresponding K_{m} values.

Protein Binding of DS-1971a in Human Liver Microsomes and S9

The free fractions of DS-1971a in human liver microsomes (0.5 mg/ml in 100 mM KPB, pH 7.4) and S9 (1 mg/ml in 100 mM KPB, pH 7.4) were determined in triplicate as fu,mic and fu,S9, respectively, based on the previously described method (Shinozuka et al., 2020).

Estimation of Fractional Metabolite Clearance and Plasma M1/P Ratio

Since no metabolism of DS-1971a into M1 in human intestinal microsomes and S9 was observed (data not shown), the M1-to-parent plasma AUC ratio (M1/P ratio) was calculated without consideration of gut metabolism based on the following equation (Nguyen et al., 2016; Obach et al., 2018):

$$\frac{AUCm}{AUCp} =$$

$$\frac{Fh, m \times fCL, m \times CLp, total}{Fh, p \times CLm, total} \times \frac{MW \text{ of metabolite } (M1 = 481)}{MW \text{ of parent } (DS-1971a = 465)}$$
 (13)

Here, AUCp, AUCm, Fh,p, Fh,m, CLp,total, CLm,total, and fCL,m represent plasma AUC of parent (DS-1971a), plasma AUC of metabolite (M1), hepatic availability of DS-1971a, hepatic availability of M1, total plasma clearance of parent (DS-1971a), total plasma clearance of metabolite (M1), and the fraction of parent (DS-1971a) clearance through hepatic metabolite (M1) formation, respectively. fCL,m in PXB-mice was obtained by measuring the total excretion (percentage of dose/100) of M1 into urine and feces after the oral administration of radiolabeled DS-1971a to PXB-mice. Meanwhile, fCL,m in humans was calculated as follows:

fCL, m in humans =
$$CL_{\text{uint}}$$
 by P450 for M1 formation in HLM \times mSF \div (CL_{uint} by P450 for DS $-$ 1971a depletion in HLM \times mSF

+
$$CL_{\text{uint}}$$
 by AO for DS $-$ 1971a depletion in human liver S9 \times sSF) (14)

Here, CL_{uint} represents unbound intrinsic clearance. CL_{uint} by P450 for M1 formation and CLuint by P450 for DS-1971a depletion were calculated by dividing respective $V_{\rm max}$ values by $K_{\rm m}$ values corrected for the fu,mic of the incubation. CLuint by AO was calculated by correcting the previously reported CLint, AO value for the fu,S9 of the incubation (Asano et al., 2021). mSF and sSF represent physiologic scaling factors for human liver microsomal proteins (46 mg protein/g liver) (Zhou et al., 2002) and S9 proteins (121 mg protein/g liver) (Houston and Galetin, 2008), respectively. Note that biliary and urinary clearance of DS-1971a were not taken into account for this calculation because of the negligible (less than 1%) excretion into bile and urine as an intact form in PXB-mouse mass balance study (described later).

Simulation of PK Profiles of DS-1971a and M1 in PXB-mice and Humans Using Dynamic PBPK Models

The minimal PBPK model (Fig. 2) was designed based on a previous report (Ito et al., 2020) with some modifications to simulate PK profiles after oral administration of DS-1971a at a dose of 0.2 mg for PXB-mice and 400 mg for humans. Development of PBPK models (Fig. 2) and simulations were all performed by the Phoenix model using WinNonlin 6.3. PBPK parameters, mass balance formulas and equations (eq. (15) - (31)), and calculation of simulated exposure were described in the Appendix.

Membrane Permeability of DS-1971a in MDCK II Cells, and Estimation of Absorption Constants-1 and -2 in Humans

Apparent permeability (P_{app}) of DS-1971a was assayed in sextuplicate using MDCK II cells, as previously reported (Furukawa et al., 2020). Absorption constants-1 and -2 (ka1 and ka2) in humans were estimated based on the following equation:

$$ka1 \text{ and } ka2 \text{ in humans} = \frac{2 \times P_{\text{eff}}}{R} \text{ (Yu and Amidon, 1999)}$$
 (32)

where R means small intestinal radius (1.75 cm) and $P_{\rm eff}$ represents effective permeability, which was calculated from $P_{\rm app}$ by SimCYPTM (Certara, Princeton, NJ) based on the following equation (Tchaparian et al., 2008):

$$log10 [P_{eff} in humans] = 1.0117 \times log10 [P_{app} in MDCK II cells] - 0.912$$
 (33)

Determination of LogD Values of DS-1971a, M1, M2, and M4

LogD values of DS-1971, M1, M2, or M4 were assayed as previously reported (Nagata et al., 2009). In brief, DS-1971a, M1, M2, or M4 (1 μ M) was added to 1-octanol-saturated PBS, and the solutions were stirred for 30 minutes at room temperature in duplicate. After centrifugation, the upper layer (1-octanol) and the lower layer (PBS) were collected, and the concentrations in each layer were determined by LC-MS/MS. The distribution coefficient (LogD) was calculated from the following equation:

$$LogD = log 10 [(peak area in the 1 - octanol layer)/(peak area in the PBS layer)]$$
 (34)

Metabolite Detection in PXB Cells

[3 H]DS-1971a (1 μ M) was incubated in triplicate with PXB cells (8×10 5 cells/ml) at 37°C and 5% CO2 for 2 hours. After the reaction had been terminated by the addition of 2 volumes of acetonitrile, the mixture was centrifuged at 21,500g for 5 minutes. The resulting supernatant was analyzed by radio-HPLC.

Radio-HPLC and LC-MS/MS Analyses

Analyses were performed as previously described (Asano et al., 2021).

Results

Metabolite Profiling in SCID and PXB-Mice

To identify DS-1971a metabolites formed in SCID and PXB-mice, plasma (at 1 hour), urinary (0–24 hours), and fecal (0–24 hours) samples were collected after the oral administration of [³H]DS-1971a (10 mg/kg), and metabolite profiling was performed. Radiochromatograms of the plasma, urinary, and fecal extracts indicated that major metabolites in

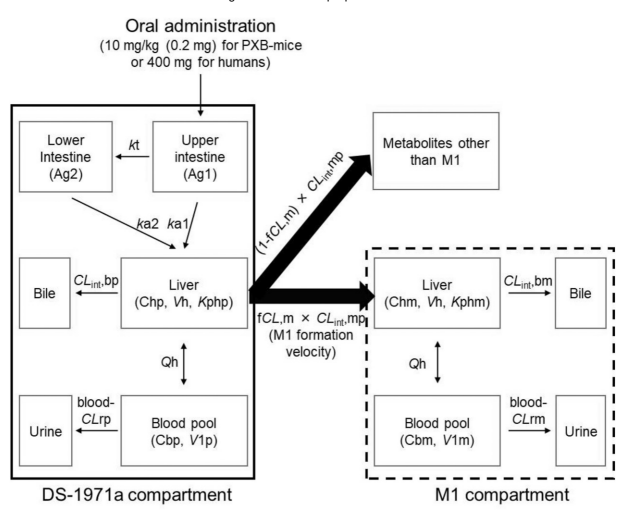


Fig. 2. The structure of the PBPK model of DS-1971 and M1 for PXB-mice and humans. Ag1, drug amount in the upper intestine; Ag2, drug amount in the lower intestine; blood-*CL*rm, renal blood clearance of M1; blood-*CL*rp, renal blood clearance of DS-1971a; Cbm, blood concentration of M1; Cbp, blood concentration of DS-1971a; Chm, hepatic concentration of M1; Chp, hepatic concentration of DS-1971a; CL_{int}, bm, biliary blood intrinsic clearance of M1; Cl_{int}, bp, biliary blood intrinsic clearance of DS-1971a; Cl_{int}, mp, metabolic blood intrinsic clearance of DS-1971a; Cl_{int}, fraction of the clearance of DS-1971a that yields M1; ka1, absorption constant-1; ka2, absorption constant-2; Kphm, liver to plasma ratio of M1; Kphp, liver to plasma ratio of DS-1971a; kt, transit constant; Qh, liver blood flow; Vh, liver volume; V1m, blood-based extrahepatic distribution volume of M1; V1p, blood-based extrahepatic distribution volume of DS-1971a.

SCID mice were AO-related metabolites, M4, M7, and M8, while those in PXB-mice were M1 and M2 (Fig. 3). The results indicated that the metabolite profile in PXB-mice was very similar to that in humans (Asano et al., 2022) but markedly distinct from that in SCID mice.

The PK profiles of major metabolites were then investigated to evaluate the exposure level of M1 and the AUC ratio of M1 to DS-1971a (M1/P ratio) in both types of mice. Based on the PK parameters (Supplementary Table 1), which were calculated from PK profiles (Supplementary Fig. 1), M1/P ratios in SCID and PXB-mice were determined to be 0.0 and 0.69, respectively, suggesting that the M1/P ratio in PXB-mice (0.69) was approximately half that in humans (1.6) (Table 1; Fig. 5). In addition, dose-normalized M1 AUC in PXB-mice (1984 ng·h/ml) was approximately one-ninth that in humans (18,400 ng·h/ml) (Table 1).

Determination of Plasma PK Parameters and Fractional Metabolite Clearance in PXB-Mice for PBPK Modeling

The obtained results showing that the M1/P ratio and M1 AUC in PXB-mice were much smaller than those in humans indicated that static and dynamic PBPK models were necessary for the quantitative

prediction of the M1/P ratio and/or M1 AUC. Therefore, various PK parameters, including plasma Vd_{ss} , total, apparent oral, metabolic, renal and biliary plasma clearance, oral bioavailability, the fraction metabolized into M1 (fCL,m), and liver-to-plasma ratio (Kph) of DS-1971a and/or M1, were obtained as follows and are summarized in Supplementary Table 3 to perform PBPK modeling.

Plasma Vd_{ss}, Total Plasma Clearance, Apparent Oral Plasma Clearance, and Oral Bioavailability in PXB-Mice. After DS-1971a or M1 was intravenously injected into PXB-mice, plasma samples were successively collected up to 24 hours later and analyzed by LC-MS/MS. Using the noncompartmental method, plasma Vd_{ss} and total plasma clearance (CLp,total and CLm,total) of DS-1971a and M1 were determined to be 0.34 and 0.23 L/kg and 20 and 11 ml/min/kg, respectively (Supplementary Table 2). After oral administration of DS-1971a, plasma concentrations of DS-1971a and M1 were quantified in the same way as for intravenous samples. The oral bioavailability of DS-1971a was calculated to be 60% in PXB-mice by dividing the oral AUC by the intravenous AUC. The apparent oral plasma clearance (CL/F) of DS-1971a and M1 were determined to be 33 and 50 ml/min/kg, respectively.

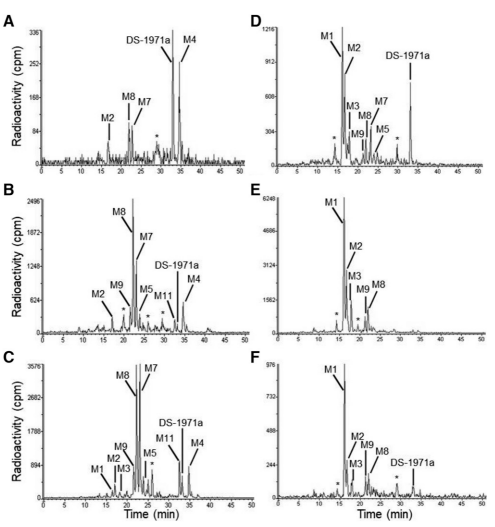


Fig. 3. Radiochromatograms of the plasma, urine, and feces from SCID mice and PXB-mice after oral dosing of radiolabeled DS-1971a at 10 mg/kg in the fasted condition. Plasma (A, D), urine (B, E), and feces (C, F) were obtained from SCID mice (A-C) and PXB-mice (D-F) at 1 h, 0-24 h, and 0-24 h, respectively, after [3H]DS-1971a administration and were analyzed using radio-HPLC. The radioactive peaks of the parent and metabolites were identified by comparing their retention times with those of authentic standards for the parent and each metabolite. * indicates unknown metabolites.

Metabolic, Renal, and Biliary Plasma Clearance in PXB-Mice.

The urinary and fecal recovery of radioactivity 0 to 24 hours after the oral administration of [3H]DS-1971a to PXB-mice at 10 mg/kg was determined to be 47% and 39%, respectively (Table 2). The 0- to 24-hour recovery of DS-1971a in urine and feces was calculated to be 0.42% and 0.82%, while that of M1 was 15% and 9.2%, respectively, by

multiplying the urinary and fecal recovery by the composition ratio of DS-1971a and M1 in the urinary and fecal radiochromatograms. The renal and biliary plasma clearance of DS-1971a (CLrp and CLbp) were determined to be 0.14 and 0.27 ml/min/kg, while for M1 (CLrm and CLbm) they were 7.1 and 4.6 ml/min/kg, respectively, based on Eq. (2) to (5). The metabolic plasma clearance of DS-1971a and M1 (CLmp

Observed plasma M1/P ratios in experimental animals and humans.

Species	Dose	DS-1971a plasma AUC (ng·h/ml)	M1 plasma AUC (ng·h/ml)	M1 dose-normalized plasma AUC ^d (ng·h/ml)	Plasma M1/P ratio
ICR mice ^a	10 mg/kg	395 ± 148	1.7 ± 0.12	0.97 ± 0.11	0.0 ± 0.0
SCID mice ^b		1736 ± 191	5.7 ± 1.6	3.3 ± 0.91	0.0 ± 0.0
PXB-mice ^b		5113 ± 548	3472 ± 253	1984 ± 145	0.69 ± 0.10
Rats ^a		3234 ± 1447	478 ± 136	273 ± 78	0.16 ± 0.038
Dogs ^a		$42,949 \pm 17,096$	0.0 ± 0.0	0.0 ± 0.0	0.0 ± 0.0
Monkeys ^a	3 mg/kg	2485 ± 473	288 ± 20	165 ± 11	0.12 ± 0.029
Humans ^c	400 mg	$12,500 \pm 3700$	$18,400 \pm 3790$		1.6 ± 0.55

^aPK parameters (mean ± SD, n = 3) are cited from Asano et al. (2021, 2022) and modified.

^dEq. (1).

^bData represent the mean ± SD of 3 mice.

^cPK parameters (mean ± SD, n = 6) are cited from Asano et al. (2021, 2022) and modified.

TABLE 2

The recovery of the radioactivity in urine and feces of SCID mice and PXB-mice after oral administration of [3H]DS-1971a at 10 mg/kg in the fasted

	Cumul	Cumulative radioactivity excretion (% of dose)				
	SCID mice		PXB-mice			
Excretion pathway	0-24 h	0-72 h	0-24 h	0–72 h		
Urine Feces Total (urine + feces)	1.6 ± 1.5 87 ± 11 88 ± 9.5	2.6 ± 1.3 91 ± 8.2 93 ± 7.0	47 ± 6.7 39 ± 2.0 86 ± 6.9	51 ± 5.9 45 ± 0.51 96 ± 5.7		

Urine and feces were collected up to 72 hours after oral administration of $[^3H]DS-1971a$ (10 mg/kg). Each value represents the mean \pm SD of 3 mice.

and *CL*mm) were calculated to be 20 and 0.0 ml/min/kg, respectively, by Eq. (6) and (7).

fCL,m in PXB-Mice. The urinary and fecal recovery of radioactivity during 0 to 72 hours after the oral administration of [³H]DS-1971a to PXB-mice at 10 mg/kg was determined to be 51% and 45%, respectively, indicating that most of the administered DS-1971a was excreted within 72 hours (Table 2). The urinary and fecal recovery of M1 during 0 to 72 hours was calculated to be 15% and 9.6%, respectively, by multiplying each recovery by the composition ratio of M1 in the radiochromatogram. The total recovery of M1 was calculated to be 25% up to 72 hours after dosing, indicating that the fraction metabolized into M1, fCL,m, was 0.25.

Kph in PXB-Mice. Plasma and liver were collected 1 hour after the oral administration of [³H]DS-1971a to PXB-mice at 10 mg/kg, and the radioactivity was quantified using a liquid scintillation counter and analyzed by radio-HPLC. **Kph** of DS-1971a and M1 (**Kphp** and **Kphm**) were calculated to be 3.2 and 1.1, respectively, by dividing the liver concentrations of DS-1971a and M1 by their plasma concentrations.

Development of a PBPK Model for PXB-Mice

Although the static model provides only M/P ratios but not metabolite AUC, PBPK models can estimate both of them. Before PBPK simulation for humans, we evaluated the validity of the constructed PBPK model (Fig. 2) by comparing the simulated and observed PK profiles in PXB-mice.

Detailed parameters for PBPK modeling in PXB-mice (Supplementary Table 4) were experimentally obtained, taken from the literature (Davies and Morris, 1993; Shinozuka et al., 2020; Asano et al., 2021, 2022), or calculated based on the observed PK parameters of DS-1971a and M1 (Supplementary Table 3). Using a PBPK model (Fig. 2) inputted with these PBPK parameters obtained with PXB-mice (Supplementary Table 4), the PK profiles of DS-1971a and M1 after the oral administration of DS-1971a at 10 mg/kg were simulated. As shown in Supplementary Fig. 2, the simulated PK profiles of DS-1971a and M1 in PXB-mice were relatively close to the observed ones. In addition, the simulated $C_{\rm max}$ and AUC values of DS-1971a and M1 were within 1.5-fold of observed data (Supplementary Table 5), and the estimated M1/P ratio was almost the same as the actual one. Thus, the PBPK model for PXB-mice was found to be established, and the disposition of DS-1971a in PXB-mice was adequately expressed by the model.

Protein Binding of DS-1971a in Human Liver Microsomes and S9

The free fractions of DS-1971a in human liver microsomes and S9 (fu,mic and fu,S9) were determined to be 0.70 \pm 0.06 and 0.77 \pm 0.06, respectively. These values were used to calculate unbound intrinsic clearance ($CL_{\rm uint,P450}$) and $CL_{\rm uint,AO}$) in vitro.

Enzyme Kinetics for DS-1971a Depletion and M1 Formation in Humans for the Determination of Human f CL_{γ} m

One of the critical parameters for estimating metabolite exposure in the static and dynamic PBPK models in PXB-mice and humans is the fraction metabolized into M1, fCL,m. To obtain this value in humans, DS-1971a depletion and M1 formation were assessed using HLM. Note that intestinal DS-1971a depletion and M1 formation were ignored because no depletion or M1 formation was observed when DS-1971a was incubated with human intestinal microsomes (data not shown).

After a 10-minute incubation of DS-1971a in HLM with the NADPH-regenerating system at concentrations ranging from 0.1 to 30 μ M, DS-1971a and M1 concentrations were quantified by LC-MS/MS. Based on the depletion and formation rates of DS-1971a and M1 at various concentrations, $K_{\rm m}$ and $V_{\rm max}$ values were determined to be 4.2 ± 2.8 μ M and 401 ± 85 pmol/min/mg microsomal protein for DS-1971a depletion and 4.4 ± 0.5 μ M and 228 ± 6.6 pmol/min/mg microsomal protein for M1 formation, respectively (Fig. 4). Considering that $CL_{\rm uint,P450}$ for DS-1971a depletion and M1 formation were calculated to be 137 and 73 μ L/min/mg microsomal protein, respectively, and $CL_{\rm uint,P450}$ for DS-1971a depletion was determined to be 3.8 μ L/min/mg S9 protein based on the previously reported $CL_{\rm int,AO}$ value of 2.9 μ L/min/mg S9 protein (Asano et al., 2021), f $CL_{\rm m}$ in humans was estimated to be 0.50 (Table 3).

Membrane Permeability of DS-1971a in MDCK II Cells and Estimation of Absorption Constants-1 and -2 in Humans

Apparent permeability ($P_{\rm app}$) of DS-1971a was determined to be $6.4 \pm 3.8 \times 10^{-6}$ cm/s. Based on the $P_{\rm app}$ value, effective permeability ($P_{\rm eff}$) and absorption constants (ka1 and ka2) were calculated to be 0.80×10^{-4} cm/s and 0.33/h, respectively, based on Eq. (33) and (32).

Prediction of M1 Exposure in Humans Based on Static and PBPK Models

Since various clearance values (renal, biliary, and metabolic clearance) of DS-1971a and M1 in humans were required for both the static and dynamic PBPK models, they were estimated from those obtained with PXB-mice using the allometric scaling method (Table 4). Obtained renal, biliary, and metabolic plasma clearance were 0.0, 0.1, and 4.3 for DS-1971a and 1.6, 1.0, and 0.0 ml/min/kg for M1, respectively. These results indicated that most of the administered DS-1971a was eliminated by hepatic metabolism (i.e., metabolic clearance > urinary and biliary clearance of DS-1971a), while M1 produced in the liver was excreted into bile and urine without further metabolism in humans (i.e., metabolic clearance of M1 = 0.0) likely due to its hydrophilicity (LogD value = -0.8), which is supported by there being no declining trend when M1 was incubated with human liver microsomes or S9 (Asano et al., 2022). Additional PBPK parameters, including intestinal availability, distribution volume, unbound fraction in plasma, liver to plasma ratio, and physiologic parameters (liver blood flow and volume), were calculated, experimentally obtained, estimated, or adopted from those in PXB-mice or taken from the literature (Davies and Morris, 1993; Shinozuka et al., 2020; Asano et al., 2021, 2022), and they are summarized in Table 5.

Based on the estimated human PK parameters and fCL,m, the M1/P ratio in humans was calculated as 1.3 (Table 4; Fig. 5) by the static model. This indicated that the static model gave a more accurate M1/P value than that observed in PXB-mice (0.69) since that in humans was 1.6 (Table 1; Fig. 5).

We further tried to predict the human PK profiles of DS-1971a and M1 using the dynamic PBPK model whose important parameters (Table 5) were derived from the estimated human PBPK parameters (Table 4). Although the absorption process in actual PK profiles of DS-1971a and M1 was more sustained and delayed than that in their

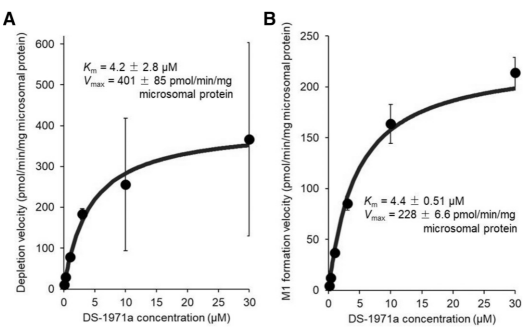


Fig. 4. $K_{\rm m}$ and $V_{\rm max}$ values for DS-1971a depletion and M1 formation in HLM. DS-1971a was incubated in HLM (0.5 mg/ml) with the NADPH-regenerating system at concentrations ranging from 0.1 to 30 μ M, and DS-1971a and M1 concentrations were determined after 10 minutes incubation. The rates of DS-1971a depletion (A) and M1 formation (B) were calculated based on the plots. Each plot represents the mean velocity \pm SD of three incubations. Using WinNonlin 6.3, $K_{\rm m}$ and $V_{\rm max}$ values were determined by fitting to the simple $E_{\rm max}$ model (model 101) [$E = E_{\rm max} \times C \div (EC_{50} + C)$] (Eq. (12)), where $E \in E_{\rm max}$, and $E \in E_{50}$ represent metabolic reaction velocity, DS-1971a concentration, $V_{\rm max}$, and $E \in E_{50}$ max as per the Michaelis–Menten equation, respectively.

simulated PK profiles (Fig. 6), the $C_{\rm max}$ values of DS-1971a and M1 (3082 and 3429 ng/ml, respectively) in humans predicted by the PBPK modeling were similar to the observed ones (2950 and 3180 ng/ml, respectively) and the estimated AUC levels of DS-1971a and M1 (13,479 and 17,116 ng·h/ml, respectively) were very close to the observed values (12,500 and 18,400 ng·h/ml, respectively) (Table 6). The M1/P ratio obtained by the dynamic PBPK model (1.3) was the same as that obtained by the static model (1.3) (Fig. 5).

Determination of LogD Values of DS-1971a, M1, M2, and M4

To consider physicochemical properties, LogD values of DS-1971a, M1, M2, and M4 at pH 7.4 were determined to be 0.8, -0.8, -0.5, and 1.5, respectively, indicating that both M1 and M2, the monooxides at the cyclohexane ring (Fig. 1), are very polar, while M4 possessing the monooxidized pyrimidine ring (Fig. 1) is more lipophilic than the intact form.

Discussion

In this study, we introduced a practical evaluation method to predict the exposure level of a human disproportionate metabolite of DS-1971a using PXB-mice and PBPK techniques. After the PK profiles of DS-1971a and M1 in PXB-mice were evaluated, a PBPK model for PXB-mice was developed based on the PK parameters obtained with PXB-mice to verify the model. Finally, human PK parameters were estimated from those in PXB-mice and inputted into the PBPK model for humans, leading to successful prediction of the metabolite exposure.

Drastic differences in the metabolite profiles between PXB-mice and SCID mice suggested the usefulness of PXB-mice for qualitative estimation of human major metabolites of mixed AO and P450 substrates. After the oral administration of DS-1971a, the most and second most abundant metabolites in PXB-mice were M1 and M2, respectively, both of which corresponded to those in humans produced by P450 (Asano et al., 2022), while major metabolites in SCID mice were AO-related metabolites, M4, M7, and M8 (Fig. 3; Supplementary Fig. 1; Supplementary Table 1). The PXB-mouse was the only animal model whose metabolite profile of DS-1971a was similar to that in humans (Asano et al., 2022), considering that the predominant metabolites in male ICR mouse, monkey, rat, and dog plasma were molecules other than M1, namely, M4, M2, M5, and M11, respectively (Asano et al., 2021; and data not shown).

TABLE 3 Calculation of fCL,m in humans.

		Human liver fraction	
Evaluation items	Parameter	Microsomes	S9
DS-1971a depletion	$CL_{\text{uint,P450}}$ (μ L/min/mg protein)	137	NA
•	$CL_{\text{uint,AO}}$ (μ L/min/mg protein)	NA	3.8
M1 formation	$CL_{\text{min}, P450}$ (μ L/min/mg protein)	73	NA
_	Protein abundance (mg protein/g liver)	46	121
The fraction of clearance of DS-1971a for M1 formation	fCL,m ^a (ratio)	0.50^{a}	

 $CL_{\text{uint,P450}}$ values were obtained by dividing the mean of V_{max} values by the mean of Km values and $f_{\text{u,mic}}$ obtained from 3 incubations. $CL_{\text{uint,AO}}$ was calculated by dividing $CL_{\text{int,AO}}$, which is cited from Asano et al. (2021), by $f_{\text{u,S9}}$. Protein abundance are cited from Zhou et al. (2002), and Houston and Galetin (2008).

NA, not applicable

Estimation of PBPK parameters in humans using the allometric scaling method and M1/P ratio calculation based on the static model

د	5	,	ε		i i Mada
Farameters	DS-19/1a	MI	Source	Static model	PBPK model
Molecular weight	465	481	T	X	X
Plasma distribution volume at steady state: Vd_{ss}^{a} (L/kg)	0.15^{a}	0.10^{a}	Allometry ^a		
Detail Blood-based extrahepatic distribution	13^b	6.8^{b}	Calculated b		×
volume: $V1p^b$ or $V1m^b$ (L)					
Liver volume: Vh (L)	1.7		Reported (Davies and Morris, 1993)		×
Liver to plasma ratio: Kp	3.2	1.1	Measured		×
fCL,m (ratio)	0.50	I	Calculated (Table 3)	×	×
Total plasma clearance: ^c CLp,total or ^c CLm,total (ml/min/kg)	4.4^{c}	2.6^c	Allometry ^c	X	
Detail Metabolic plasma clearance: CLmp ^c	4.3^c	0.0^c		×	×
or CLmm ^c (ml/min/kg)					
Biliary plasma clearance: CLbpc or	0.059^c	1.0^{c}		X	×
CLbm' (ml/min/kg)					
Renal plasma clearance: CLrp ^c or	0.030^c	1.6^c			×
$CLrm^{c}$ (ml/min/kg)					
Hepatic availability: Fh,p^d or Fh,m^e	0.62^{d}	0.92^e	Calculated ^{d,e}	X	
Plasma MI/P ratio ^f by static model	1.3^{f}		Calculated f	×	

X, used.
^aEq. (9).
^bEq. (31).
^cEq. (8).
^dEq. (10).
^eEq. (10).
^eEq. (11).
^fEq. (13).

The PXB-mouse was also found to be a predictive animal model of human disproportionate metabolite formation by CYP2C8. Our recent work demonstrated that humans possess much higher activity for M1 formation, which is predominantly mediated by CYP2C8, than any other animals (ICR mice, rats, dogs, and monkeys) (Asano et al., 2022). Even though monkeys possess CYP2C8 (previously termed CYP2C20) with high amino acid sequence homology (92%) to human CYP2C8, much lower activity for 6α-hydroxy paclitaxel formation by CYP2C8 in monkeys than in humans is also known (Yoda et al., 2012; Emoto et al., 2013). Owing to such species dependence in CYP2C8 metabolism, M1 exposure in humans was not covered by experiments with safety animals, leading to M1 becoming a human disproportionate metabolite (Asano et al., 2022). The highest exposure to M1 of all the DS-1971a metabolites in PXB-mice (Fig. 3; Supplementary Fig. 1; Supplementary Table 1) indicated the dominant contribution of CYP2C8 to DS-1971a metabolism in these mice. Given the additional fact that 3-hydroxydesloratadine-glucronide, which is another human disproportionate metabolite produced sequentially by UGT2B10, CYP2C8, and then UGTs (Kazumi et al., 2015), was also detected as the predominant metabolite in the plasma of PXB-mice after oral administration of desloratadine (Kato et al., 2020), PXB-mice can provide reliable qualitative information for CYP2C8-mediated formation of human disproportionate metabolites of mixed non-P450 and P450 substrates.

Meanwhile, from a quantitative perspective, the underestimation of the human M1 to DS-1971a AUC ratio (M1/P ratio) in PXB-mice deserves attention, although the M1/P ratio in PXB-mice (0.69) was closer to that in humans (1.6) than those in ICR mice (0.0), SCID mice (0.0), rats (0.16), dogs (0.0), and monkeys (0.12) (Table 1; Fig. 5). Underestimation of the M/P ratio in PXB-mice was also observed with various compounds and their metabolites: AUC ratios of troglitazone and its sulfate, of zaleplon and its 5-oxo metabolite, and of GDC-0834 and its hydrolyzed metabolite in humanized mice were 0.5 (Schulz-Utermoehl et al., 2012), 1.7 (Tanoue et al., 2013), and 0.7 (Liu et al., 2011), respectively, and they were lower than those in humans [6.8-8.1 (Young et al., 1998), 2.9 (Smith and Obach, 2009), and an incalculably large value due to no exposure to the parent (Liu et al., 2011), respectively]. Though there is a possibility that innate functions of tissues other than liver (e.g., kidney) in PXB-mice might make a difference in M1/P AUC ratios between PXB-mice and humans, we found that the leading reason for the smaller M1/P AUC ratios in PXB-mice than in humans is the smaller fraction of clearance of DS-1971a for metabolite (M1) formation (fCL,m) in PXB-mice than in humans. In fact, the fCL,m in PXB-mice (0.25) (Supplementary Table 4) was actually half that in humans (0.50) (Table 4), probably due to metabolism by remaining mouse hepatocytes in PXB-mice (Kamimura et al., 2010). This assumption is supported by more abundant formation of M4 than M1 (Supplementary Fig. 3A) and the fCL,m value of 0.27 (data not shown) in hepatocytes from PXB-mice (PXB cells). Although there was negligible M4 in the plasma, urine, and feces from PXB-mice (Fig. 3), M4 was detected in the liver from PXB-mice (Supplementary Fig. 3B), indicating that M4, which was more hydrophobic (LogD = 1.5) than the intact form and M1 (LogD = 0.8 and -0.8), was likely metabolized into further metabolites including M8 and M9 due to its poor metabolic stability (Asano et al., 2022). The slight difference of M1/P ratios between PXB-mice (0.69) and humans (1.6) directed us to the additional research on estimation of the human M1/P ratio by static and dynamic PBPK methods.

Prior to the estimation of the M1/P ratio in humans, a robust PBPK model for PXB-mice (Fig. 2) was constructed. In this research, we focused on the PK parameters of $C_{\rm max}$ and AUC because these parameters are used as criteria for disproportionate metabolite and safety

TABLE 5
Inputted parameters in the PBPK model for humans.

Inputted parameter	Explanation	Value	Unit	Source
Dose	Clinical dose	400	mg	Reported (Asano et al., 2022)
Fa	Fraction absorbed	1.0		Reported (Asano et al., 2021)
Fg	Intestinal availability	1.0		Calculated
Tlag	Residence time in the stomach in a fed condition	1.3	h	Reported (Kenyon et al., 1995)
Ag1	Drug amount in upper intestine	Variable	mg	· —
Ag2	Drug amount in lower intestine	Variable	mg	
Vh	Liver volume	1.7	L	Reported (Davies and Morris, 1993)
Qh	Liver blood flow	21	ml/min/kg	, ,
ka1 ^a	Absorption constant in upper intestine	0.33^{a}	/h	Calculated ^a
ka2 ^a	Absorption constant in lower intestine	0.33^{a}	/h	Calculated ^a
kt ^b	Transit constant in gastrointestinal tract	0.57^{b}	/h	Calculated ^b
MWp and MWm	Molecular weight	465, 481	g/mole	Calculated
Rbp and Rbm	Blood-to-plasma concentration ratio of DS- 1971a and M1	0.55, 0.56	_	Measured
fup and fum	Unbound fraction of DS-1971a and M1 in plasma	0.010, 0.020	_	Reported (Shinozuka et al., 2020; Asano et al., 2022)
Chp and Chm	Hepatic concentration of DS-1971a and M1	Variable	ng/ml	· <u> </u>
Kphp and Kphm	Liver-to-plasma ratio of DS-1971a and M1	3.2, 1.1	_	Measured
Cbp and Cbm	Blood concentration of DS-1971a and M1	Variable	ng/ml	_
$V1\hat{p}$ and $V1m$	Blood-based extrahepatic distribution volume of DS-1971a and M1	13, 9.8	L	Calculated (Table 4)
$CL_{\rm int}$, bp ^c and $CL_{\rm int}$, bm ^d	Biliary blood intrinsic clearance of DS- 1971a and M1	$25,^{c} 229^{d}$	L/h	Calculated ^{c,d}
blood-CLrp ^e and -CLrm f	Renal blood clearance of DS-1971a and M1	$0.23^{e}_{,}$ $12^{f}_{,}$	L/h	Calculated ^{e,f}
f <i>CL</i> ,m	Fraction of the clearance of DS-1971a that yields M1	0.50	_	Calculated (Table 3)
$CL_{\rm int}, { m mp}^g$	Metabolic blood intrinsic clearance of DS- 1971a	2923 ^g	L/h	Calculated ^g

^aEq. (32).

assessment in the US Food and Drug Administration guidance for industry on safety testing of drug metabolites (US Food and Drug Administration, 2020) and/or International Conference on Harmonization (2010) guidance. Since the simulated $C_{\rm max}$, AUC, and M1/P ratios in

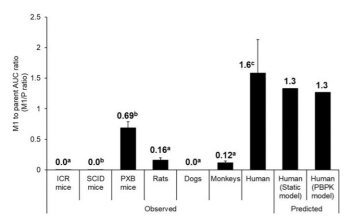


Fig. 5. Observed and predicted plasma AUC ratios of M1 to DS-1971a in various species. a, c: M1/P ratios (mean \pm SD of n = 3^a or 6^c) are cited and modified from Asano et al. (2021, 2022). b: Data represent the mean \pm SD (n = 3). Static model-based prediction was performed according to Eq. (13). PBPK model-based estimation was performed by dividing the predicted M1 AUC by the predicted DS-1971a AUC.

PXB-mice were close to the observed values (Supplementary Table 5), we moved forward with human PK prediction.

The human PBPK model was developed based on the same model structure (Fig. 2) as that for PXB-mice with the replacement of parameters in PXB-mice by those in humans, enabling the prediction of exposure to DS-1971a and M1 in humans. Important PK parameters in humans such as Vd_{ss} and various clearance (Table 4) were estimated by the allometric scaling of PK parameters in PXB-mice as the only animal model showing as high exposure to M1 as humans (Fig. 3). The human PBPK model, in which various parameters (Table 5) converted from those in Table 4 were inputted, successfully provided us with $C_{\rm max}$ and AUC (3082 ng/ml and 3429 ng/ml, and 13,479 and 17,116 ng·h/ml) of DS-1971a and M1, which were very close to the observed values (2950 and 3180 ng/ml, and 12,500 and 18,400 ng·h/ml) (Table 6). Meanwhile, the simulated PK profile lacked the prolonged absorption profile of the actual one (Fig. 6.). A potential reason for this is the delayed absorption in the fed condition because enterohepatic circulation of DS-1971a was not observed in mice and monkeys (Asano et al., 2021), and DS-1971a was administered to humans after ingestion of high-calorie food. Food has a strong influence on absorption process of drugs based on previous research reporting that gastric transit time was prolonged up to 5 hours by the presence of food in the gastrointestinal tract (Weitschies et al., 2010). The lack of adequate simulation of the absorption process under fed conditions is

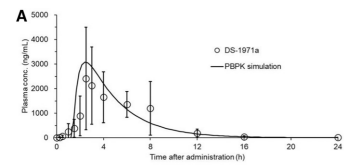
^bEq. (30).

Eq. (26).

^dEq. (25).

^eEq. (23).

^fEq. (24). ^gEq. (27).



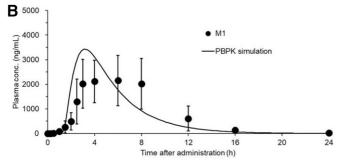


Fig. 6. The predicted plasma PK profiles of DS-1971a (A) and M1 (B) in humans after oral administration at 400 mg in the fed condition using the dynamic PBPK model. PBPK modeling was performed based on the customized model (Fig. 2) with the parameters shown in Table 5 using WinNonlin 6.3. Open and closed circles represent the observed plasma concentrations of DS-1971a and M1 (mean ± SD of six humans), respectively (Asano et al., 2022). Solid line indicates the simulated PK profiles of DS-1971a and M1 by the dynamic PBPK model.

a limitation of this research, as illustrated by the better simulated PK results using optimized absorption and transit constants (Supplementary Fig. 4). Another limitation was the lack of inclusion of transporters in the model. Although we quantified various clearance based on PK results, the contribution of transporters to these parameters was not estimated.

Both static and dynamic models provided us with the same M1/P ratio (1.3), which is close to the observed value (1.6). Although the static model appears to be more useful due to the smaller number of PK parameters required for the calculation, the accuracy of the parameters should be examined using a dynamic model. For example, the plasma $C_{\rm max}$ and AUC of DS-1971a and M1 simulated by the dynamic model were comparable to those observed (Supplementary Fig. 2; Supplementary Table 5), thereby confirming the validity of the obtained parameters, such as hepatic and systemic clearance, which are also used to perform static calculation. Therefore, it is better to use both static and dynamic models in combination to reproduce the observed data, and which method is selected would depend on a case-by-case basis. Although the construction of a dynamic model requires numerous parameters, a dynamic model has a wide range of

applications, such as in drug-drug interaction simulation (Yamada et al., 2020), pharmacokinetics-pharmacodynamics (Loisios-Konstantinidis et al., 2020), and model-informed drug discovery and development (Marshall et al., 2016).

In conclusion, we have demonstrated that PXB-mice are useful to predict the PK of and exposure to human disproportionate metabolites generated by CYP2C8-mediated primary metabolism. Although the metabolite profiles of DS-1971a and M1 in PXB-mice were very similar to those in humans, the observed M1/P ratio in PXB-mice underestimated that in humans. The present results suggest that PBPK modeling based on not only estimated human PK parameters obtained with PXB-mice but also metabolite formation kinetics in human liver fractions is a practical method to predict the human PK profile of M1 and its exposure level in a quantitative manner.

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Authorship Contributions

Participated in research design: Asano Conducted experiments: Asano, Hamada

Performed data analysis: Asano, Nakamura, Nishiya, Yahara

Wrote or contributed to the writing of the manuscript: Asano, Nakamura, Nishiya, Shiozawa, Takakusa, Shibayama, Inoue, Shinozuka, Yahara, Watanabe, Yoshinari

Appendix

PBPK Parameters

Ag1, drug amount in the upper intestine; Ag2, drug amount in the lower intestine; blood-CLrm, renal blood clearance of M1; blood-CLrp, renal blood clearance of DS-1971a; Cbm, blood concentration of M1; Cbp, blood concentration of DS-1971a; Chm, hepatic concentration of M1; Chp, hepatic concentration of DS-1971a; CLint, bm, biliary blood intrinsic clearance of M1; CL_{int}, bp, biliary blood intrinsic clearance of DS-1971a; CL_{int}, mp, metabolic blood intrinsic clearance of DS-1971a; fCL,m, fraction of the clearance of DS-1971a that yields M1; Fg, intestinal availability; fum, unbound fraction of M1 in plasma; fup, unbound fraction of DS-1971a in plasma; ka1, absorption constant-1; ka2, absorption constant-2; Kphm, liverto-plasma ratio of M1; Kphp, liver-to-plasma ratio of DS-1971a; kt, transit constant; MWm, molecular weight of M1; MWp, molecular weight of DS-1971a; Qh, hepatic blood flow; Rbm, blood-to-plasma concentration ratio of M1; Rbp, blood-to-plasma concentration ratio of DS-1971a; Vh, liver volume; V1m, blood-based extrahepatic distribution volume of M1; V1p, bloodbased extrahepatic distribution volume of DS-1971a

Mass Balance Formulas for PBPK model

The following mass balance formulas were simultaneously solved by Phoenix model using WinNonlin 6.3. The inputted value of each parameter for PXB-mice and humans are described in Supplementary Table 4 and Table 5, respectively.

TABLE 6

Observed and PBPK model-predicted plasma PK parameters for humans after 400 mg oral dose of DS-1971a in the fed condition.

Compound	Method	Plasma $C_{ m max}$ (ng/ml)	Plasma AUC (ng·h/ml)	Plasma M1/P ratio
DS-1971a	Dynamic PBPK simulation	3082	13,479	_
	Observed ^a	2950 ± 1810^a	$12,500 \pm 3700^a$	_
M1	Dynamic PBPK simulation	3429	17,116	1.3
	Observed ^a	3180 ± 465^a	$18,400 \pm 3790^a$	1.6 ± 0.55^a

 $^{^{}a}PK$ parameters (mean \pm SD of n = 6) are cited and modified from Asano et al. (2022).

Upper intestinal compartment of DS-1971a

$$\frac{\mathrm{dAg1}}{\mathrm{dt}} = -ka1 \times \mathrm{Ag1} - kt \times \mathrm{Ag1} \tag{15}$$

Lower intestinal compartment of DS-1971a

$$\frac{\mathrm{dAg2}}{\mathrm{dt}} = kt \times \mathrm{Ag1} - ka2 \times \mathrm{Ag2} \tag{16}$$

Liver compartment of DS-1971a

$$\frac{dChp}{dt} \times Vh = Ag1 \times Fg \times ka1 + Ag2 \times Fg \times ka2 - CL_{int}, mp$$

$$\times \left(\frac{Chp}{Kphp} \times fup\right) - Qh \times \left(\frac{Chp}{Kphp} \times Rbp - Cbp\right) - CL_{int}, bp$$

$$\times \left(\frac{Chp}{Kphp} \times fup\right)$$

$$(17)$$

Blood compartment of DS-1971a

$$\frac{\text{dCbp}}{\text{dt}} \times V1p = Qh \times \left(\frac{\text{Chp}}{Kphp} \times Rbp - \text{Cbp}\right)$$

$$- \text{blood-}CLrp \times \text{Cbp}$$
(18)

Liver compartment of M1

$$\frac{\text{dChm}}{\text{dt}} \times Vh = fCL, m \times CL_{\text{int}}, mp \times \left(\frac{\text{Chp}}{Kphp} \times fup\right) \times \frac{\text{MWm}}{\text{MWp}}$$
$$-Qh \times \left(\frac{\text{Chm}}{Kphm} \times Rbm - Cbm\right) - CL_{\text{int}}, bm \times \left(\frac{\text{Chm}}{Kphm} \times fum\right)$$
(19)

Blood compartment of M1

$$\frac{\text{dCbm}}{\text{dt}} \times V1\text{m} = Q\text{h} \times \left(\frac{\text{Chm}}{K\text{phm}} \times R\text{bm} - \text{Cbm}\right) - \text{blood-}CL\text{rm} \times \text{Cbm}$$
(20)

Note that hepatic metabolism of M1 is not included in Eq. (19) because M1 was reported to be metabolically stable against human liver microsomes and S9 (Asano et al., 2022), indicating that the only elimination pathways of M1 are its biliary and urinary excretion in its unchanged form.

Plasma concentrations, C_{max} and AUC simulated by PBPK model

Since mass balance formulas were described as blood-based parameters, plasma concentrations of DS-1971a and M1 were calculated as follows:

Plasma concentration of DS-1971a =
$$\frac{\text{Cbp}}{R\text{bp}}$$
 (21)

Plasma concentration of M1 =
$$\frac{\text{Cbm}}{R\text{bm}}$$
 (22)

PK parameters were calculated by noncompartmental analysis of the simulated plasma PK profiles of DS-1971a and M1 in PXB-mice and humans using WinNonlin 6.3.

Acquisition of parameters used in mass balance formulas for PBPK model

The values of fum, fup, Qh, and Vh are cited from previous reports (Davies and Morris, 1993, Shinozuka et al., 2020; Asano et al., 2022). fCL,m, Kphm, Kphp, Rbm, and Rbp values were experimentally obtained (described in the Materials and Methods). Other parameters besides certain variables (Ag1, Ag2, Cbm, Cbp, Chm, and Chp) were acquired as follows. It is noted that BW represents BW of 70 kg for humans (Davies and Morris, 1993) and 0.020 kg for PXB-mice (the measured value of the purchased PXB-mice).

lack blood-CLrp and blood-CLrm

Renal plasma clearance (CLrp and CLrm) were divided by Rb values (Rbp and Rbm) with unit conversion.

blood-
$$CL$$
rp (L/h) = $\frac{CL$ rp \times BW \times 60 min \div 1000 (23)

blood-
$$CL$$
rm $(L/h) = \frac{CL$ rm $Rbm \times BW \times 60 min \div 1000$ (24)

• $\blacksquare \blacksquare \blacksquare CL_{int}$, bm, CL_{int} , bp, and CL_{int} , mp

 CL_{int} ,mp, CL_{int} ,bp, and CL_{int} ,bm were calculated based on the well-stirred model with unit conversion as follows:

$$CL_{\text{int}}, \text{bm} (\text{L/h}) = \frac{CL\text{bm}}{f\text{um}} \times \frac{Q\text{h}}{(Q\text{h} - CL\text{bm}/R\text{bm})} \times \text{BW} \times 60 \text{ min} \div 1000$$
 (25)

$$CL_{\rm int}$$
, bp (L/h) =

$$\frac{\textit{CLbp}}{\textit{fup}} \times \frac{\textit{Qh}}{(\textit{Qh-CLbp/Rbp})} \times \text{BW} \times 60 \, \text{min} \, \div \, 1000 \tag{26}$$

$$\frac{\mathit{CL}_{int}, mp \left(L/h \right) =}{\mathit{fup}} \times \frac{\mathit{Qh}}{(\mathit{Qh} - \mathit{CLmp}/\mathit{Rbp})} \times BW \times 60 \, \text{min} \, \div \, 1000 \tag{27}$$

• $\blacksquare \blacksquare \blacksquare Fa \text{ and } Fg$

Our previous work (Asano et al., 2021) indicated that 95% and 4.2% of the doses were recovered in bile and urine, respectively, after the oral administration of radiolabeled DS-1971a to mice, suggesting that the oral absorption of DS-1971a is more than 95% in mice. In addition, oral absorption in monkeys can be calculated to be 100% based on the urinary (35% and 38%) and total recovery (91% and 95%) after intravenous and oral administration (Asano et al., 2021) using a previously reported equation (Pleiss, 2005) with modification.

$$Oral absorption = \frac{Urinary recovery after oral administration}{Urinary recovery after intravenous administration} \times \frac{Total recovery after intravenous administration}{Total recovery after oral administration} \times 100$$
 (28)

Based on this evidence and the fact that no metabolism of DS-1971a with human intestinal microsomes and S9 was observed (data not shown), both fraction absorbed (Fa) and intestinal availability (Fg) were assumed to be 1.0. These assumptions were also applicable to Fa and Fg values in PXB-mice both because Fh,p in PXB-mice can be calculated to be 0.59 by Eq. (10) based on the observed CLmp of 20 ml/min/kg, CLbp of 0.27 ml/min/kg,

*R*b value of DS-1971a of 0.55 (Supplementary Tables 3 and 4), and *Q*h value of 90 ml/min/kg in mice (Davies and Morris, 1993), and because the $Fa \times Fg$ value can be calculated to be 1.0 by the following equation:

$$Fa \times Fg \text{ in PXB-mice} = \frac{\text{Oral bioavailability, pxb}}{100 \times Fh, p, pxb}$$
 (29)

where observed oral bioavailability and Fh,p in PXB-mice was 60% (Supplementary Table 2) and 0.59 (calculated above), respectively.

Thus, both Fa and Fg values were assumed to be 1.0 in the PBPK model, and the initial value of Ag1 (Ag1 value at time 0 hour) was equal to the oral doses (0.2 mg for PXB-mice and 400 mg for humans) by multiplying the oral doses by an Fa value of 1.0.

• ■■■ ka1, ka2, and kt

The values of ka1, ka2, and kt for PXB-mice were derived by fitting the plasma concentration of DS-1971a in PXB mice to the previously described PBPK model with multiplicative error using WinNonlin 6.3. ka1 and ka2 values for humans were estimated from the permeability value in MDCK II cells (described in the Materials and Methods). kt value for humans was calculated based on the following equation (Yu and Amidon, 1999):

$$kt \text{ in humans} = \frac{\text{Number of intestinal compartment } (N = 2)}{\text{Intestinal transit time } (3.5 \text{ h})}$$
(30)

Supplementary ka1, ka2, and kt values for humans were obtained in the same way as those for PXB-mice.

■■■ MWm and MWp

Molecular weights of M1 and DS-1971a (MWm and MWp) were calculated to be 481 and 465 g/mole, respectively.

• $\blacksquare \blacksquare \blacksquare V1m \text{ and } V1p$

Blood distribution volumes (Vd) other than liver (Vd of blood components were called V1p for DS-1971a and V1m for M1) were calculated as in a previous report (Ito et al., 2020) using the following equation:

$$\frac{V \lg \operatorname{or} V \lg \operatorname{m} (L) =}{\frac{V \lg \operatorname{or} V \lg \operatorname{m} (L)}{(R \lg \operatorname{por} R \lg \operatorname{m})}} - \frac{V \operatorname{h} \times (K \operatorname{php} \operatorname{or} K \operatorname{phm})}{(R \lg \operatorname{por} R \lg \operatorname{m})} \times (F \operatorname{h}, \operatorname{por} F \operatorname{h}, \operatorname{m})$$
(31)

The values of BW, liver volume (Vh), and Qh used were 0.02 kg, 0.0013 L, and 90 ml/min/kg for PXB-mice and 70 kg, 1.7 L, and 21 ml/min/kg for humans, respectively (Davies and Morris, 1993).

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