Building and evaluation of a PBPK model for inulin in rats

|  |  |
| --- | --- |
| Version | x.x-OSPy.y |
| based on *Model Snapshot* and *Evaluation Plan* | https://github.com/Open-Systems-Pharmacology/Inulin-Model/releases/tag/vx.x |
| OSP Version | y.y |
| Qualification Framework Version | z.z |

This evaluation report and the corresponding PK-Sim project file are filed at:

https://github.com/Open-Systems-Pharmacology/OSP-PBPK-Model-Library/

Table of Contents

# Introduction

Inulin is a highly hydrophilic polysaccharide which does not distribute into cells and is cleared via glomerular filtration.

Inulin has a considerably smaller solute radius than the proteins which had been used to develop the generic large molecule physiologically based pharmacokinetic (PBPK) model in PK-Sim ([Niederalt 2018](#Xa559bd5247f9128e74e6edcc17de37648e8cde6)).

The herein presented evaluation report evaluates the performance of the PBPK model for inulin in rats using the large molecule model in PK-Sim.

The presented inulin PBPK model as well as the respective evaluation plan and evaluation report are provided open-source (https://github.com/Open-Systems-Pharmacology/Inulin-Model).

# Methods

## Modeling Strategy

The development of the large molecule PBPK model in PK-Sim® has previously been described by Niederalt et al. ([Niederalt 2018](#Xa559bd5247f9128e74e6edcc17de37648e8cde6)). In short, the model was built as an extension of the PK-Sim® model for small molecules incorporating (i) the two-pore formalism for drug extravasation from blood plasma to interstitial space, (ii) lymph flow, (iii) endosomal clearance and (iv) protection from endosomal clearance by neonatal Fc receptor (FcRn) mediated recycling.

For model development and evaluation, PK data were used from compounds with a wide range of solute radii and from different species. The PK data used for parameter estimation were from the following compounds: antibody–drug conjugate BAY 79-4620 in mice (Bayer in house data), antibody 7E3 in wild-type and FcRn knockout mice ([Garg 2007](#Xa559bd5247f9128e74e6edcc17de37648e8cde6), [Garg2009](#Xa559bd5247f9128e74e6edcc17de37648e8cde6)), domain antibody dAb2 in mice ([Sepp 2015](#Xa559bd5247f9128e74e6edcc17de37648e8cde6)), antibodies MEDI-524 and MEDI-524-YTE in monkeys ([Dall’Acqua 2006](#Xa559bd5247f9128e74e6edcc17de37648e8cde6)), and antibody CDA1 in humans ([Taylor 2008](#Xa559bd5247f9128e74e6edcc17de37648e8cde6)). The PK data used for model evaluation were from inulin in rats ([Tsuji1983](#Xa559bd5247f9128e74e6edcc17de37648e8cde6)) and tefibazumab in humans ([Reilly 2005](#Xa559bd5247f9128e74e6edcc17de37648e8cde6)).

The PBPK model including the estimated physiological parameters as described by Niederalt et al. ([Niederalt 2018](#Xa559bd5247f9128e74e6edcc17de37648e8cde6)) is available in the Open Systems Pharmacology Suite from version 7.1 onwards.

This evaluation report focuses on the PBPK model for inulin.

Details about input data (physicochemical, *in vitro* and PK) can be found in [Section 2.2](#X681d1fae3f233cf07e1ee33c73d775b8586b96f).

Details about the structural model and its parameters can be found in [Section 2.3](#X920e58f6eb345d17092abc826058b6950015cf7).

## Data

### In vitro / physico-chemical Data

A literature search was performed to collect available information on physicochemical properties of Inulin. The obtained information from literature is summarized in the table below.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **Parameter** | **Unit** | **Value** | Source | **Description** |
| MW | g/mol | 5000-5500 | [Ohno 1978](#Xa559bd5247f9128e74e6edcc17de37648e8cde6) | Molecular weight |
| r | nm | 1.39 | [Ghandehari 1997](#Xa559bd5247f9128e74e6edcc17de37648e8cde6) | Hydrodynamic solute radius |
| logP | µM | < -10 | [Dubbelboer 2022](#Xa559bd5247f9128e74e6edcc17de37648e8cde6) | Lipophilicity (octanol/water partition coefficient). Inulin is highly hydrophilic. A logP = -10 is insensitively small in the PBPK model. |
| Kd (FcRn) | µM | 999,999 |  | Dissociation constant for binding to FcRn. High value representing no FcRn binding. |

### PK Data

Published plasma and tissue PK data on inulin in rats were used.

|  |  |
| --- | --- |
| Publication | Description |
| [Tsuji 1983](#Xa559bd5247f9128e74e6edcc17de37648e8cde6) | Plasma and tissue concentrations after i.v. application of 20 and 200 mg/kg inulin in rats (for 200 mg/kg plasma only). |

## Model Parameters and Assumptions

### Absorption

There is no absorption process since inulin was administered intravenously

### Distribution

The standard vascular properties of the different tissues (hydraulic conductivity, pore radii, fraction of flow via large pores) and standard lymph and fluid recirculation flow rates from PK-Sim were used ([Niederalt 2018](#Xa559bd5247f9128e74e6edcc17de37648e8cde6)).

### Metabolism and Elimination

Inulin is renally excreted via glomerular filtration. The standard glomerular filtration rate from the PK-Sim library was used (GFR fraction = 1).

### Tissue Concentrations

For the comparison with experimental data the parameters Fraction of blood for sampling used in the Observer for the tissue concentrations were set for all organs to 0.18. This value is based on the parameter identification for different compounds reported in Ref. ([Niederalt 2018](#Xa559bd5247f9128e74e6edcc17de37648e8cde6)) for comparison with tissue dissection data. (The parameter Fraction of blood for sampling specifies residual blood in tissue as ratio of blood volume contributing to the measured tissue concentration to the total in vivo capillary blood volume.)

|  |  |  |
| --- | --- | --- |
| Model Parameter | Value | Unit |
| Fraction of blood for sampling (all organs) | 0.18 |  |

Experimentally, gut concentrations (from duodenum to the cecum) were measured ([Tsuji 1983](#Xa559bd5247f9128e74e6edcc17de37648e8cde6)). In the present evaluation report, the experimental gut concentrations were compared to simulated organ concentrations for small and large intestine separately in the goodness of fit plots as well as in the concentration-time profile plot.

### Automated Parameter Identification

No drug specific parameters were fitted.

# Results and Discussion

The PBPK model for inulin was evaluated with plasma and tissue PK data from rats.

The next sections show:

1. the final model parameters for the building blocks: [Section 3.1](#final-input-parameters).
2. the overall goodness of fit: [Section 3.2](#diagnostics-plots).
3. simulated vs. observed concentration-time profiles for the clinical studies used for model building and for model verification: [Section 3.3](#ct-profiles).

## Final input parameters

The compound parameter values of the final PBPK model are illustrated below.

### Compound: Inulin

#### Parameters

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| Name | Value | Value Origin | Alternative | Default |
| Solubility at reference pH | 9999 mg/l | Other-/Dummy value not used in the simulation | Measurement | True |
| Reference pH | 7 | Other-/Dummy value not used in the simulation | Measurement | True |
| Lipophilicity | -10 Log Units | Other-Highly hydrophilic | Measurement | True |
| Fraction unbound (plasma, reference value) | 1 | Other-Assumption | Measurement | True |
| Is small molecule | Yes |  |  |  |
| Molecular weight | 5500 g/mol | Publication-Ohno1978 |  |  |
| Plasma protein binding partner | Unknown |  |  |  |
| Radius (solute) | 0.00139 µm | Publication-Ghandehari1997 |  |  |

#### Calculation methods

|  |  |
| --- | --- |
| Name | Value |
| Partition coefficients | PK-Sim Standard |
| Cellular permeabilities | PK-Sim Standard |

#### Processes

##### Systemic Process: Glomerular Filtration-GFR

Species: Human

###### Parameters

|  |  |  |
| --- | --- | --- |
| Name | Value | Value Origin |
| GFR fraction | 1 |  |

## Diagnostics Plots

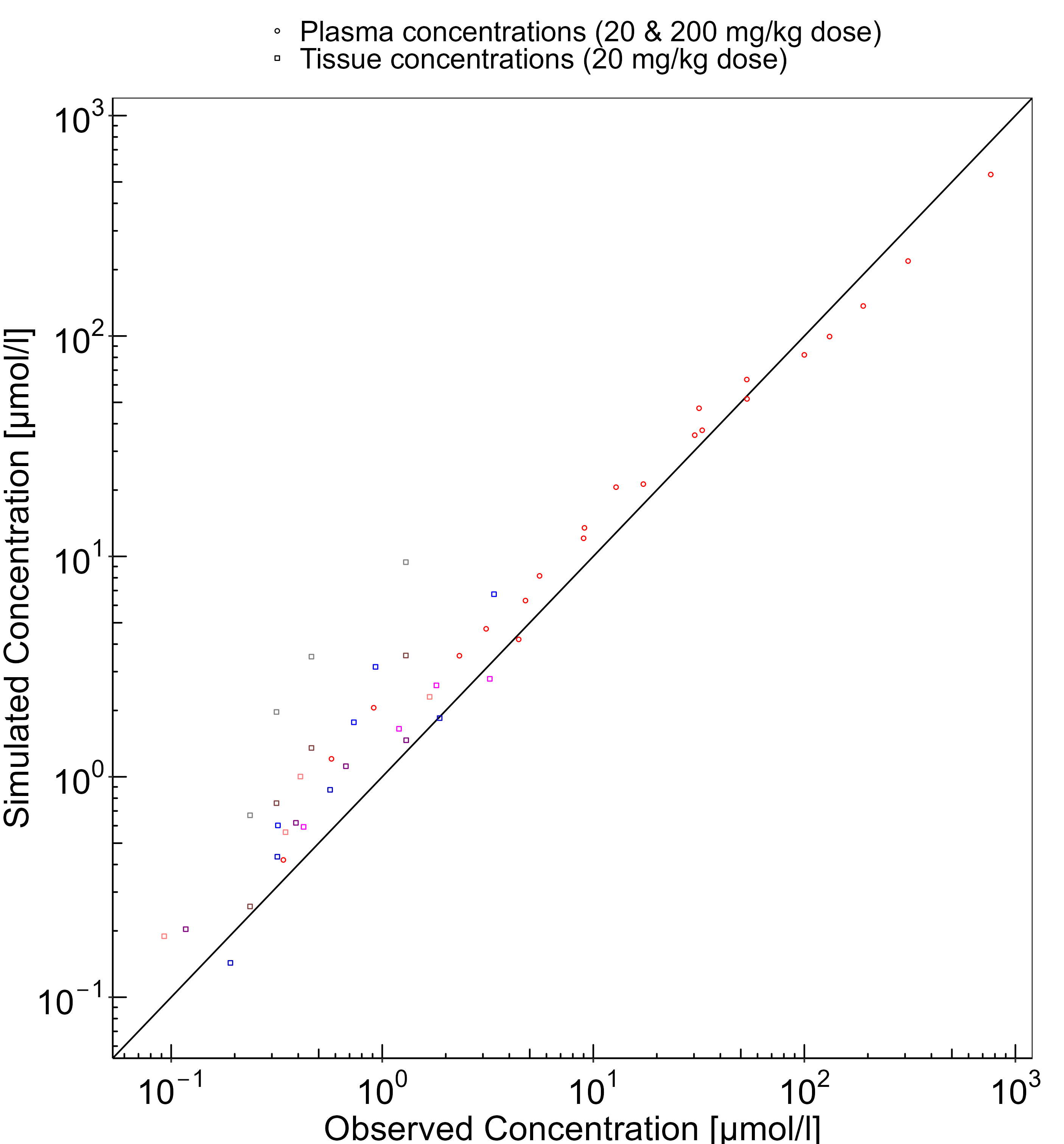
Below you find the goodness-of-fit visual diagnostic plots for the PBPK model performance of all data used presented in [Section 2.2.2](#PK-data).

The first plot shows observed versus simulated plasma concentration, the second weighted residuals versus time.

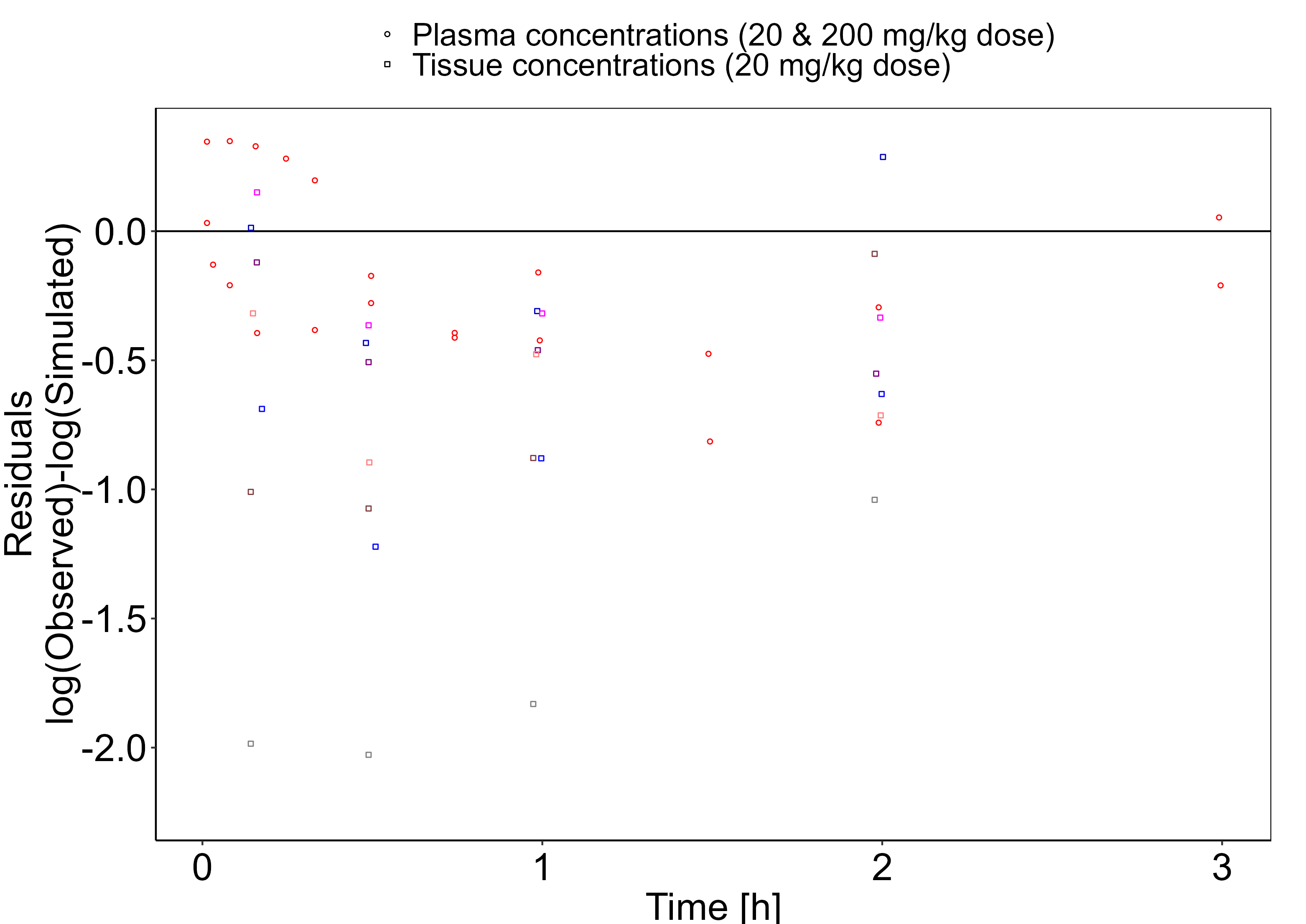
**Table 3-1: GMFE for Goodness of fit plot for concentration in plasma and tissues**

|  |  |
| --- | --- |
| Group | GMFE |
| Plasma concentrations (20 & 200 mg/kg dose) | 1.38e+00 |
| Tissue concentrations (20 mg/kg dose) | 2.01e+00 |
| All | 1.71e+00 |

**Figure 3-1: Goodness of fit plot for concentration in plasma and tissues**



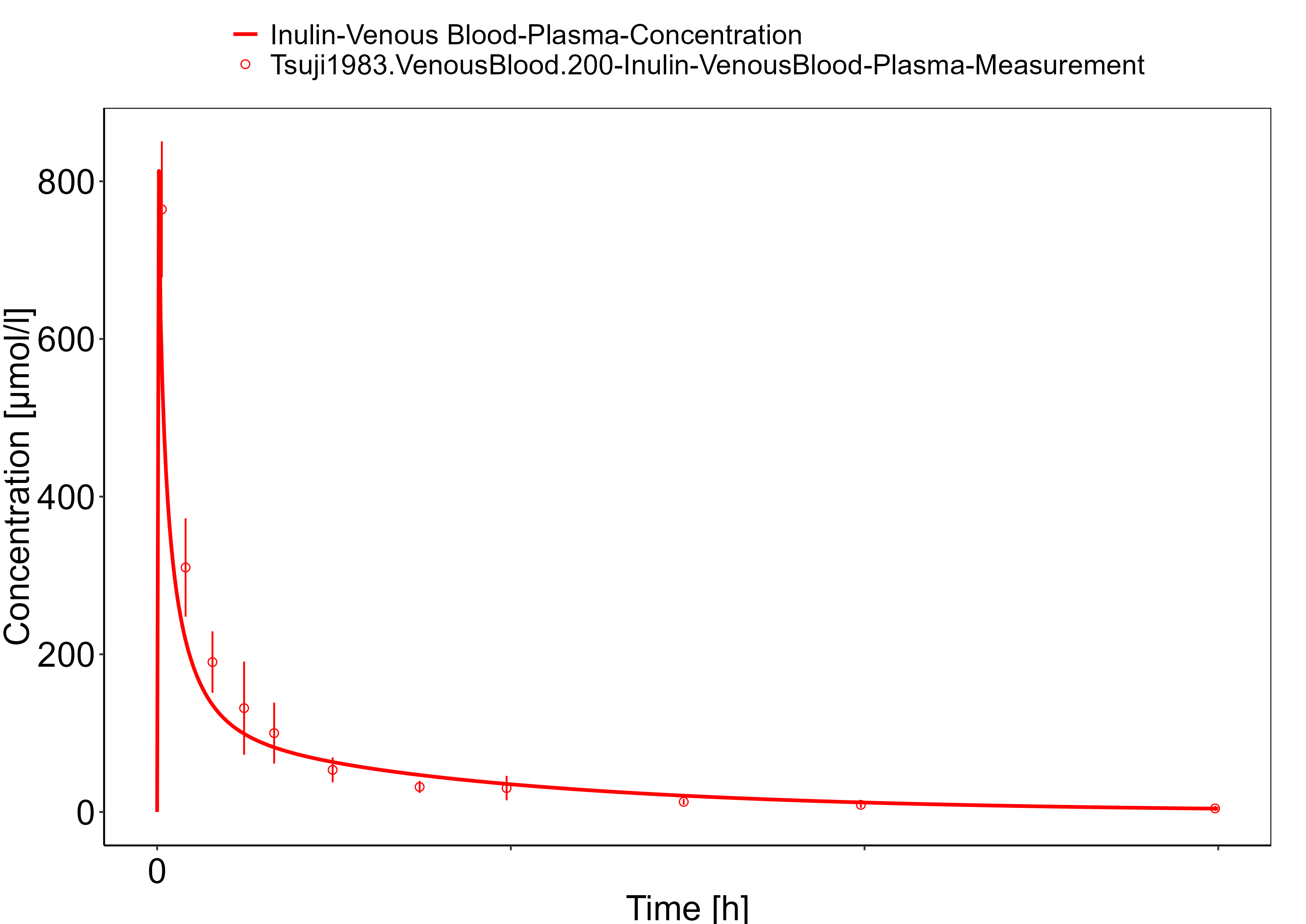
**Figure 3-2: Goodness of fit plot for concentration in plasma and tissues**



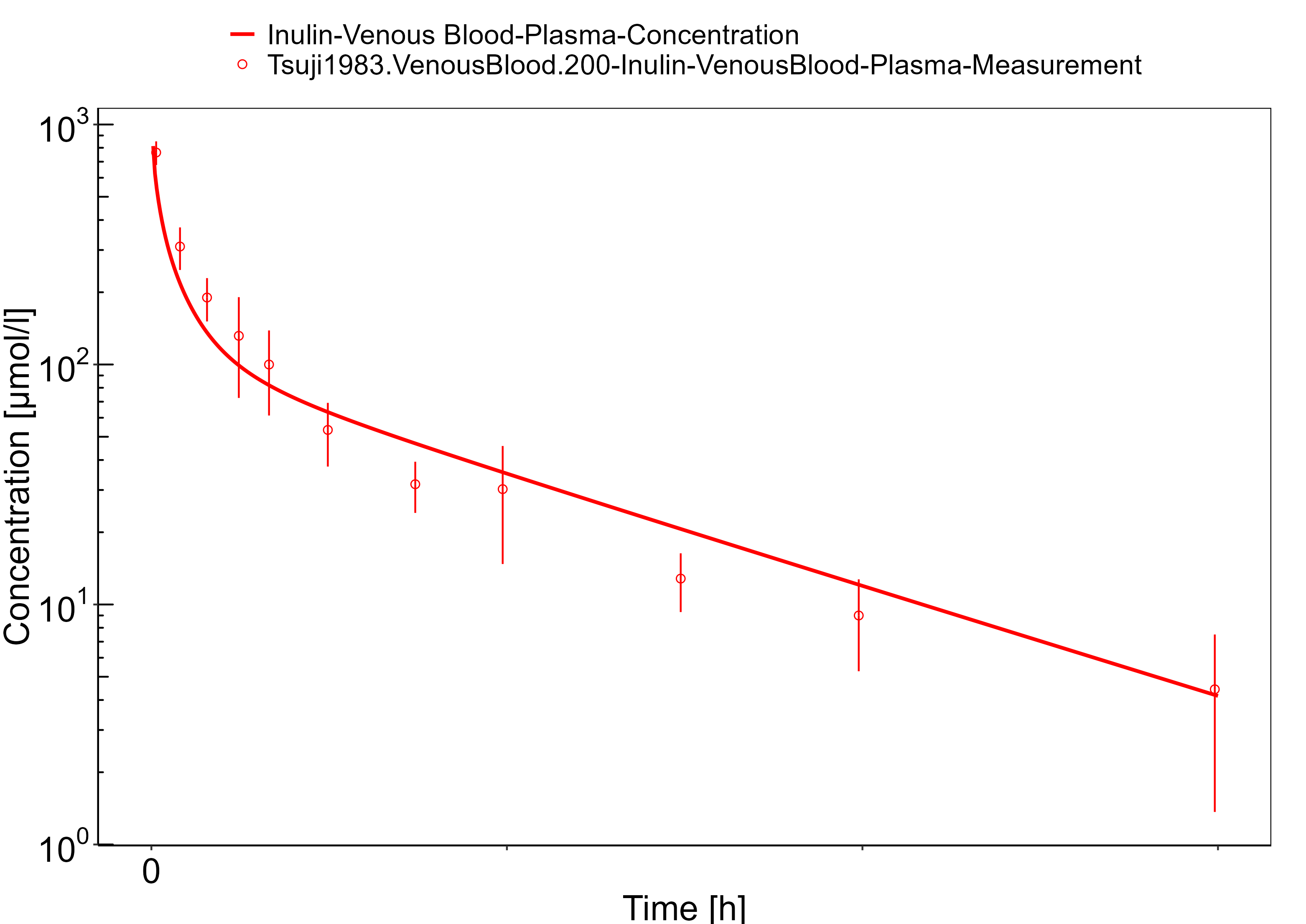
## Concentration-Time Profiles

Simulated versus observed concentration-time profiles of all data listed in [Section 2.2.2](#PK-data) are presented below.

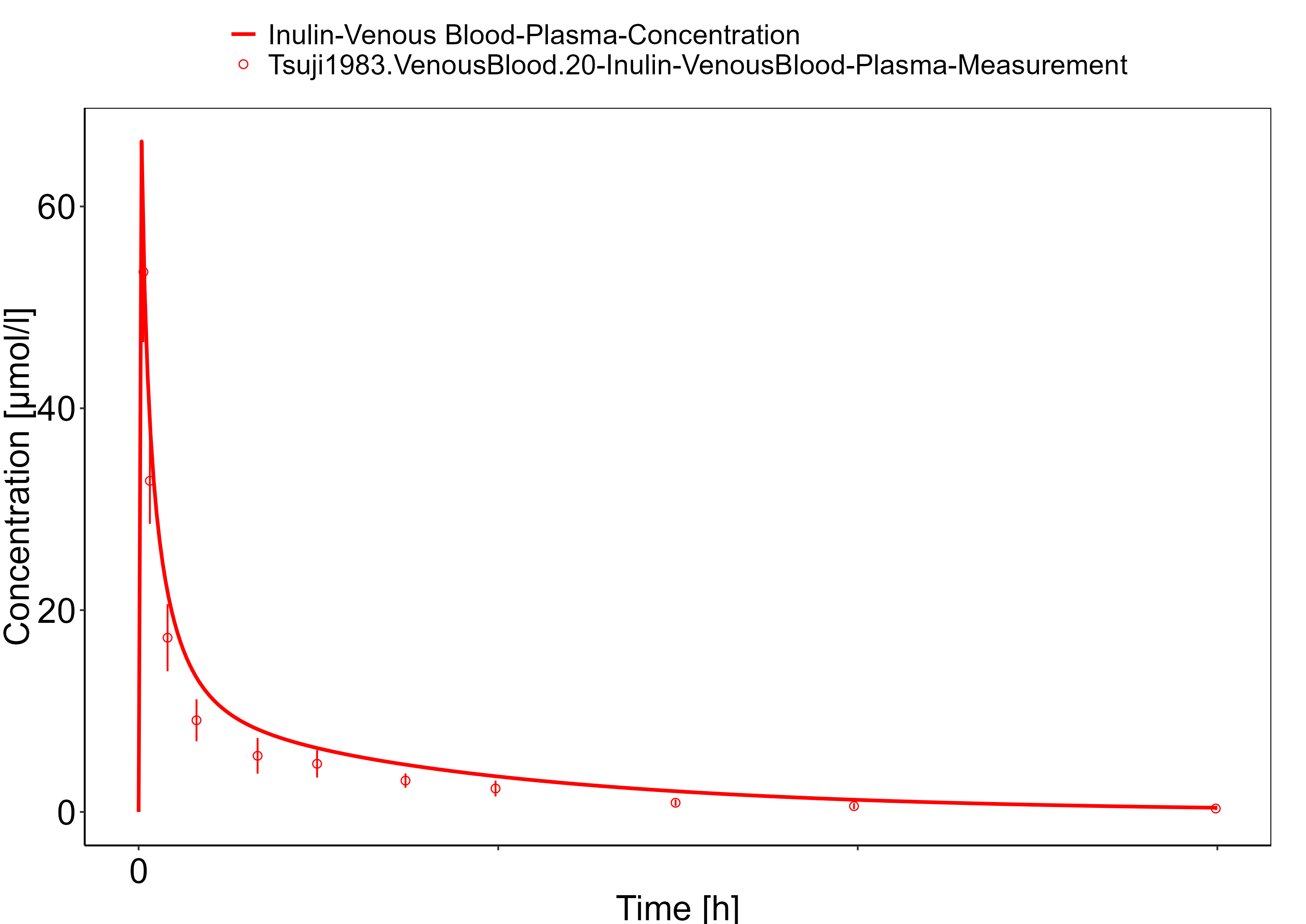
**Figure 3-3: Plasma concentration (linear scale)**



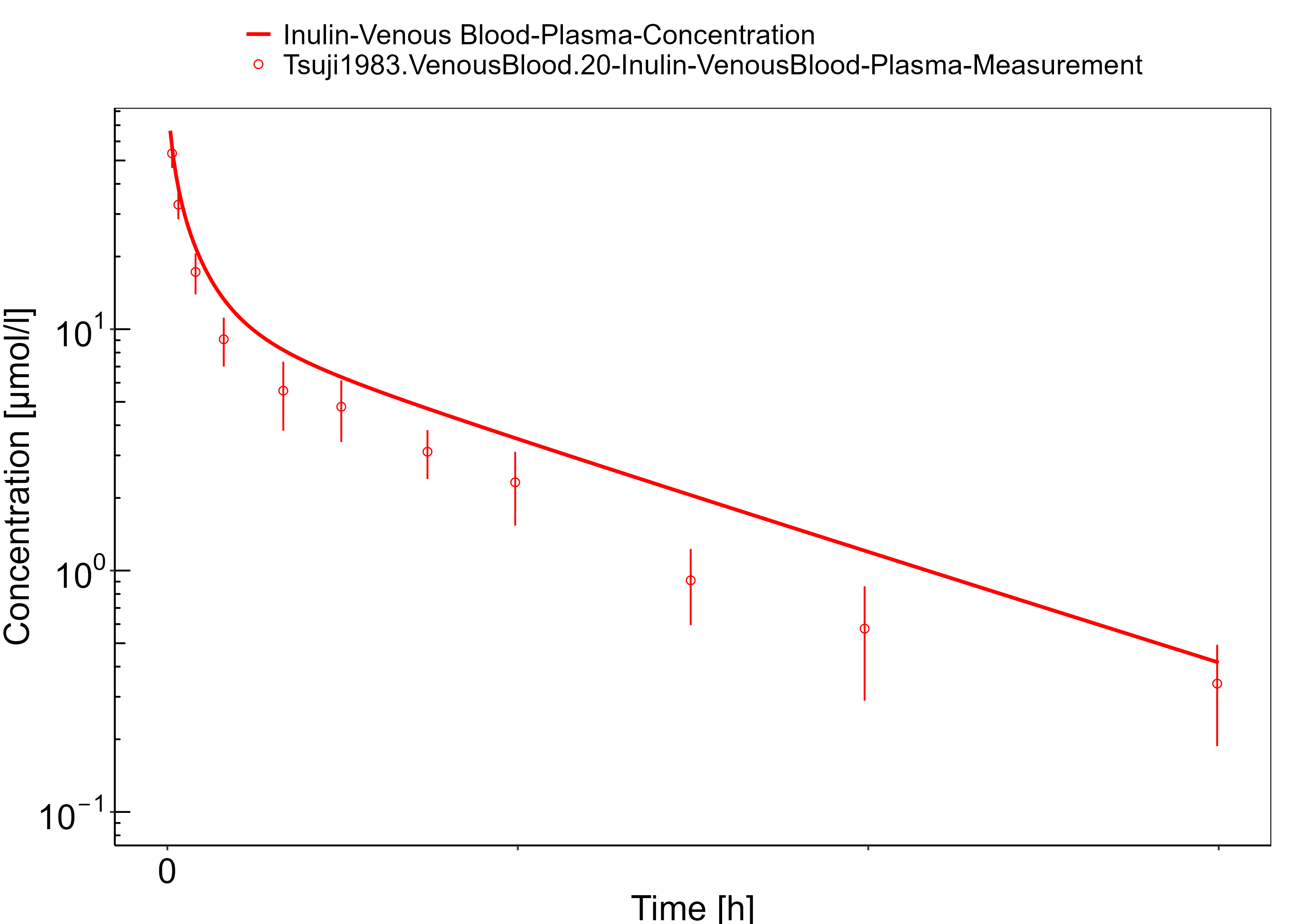
**Figure 3-4: Plasma concentration (log scale)**



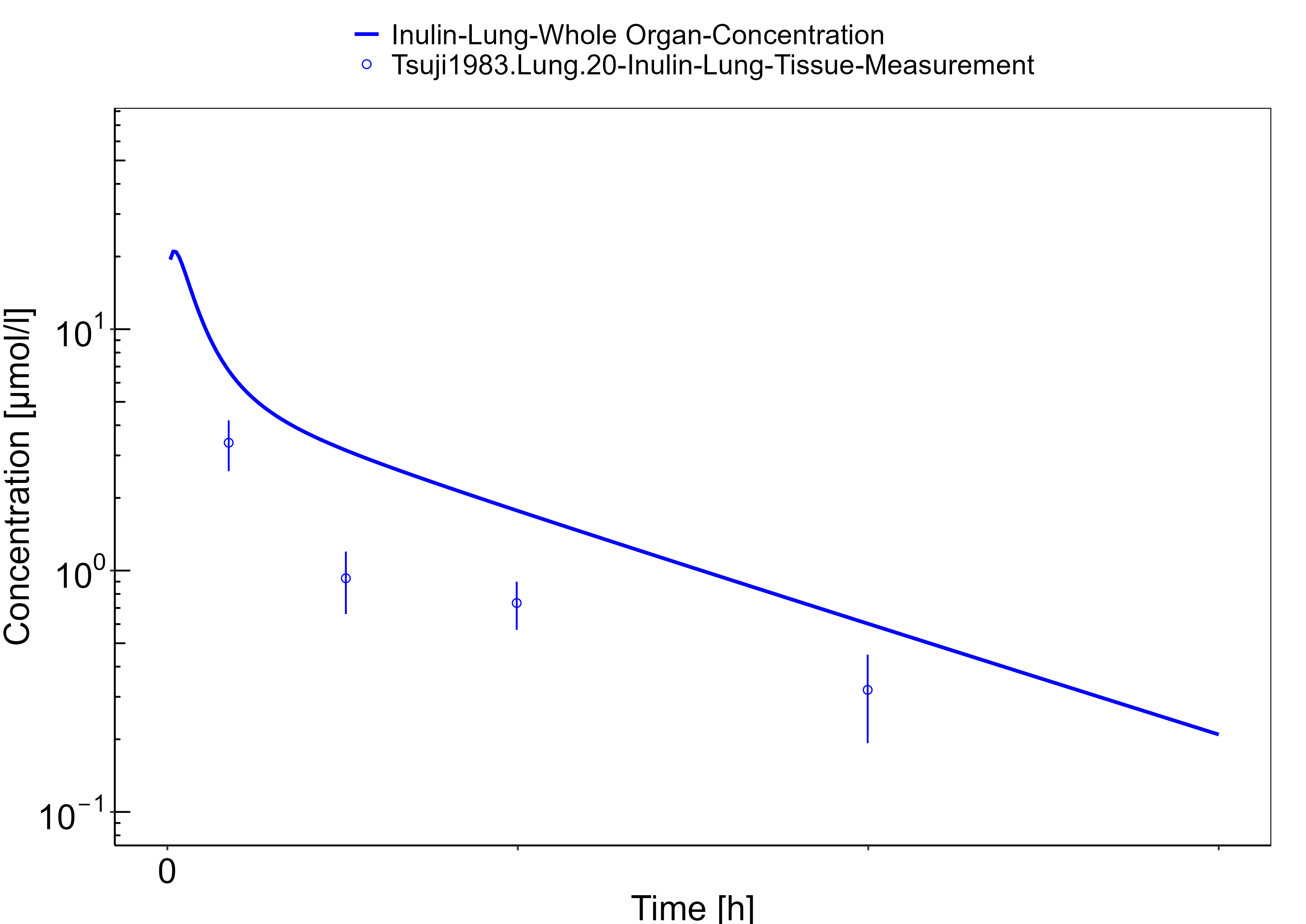
**Figure 3-5: Plasma (linear scale)**



**Figure 3-6: Plasma (log scale)**



**Figure 3-7: Lung**



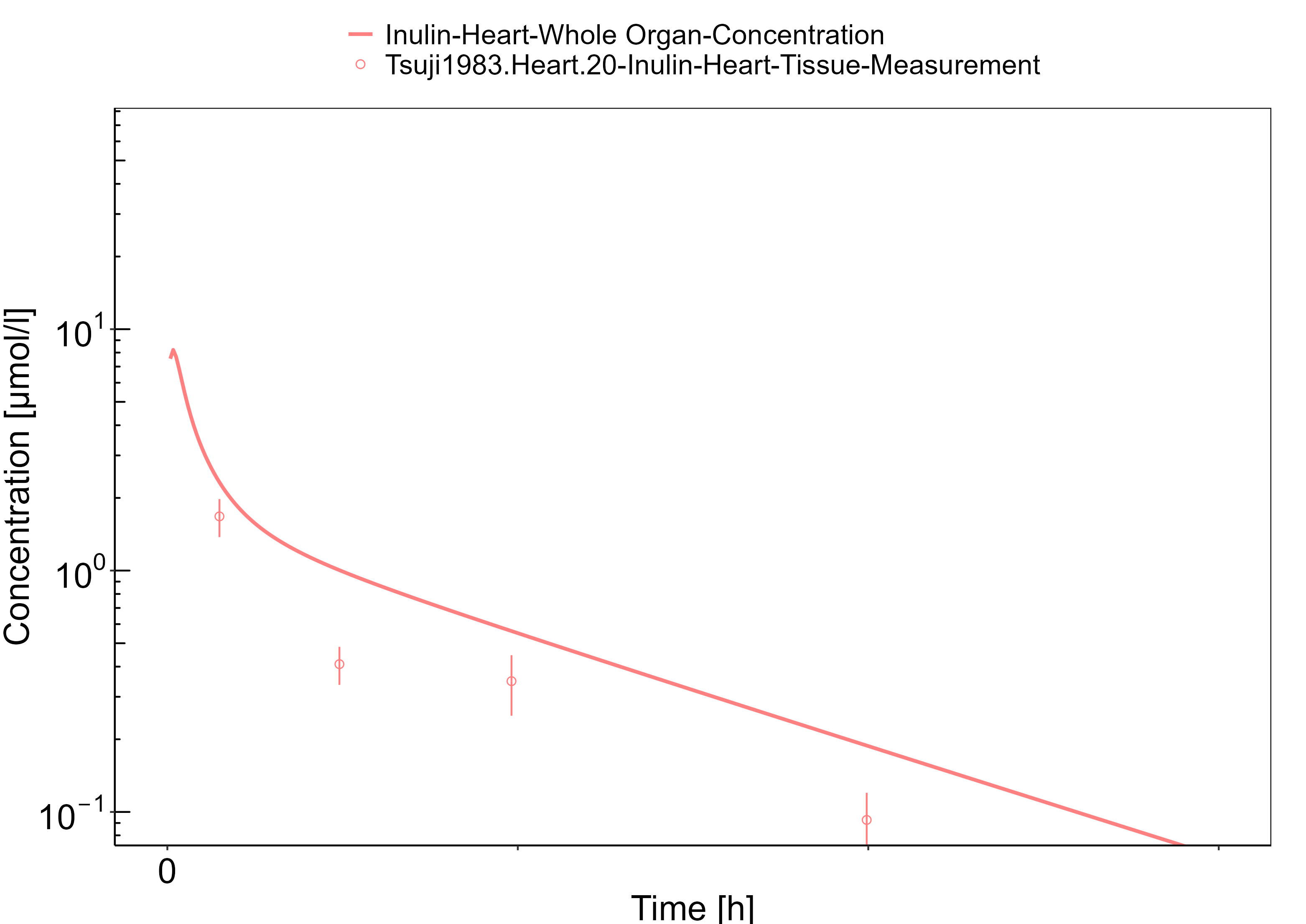
**Figure 3-8: Muscle**



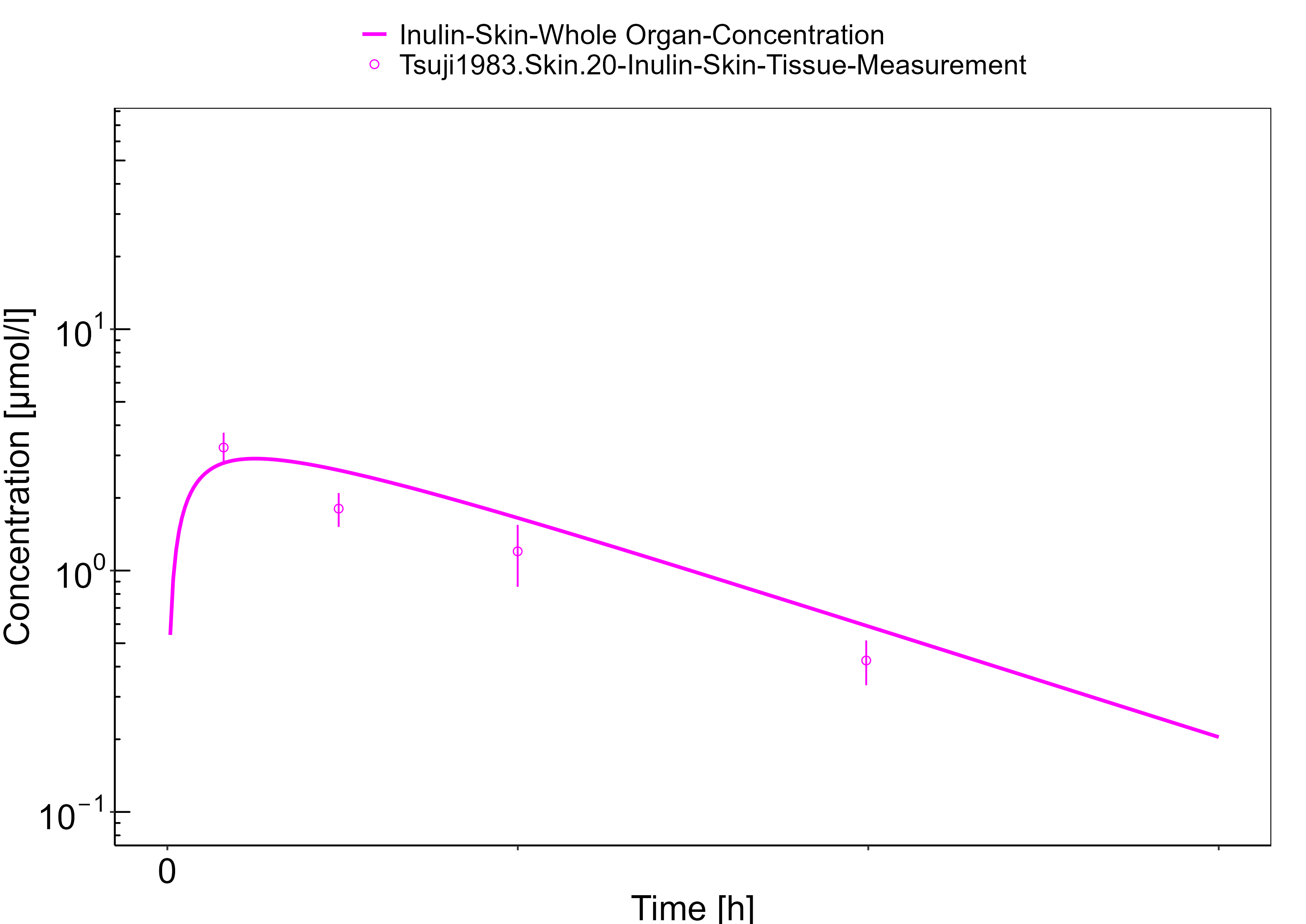
**Figure 3-9: Bone**



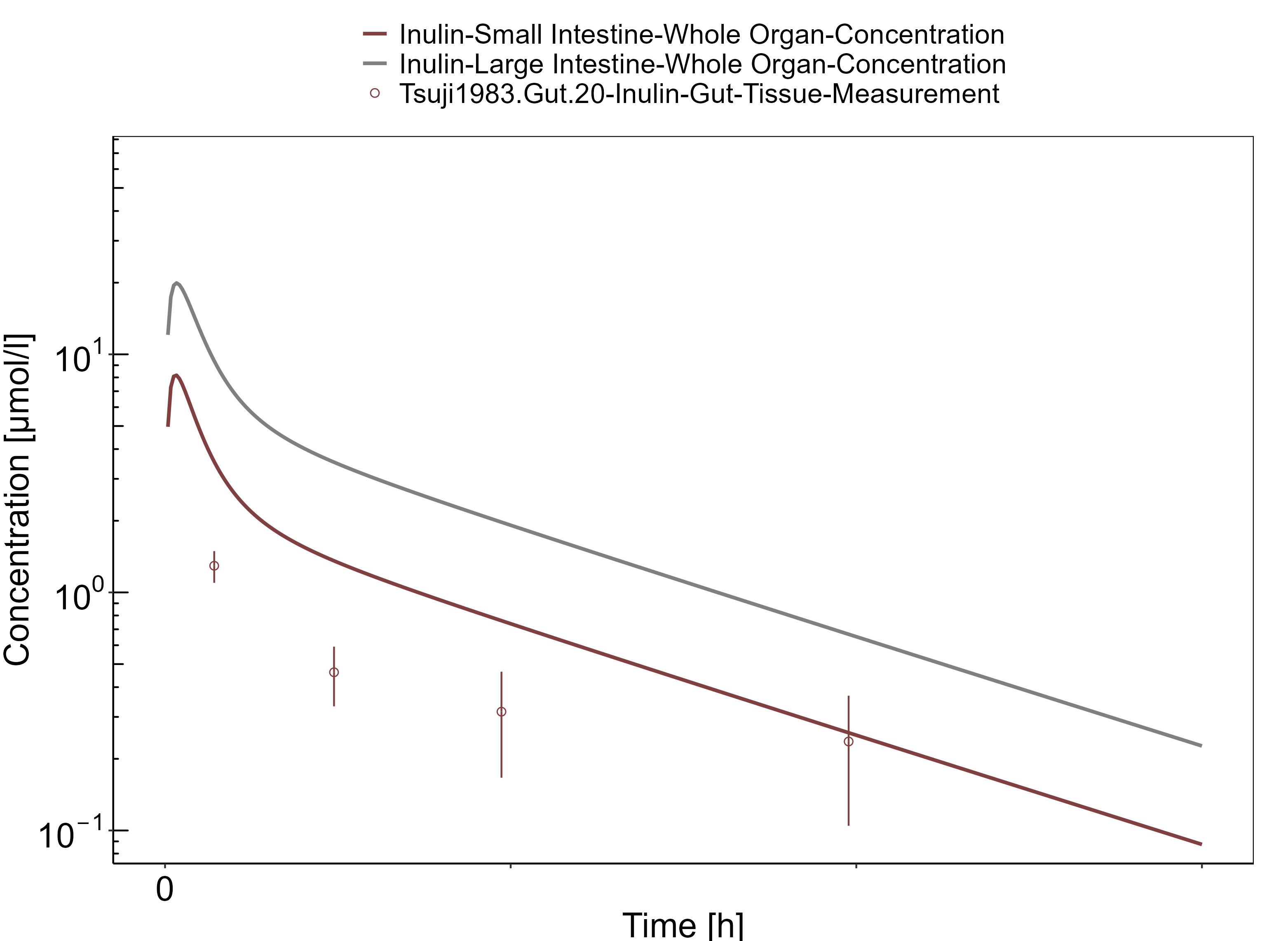
**Figure 3-10: Heart**



**Figure 3-11: Skin**



**Figure 3-12: Gut**



# Conclusion

The herein presented PBPK model overall adequately describes the plasma pharmacokinetics of inulin in rats. Tissue concentrations tend to be overestimated by the model, the largest deviations being observed for lung, gut, and heart concentrations. Simulations are predictions without adjusting any compound specific parameter (i.e., using a literature value for the solute radius of inulin and the default physiological parameters in PK-Sim especially for the glomerular filtration rate, vascular properties and compartment volumes).

# References

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