Protein binding and hepatic clearance: Re-examining the discrimination between models of hepatic clearance with diazepam in the isolated perfused rat liver preparation.

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Abbreviation list

$C_{in}$, entering concentration; $CL$, clearance; $CL_{int}$, intrinsic clearance; $C_{out}$, exiting concentration; $DM$, dispersion model; $DZP$, diazepam; $ER$, extraction ratio; $F_{est}$, estimated availability; $F_{obs}$, observed availability; $f_{u,B}$, fraction unbound in blood; $HSA$, human serum albumin; $IPRL$, isolated perfusate rat liver; $LC$–$MS/MS$, high performance liquid chromatography-tandem mass; $PTM$, parallel-tube model; $Q$, organ blood flow; $RED$, Rapid Equilibrium Dialysis; $WSM$, well-stirred model;
Abstract

Here we re-examine the hepatic extraction for diazepam, the only drug for which isolated perfused rat liver (IPRL) studies have been reported to not be consistent with the well-stirred model of organ elimination when only entering and exiting liver concentration measurements are available. First we examine the time dependency of diazepam equilibrium fraction unbound measurements from 4 to 24 hours, reporting the continuing increases with time. We conclude that the time dependency of equilibrium protein binding measurements for very highly bound drugs may be an issue that is not readily overcome. Examining $C_{\text{out}}/C_{\text{in}} (F_{\text{obs}})$ measurements for diazepam when no protein is added to the incubation media, we observe IPRL outcomes consistent with previous reports of Rowland and co-workers. Recent studies have shown the marked under-predictability of in vivo clearance from in vitro measures of elimination in the absence of protein for very highly bound drugs, which is markedly diminished in the presence of albumin. Thus, we measured $F_{\text{obs}}$ for diazepam at additional low concentrations of protein that would allow discrimination of the models of hepatic elimination and our results are not consistent with the dispersion and parallel-tube models. Therefore, although we can confirm similar outcomes to that reported by Rowland and co-workers when no protein is added to perfusion media, our IPRL results for diazepam, as well of that of Rowland and co-
workers, cannot be reasonably interpreted as proving that hepatic organ elimination is model independent or supporting the dispersion and parallel-tube models of organ elimination.
Significance Statement

The only drug experiments for which isolated perfusion rat liver studies do not support hepatic clearance being best described by the well-stirred model were carried out with diazepam at zero protein concentration. We repeat those studies confirming the previous results at zero protein concentration, but the addition of low protein binding conditions capable of differentiating the various models of hepatic elimination are more consistent with the well-stirred model of hepatic elimination.

These experimental studies do not support the preference for alternate models of hepatic elimination, nor the proposal that hepatic organ clearance is model independent.
**Introduction**

In 2018, Benet et al. (2018) proposed that calculating organ clearance \((CL)\) as the product of the extraction ratio \((ER)\) and the blood flow to the organ \((Q)\) as given in Eq. 1, as first proposed by Rowland (1972), was only consistent with the well-stirred model in pharmacokinetics. They maintain that this is true since the driving force concentration for elimination is assumed to be the concentration entering the organ, \(C_{in}\), rather than the concentration within the organ, i.e., \(Rate\ Out = Q \cdot (C_{in} - C_{out}) = CL \cdot C_{in}\). That is, there is no incremental clearance within the organ and all elimination is driven only by the entering concentration.

\[
CL = Q \cdot ER = Q \cdot \frac{C_{in} - C_{out}}{C_{in}} \tag{Eq. 1}
\]

Rowland and Pang (2018) questioned this concept and argue that Eq. 1 “simply expresses [a] proportionality between the observed rate of elimination and a reference concentration” and is not model dependent. That is, clearance as calculated by Eq. 1 is the organ clearance for all models of organ elimination, whether drug elimination in the organ follows the parallel tube model, various axial dispersion models or the well-stirred model.
It is universally agreed that the various models of organ clearance can only be
differentiated for high clearance (extraction ratio) substrates, and there are only three
published studies that directly test the difference in models for high extraction ratio
drugs in rat isolated perfused organs. All three of those studies, including two from
the Rowland laboratory, conclude that the data are consistent with the well-stirred
model, not alternate hepatic clearance models. Pang and Rowland (1977) evaluated
the effect of changing organ blood flow on the extraction ratio of lidocaine; the
Rowland laboratory also evaluated the effect of changing blood flow on the extraction
ratio of meperidine (Ahmad et al., 1983); while Jones et al. (1984) evaluated the
effect of changing protein binding on the extraction ratio of propranolol. Yet, these
experimental results are generally ignored.

In each of those studies for high clearance (extraction ratio) drugs, the investigators
concluded that the outcomes were consistent with the well-stirred model as given in
Eq. 1, as represented by Eq. 2, as opposed to the parallel tube model given in Eq. 3

\[
CL_H = \frac{Q_H \cdot f_{u,B} \cdot CL_{int,u}}{Q_H + f_{u,B} \cdot CL_{int,u}} \quad \text{(Eq. 2)}
\]

\[
CL_H = Q_H(1 - e^{-\frac{-f_{u,B} \cdot CL_{int,u}}{Q_H}}) \quad \text{(Eq. 3)}
\]
where the hepatic clearance, $CL_H$, is calculated from the hepatic blood flow, $Q_H$, the fraction unbound in blood, $f_{u,B}$ and the unbound intrinsic clearance, $CL_{int,u}$.

However, there are two experimental organ clearance studies with the low hepatic clearance drug, diazepam, in which diazepam has been manipulated to be high clearance in the absence of plasma proteins, as reported by the Rowland laboratory (Rowland et al., 1984, Diaz-Garcia et al., 1992), that demonstrate preference for the parallel tube and axial dispersion models versus the well-stirred model. Yet, the only experimental data values detailed in these two papers showing preference for the alternate models were carried out when no protein is present in the perfusion media and the outcomes, $C_{out}/C_{in}$ measures, differ by about 4-fold: 0.042 (CV = 76 %) reported by Rowland et al. (1984) using a gas liquid chromatographic analytical method and 0.011 (CV = 45 %) reported by Diaz-Garcia et al. (1992) using radiolabel measurements. Although the authors report that “diazepam did not bind to the perfusion apparatus” (Diaz-Garcia et al., 1992), we and other drug metabolism scientists are very wary of carrying out studies in the absence of any binding protein in the drug containing media, especially since diazepam in human plasma is so highly protein bound (98.5 %; Allen and Greenblatt, 1980). There is also marked uncertainty as to how protein binding measurements should be interpreted and utilized in
metabolism/transport studies as we and others have reviewed (Poulin et al., 2016; Bowman and Benet, 2018; Bteich et al., 2019). In vitro measures in the absence of protein (albumin) markedly under-predict in vivo clearance, where this difference is diminished in the presence of albumin (Poulin and Haddad, 2018; Kim et al., 2019). Therefore, in the present work, we have re-examined the rat liver perfusion clearance of diazepam in the absence of protein, but also at additional protein concentrations that discriminate, using mass spectrometric assay methods, which hepatic disposition model describes the hepatic clearance of diazepam.

Materials and Methods

Chemicals. With approval to use a controlled substance by the University of California San Francisco (UCSF), diazepam (DZP) and its internal standard diazepam-d5 were obtained from Spectrum (Gardena, CA) and Cerilliant (Round Rock, TX), respectively. The buffer reagents used in the IPRL experiments including taurocholate, sodium bicarbonate, and Krebs-Ringer powder, were all purchased from Sigma-Aldrich (St. Louis, MO). Fatty acid-free human serum albumin (HSA) was supplied commercially from SeraCare (Milford, MA). Rapid equilibrium dialysis (RED) device inserts were purchased from Thermo Scientific (Rockford, IL). All other chemicals and solvents for the analysis were of analytical or liquid
chromatography-mass spectrometry (LC–MS) grade.

**Diazepam Protein Binding**

Equilibrium dialysis experiments with DZP were carried out using RED device inserts (Waters et al., 2008). DZP was prepared at 1 μg/ml in Kreb’s buffer containing varied HSA concentrations (W/V) of 0%, 0.025%, 0.05%, 0.1%, 0.5%, 1%, 2% and 4%. A 500 μl volume of DZP testing solution was spiked into the sample chamber of the insert and followed by adding 750 μl blank buffer in the adjacent chamber. The loaded inserts were assembled in the Teflon based plate and the plate was sealed with a self-adhesive lid, swayed at 20 strokes per min under 37°C on an incubator shaker (Feasterville, PA) for the required experimental times of 4, 8, 12 or 24 h.

**Surgery and Perfusion of Livers.** Male Sprague-Dawley rats (350–400 g; Charles River Lab., Hollister, CA) were housed in the UCSF animal care facility with a 12-h light/dark cycle and allowed free access to water and food. Approval for their use in experiments was obtained from the Committee on Animal Research, UCSF. Anesthesia was conducted by intraperitoneal injection with a 1 ml/kg dose of ketamine/xylazine (91 mg, 9.1 mg/ml) before surgery. Livers were isolated for perfusion *ex situ*, as described previously with slight modifications (Wu and Benet, 2003; Lau et al., 2004). Oxygenated Krebs-Ringer bicarbonate buffer (pH 7.4), supplemented with different concentrations of HSA (ranging from 0% to 2%), sodium
bicarbonate (15 mM), and sodium taurocholate (10 μM), was pumped through the liver at a flow rate of 15 ml/min via a catheter inserted in the portal vein. Perfusion was performed in 8 rats at 37°C in a single pass manner from a reservoir containing 500 ml of medium, through a 1 μm pore size glass fiber filter, oxygenator, and bubble trap placed before the liver. The perfusate was oxygenated by carbogen (95% O₂/5% CO₂) as it passed through the semipermeable tube prior to entering the liver. Liver viability was judged on the basis of its appearance (uniformly pink to brown), portal vein pressure (8–10 mm Hg), and the pH of perfusate (in the range of 7.35–7.45), as well as metabolic capability. After a 20 min stabilization period, single-pass perfusions were started sequentially with the perfusates containing 1 μg/ml DZP and different HSA concentrations (0%, 0.025%, 0.05%, 0.1%, 0.5% and 2.0%). The order of the perfusate concentrations randomly followed one of the two sequences depicted in Fig. 1. During each rat perfusion experiment, 2 ml samples from the inferior vena cava (C_{out}) were taken at 16, 17, 18, 19 and 20 min. At the end of each perfusion period a 2 ml influx sample (C_{in}) was obtained. Due to fluctuating bile flow rate from the cannulated bile duct after liver isolation, no attempt was made to quantitate DZP in the bile.

**Sample Preparation and Analysis.** A simple and quick method of protein precipitation was employed in sample preparation. Each 200 μl perfusate sample was
combined with 40 μl internal standard solution (500 ng/ml in acetonitrile) as well as
560 μl cold acetonitrile. The final sample/internal standard/organic solvent mixtures
were then mixed briefly followed by centrifugation at 13,000 g for 10 min. Then 200
μl of the supernatant was transferred into HPLC screw cap vials. Preliminary studies
indicated no matrix effects as a function of different HSA concentrations, therefore
calibration curves for DZP were constructed over the range 1 ng/ml to 1000 ng/ml in
perfusate with 2% HSA.

DZP and its internal standard DZP-d5 were analyzed using a Shimadzu LC-20AD
high performance liquid chromatography (Kyoto, Japan) coupled to a Biosystems-
Sciex API 4000 series triple-quadrupole mass spectrometer (Foster City, CA) with an
electrospray ionization interface. Chromatographic separation was carried out on a
Thermo BDS Hypersil C18 column (4.6 × 100 mm, 5 μm) using a mobile phase
composed of solvent A (2 mM ammonium acetate and 0.1% formic acid in water) and
solvent B (2 mM ammonium acetate and 0.1% formic acid in methanol) under an
isocratic program with 3 min run time and 0.7 ml/min flow rate. The autosampler was
conditioned at 10 °C and the injection volume was set at 10 μl. Acquisition for tandem
mass spectrometry was performed in positive ionization mode, and the multiple
reaction monitoring mode was selected for quantification of the analytes. The
precursors to production ion transitions in the analytes were 285.2→154.1 and 285.2
→193.0 for DZP as well as 290.2→154.1 for DZP-d5 (Marin et al., 2012). The ion-source temperature was maintained at 550 °C, and the ion spray voltage was 5.0 kV. Analyst 1.4.2 software (Applied Biosystems-Sciex; Foster City, CA) was used to collect and process data.

**Models of Hepatic Drug Clearance**

Three models of hepatic elimination are considered in the figures comparing the outcomes by Diaz-Garcia et al. (1992): the well-stirred model, the parallel tube model and the axial dispersion model with a dispersion number of 0.34. The theoretical figures comparing the various hepatic models with changing protein binding or changing flow rate are based on the assumption that Eq. 1 is model independent. Although we disagree with that assumption, and that only the well-stirred model simulation is relevant, we calculated the theoretical curves here, following the procedure in Diaz-Garcia et al. (1992), so that a comparison with previous publications can be made. This was done by utilizing the mean observed \( f_u \) and \( C_{out}/C_{in} \) (\( F_{obs} \)) values for all protein concentrations and estimating the \( CL_{int,u} \) for each of the models. This value was then used to determine the estimated \( C_{out}/C_{in} \) (\( F_{est} \)) values at each value of \( f_u \), from which the hypothetical curve representing each of the models could be drawn.
Results

Equilibrium Dialysis. Variations of diazepam protein binding under different HSA concentrations are shown in Fig. 2 and Table 1 reflecting the time dependent changes in diazepam protein binding as a function of equilibration times between 4 and 24 hours. The time dependency of equilibrium protein binding measurements is well recognized. For example, Obach (1997) demonstrated this as a potential error source for IVIVE predictions for warfarin, imipramine and propranolol. Table 1 presents the mean values of percent fraction unbound for the four different dialysis times at the eight different HSA concentrations. The table also presents the standard deviations at each HSA concentration and the coefficient of variation of these measures. Although there are differences in variability (coefficient of variation) at different dialysis times and different HSA concentrations, there appears to be no discernable pattern. That is, the coefficients of variation appear to be randomly distributed across the 32 different mean values. Of particular relevance is the ratio of mean protein binding values at each of the HSA concentration for the 24 hour measurements versus the 4 hour measurements. The change in the extent of binding for zero HSA concentration, where fraction unbound percentages increase from 78.9 to 96.0 % over the 18 hour time period (ratio of 1.22), was very similar to the proportional change in the extent of protein binding at 4 % HSA, where percent unbound increased from 0.65 % to 0.84%
The protein binding change for all HSA concentrations changed similarly, ranging from 1.22 to 1.82. Thus, although it may appear in Fig. 2A that significant changes in protein binding with respect to incubation time only occur at low HSA concentration, the actual data as presented in Table 1 demonstrate that the relative change in the extent of binding from 4 to 24 hours is approximately the same for all HSA concentrations.

**IPRL Experiments.** We conducted IPRL with 8 male rat livers utilizing two different perfusion sequences varying HSA concentrations (Fig. 1). Figure 3 depicts the observed $\frac{C_{out}}{C_{in}}$ ratios for all 8 rats at the 6 different HSA concentrations, each with 5 measurements taken at minutes 16-20 as indicated. Two perfusion sequences were used, therefore $\frac{C_{out}}{C_{in}}$ ratios were averaged for the 2 sequences, however the final 0% HSA measurement in each rat (depicted as 0') is shown separately from the average of the earlier 0% HSA perfusions. Compared to the first period of HSA-free perfusion (0-20 min or 40-60 min), the availability of DZP at 120-140 min was increased approximately 2-fold ($0.033 \pm 0.022 \%$ vs $0.073 \pm 0.043\%$, p<0.05). The individual results for the 8 rats in the IPRL studies are depicted in Fig. 4, where the 0% HSA results are for the initial measurements only. The average $F_{obs}$ values are presented in Table 2 together with the calculated $F_{est}$ for each model using the average 8h equilibrium fraction unbound measurements for each HSA concentration.
(Although $F_{est}$ values would differ slightly for the different “equilibrium times”
chosen, the differentiation between models would be similar.) The mean $F_{obs}$ values
were then used for each model to generate the theoretical curves as depicted in Figs. 4
and 5. While Fig. 4 displays the individual data, Fig. 5 depicts the average of the
experimental $F_{obs}$ values for each HSA concentration versus fraction unbound and the
theoretical curves for the three models of hepatic elimination. The last three columns
of Table 2 give the estimated F values for each model for each HSA concentration.

Discussion

The motivation for the present study resulted from our interest in reexamining the
data from apparently the only isolated rat liver perfusion drug studies that provided
results that were not best explained by the well-stirred model of organ clearance.

Those studies (Rowland et al., 1984; Diaz-Garcia et al., 1992) examined diazepam
 clearance and protein binding at HSA concentrations of 0, 0.1, 0.5, 1 and 2%. The
authors reported that “Preliminary experiments showed that equilibrium was reached
within 3 hr, that volume shifts were negligible, and that diazepam did not bind to the
experimental system.” The mean experimental percent fraction unbound values from
these studies are included in Fig. 2b and shown at 3 hr. Our study using LC-MS/MS
measurements in the RED device yield lower results.
The variation of diazepam protein binding under different HSA concentrations is shown in Fig. 2. The fraction unbound of diazepam in the HSA-free medium is 78.9% after 4 h incubation, not reaching the theoretical value of 100% even after a 24 h incubation, as opposed to the 100% free fraction reported by Diaz-Garcia et al. (1992) after a 3 hour incubation. As seen in Table 1 the fractions unbound at 0.1% HSA ranged from 27.6% after 4 h incubation to 50.1% after 24 h incubation, versus the reported 37.6% after a 3 h incubation by Diaz-Garcia et al. (1992). None of the protein binding measurements at various HSA concentrations reached equilibrium at 24 h, all of which exhibit a similar degree of non-equilibrium, independent of protein concentration. Further, the variability of the measurement and the difference of the measurement between 4 and 24 hours appears to be independent of protein concentration. We conclude that the time dependency of equilibrium protein binding measurements for very highly bound drugs may be an issue that is not readily overcome.

For the isolated perfused diazepam liver studies, we show similar results as Rowland and co-workers for the initial measurements in the absence of added protein. That is, the mean data for this point (0% HSA; 91.6% average fraction unbound; $F_{obs}=0.033$)
appears to be consistent with the parallel-tube model (see Figs. 4 and 5). As depicted in Fig. 4, there is great variability in the 8 individual measurements at zero protein addition ($F_{obs} = 0.033; CV = 67\%$), with this high variability ($F_{obs} = 0.042; CV = 76\%$) also reported by Rowland et al. (1984) using a gas liquid chromatographic analytical method and ($F_{obs} = 0.011; CV = 45\%$) reported by Diaz-Garcia et al. (1992) using radiolabel measurements. Our second measures at zero protein addition of $F_{obs}$ at 120-140 min ($F_{obs} = 0.073\% ; CV = 59\%$) were statistically higher than the earlier first measurements, probably reflecting the decrease in liver function with time, as this parameter is much more sensitive than traditional measures of liver viability. When our second measure at zero protein is plotted in Fig. 5, this mean value would appear to be more consistent with the dispersion model (0% HSA; 91.6% average fraction unbound; $F_{obs}$=0.073). The 2-fold difference here in our measurements should be considered in light of the 4-fold difference reported in the two Rowland and co-worker studies.

The Rowland and co-workers investigations did not study any other protein concentrations that would readily allow comparison of the different models. Thus, we investigated 0.025% and 0.05% protein concentrations in our IPRL studies. As seen in Figs. 4 and 5 these studies appear to be more consistent with the well-stirred
model. In Table 2 we examined the estimated availability for each of the models (under the assumption that clearance was model independent). At lower protein concentrations where the diazepam clearance (extraction ratio) is high, differentiation between the models is obvious. This differentiation dissipates as protein binding increases and clearance (extraction ratio) decreases. Comparing the least square differences between the three models as given in Table 2 and shown in Fig. 5, the initial average $F_{obs}$ measurement when no protein is added to the perfusion media is consistent with the theoretical parallel-tube curve and the later 120-140 min measurement is consistent with the dispersion model theoretical curve. Thus, it appears that we confirm the experimental results observed by Rowland et al. (1984) and Perez-Garcia et al. (1992), respectively, when no protein is added to the perfusion media. However, when we studied low concentrations of protein that could differentiate the various theoretical models our results appear to be consistent with the well-stirred model. As noted above, today there is increasing attention being paid to how fraction unbound is interpreted in predicting metabolic interactions and the effect that albumin can have on these protein binding and metabolism measurement as measures in the absence in protein markedly under-predict in vivo clearance as recently reviewed (Bowman and Benet, 2018; Bteich et al., 2019). Recent studies with hepatocytes report markedly better IVIVE predictability in the presence of albumin
than in its absence (Poulin and Haddad, 2018; Kim et al., 2019). Thus, although we

can confirm similar outcomes to that reported by Rowland and co-workers when no

protein is added to perfusion media, those results cannot be reasonably interpreted as

proving that Eq. 1 is model independent.
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Authorship Contributions

Participated in research design: Wang, Benet

Conducted experiments: Wang

Performed data analysis: Wang, Benet

Wrote or contributed to the writing of the manuscript: Wang, Benet
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dispersion model of hepatic drug elimination to the kinetics of diazepam in the

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clearance from in vitro uptake clearance by suspended human hepatocytes for
anionic drugs with high binding to human albumin: Improvement of in vitro-
to-in vivo extrapolation by considering the "albumin-mediated" hepatic uptake
mechanism on the basis of the "facilitated-dissociation model". *Drug Metab
Dispos* **47**: 94-103.

and efflux significantly changes metabolism: hepatic enzyme-transporter

assay for 21 benzodiazepine drugs and metabolites, zolpidem and zopiclone in


Legends for Figures

Figure 1. Drug loading design of diazepam (1 μg/ml) combined with varied HSA concentrations in IPRL studies. Two sequences of HSA concentrations were utilized and individual rats were randomly assigned to a sequence. The Arabic number superscripts indicate where the individual HSA concentrations were in the testing sequence (i.e., in sequence 1, 0.1% HSA was studied between 100 and 120 min, while in sequence 2, this HSA concentration was studied between 60 and 80 min.)

Figure 2. (A) Plot of the free fraction of diazepam in the perfusate as a function of logarithmic HSA concentration for incubation times ranging from 4 to 24h. (B) Solid lines connect the fraction unbound as a function of equilibration time reported here. Data points at 3 h represent fraction unbound reported by Diaz-Garcia et al. (1992).

Figure 3. The availabilities of diazepam in IPRL studies under varied HSA concentrations in total seven periods with constant perfusate flow rate (15 ml/min) and drug concentration (1 μg/ml). Flow-out samples were collected every minute from 16 to 20 min in each period, including the flow-in perfusate at the last sampling time.
Figure 4. Estimated results of well-stirred (WSM), parallel-tube (PTM), and dispersion (DM) models for the relationship between availability and fraction unbound of diazepam. Symbols represent the individual results in each of the IPRL studies. Values for the zero protein added to the incubation are for the initial measurements.

Figure 5. Estimated results of well-stirred (WSM), parallel-tube (PTM), and dispersion (DM) models for the relationship between availability and fraction unbound of diazepam. Symbols represent the mean results (n=8) in all of the IPRL studies. For zero protein additions, ● depicts the initial measurements between 0-20 min and 40-60 min, while ◊ depicts the second measurements in each perfusion at 120-140 min.
Table 1. Diazepam Fraction Unbound (%) as a Function of Incubation Time and HSA Concentration

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<th>Dialysis Time</th>
<th>HSA (%) W/V</th>
<th>Mean</th>
<th>SD</th>
<th>CV%</th>
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Table 2. Observed ($F_{obs}$) and estimated availability ($F_{est}$) of diazepam in the IPRL under varying protein (HSA) binding conditions and measured 8h fraction unbound for three different hepatic drug clearance models.

<table>
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<th>HSA (%)</th>
<th>$f_{u,8h}$ (%)</th>
<th>$F_{obs}$ (%)</th>
<th>Estimated availability; $F_{est}$ (%)</th>
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<td>92.7</td>
<td>88.3</td>
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</table>

$^a$ well-stirred model  
$^b$ parallel-tube model  
$^c$ dispersion model
Fig 1

Cannulation

-20 0 20 40 60 80 100 120 140 Time (min)

0: Stabilized by DZP-free perfusate

1: 0% HSA\(^1\) or 0.3% HSA\(^2\)

2: 0.05% HSA\(^1\ & 2\)

3: 0.5% HSA\(^1\) or 0% HSA\(^2\)

4: 0.25% HSA\(^1\) or 0.1% HSA\(^2\)

5: 2% HSA\(^1\ & 2\)

6: 0.1% HSA\(^2\) or 0.025% HSA\(^2\)

7: 0% HSA\(^1\ & 2\) retest
(A) Fraction unbound (%) vs. HSA concentration (% W/V) for 4, 8, 12, and 24 h incubation (n=4).

(B) Fraction unbound (%) vs. Duration of equilibrium (h) for HSA concentrations 0%, 0.025%, 0.05%, 0.1%, 0.5%, 1%, 2%, and 4%.

Fig 2
Fig 3

(A): Sampling time in each period (min)
(B): HSA concentration (%)
Fig 4
Fig 5