Preclinical Pharmacokinetics of MI-219, a Novel Human Double Minute 2 (HDM2) Inhibitor and Prediction of Human Pharmacokinetics

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ABSTRACT – Purpose. The two purposes of this study were evaluating preclinical pharmacokinetics of MI-219 and predicting clearance (CL) and volume of distribution at steady-state (Vdss) of MI-219 in humans. Methods. Pharmacokinetic studies were conducted on mice, rats, dogs, and monkeys. Human CL of MI-219 was predicted using allometric scaling (SA), multi-exponential allometric scaling (ME), rule of exponents (RoE), single species scaling, two-term power equation (TTPE), physiologically based in vitro-in vivo extrapolation (IVIVE), and fu corrected intercept method (FCIM). In vitro assays were conducted to determine in vitro intrinsic CL, protein binding, and blood-plasma partition coefficients. To estimate half-life of MI-219, plasma concentration-time profile in humans was predicted using kallynochron and apolysichron time transformation (Dedrick plots) and normalization with MRT and Vdss (Wajima’s method). In addition, simultaneous interspecies scaling of CL, Vdss and concentration-time profile were performed by using Nonlinear Mixed Effects Modeling (NONMEM). Results. Preclinical studies showed that the elimination of MI-219 was mainly through metabolism. The validation using observed monkey CL and Vdss showed that MA, IVIVE and Oie-Tozer methods were accurately than the other methods. Human CL of MI-219 predicted by ME and IVIVE was between 0.237-0.342 L*h⁻¹*kg⁻¹. Human Vdss predicted by Oie-Tozer method and allometric scaling of unbound volume of distribution of tissues (Vt/fuT) method was between 0.93-1.40 L*kg⁻¹. Superimposition of rat, monkey and dog data was observed in Dedrick plots and Wajima’s transformations. Conclusions. The predicted human pharmacokinetics is useful for the design of first-in-human study.

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INTRODUCTION

MI-219 is a potent, highly selective and orally active small-molecule inhibitor of human double minute 2 (HDM2)–p53 interaction (Figure S1) (1-3). MI-219 stimulates rapid but transient p53 activation in tumor tissues, resulting in inhibition of cell proliferation, selective induction of cancer cell apoptosis, and complete tumor growth inhibition in xenograft models. Preclinical studies have shown that MI-219 is a promising lead candidate for p53 wild-type cancer therapy.

Prediction of human pharmacokinetics is an essential element in clinical development of a potential drug molecule. Over the past few decades, numerous empirical or physiological approaches have been developed for quantitative predictions of human clearance (CL) (4, 5), volume of distribution at steady-state (Vdss) (6, 7) and human plasma concentration-time profile from preclinical animal data (8, 9). These predictive approaches typically use in vivo preclinical animal data and/or in vitro metabolism and disposition data obtained from human tissue such as microsomes or hepatocytes. However, there is not a standard method superior to other methods for each individual compound.

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Each approach has its unique advantages and disadvantages in its application to predicting human pharmacokinetics (5).

In current study, we first delineated in vitro and in vivo pharmacokinetics of MI-219 in mice, rats, dogs and monkeys and quantitatively predicted pharmacokinetics of MI-219 in humans. Preclinical studies showed that the elimination of MI-219 was mainly due to drug metabolism. Various methods were utilized to predict CL and Vdss values of MI-219 in human. The plasma concentration–time profile in humans was also predicted using Dedrick plots and Wajima’s method (9-12).

**METHODS**

**Reagents**
MI-219 and MI-343 (internal standard) were synthesized using a previously published method (2, 13). Mouse, rat, dog, monkey and human liver microsomes were supplied by Sigma (St. Louis, MO, USA) or Xenotech (Lenexa, KS, USA). All other chemicals and reagents were purchased from commercial suppliers and were of HPLC grade.

**Metabolic stability assay**
The metabolic stability of MI-219 was assessed in human (a pool of fifty subjects), CD-1 mouse (a pool of ten male mice), Sprague–Dawley rat (a pool of fifteen male rats), beagle dog (a pool of eight male dogs), and cynomolgus monkey (a pool of six male monkeys) liver microsomes. The enzymes were activated by β-NADPH (reduced form). The incubation solution was diluted with 0.1 M phosphate buffer (containing MgCl2) to 0.4 mL. The final concentrations of MI-219, liver microsomes, β-NADPH, and MgCl2 were 1 µM, 1 mg/mL, 1 mM, and 3.3 mM respectively. The solution was incubated at 37°C. An aliquot of 40 µl of mixture was collected at 0, 5, 10, 15, 30, 45 and 60 min, and then proteins were precipitated with 120 µl of ice-cold acetonitrile containing an internal standard (100 ng/mL). The processed samples were injected on a Zorbax Bonus-RP column (2.1 mm × 50 mm, 5 µm). The system was run in isocratic mode with mobile phase consisting of methanol and water in the ratio of 85:15 (v/v) at a flow rate of 0.3 mL/min. An electrospray ion source in the negative-ion multiple reaction monitoring (MRM) mode was used for detection. The MRM transition channel was m/z 550 to m/z 306 for MI-219 and m/z 596 to m/z 352 for IS. The ion spray voltage was set at -2700 V. Ionization temperature was set as 700 °C. Data acquisition and quantitation were performed using analyst software version 1.4.1 (Applied Biosystems, MDS Sciex Toronto, Canada). The peak area ratios (MI-219 peak area/ internal standard peak area) were plotted against time and the data were fitted with a two compartment model using WinNonlin version 3.2 (Pharsight Corporation, Mountain View, CA, USA) to calculate the initial elimination rate constants (Kc).

**Animals**
The animal procedures were performed according to a protocol approved by the University Committee for the Use and Care of Animals (UCUCA) at the University of Michigan. Male CD-1 (ICR) mice (n=33, body weight 18 ± 1.7 g), male Sprague Dawley rats (n=3, body weight 195 ± 15 g), male cynomolgus monkeys (n=6, body weight 2.5 ± 0.26 kg), and male beagle dogs (n=3, bodyweight 8.0 ± 1.1 kg) were used for pharmacokinetic studies. Animals were maintained under a 12-h light/ dark cycle in cages with free access to commercial food pellets. Water was supplied ad libitum. All blood samples were collected into heparinized polypropylene microcentrifuge tubes.

**Mouse pharmacokinetic study**
Thirty three male CD-1 mice were given an intravenous dose of 10 mg/kg of MI-219 dissolved in saline containing 20% polyethylene glycol (PEG 400) and 3% cremophor via the tail vein (MI-219 concentration 2 mg/mL and injection volume 5 mL/kg). Approximately 0.2 mL of blood was collected from each mouse by terminal cardiac puncture under isoflurane. Blood samples (n = 3 mice per time point) were taken from intravenously dosed mice at the following time points: before dosing and subsequently at 5, 15, 30, 45 min, 1, 2, 3, 5, 9, and 24 h after i.v administration. Blood samples were centrifuged at 21913 g × 5 min and 10 µl of supernatant was injected into LC-MS/MS. All incubations were performed in triplicate.

An Agilent 1200 HPLC system coupled with a Qtrap 3200 mass spectrometer (Applied Biosystems, MDS Sciex Toronto, Canada) was used to quantify the remaining MI-219 in the liver microsome samples. The processed samples were injected on a Zorbax Bonus-RP column (2.1 mm × 50 mm, 5 µm). The system was run in
**Rat pharmacokinetic study**

Two days before the initiation of the study, Sprague–Dawley rats were carotid cannulated using anaesthesia consisting of intraperitoneal administration of ketamine (100 mg/kg) and xylazine (10 mg/kg). Three male Sprague–Dawley rats were given an intravenous dose of 5 mg/kg of MI-219 in 10% DMSO and 90% saline (MI-219 concentration 5 mg/mL and injection volume 1 mL/kg) via a tail vein. Blood samples (approximately 0.2 mL per sample) were collected from each animal via a jugular vein catheter at pre-dose and 5, 15, 30 min, 1, 2, 4, 6, 8, and 24 hours post-dose. Blood samples were centrifuged within 30 min of collection and plasma was harvested. Plasma samples were stored at approximately -70°C until analysis.

**Monkey pharmacokinetic study**

Six male cynomolgus monkeys were given 10 mg/kg of MI-219 in 10% DMSO and 90% saline (MI-219 concentration 10 mg/mL and injection volume 1 mL/kg) via a saphenous vein. Blood samples (approximately 1 mL per sample) were collected from the femoral vein of each animal at pre-dose, 5, 15, 30 min, 1.5, 2, 3, 4, 6, 8, and 24 h post-dose. All blood samples were centrifuged at 13,000 rpm for 5 min and stored at approximately -70°C until analysis.

**Dog pharmacokinetic study**

Three male beagle dogs were given a single intravenous dose of 2 mg/kg of MI-219 in 10% DMSO and 90% saline (MI-219 concentration 2 mg/mL and injection volume 1 mL/kg) via a cephalic vein. Blood samples (approximately 2 mL per sample) were collected from the jugular vein of animals at pre-dose, 7, 12 and 30 min post-dose; and 1, 3, 4, 6, 8, 12 and 24 h post-dose. All blood samples were centrifuged in tubes containing heparin. Blood samples were centrifuged and plasma was harvested within 1 h of collection. Plasma samples were stored at approximately -70°C until analysis.

**Plasma sample preparation and LC-MS/MS analysis**

To prepare the plasma sample for LC-MS/MS analysis, ice-cold acetonitrile containing 100 ng/mL internal standard (MI-343) was added to 50 µl mouse plasma samples or 100 µl of plasma samples of rats, monkeys or dogs to precipitate proteins. After centrifugation at 13,000 rpm for 5 min, 10 µL of the supernatant was introduced into the LC-MS/MS system. LC-MS/MS conditions are similar to that used for microsomal sample analysis. The lower limits of quantitation (LOQs) in plasma are 2.1 ± 11% ng/mL (mouse), 1.0 ng/mL ± 13% (rat), 1.0 ng/mL ± 9% (monkey), and 1.5 ng/mL ± 10% (dog).

**Pharmacokinetic analysis**

All pharmacokinetic parameters were calculated by noncompartmental methods using WinNonlin version 3.2 (Pharsight Corporation, Mountain View, CA, USA). Parameters are presented as a mean ± standard deviation (SD).

**Plasma and liver microsome protein binding.**

Plasma protein binding was measured using a previously published method (14). Plasma was spiked with MI-219 to yield a final concentration of 5 µM. After incubation at 37°C for 1 h, an aliquot was transferred to a 10 kD cut-off ultrafiltration device (Millipore Corporation, Billerica, MA, USA), which was centrifuged at 2000 g for 3 h at 37°C. Samples were analyzed by LC-MS/MS as described above. The concentrations of MI-219 in the filtrate were determined by comparing with those of a standard curve containing known amounts of MI-219 dissolved in protein-free plasma (Macromolecules had been removed from the plasma by 10 kD cut-off ultrafiltration). The unbound fraction was estimated from the ratio of MI-219 concentration in the filtrate to that in the original plasma samples. Parallel studies using protein-free plasma instead of plasma indicated that MI-219 was bound minimally to the ultrafiltration device.

To determine the protein binding of MI-219 in liver microsomes, liver microsomes were diluted with 0.1 M phosphate buffer to 1 mg/mL and spiked with MI-219 to yield a final concentration of 1 µM. After incubation at 37°C for 1 h, an aliquot was transferred to a 10 kD cut-off ultrafiltration device, which was centrifuged at 2000 g for 3 h at 37°C. The concentrations of MI-219 in the filtrate were determined by comparing with those of a standard curve containing known amounts of MI-219 dissolved in ultrafiltrated 1 mg/mL liver microsome solution. The unbound fraction was estimated from the ratio of MI-219 concentration in the filtrate to 1 µM.
Blood-plasma partitioning (BP ratio)
BP ratios was measured using a previously published method (14). MI-219 was added to whole blood to obtain a final concentration of 5 µM and incubated at 37°C for 30 min. Plasma was separated from blood samples and concentration of MI-219 was determined by comparing with a standard curve prepared in blank plasma. The blood/plasma concentration ratio (BP) was determined by dividing 5 µM by that found in plasma separated from blood samples.

The concentration of MI-219 in blood cells is assumed to be equal to its unbound concentration in plasma. The theoretical BP ratio was calculated using equation 1 (14).

\[
BP = 1 + H \times (f_u - 1) \quad [1]
\]

where \(f_u\) is the fraction of unbound in plasma, and \(H\) is the hematocrit (mouse, 0.45; rat, 0.46; dog, 0.42; monkey, 0.41; rabbit, 0.36; human, 0.44)(15).

Prediction of pharmacokinetics of MI-219 in human

Simple Allometric scaling (SA)
The CL (L*h\(^{-1}\)) and Vss (L) values of MI-219 in animals and body weight W (kg) of animals after logarithmic conversion were fitted to log-transformed form of the following allometric equations (16) by linear regression:

\[
CL = a \times W^b \quad Vd_{ss} = a' \times W^c \quad [2]
\]

Where \(a\) and \(a'\) are the coefficients; \(b\) and \(c\) are the exponents of simple allometry. Human CL was extrapolated using the fitted line and an assumed body weight of 70 kg.

Allometric scaling after normalized by in-vitro metabolic data
Human and animal in vitro intrinsic CL (CL\(_{int,in\text{ vitro}}\)) was used to correct in vivo CL prior to allometric scaling. CL\(_{int,in\text{ vitro}}\) of MI-219 was derived from in vitro elimination rate constant in various liver microsomes.

\[
CL_{int,in\text{ vitro}} = \frac{0.693}{T_{1/2} \times C_{\text{protein}}} = \frac{K_e}{C_{\text{protein}}} \quad [3]
\]

Where \(K_e\) and \(C_{\text{protein}}\) (1mg/mL) were the rate constant of drug elimination in liver microsomes and the concentration of liver microsomes respectively.

Two-term power equation (TTPE)
CL was scaled allometrically by using an empirical equation which contains two power terms.

\[
CL = a \times W^{a_1} \times BrW^{b_1} \quad [5]
\]

where BrW is the brain weight.

Multi-exponential Allometric scaling (ME)
To avoid the uncertainty around the selection of correction factor used with SA, multi-exponential allometric scaling equation (17) was used to predict CL of MI-219 in human.

\[
CL = a \times W^{b_1} + \frac{1 - \frac{3}{2}b_2}{2} \times a \times W^{0.9} \quad [6]
\]

where \(a\) and \(b\) are the coefficient and exponent of SA; \(W\) is human body weight (70 kg).

Single species scaling
The following single species scaling equations (18) were employed to predict CL of MI-219 in human:

\[
\frac{CL_{\text{human}}}{kg} = 0.41 \times CL_{\text{dog}} / kg \\
\frac{CL_{\text{human}}}{kg} = 0.152 \times CL_{\text{rat}} / kg \\
\frac{CL_{\text{human}}}{kg} = 0.407 \times CL_{\text{monkey}} / kg \quad [7]
\]

\(f_u\) corrected intercept method (FCIM)

\[
CL = 33.35 ml/min \times \left( \frac{a}{Rf_u} \right)^{0.77} \quad [8]
\]

where \(Rf_u\) is the ratio of unbound fraction in plasma between rats and humans and \(a\) is the coefficient of SA.
Physiologically based in vitro-in vivo extrapolation (IVIVE)

CL of MI-219 in humans was predicted using the physiologically based in vitro-in vivo extrapolation (IVIVE). To assess the prediction performance of this method, CL of MI-219 in mice and rats were also predicted using IVIVE. The in vitro intrinsic CL of MI-219 in mouse, rat and human liver microsomes were scaled up to whole-organ in vivo intrinsic CL by using the following scaling factors: microsomal protein yield per gram of liver (mg/g): mouse, rat and monkey, 45; and human, 32; and liver weight (g/kg body weight): mouse, 87.5; rat, 40; monkey, 30; and human, 26 (14, 15). The in vivo intrinsic CL was corrected with the fractions unbound in plasma and liver microsomes. Then, the corrected in vivo intrinsic CL was used to calculate hepatic CL in mice, rats and humans by using well-stirred model and parallel-tube model (19). The values of liver blood flow (L*h⁻¹*kg⁻¹) were as follows: mouse 5.4; rat, 4.2; monkey, 2.7; and human, 1.24(14, 15). The calculated hepatic CL was then converted to plasma CL by multiplying BP ratio of MI-219 (MI-219 was assumed to be cleared exclusively by hepatic metabolism).

\[ CL_{in vivo} = \frac{K_e}{C_{protein}} \times \frac{32mg \text{ protein}}{1g \text{ liver weight}} \times \frac{26g \text{ liver weight}}{kg \text{ body weight}} \]  \[ CL \]  

Well-stirred model

\[ CL_{int,int, vivo} = \frac{fu_p \times CL_{int,int, vivo}}{fu_m} \]  \[ CL \]

Parallel-tube model

\[ CL_{H} = Q_H \times \left( 1 - e^{-\frac{CL_{int,int, vivo}}{Q_H}} \right) \]  \[ CL \]

\[ CL_{P} = CL_{H} \times BP \]  \[ CL \]

Allometric scaling of unbound volume of distribution of tissues (V_{f/uT})

The correlation between human and rat unbound volume of distribution of tissues (\( V_{f/uT} \)) was utilized to estimate human V_{ss}.(20)

\[ \left( \frac{V_T}{fu_T} \right)_{rat} = \frac{V_{dss, rat} - V_{b, rat} \times BP_{rat}}{fu_{P, rat}} \]

\[ \left( \frac{V_T}{fu_T} \right)_{human} = \left( \frac{V_T}{fu_T} \right)_{rat}^{0.95} \]  \[ 14 \]

\[ V_{dss,human} = fu_{P, human} \times \left( \frac{V_T}{fu_T} \right)_{human} + V_{b, human} \times BP_{human} \]  \[ 15 \]

where \( V_T \) was volume of tissue and \( fu_T \) was the drug unbound fraction in tissue; \( V_b \) was the blood volume (human, 0.0743 L/kg; rat, 0.054 L/kg).

Oie-Tozer equation

Oie-Tozer equation was used to calculate human V_{ds} from unbound fraction in plasma (\( fu_p \)) and tissue (\( fu_T \)), the plasma volume (\( V_p \): 0.0313, 0.0448, 0.0515 and 0.0436 L/kg for rat, monkey, dog and human), the extracellular fluid volume (\( V_e \): 0.265, 0.208, 0.216 and 0.151 L/kg for rat, monkey, dog and human), the ratio of extravascular to intravascular proteins (\( R_e/i : 1.4 \) for all the species), the tissue volume minus extracellular space (\( V_r \): 0.364, 0.485, 0.450 and 0.38 L/kg)(21). The unbound fractions in tissue \( fu_T \) of rats, monkey and dogs were calculated from corresponding V_{dss} and the average value of \( fu_T \) was used to calculate human V_{ds}.

\[ fu_{T, animal} = \frac{V_{dss} - V_p - fu \times V_e - (1 - fu_p) \times R_{e/i} \times V_p}{V_e \times (1 - fu_p)} \]  \[ 16 \]

\[ V_{dss,human} = V_p + fu_{P, human} \times V_e + (1 - fu_{T, human}) \times R_{e/i} \times V_p + V_e \times \frac{fu_{P, human}}{fu_{average}} \]  \[ 17 \]

Dedrick plots

The plasma concentration–time profile of MI-219 in humans (assume an i.v administration of 5mg/kg of MI-219) was simulated by using two Dedrick plot methods (based on Kallynochrons and
Apolysichrons)\(8, 11, 22\). The equations of the Dedrick plot methods were listed as below.

Kallynochrons (elementary Dedrick plot):

\[
\text{Time}_{\text{a}} = \text{Time}_{\text{a}} \times \left( \frac{W_{\text{a}}}{W_{\text{i}}} \right)^{1-b} \tag{18}
\]

\[
\text{Conc}_{\text{a}} = \text{Conc}_{\text{a}} \times \frac{\text{Dose}_{\text{i}}}{\text{Dose}_{\text{i}}} \times \left( \frac{W_{\text{a}}}{W_{\text{i}}} \right)^{b} \tag{19}
\]

Apolysichrons:

\[
\text{Time}_{\text{a}} = \text{Time}_{\text{a}} \times \left( \frac{W_{\text{a}}}{W_{\text{i}}} \right)^{c} \tag{20}
\]

\[
\text{Conc}_{\text{a}} = \text{Conc}_{\text{a}} \times \frac{\text{Dose}_{\text{i}}}{\text{Dose}_{\text{i}}} \times \left( \frac{W_{\text{a}}}{W_{\text{i}}} \right)^{c} \tag{21}
\]

Where \(W\) (kg) was the average body weight of animal or humans; \(\text{Dose}\) was in milligram. \(b\) and \(c\) were the exponents of simple allometry of \(\text{CL}\) and \(\text{Vdss}\). The \(\text{CLs}\) in different animal species were multiplied by their respective MLP and plotted as a function of the body weight on a log-log scale to generate exponent \(d\). The superimposition of the transformed animal concentration-time curves was expected. A two-compartment model (WinNonlin, version 5.2.1) was used to fit the superimposed curves. The goodness of fit was evaluated by correlation of observed and predicted values (\(r^2\)) and Akaike Information Criteria (AIC)(10).

Wajima’s method

The plasma concentration–time profile of MI-219 in humans was predicted by normalizing concentration–time profiles in animals with \(\text{MRT} = \text{Vdss}/\text{CL}\) and \(\text{C}_{\text{ss}} = \text{Dose}/\text{Vdss}\) (9). The superimposition of the normalized animal concentration-time curves was expected. A two-compartment model (WinNonlin, version 5.2.1) was used to fit the superimposed curves. Similar to Dedrick plot analysis, a two-compartment model (WinNonlin, version 5.2.1) was used to fit the predicted human plasma concentration–time profile of MI-219.

RESULTS

Metabolic stability assay

The percentages of MI-219 remaining in mouse, rat, dog, monkey and human liver microsomes (1 mg/mL) after 5, 10, 15, 30, 45 and 60 min incubation are presented in Figure 1. In all species tested, more than 50% of MI-219 was metabolized after incubation for 1 h, suggesting MI-219 was extensively metabolized by CYP enzymes. The elimination rate constants and half-lives of MI-219 in mouse, rat, monkey, dog and human liver microsomes were shown in Table 1.

Plasma and liver microsome protein binding and blood-plasma partitioning (BP ratio)

As shown in Table 2, the unbound fraction of MI-219 in mouse, rat, monkey and human plasma at a concentration of 5 \(\mu\)M ranged from 1.39% to 4.45%. The unbound fractions of MI-219 (1\(\mu\)M) in 1 mg/mL of mouse, rat, monkey and human liver microsomes are 2.03%, 2.44%, 4.40% and 6.16% respectively. The BP ratios of MI-219 in mice, rats and humans are experimentally measured and range from 0.572 to 0.597, which agree with the BP ratio values calculated from corresponding \(f_{\text{up}}\).

Pharmacokinetics of MI-219 in rat, mouse, dog and monkey

Figure 2 is a semi-log plot of MI-219 plasma concentration versus time for mouse, rat, dog, and monkey following intravenous administration. The concentrations of MI-219 in mouse plasma at 24 h are lower than the limit of quantification (2 ng/mL) of the LC-MS/MS assay. The pharmacokinetic parameters of MI-219 in the four species are summarized in Table 3.
Figure 1. Metabolic stability of MI-219 in animal and human liver microsomes (n = 3)

Table 1. Elimination rate constants and half-lives of MI-219 in liver microsomes

| Species   | Mouse | Rat  | Monkey | Dog   | Human |
|-----------|-------|------|--------|-------|-------|
| Elimination rate constant (min⁻¹) | 0.04449 | 0.04935 | 0.07842 | 0.06535 | 0.04747 |
| Half-life (min) | 15.6 | 14.0 | 8.8   | 10.6  | 14.6  |

Table 2. Unbound fraction of MI-219 in plasma and liver microsomes and blood-plasma partition

| Species | Unbound fraction in plasma (fuₚ) | Unbound fraction in liver microsome (fuₘ) | Blood-Plasma partition (calculated) | Blood-Plasma partition (measured) |
|---------|----------------------------------|------------------------------------------|-----------------------------------|-----------------------------------|
| Mouse   | 4.45%±0.04%                      | 2.03%±0.29%                              | 0.570                             | 0.581±0.010                      |
| Rat     | 3.87%±0.79%                      | 2.44%±0.73%                              | 0.558                             | 0.597±0.008                      |
| Monkey  | 1.96%±0.65%                      | 4.40%±0.94%                              | 0.598                             | N.A.                             |
| Dog     | 1.39%±0.48%                      | N.A.                                     | 0.586                             | N.A.                             |
| Human   | 1.65%±0.27%                      | 6.16%±0.58%                              | 0.567                             | 0.572±0.006                      |

Figure 2. Mean plasma concentration–time following single intravenous administration of MI-219. Mice (three animals per time point), rats (n = 3), dogs (n = 3), and monkeys (n = 6). * The concentrations of MI-219 in mouse plasma at 24 hr post-injection were lower than LOQ.
MI-219 has high plasma CLs in mice and rats at 7.15 and 2.13 L*h$^{-1}$*kg$^{-1}$, respectively. The corresponding blood CLs in mice and rats are 12.31 and 3.57 L*h$^{-1}$*kg$^{-1}$, respectively (more than 80% of hepatic blood flow). Especially, the blood CL exceeds hepatic blood flow in mice. In monkeys, MI-219 had a moderate plasma CL of 0.85 L*h$^{-1}$*kg$^{-1}$ and blood CL of 0.61 L*h$^{-1}$*kg$^{-1}$ (approximately 34% of hepatic blood flow). The terminal half-life ($t_{1/2}$) ranged from 1.89 h in mice to 3.90 h in dogs. The volume of distribution at steady-state ($V_{dss}$) ranged from 0.60 L*kg$^{-1}$ in dogs to 6.37 L*kg$^{-1}$ in mice.

Prediction of CL of MI-219 in humans

To validate various approaches for CL prediction, monkey CL was predicted from in vitro and/or in vivo data obtained from human liver microsomes, mice, rats and monkeys. As shown in Table 4, the predicted monkey CLs by simple allometric scaling (SA), allometric scaling normalized by microsomal data, TTPE, ME and IVIVE were 0.63, 0.73, 0.65, 0.91, 0.82 (IVIVE, well-stirred model) and 1.04 L*h$^{-1}$*kg$^{-1}$ (IVIVE, parallel-tube model), respectively. ME and IVIVE provided more accurate predictions than the other approaches. Single species scaling and FCIM are not applicable to monkey CL prediction.

The predicted human CL by SA was 0.146 L*h$^{-1}$*kg$^{-1}$. The coefficient and exponent for SA of CL were 1.076 and 0.5301, respectively (Figure 3A). According to Rule of Exponents (RoE), the CL (0.146 L*h$^{-1}$*kg$^{-1}$) is likely to be underestimated since the exponent is less than 0.55 (4). Corrections with maximum lifespan potential (MLP) or brain weight (BrW) would not apply. To eliminate the interspecies differences in hepatic metabolism, animal CL was corrected by multiplying the ratio of elimination rate constants in human and animal liver microsomes ($K_{e,human}/K_{e,animal}$). Allometric scaling of the corrected animal CL resulted in a lower predicted CL (0.08 L*h$^{-1}$*kg$^{-1}$) (Figure 3B).

Figure 4 shows the plot of CL against body weight (W) and brain weight (BrW) by using two-term power equation (TTPE). TTPE showed a good correlation among CL, W and BrW and the coefficient of determination ($r^2$) was 0.9988. The predicted human CL was 0.184 L*h$^{-1}$*kg$^{-1}$.

### Table 3. Pharmacokinetics (mean ± standard deviation) of MI-219 in animals

| Species | Body weight (kg) | Dose (mg*kg$^{-1}$) | AUC$_{(0-\infty)}$ (h*ng*L$^{-1}$) | MRT$_{(0-\infty)}$ (h) | Vd$_{z}$ (L*kg$^{-1}$) | T$_{1/2}$ (h) | Vd$_{ss}$ (L*kg$^{-1}$) | CL (L*h$^{-1}$*kg$^{-1}$) |
|---------|----------------|---------------------|-------------------------------|-----------------|----------------------|--------------|----------------------|------------------|
| Mouse   | 0.018          | 10                  | 7046±760                      | 0.9±0.04        | 19.53±2.12           | 1.89±0.04   | 6.37±0.40           | 7.15±0.76       |
| Rat     | 0.2            | 5                   | 2349±30                       | 1.38±0.12       | 6.76±1.14            | 2.20±0.34   | 2.95±0.31           | 2.13±0.03       |
| Monkey  | 2.5            | 10                  | 11884±1714                    | 2.41±0.28       | 4.13±0.56            | 3.35±0.09   | 2.07±0.35           | 0.85±0.12       |
| Dog     | 8              | 2                   | 6236±2312                     | 1.63±0.36       | 1.89±0.39            | 3.90±0.78   | 0.60±0.32           | 0.36±0.14       |

### Table 4. Predicted CL of MI-219 in human and monkey

| Methods                                | Predicted monkey CL (L*h$^{-1}$*kg$^{-1}$) | Predicted human CL (L*h$^{-1}$*kg$^{-1}$) |
|----------------------------------------|--------------------------------------------|--------------------------------------------|
| Simple allometry                       | 0.63                                       | 0.146                                      |
| Allometric scaling normalized by in vitro microsomal data | 0.73                                       | 0.080                                      |
| Two term power equation                | 0.65                                       | 0.184                                      |
| Multi-exponential allometric scaling   | 0.91                                       | 0.342                                      |
| Single species scaling                 | N.A.                                       | 0.324 (rat), 0.148 (dog), 0.346 (monkey)   |
| Physiologically based in vitro-in vivo scaling | 0.82 (well-stirred model)                  | 0.237 (well-stirred model)                 |
| f$_c$ corrected intercept method       | N.A.                                       | 0.168                                      |
Figure 3. Allometric scaling of CL (A); CL corrected by microsomal data (B); Vdss (C); unbound Vdss (D).

Figure 4. Human CL prediction using two-term power equation
The coefficient (1.076) and exponent b (0.5301) obtained from SA of CL were applied to the multi-exponential allometric scaling (ME) equation to estimate CL in human. The predicted human CL was 0.342 L*h⁻¹*kg⁻¹.

To simplify SA, single-species scaling methods proposed by Tang et. al. (18) were also applied to the prediction of CL of MI-219 in humans. A big variation was observed when the CL in humans was derived from rats (0.324 L*h⁻¹*kg⁻¹), dogs (0.148 L*h⁻¹*kg⁻¹), and monkeys (0.346 L*h⁻¹*kg⁻¹).

Considering the extremely high plasma protein binding of MI-219, the fu_corrected intercept (0.0387/0.0165 = 2.345) and coefficient of simple allometry (0.0317) were applied to FCIM equation. The resulted CL in humans was 0.168 L*h⁻¹*kg⁻¹.

CL of MI-219 was scaled from in vitro intrinsic CL determined in human liver microsome incubation using physiologically based in vitro-in vivo extrapolation method. The elimination rate constant of MI-219 in 1 mg/mL human liver microsome was 0.04747 min⁻¹. fu,m in human liver constant of MI-219 in 1 mg/mL human liver microsome was 0.342 L*h⁻¹. fu,m in human liver was estimated to be 0.342 L*h⁻¹.

The predicted human CL was extrapolated from rat (0.00475) were derived from corresponding Vdss.

To validate various approaches for Vdss prediction, monkey Vdss by allometric scaling, allometric scaling of undound Vdss, and Oie-Tozer method were 1.32, 0.77 and 2.05 L*kg⁻¹, respectively. Oie-Tozer equation provided the best predictions of monkey Vdss. Interspecies scaling of unbound volume of distribution of tissues (V_T/fu_T) method is not applicable to monkey Vdss prediction.

Estimates of Vdss (L) from mouse, rat, and monkey pharmacokinetic studies were plotted against body weight (kg) to predict human Vdss. Human Vdss predicted by SA was 0.418 L*kg⁻¹ (Figure 3C). As a comparison, human Vdss extrapolated from unbound Vdss (V_d/fu_d) of four animal species was 0.694 L*kg⁻¹ (Figure 3D). When human Vdss was extrapolated from mouse, rat, and monkey data, the predicted values by SA and allometric scaling of unbound Vdss were 0.90 and 1.22 L*kg⁻¹, respectively.

Interspecies scaling of V_T/fu_T method was used to estimate Vdss of MI-219 in human. V_T/fu_T in rats was derived from rat Vdss as 75.4 L*kg⁻¹. V_T/fu_T in human was 61.0 L*kg⁻¹, resulting in a Vdss in human of 1.05 L*kg⁻¹. Physiologically based Oie-Tozer equation was used to predict Vdss in human. fu_T of rats (0.00491), dogs (0.0131) and monkeys (0.00475) were derived from corresponding Vdss. The average fu_T of the three species was used to calculate Vdss of MI-219 in human, which was determined as 0.93 L*kg⁻¹. When the average fu_T of rats and monkeys was used as human fu_T, the predicted human Vdss was 1.40 L*kg⁻¹.

**Dedrick plot analysis**

Two Dedrick plot methods were applied for the prediction of the plasma concentration–time profile and half-life (t½) of MI-219 in human. The plasma concentration–time curves obtained from mice, rats, monkeys and dogs were normalized according to the pharmacokinetic times of kallynochrons (elementary Dedrick plot) and apolysichrons (complex Dedrick plot). As shown in Figure 5, reasonable superimposition was observed among the curves transformed from rat, monkey and dog plasma concentration–time profiles. However, the curve transformed from mouse plasma concentration–time profile was not superimposable to other curves. Hence, only the curves derived from rats, monkeys and dogs were used to predict human pharmacokinetics. A two-compartment model was employed to fit the superimposed curves using WinNonlin. The predicted human pharmacokinetic parameters (alpha, beta, A, B, CL, Vdss, MRT, and t½), coefficients of correlation and...
AIC were shown in Table 5. The kallynochron transformation resulted in CL (0.148 L*h\(^{-1}\)*kg\(^{-1}\)) and Vd\(\text{ss}\) (0.406 L*kg\(^{-1}\)) comparable with that derived from SA. The human CL predicted by apolysichron time transformation (0.164 L*h\(^{-1}\)*kg\(^{-1}\)) was more comparable with those obtained by TTPE and FCIM methods. The t\(\frac{1}{2}\) values predicted by kallynochrons and apolysichrons transformations were 3.08 h and 2.53 h respectively.

Wajima’s method
To estimate t\(\frac{1}{2}\), the concentration–time profiles in mice, rats, dogs and monkeys were normalized by MRT and C\(\text{ss}\). Reasonable superimposition was observed among the normalized curves from rats, monkeys and dogs (Figure 6A). The normalized mouse curve was not superimposable to other curves. The normalized concentrations and times from rats, monkeys and dogs were transformed into human concentrations and times by multiplying the predicted human C\(\text{ss}\) and MRT. Since allometric scaling tends to underpredict CL and Vd\(\text{ss}\), the average CL predicted from TTPE (0.184 L*h\(^{-1}\)*kg\(^{-1}\)), ME (0.342 L*h\(^{-1}\)*kg\(^{-1}\)), FCIM (0.168 L*h\(^{-1}\)*kg\(^{-1}\)) and IVIVE methods (0.237 and 0.282 L*h\(^{-1}\)*kg\(^{-1}\)) and the Vd\(\text{ss}\) predicted from Oie-Tozer equation (0.93 L*kg\(^{-1}\)) were utilized to generate MRT (3.85 h) and C\(\text{ss}\) (5.38 mg*L\(^{-1}\)) in human. The transformed human concentrations and times were plotted (Figure 6B) and fitted with a two-compartment model using WinNonlin. The predicted human pharmacokinetic parameters (alpha, beta, A, B, CL, Vd\(\text{ss}\), MRT, and t\(\frac{1}{2}\)) were listed in Table 5. The r\(^2\) (0.979) and AIC (543) values suggested a good curve-fitting.

Interspecies scaling using NONMEM
Finally, we coupled allometric scaling with NONMEM approach, which can model multiple PK parameters simultaneously and predict human plasma concentration-time profile. Pharmacokinetic parameters of rats, dogs and monkeys were estimated from measured plasma concentrations using individual or combined NONMEM models (Table S1). The estimated animal parameters were scaled to human parameters by the derived NONMEM interspecies scaling model. Human PK profile after i.v. administration was simulated (Figure 7) and human PK parameters were predicted (Table 5). The developed model was validated in all the animal species. The observed animal PK profile falls in the 5th and 95th quantile of model simulations, in all animal species. Figure S2 showed the validation in dogs.
Table 5. Predicted pharmacokinetic parameters of MI-219 in humans by Detrick plot, Wajima’s method and NONMEM modeling

| Parameters       | Kallynochr-ons | Apolysichrons | Wajima’s method | NONMEM |
|------------------|----------------|---------------|-----------------|--------|
| alpha            | 5.671          | 4.521         | 4.644           | 2.839  |
| beta             | 0.225          | 0.274         | 0.253           | 0.161  |
| A                | 76308          | 56043         | 32457           | 21714  |
| B                | 4554           | 4957          | 3017            | 3109   |
| CL (L*h⁻¹*kg⁻¹)  | 0.148          | 0.164         | 0.265           | 0.186  |
| Vdₑ (L*h⁻¹)      | 0.406          | 0.370         | 0.679           | 0.843  |
| MRT (h)          | 2.74           | 2.25          | 2.57            | 4.53   |
| T₁/₂             | 3.08           | 2.53          | 2.73            | 4.30   |
| R²               | 0.958          | 0.952         | 0.979           | N.A.   |
| AIC              | 556            | 619           | 543             | N.A.   |

Figure 6. Superimposition of normalized concentration-time profiles of MI-219 in animals using Css and MRT (Wajima’s method) (A) and predicted human concentration-time profile following a single i.v. injection of MI-219 (5 mg*kg⁻¹) (B)
Effective inhibition of p53 wild-type tumors in preclinical models makes MI-219 an attractive candidate for anti-cancer drug development (1). Accurate prediction of human pharmacokinetics of MI-219 is critical for the assessment of its potential to succeed as a drug. Numerous approaches based on in vitro and in vivo preclinical data for human pharmacokinetics are available but there is no universal approach that will work for every compound. Although allometric scaling techniques are simple and most widely used in industry, the use of allometric scaling should be cautious due to its empirical nature. In current study, we intend to utilize and evaluate various approaches including both empirical and physiologically based approaches for predicting pharmacokinetic parameters of MI-219 in humans.

The results from animal pharmacokinetic studies show that MI-219 has moderate to high plasma CL in all species (E_{H} >0.3). Especially, the blood CL in mice exceeds the hepatic blood flow, implying the existence of non-hepatic CL of MI-219 in mice. Since renal CL does not contribute significantly to the systemic CL of MI-219, the high CL of MI-219 is likely due to both hepatic and extrahepatic metabolism. Rapid metabolic turnover of MI-219 in mouse, rat, monkey, dog and human liver microsome incubation suggests extensive hepatic metabolism. High volume of distribution in mice, rats and monkeys suggests extensive distribution of MI-219 in peripheral tissues.

Considering the moderate-to-high hepatic extraction ratios of MI-219 in animals, it is not surprising that simple allometric scaling (SA) of CL showed a good correlation between CL and body weights since CL of compounds with high hepatic extraction ratios is limited by liver blood flow, which is well correlated with body weights across species (24). However, it has been suggested that a high r^2 value does not warrant an accurate CL prediction(25). Incorporation of elimination rate constants determined in live microsome incubations generated a lower CL prediction compared with SA. As declared in the Rule of Exponent (RoE), human CL of MI-219 is likely underestimated by SA since the exponent is less than 0.55 and the application of correction factors such as MLP and BrW will not improve the prediction (4). Consistently, CL of MI-219 in monkey was underpredicted by allometric scaling with or without correction factors.

The CL values of MI-219 estimated using single-species scaling exhibited great variation when scaled from different species. The interspecies difference limits its application in human CL prediction. Human CLs predicted by two other empirical approaches, TTPE and FCIM, are higher than that obtained from SA. In TTPE, CL is allometrically correlated with both body weight and
brain weight, which probably performs better than SA for some compounds. Unfortunately, TTPE provided an underestimation of monkey CL. FCIM was developed to correct vertical allometry. Compared with SA, FCIM minimizes the prediction errors caused by the interspecies differences in plasma protein binding between rats and human(26). However, FCIM ignores the differences in metabolic activities between rats and humans and it cannot be validated with animal data.

The assumption of multi-exponential allometric scaling (ME) is that CL is correlated with both body weight and volumes of elimination organs such as liver and kidneys (17). ME gave a good prediction of monkey CL, suggesting that this method can correct the underprediction caused by SA. The human CL (0.342 L*h⁻¹*kg⁻¹) obtained from ME is 2-fold higher than that predicted by SA (0.146 L*h⁻¹*kg⁻¹). ME utilizes both the exponent of SA (0.5301) and a fix exponent 0.9 to correlate human CL and body weight. The first exponent reflects the relationship between CL and body weight while the fix exponent reflects the relationship between hepatic CL and liver weight across species.

Interspecies differences in protein binding and metabolic enzymes are likely to result in inaccurate predictions of human CL by SA. Physiologically based IVIVE method is designed to minimize the errors caused by these interspecies differences by incorporating in vitro intrinsic CL, unbound fraction in plasma (fuₚ) and unbound fraction in liver microsomes (fuₘ) into the equations (27). To verify the predictability of IVIVE, CLs of MI-219 in mouse, rat and monkey were predicted using this approach. Although an underprediction of CL in mouse was observed, the predicted CLs in rat and monkey agreed well with the observed CLs. The underprediction of CL in mouse is probably due to extrahepatic clearance. Hence, to accurately predict human CL, more in vitro investigation is required to detect the possible extrahepatic clearance of MI-219 in humans.

The observed dog CL and predicted human CL are lower than that of the other species. The low CLs in dogs and humans are probably due to the interspecies differences in protein binding, hepatic physiological parameters, and/or biliary excretion. High protein binding in dog and human plasma results in low Vdss and decreased hepatic metabolism. Compared with rats and mice, humans and dogs have lower hepatic blood flow, lower microsomal proteins per gram of liver, and lower liver weight per kg of body weight. In addition, biliary excretion and enterohepatic circulation can decrease the CL in dogs and human, which might explain the difference in CL between dog and monkey.

Similar to CL prediction, underestimation of monkey Vdss by SA and allometric scaling of unbound Vdss were observed. Oie-Tozer method provided better prediction of monkey Vdss than allometric scaling (28). The underestimation of monkey Vdss is mainly caused by the low dog Vdss. The removal of dog Vdss from the data set resulted in a good prediction of monkey Vdss by Oie-Tozer method. Consistent with monkey Vdss prediction, Oie-Tozer method provided a higher prediction of human Vdss (0.93 L*kg⁻¹) than allometric scaling methods. When dog Vdss was removed, the predicted human Vdss was 1.40 L*kg⁻¹. Both the two predictions are comparable to the human Vdss predicted by allometric scaling of V₉/fu₉ from rat data (1.05 L*kg⁻¹). One advantage of Oie-Tozer method and allometric scaling of V₉/fu₉ method is that they can minimize the errors caused by interspecies differences in plasma protein binding and tissue binding by incorporating unbound fractions in plasma and tissues into the equations. In contrast, allometric scaling of unbound Vdss ignores tissue binding and SA ignores both tissue binding and plasma protein binding.

Detrick plot method was used to predict the plasma concentration-time profile of MI-219 in human. After kallynochron and apolysichron transformation, the data from rats, monkeys and dogs were superimposable. Essentially, Detrick plot method originated from the allometric scaling approach and its prediction performance depends on the allometric relationships of the pharmacokinetic parameters across species. Therefore, the CL and Vdss estimated by kallynochron and apolysichron methods are comparable to those obtained from SA. The advantage of Detrick plots is that it can be used to estimate half-life and AUC which are important for the design of human doses.

To avoid the possible underestimation of both CL and Vdss from allometric scaling and Detrick plots, Wajima’s normalization method was also employed to predict the concentration-time profile of MI-219 in humans. CL and Vdss derived from physiologically-based methods (IVIVE, FCIM, and Oie-Tozer equation) instead of simple allometric scaling were used to estimate human concentration-time profile, which fitted the two-compartment model better than the data transformed from Detrick plots.
In summary, the pharmacokinetics of MI-219 was studied in mice, rats, monkeys and dogs after i.v. injection. The pharmacokinetic parameters of MI-219 in humans were predicted using various empirical and physiologically based methods. The plasma concentration-time profile in humans was predicted using Detrick plot method and Wajima’s Css-MRT method. The validation using monkey in vivo PK parameters showed that ME and IVIVE approaches provided better CL predictions than the other approaches. Oie-Tozer method was superior to the other approaches for Vdss prediction. Monkey CL and Vdss were underpredicted by allometric scaling methods. When applied to human PK prediction, ME and IVIVE predicted comparable human CLs (0.237, 0.282 and 0.342 L*h^-1*kg^-1). Oie-Tozer method and allometric scaling of Vf/fuT method also provided similar predictions of human Vdss (0.93 or 1.40 and 1.05 L*kg^-1). Superimposition of rat, monkey and dog data and good curve fitting were observed after apolysichron time transformation and normalization with MRT and Vdss. The validation using monkey in vivo data and a narrow range of the predicted human PK parameters provides us a confidence in the human PK predictions of MI-219.

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