

# Building and evaluation of a PBPK model for erythromycin in healthy adults

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based on <i>Model Snapshot and Evaluation Plan</i>	<a href="https://github.com/Open-Systems-Pharmacology/Erythromycin-Model/releases/tag/v1.4">https://github.com/Open-Systems-Pharmacology/Erythromycin-Model/releases/tag/v1.4</a>
OSP Version	12.0
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This evaluation report and the corresponding PK-Sim project file are filed at:

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

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# 1 Introduction

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The presented model building and evaluation report evaluates the performance of a PBPK model for erythromycin in healthy adults.

Erythromycin a macrolide antibiotic used for the treatment of a number of bacterial infections, including respiratory tract infections, skin infections, chlamydia infections, and others. Erythromycin is available in the form of various salts and formulations, for example as:

- erythromycin lactobionate for intravenous injection
- erythromycin base in enteric-coated capsules or tablets for oral administration
- erythromycin stearate in filmcoated tablets for oral administration
- erythromycin ethylsuccinate in suspension or in filmcoated tablets for oral administration

In its free form as base, erythromycin is easily hydrolyzed in acidic aqueous solution ([Mordi 2000](#)).

Therefore, orally administered erythromycin is given in the form of enteric-coated formulations or as more acid-stable salts or esters of erythromycin (e.g. erythromycin ethylsuccinate). Once in the small intestine, erythromycin is rapidly absorbed displaying a highly variable bioavailability ([Chun 1977](#), [Mather 1981](#)). Erythromycin diffuses in most tissues and accumulates in leukocytes and phagocytes ([Miller 1984](#), [Carlier 1987](#)). About 70% of erythromycin is bound to plasma proteins ([Barre 1987](#)).

Erythromycin has been shown to be a substrate for various transporters including P-gp and OATP1B1. The latter has been shown to critically affect erythromycin disposition ([Lancaster 2012](#)). Erythromycin is extensively metabolized through N-demethylation catalyzed by CYP3A. Metabolism via CYP4F11 has also been suggested ([Kalsotra 2004](#)). Biliary excretion also appears to play an important role in erythromycin clearance ([Acocella 1968](#), [Chelvan 1979](#)), but its contribution to total elimination remains unknown. The dose fraction excreted unchanged in urine is minimal and highly variable; reported fractions after IV administration range from  $0.018 \pm 0.005$  to  $0.171 \pm 0.11$  (mean  $\pm$  SD) ([Pasic 1987](#), [Austin 1980](#)). There is abundant evidence from *in vitro* studies that erythromycin irreversibly inhibits CYP3A (e.g. [Larrey 1983](#)) and the FDA lists erythromycin as moderate index inhibitor for CYP3A. Findings from *in vivo* studies investigating the dose linearity of erythromycin pharmacokinetics are not fully conclusive, but some studies observed a slight dose dependency ([Austin 1980](#), [Josefsson 1982](#)).

# 2 Methods

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## 2.1 Modeling Strategy

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The general workflow for building an adult PBPK model has been described by Kuepfer et al. ([Kuepfer 2016](#)). Relevant information on the anthropometry (height, weight) was gathered from the respective clinical study, if reported. Information on physiological parameters (e.g. blood flows, organ volumes, hematocrit) in adults was gathered from the literature and has been incorporated in PK-Sim® as described previously ([Willmann 2007](#)). The applied activity and variability of plasma proteins and active processes that are integrated into PK-Sim® are described in the publicly available 'PK-Sim® Ontogeny Database Version 7.3' ([PK-Sim Ontogeny Database Version 7.3](#)).

The PBPK model was developed based on clinical data of healthy adult subjects obtained from the literature, covering different formulation types and erythromycin salts. Multiple doses and dosing schedules following intravenous (IV) and oral (PO) administration were included in model building. Mass balance information on urinary excretion of unchanged erythromycin after IV administration was also accounted for during the model building process.

Unknown parameters were simultaneously optimized using all available PK data, in particular:

- 4 data sets following single IV administration of 4 different doses of erythromycin (125 mg, 250 mg, 300 mg, 500 mg) as erythromycin lactobionate
- 6 data sets following single and multiple PO administration of 3 different doses of erythromycin (250 mg, 500 mg, 1000 mg) as film-coated tablets containing erythromycin stearate
- 2 data sets following single PO administration of 500 mg erythromycin as enteric-coated tablets containing erythromycin as base
- 2 data sets following single and multiple PO administration of 2 different doses of erythromycin (250 mg, 500 mg) as enteric-coated capsules containing pellets of erythromycin as base

Structural model selection was mainly guided by visual inspection of the resulting description of data and biological plausibility. The following parameters were identified using the Parameter Identification module provided in PK-Sim® and MoBi® ([Open Systems Pharmacology Documentation](#)):

- `Specific intestinal permeability (transcellular)`
- `Transport Protein - OATP1B1 - kcat`
- `Metabolizing Enzyme - CYP3A4 - kcat`
- `Total Hepatic Clearance - Specific clearance`
- `GFR fraction`
- `K_kinact_half`
- `kinact`
- `Dissolution shape` (separately for the film-coated tablet containing erythromycin stearate, the enteric-coated tablet containing erythromycin as base, and the enteric-coated capsule containing pellets of erythromycin as base)
- `Dissolution time (50% dissolved)` (separately for the film-coated tablet containing erythromycin stearate, the enteric-coated tablet containing erythromycin as base, and the enteric-coated capsule containing pellets of erythromycin as base)

- `Dissolution lag time` (separately for the enteric-coated tablet containing erythromycin as base and the enteric-coated capsule containing pellets of erythromycin as base)
- `Solubility at reference pH` (only for the enteric-coated tablet containing erythromycin as base)

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

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

## 2.2 Data

### 2.2.1 In vitro / physicochemical data

A literature search was carried out to collect available information on physicochemical properties of erythromycin. The obtained information from the literature is summarized in the table below and is used for model building.

Parameter	Unit	Literature	Description
Molecular weight	g/mol	733.9 ( <a href="#">drugbank.ca</a> )	Molecular weight
pK <sub>a</sub> (basic)		8.8 ( <a href="#">Lien 1974</a> ); 8.88 ( <a href="#">McFarland 1997</a> )	Acid dissociation constant
logP		2.48 ( <a href="#">Lien 1974</a> ); 2.92 ( <a href="#">Capobianco 1994</a> ); 3.06 ( <a href="#">McFarland 1997</a> )	Partition coefficient between octanol and water
f <sub>u</sub>		0.27 ± 0.05 <sup>a</sup> ( <a href="#">Sun 2010</a> ); 0.28 ± 0.04 <sup>a</sup> ( <a href="#">Iliopoulos 1982</a> ); 0.305 ± 0.028 <sup>a</sup> ( <a href="#">Barre 1987</a> ); 0.326 ( <a href="#">Xu 2009</a> )	Fraction unbound in human plasma
Water solubility (erythromycin lactobionate)	mg/L	200000 ( <a href="#">U.S. Patent 2,761,859</a> )	Solubility of erythromycin lactobionate in water
Water solubility (erythromycin stearate)	mg/L	182 ( <a href="#">Jones 1969</a> )	Solubility of erythromycin stearate in water
Water solubility (erythromycin base)	mg/L	2100 ( <a href="#">Manna 1998</a> )	Solubility of free erythromycin as base in water

<sup>a</sup> mean ± SD

### 2.2.2 In vitro data on mechanism-based inhibition of CYP3A

A literature search was carried out to collect quantitative information the kinetic parameters describing the mechanism-based inhibition of CYP3A by erythromycin, K<sub>i</sub> and k<sub>inact</sub>. Ample data was collected from the literature and is summarized in the table below.

Victim compound	In vitro system	Enzyme	$K_I$ [μM]	$k_{inact}$ [min <sup>-1</sup> ]	Concentration range of erythromycin [μM]	Concentration of victim compound [μM]	Reference
testosterone	rec cDNA CYP3A4	CYP3A4	1.04	0.0293	0 - 20	100	Akiyoshi 2013
testosterone	rec cDNA CYP3A4	CYP3A4	1.21	0.0164	0 - 20	100	Akiyoshi 2013
testosterone	rec cDNA CYP3A4	CYP3A4	0.415	0.0159	0 - 20	100	Akiyoshi 2013
testosterone	rec cDNA CYP3A4	CYP3A4	2.24	0.0293	0 - 20	100	Akiyoshi 2013
testosterone	rec cDNA CYP3A4	CYP3A4	0.753	0.0248	0 - 20	100	Akiyoshi 2013
midazolam	rec cDNA CYP3A4	CYP3A	9.5	0.16	50-fold range	10	Atkinson 2005
midazolam	rec cDNA CYP3A4	CYP3A	8.82	0.12	50-fold range	10	Atkinson 2005
midazolam	rec cDNA CYP3A4	CYP3A	8.3	0.09	50-fold range	10	Atkinson 2005
testosterone	rec cDNA CYP3A4	CYP3A	9.5	0.06	4 - 200	16.7	Atkinson 2005
triazolam	HLM	CYP3A	5.4	0.069		400	Aueviriyavit 2010
testosterone	rec cDNA CYP3A4	CYP3A4	5.3	0.12	5 - 20	200	Chan 2000
midazolam	cryopreserved HEP	CYP3A	60	0.081	3 - 300	30	Chen 2011
midazolam	cryopreserved HEP	CYP3A	67.9	0.079	3 - 300	30	Chen 2011
midazolam	rec cDNA CYP3A4	CYP3A4	0.762	0.0648	0.1 - 30	4	Ishikawa 2017
testosterone	rec cDNA CYP3A4	CYP3A4	1.00	0.0604	0.1 - 30	150	Ishikawa 2017
nifedipine	rec cDNA CYP3A4	CYP3A4	0.794	0.0766	0.1 - 30	6	Ishikawa 2017
triazolam	HLM	CYP3A4	15.9	0.062	3 - 100	300	Kanamitsu 2000
triazolam	HLM	CYP3A4	17.4	0.055	3 - 100	300	Kanamitsu 2000
triazolam	rec cDNA CYP3A4	CYP3A4	19.1	0.173	3 - 100	300	Kanamitsu 2000
triazolam	rec cDNA CYP3A4	CYP3A4	18.9	0.097	3 - 100	300	Kanamitsu 2000
testosterone	HLM	CYP3A4	29.4	0.0271		250	Kosaka 2017
testosterone	HLM	CYP3A4	30	0.040	0.3 - 300		Kozakai 2013
midazolam	HLM	CYP3A4	12	0.035	0.3 - 300		Kozakai 2013
midazolam	HLM	CYP3A4	20	0.033	0.3 - 300		Kozakai 2013
midazolam	cryopreserved HEP	CYP3A	25.15	0.08	0.13 - 100	30	Mao 2011
midazolam	HLM	CYP3A	10.8	0.032			Mao 2016
midazolam	cryopreserved HEP	CYP3A	30.7	0.05	0 - 300	20	Mao 2016
midazolam	cryopreserved HEP	CYP3A	59.2	0.062	0 - 100	5	Mao 2016
midazolam	cryopreserved HEP	CYP3A	80.3	0.052	0 - 100	20	Mao 2016
midazolam	rec cDNA CYP3A4	CYP3A4	7.47	0.042	2 - 50	8	McConn 2004
midazolam	HLM	CYP3A4	10.9	0.046	2 - 100	8	McConn 2004

Victim compound	In vitro system	Enzyme	$K_I$ [μM]	$k_{inact}$ [min <sup>-1</sup> ]	Concentration range of erythromycin [μM]	Concentration of victim compound [μM]	Reference
midazolam	primary HEP	CYP3A	11	0.07	0.1 - 10	3	McGinnity 2006
midazolam	HLM	CYP3A4	10	0.036	0 - 25		Obach 2007
testosterone	HLM	CYP3A4	9.8	0.039	0 - 25	500	Obach 2007
testosterone	rec cDNA CYP3A4	CYP3A4	0.92	0.058	5 - 100	250	Polasek 2006
testosterone	HLM	CYP3A4	12.8	0.037	5 - 100	250	Polasek 2006
midazolam	rec cDNA CYP3A4	CYP3A4	5.1	0.30	0.5 - 50	100	Ring 2005
testosterone	rec cDNA CYP3A4	CYP3A4	0.92	0.058			Teng 2010
testosterone	HLM	CYP3A4	4.579	0.0115			Teng 2010
domperidone	HLM	CYP3A4	18.4	0.022	2.5 - 200	500	Ung 2009
domperidone	rec cDNA CYP3A4	CYP3A4	4.1	0.026	2.5 - 200	500	Ung 2009
midazolam	HLM	CYP3A4	12.1	0.0215	0 - 100	25	Watanabe 2007
nifedipine	HLM	CYP3A4	11.3	0.0295	0 - 100	50	Watanabe 2007
testosterone	HLM	CYP3A4	10.9	0.0352	0 - 100	200	Watanabe 2007
midazolam	HLM	CYP3A	1.48	0.017	0.5 - 500	20	Xu 2009
midazolam	cryopreserved HEP	CYP3A	20.0	0.016	0.5 - 500	20	Xu 2009
midazolam	cryopreserved HEP	CYP3A	109	0.055	0.5 - 500	20	Xu 2009
midazolam	HLM	CYP3A	81.8	0.0665	20 - 400	10	Yamano 2001
midazolam	HLM	CYP3A	15.7	0.1			Zhang 2006
testosterone	rec cDNA CYP3A4	CYP3A4	5	0.34	1 - 50	200	Zhang 2009
testosterone	HLM	CYP3A4	15.7	0.09	1 - 50	200	Zhang 2009
midazolam	HLM	CYP3A4	26.5	0.041	2.5 - 50	10	Zimmerlin 2011

**Note:** Abbreviations: HEP: human hepatocytes; HLM: human liver microsomes; rec cDNA CYP3A4: human recombinant c-DNA CYP3A4 enzymes (e.g. supersomes, baculovirus-insect cell system, E. coli transfected cells)

The data listed in the Table above can be statistically summarized as follows:

Parameter [unit]	Min	Q1	Geometric mean	Median	Arithmetic mean	Q3	Max
$K_I$ [μM]	0.420	4.89	8.71	10.9	18.4	19.3	109
$k_{inact}$ [min <sup>-1</sup> ]	0.0115	0.0314	0.0504	0.0535	0.0664	0.0772	0.340

## 2.2.3 Clinical data

A literature search was carried out to collect available PK data on erythromycin in healthy adults. The following data from the publications listed below were used for model building and evaluation:

<b>Publication</b>	<b>Study description</b>
Austin 1980	IV administration of 125 mg, 250 mg, 500 mg, and 900 mg as erythromycin lactobionate; single dose
Barre 1987	IV administration of 500 mg as erythromycin lactobionate; single dose
Berend 1979	PO administration of 500 mg erythromycin stearate; multiple dose
Birkett 1990	PO administration of 250 mg as enteric-coated capsules containing pellets of erythromycin base; single and multiple dose
Brannan 1995	PO administration of 500 mg erythromycin stearate; multiple dose
DiSanto 1981	PO administration of 500 mg as unprotected tablets containing erythromycin base, as film-coated tablets containing erythromycin base, as enteric-coated tablets containing erythromycin base and as film-coated tablets containing erythromycin stearate; single and multiple dose
Henry 1980	PO administration of 500 mg erythromycin stearate; single dose
Huppertz 2011	IV administration of 1000 mg as erythromycin lactobionate; single dose
Iliopoulou 1982	PO administration of 500 mg as film-coated tablet containing erythromycin stearate; multiple dose
Josefsson 1982	PO administration of 500 mg as film-coated tablet containing erythromycin stearate and 250 mg, 500 mg, 1000 mg as enteric coated capsules containing pellets of erythromycin base; single dose
Kavi 1988	PO administration of 500 mg as film-coated tablet containing erythromycin stearate; single dose
Kivistö 1997	PO administration of 500 mg as enteric-coated capsules containing erythromycin base; multiple dose
Kroboth 1982	PO administration of 500 mg as enteric-coated tablets containing erythromycin base; single dose
Malmborg 1978	PO administration of 500 mg as film-coated tablet containing erythromycin stearate; multiple dose
Miglioli 1990	PO administration of 1000 mg erythromycin stearate; multiple dose
Olkkola 1993	Midazolam-erythromycin interaction study; PO administration of 500 mg as enteric-coated tablet containing erythromycin base; multiple dose
Parsons 1977	PO administration of 500 mg erythromycin stearate; single dose
Pasic 1987	IV administration of 300 mg as erythromycin lactobionate; single dose
Posti 1983	PO administration of 500 mg as film-coated tablets containing erythromycin stearate, as enteric-coated tablets containing erythromycin base, and as enteric-coated tablets containing erythromycin stearate; single dose
Schreiner 1984	PO administration of 500 mg as film-coated tablets containing erythromycin stearate and as enteric-coated capsule containing pellets of erythromycin base; single dose
Shanson 1984	PO administration of 1500 mg erythromycin stearate; single dose
Simon 1980	IV administration of 500 mg erythromycin lactobionate and PO administration of 500 mg erythromycin stearate; single dose
Sun 2010	IV administration of 125 mg and PO administration of 250 mg (salt and formulation type not specified); single dose
Yakatan 1979	PO administration of 250 mg as film-coated tablet containing erythromycin stearate; single dose

Publication	Study description
Yakatan 1980	PO administration of 250 mg as film-coated tablet containing erythromycin stearate and as enteric-coated tablet containing erythromycin base; single and multiple dose

## 2.3 Model Parameters and Assumptions

### 2.3.1 Dissolution and absorption

Development of an adequate absorption model for erythromycin was complicated by the large intersubject variability in the absorption kinetics of erythromycin (Chun 1977, Mather 1981). Additionally, multiple formulation types are available entailing different dissolution and absorption kinetics (Chun 1977, Yakatan 1980, Mather 1981). The herein presented model was developed for the following oral dosage forms:

- film-coated tablets containing erythromycin stearate
- enteric-coated tablets containing erythromycin as free base
- enteric-coated capsules containing pellets of erythromycin as free base

The following sections briefly address how each of these formulations was modeled in presented PBPK models.

#### 2.3.1.1 Film-coated tablet containing erythromycin stearate

Comparison of the reported erythromycin PK following administration of different doses of film-coated tablets containing the stearate salt indicated that the lowest dose (250 mg) yields a larger  $t_{max}$  than the higher doses (500, 1000 and 1500 mg). Different dissolution kinetics were therefore used for this formulation type; relatively slow dissolution kinetics was used for the 250 mg dose, whereas dissolution of the higher doses (500, 1000 and 1500 mg) was described by immediate-release kinetics. Both types of dissolution kinetics were described by the Weibull function. The parameters `Dissolution shape` and `Dissolution time (50% dissolved)` were optimized to optimally describe the clinical data. Additionally, solubility was also optimized as it was found during model building that solubility-limited absorption of the high doses captured the observed PK profiles better.

#### 2.3.1.2 Enteric-coated tablet containing erythromycin as free base

Relatively few PK data were available for enteric-coated tablets containing erythromycin as free base. For this formulation type, PK profiles were only available for the 250 mg and 500 mg dose. For these two doses, dissolution was described by the Weibull function. To account for the enteric coating (protecting the tablet from dissolving in the stomach), a `Lag time` > 0 min was applied in the model. To optimally describe the clinical data, relevant parameters of the Weibull function were optimized, namely the `Dissolution shape`, `Dissolution time (50% dissolved)` and `Lag time`. Similar to the film-coated tablet containing the stearate salt, it was found that the observed PK profiles of the enteric-coated tablet containing erythromycin as free base could be better described when optimizing solubility.

#### 2.3.1.3 Enteric-coated capsule containing pellets of erythromycin as free base

Comparison of the reported erythromycin PK following administration of different doses of enteric-coated capsules containing pellets of erythromycin base showed large variability in  $C_{max}$  and  $t_{max}$  both within and between different doses (dose range: 250 mg - 1000 mg). Here, dissolution kinetics was also described by the Weibull function with relevant parameters (`Dissolution shape`, `Dissolution time`,

(50% dissolved) and lag time) being optimized to better describe clinical data. However, optimizing solubility did not result in a better description of the observed PK profiles. Consequently, solubility of erythromycin in this formulation type was fixed to a high value (500 mg/L) to avoid any solubility-related limitations in dissolution and absorption.

For all three formulations, the same specific intestinal permeability (transcellular) was used in the PBPK model. This parameter was also optimized to better describe the clinical PK data.

### 2.3.2 Distribution

With an average fraction unbound in human plasma of approximately 0.30, erythromycin is moderately protein-bound. In the developed model, the fraction unbound (plasma, reference value) was set to 0.305 which is the value reported by Barre et al. (Barre 1987). Lipophilicity was fixed to a value of 2.82 which is the average of three experimentally measured values reported in the literature (Lien 1974, Capobianco 1994, McFarland 1997). The observed PK data were found to be best described using the model for estimating intracellular-to-plasma partition coefficients according to the method by Rodgers and Rowland (Rodgers 2005, Rodgers 2006). Cellular permeabilities were automatically calculated using the method charge dependent schmitt (Open Systems Pharmacology Documentation). Active transfer of erythromycin by OATP1B1 was modeled as Michaelis-Menten kinetics using an Influx transporter type; the Km value of 13.2  $\mu$ M was taken from the literature (Lancaster 2012) and kcat was optimized to best match observed clinical data. The gene expression profile of OATP1B1 (default symbol for the gene: LST-3TM12) was loaded from the internal PK-Sim® database using the expression data quantified by RT-PCR (Open Systems Pharmacology Documentation).

### 2.3.3 Elimination

Erythromycin is extensively metabolized via N-demethylation catalyzed by CYP3A. Kinetics of this biotransformation was described by a Michaelis-Menten process. The following kinetic parameters erythromycin N-demethylation have been measured in human liver microsomes (HLM) and reported in the literature (mean  $\pm$  standard error):

Vmax [nmol/min/mg mic protein]	Km [ $\mu$ M]	Microsomal preparation	Reference
2 $\pm$ 0.09	78 $\pm$ 9	HLM from donor HL 3926	Wang 1997
0.41 $\pm$ 0.02	44 $\pm$ 7	HLM from donor HL 24493	Wang 1997
0.345 $\pm$ 0.013	88 $\pm$ 10	mixed HLM pool	Riley 1997

In the PBPK model, Vmax and Km were fixed to the mean of the values tabulated above (70  $\mu$ M and 0.918 nmol/min/mg mic protein). The gene expression profile of CYP3A4 was loaded from the internal PK-Sim® database using the expression data quantified by RT-PCR (Open Systems Pharmacology Documentation).

Although it has also been observed that erythromycin is metabolized via CYP4F11 *in vitro* (Kalsotra 2004), this elimination pathway was not accounted for in the model because its contribution to overall elimination was assumed to be low. In humans, CYP4F11 is mainly expressed in the liver and to a much lesser extent in the kidney (Cui 2000) and the CYP4F family makes up approximately 15% of all hepatic CYP enzymes (Michaels 2014). The K<sub>m</sub> and V<sub>max</sub> values for the CYP4F11-mediated biotransformation reported by Kalsotra 2004 are similar to those measured for CYP3A4 (Riley 1997, Wang 1997), suggesting that the relative mass balance of these two metabolism pathways mainly

depends on the absolute amount of each enzyme in the liver. While no information on total CYP4F11 in the human liver could be found in the literature, CYP4F11 expression in the liver of cynomolgus monkeys was observed to be approximately 6-fold lower than that of CYP3A4 (Uehara 2015). Hence, it was assumed that CYP4F11-mediated metabolism of erythromycin can be neglected in humans.

Additional elimination pathways suggested for erythromycin are acid-catalyzed degradation (hydrolysis) in the acidic milieu of the stomach (Mordi 2000) and biliary excretion (Acocella 1968, Chelvan 1979), but no quantitative information on the mass balance of these pathways could be found in the literature. Additionally, mechanism-based inhibition of CYP3A4 by erythromycin might constitute another clearance process which was neither considered in the model. However, a `total hepatic clearance` process was implemented in the model which could at least partly account for other elimination pathways not explicitly accounted for in the model. Of note, despite the name `total hepatic clearance`, this clearance pathway was implemented as dummy clearance accounting for additional elimination processes that are not covered by CYP3A-mediated clearance and unchanged renal excretion and it should hence rather be regarded as a partial than a total clearance.

The reported dose fractions of erythromycin undergoing unchanged renal excretion after IV administration range from  $0.018 \pm 0.005$  to  $0.171 \pm 0.11$  (mean  $\pm$  SD) (Pasic 1987, Austin 1980). This information was accounted for in the model by implementing a glomerular filtration process and optimizing the `GFR fraction` to match the observed dose fractions excreted unchanged in urine.

### 2.3.4 Autoinhibition via CYP3A4

In the scientific literature, large ranges have been reported for  $K_I$  and  $k_{inact}$  (Section 2.2.2). Since the exact values are unknown, `K_kinact_half` and `kinact` were both optimized within the observed range (see Section 2.2.2) during model building to best match the observed clinical data.

To better inform optimization of these two parameters, clinical data of a midazolam-erythromycin interaction study conducted by Olkkola et al. (Olkkola 1993) were included in the parameter optimization during model building. Therefore, the midazolam PBPK model v0.9 available on OSP GitHub (<https://github.com/Open-Systems-Pharmacology/Midazolam-Model/releases/tag/0.9>) was loaded in the PK-Sim® erythromycin file and the study by Olkkola et al. (Olkkola 1993) was simulated. However, instead of using the reported midazolam plasma concentrations as observed data in the parameter identification, the AUC of midazolam was used. More specifically, a midazolam target AUC after IV and oral administration was calculated by multiplying the simulated midazolam AUC (24.3 and 54.0  $\mu\text{mol min/L}$  and after IV and oral administration, respectively) with the observed geometric mean AUC ratio (1.96 and 4.07 after IV and PO administration, respectively) (Olkkola 1993) resulting in target AUCs of 47.4 and 220  $\mu\text{mol min/L}$  after IV and oral administration of midazolam, respectively. These values were included as observed data values in the parameter identification during model building. Since the AUC is not a default output that can directly be used in the parameter identification, the PBPK model structure was modified prior to running the parameter identification as described in the following. After exporting the model to MoBi®, an artificial reaction of a dummy molecule was created. The reaction rate was defined as the simulated peripheral venous blood plasma concentration of midazolam, hence yielding the AUC at any specific time point. Thereafter, the model was imported in PK-Sim® and included in the parameter identification. After being used in the parameter identification during model building, the model was not used any further.

## 2.3.5 Inhibition of P-gp

The model also includes competitive inhibition of P-gp with a  $K_i$  of 22.7  $\mu\text{mol/L}$  (Eberl 2007).

## 2.3.6 Automated Parameter Identification

This is the result of the final parameter identification:

Model Parameter	Formulation type/salt form	Optimized Value	Unit
Dissolution shape	Enteric coated pellets	1.0564916105	
Dissolution shape	Enteric coated tablet	1.0838799888	
Dissolution shape	Filmcoated tablet (except 250 mg dose)	1.0960212213	
Dissolution shape	Filmcoated tablet, 250 mg	3.2811974117	
Dissolution time (50% dissolved)	Enteric coated pellets	1.7462743767	min
Dissolution time (50% dissolved)	Enteric coated tablet	79.6337524677	min
Dissolution time (50% dissolved)	Filmcoated tablet (except 250 mg dose)	1.7038947098	min
Dissolution time (50% dissolved)	Filmcoated tablet, 250 mg	83.6562552486	min
GFR fraction		1.1591081815	
$K_{kinact,half}$ (CYP3A4)		7.6007360452	$\mu\text{mol/L}$
$k_{cat}$ (OATP1B1)		2.0201069202*	1/min
$kinact$ (CYP3A4)		0.0296261146	1/min
$km$ (OATP1B1)		0.735836485	$\mu\text{mol/L}$
Lag time	Enteric coated pellets	54.3490442506	min
Lag time	Enteric coated tablet	78.7967495765	min
Solubility at ref pH	Enteric coated tablet, erythromycin base	8.3990771997	mg/L
Solubility at ref pH	Filmcoated tablet, erythromycin stearate	28.0708790976	mg/L
Specific clearance		4.1462183378	1/min
Specific intestinal permeability		0.00038668371665	cm/min

\* The value in the model was updated to 1.350032201 with the release of PK-Sim 10 to account for the updated calculation method of interstitial concentrations (please refer to the respective [release notes of version 10](#)).

# 3 Results and Discussion

The PBPK model for erythromycin was developed and verified with clinical pharmacokinetic data.

The next sections show:

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

## 3.1 Final input parameters

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

### Compound: Erythromycin

#### Parameters

Name	Value	Value Origin	Alternative	Default
Solubility at reference pH	28.0708790976 mg/l	Parameter Identification-Parameter Identification-Value updated from '003 - BP conc ratio FIX' on 2019-11-14 14:43	Erythromycin stearate film-coated tablet	True
Reference pH	7	Parameter Identification-Parameter Identification-Value updated from '001' on 2019-10-01 13:46	Erythromycin stearate film-coated tablet	True
Solubility at reference pH	500 mg/l	Parameter Identification-Parameter Identification-Value updated from '001' on 2019-10-01 13:46	Erythromycin base enteric coated pellets	False
Reference pH	7	Parameter Identification-Parameter Identification-Value updated from '001' on 2019-10-01 13:46	Erythromycin base enteric coated pellets	False
Solubility at reference pH	200 mg/ml	Publication-In Vitro-Hoffhine, Jr Charles E. "Aqueous soluble salts of erythromycin." U.S. Patent 2,761,859, issued September 4, 1956.	Erythromycin lactobionate	False
Reference pH	7	Publication-In Vitro-Hoffhine, Jr Charles E. "Aqueous soluble salts of erythromycin." U.S. Patent 2,761,859, issued September 4, 1956.	Erythromycin lactobionate	False
Solubility at reference pH	8.3990771997 mg/l	Parameter Identification-Parameter Identification-Value updated from 'Parameter Identification 1' on 2020-01-13 15:01	Erythromycin base enteric coated tablet	False
Reference pH	7		Erythromycin base enteric coated tablet	False

Name	Value	Value Origin	Alternative	Default
Lipophilicity	2.82 Log Units	Parameter Identification-Parameter Identification-Value updated from 'Parameter Identification 1' on 2020-01-13 15:01	Literature (average value)	True
Fraction unbound (plasma, reference value)	0.305	Publication-In Vivo-PMID: 3606934	Barre 1987	True
Specific intestinal permeability (transcellular)	0.00038668371665 cm/min	Parameter Identification-Parameter Identification-Value updated from 'Parameter Identification 1' on 2020-01-13 15:01	Fitted	True
Is small molecule	Yes			
Molecular weight	733.927 g/mol	Internet-drugbank.ca		
Plasma protein binding partner	Unknown			

## Calculation methods

Name	Value
Partition coefficients	Rodgers and Rowland
Cellular permeabilities	Charge dependent Schmitt

## Processes

### Metabolizing Enzyme: CYP3A4-Biotransformation\_fitted

Molecule: CYP3A4

#### Parameters

Name	Value	Value Origin
In vitro Vmax for liver microsomes	918.33333 pmol/min/mg mic. protein	Publication-In Vitro-PMID: 9566442
Km	70 µM	Publication-In Vitro-Average of reported values in the literature (PMID: 9107550 and PMID: 9566442)

### Systemic Process: Glomerular Filtration-fitted

Species: Human

#### Parameters

Name	Value	Value Origin
GFR fraction	1.1591081815	Parameter Identification-Parameter Identification-Value updated from 'Parameter Identification 1' on 2020-01-13 15:01

## Inhibition: CYP3A4-MBI

Molecule: CYP3A4

### Parameters

Name	Value	Value Origin
kinact	0.0296261146 1/min	Parameter Identification-Parameter Identification-Value updated from 'Parameter Identification 1' on 2020-01-13 15:01
K_kinact_half	7.6007360452 μmol/l	Parameter Identification-Parameter Identification-Value updated from 'Parameter Identification 1' on 2020-01-13 15:01

## Transport Protein: OATP1B1-fitted

Molecule: OATP1B1

### Parameters

Name	Value	Value Origin
Transporter concentration	1 μmol/l	
Vmax	11.66 pmol/ml/min	Publication-In Vitro-PMID: 22990751
Km	0.735836485 μmol/l	Parameter Identification-Parameter Identification-Value updated from 'Parameter Identification 1' on 2020-01-13 15:01
kcat	1.350032201 1/min	Parameter Identification-Parameter Identification-Value updated from 'Parameter Identification 1' on 2020-01-13 15:01

## Systemic Process: Total Hepatic Clearance-fitted

Species: Human

### Parameters

Name	Value	Value Origin
Fraction unbound (experiment)	0.305	
Lipophilicity (experiment)	2.48 Log Units	
Plasma clearance	0 ml/min/kg	
Specific clearance	4.1462183378 1/min	Parameter Identification-Parameter Identification-Value updated from 'Parameter Identification 1' on 2020-01-13 15:01

## Inhibition: P-gp-Eberl2007

Molecule: P-gp

## Parameters

Name	Value	Value Origin
Ki	22.7 μmol/l	Publication-Eberl S, Renner B, Neubert A, Reisig M, Bachmakov I, König J, Dörje F, Mürdter TE, Ackermann A, Dormann H, Gassmann KG, Hahn EG, Zierhut S, Brune K, Fromm MF. Role of p-glycoprotein inhibition for drug interactions: evidence from in vitro and pharmacoepidemiological studies. Clin Pharmacokinet. 2007;46(12):1039-49. doi: 10.2165/00003088-200746120-00004. PMID: 18027988.

## Formulation: Erythromycin\_Weibull\_enteric-coated-pellets

Type: Weibull

## Parameters

Name	Value	Value Origin
Dissolution time (50% dissolved)	1.7462743767 min	Parameter Identification-Parameter Identification-Value updated from 'Parameter Identification 1' on 2020-01-13 15:01
Lag time	54.3490442506 min	Parameter Identification-Parameter Identification-Value updated from 'Parameter Identification 1' on 2020-01-13 15:01
Dissolution shape	1.0564916105	Parameter Identification-Parameter Identification-Value updated from 'Parameter Identification 1' on 2020-01-13 15:01
Use as suspension	Yes	

## Formulation: Erythromycin\_Weibull\_enteric-coated-tablet

Type: Weibull

## Parameters

Name	Value	Value Origin
Dissolution time (50% dissolved)	79.6337524677 min	Parameter Identification-Parameter Identification-Value updated from 'Parameter Identification 1' on 2020-01-13 15:01
Lag time	78.7967495765 min	Parameter Identification-Parameter Identification-Value updated from 'Parameter Identification 1' on 2020-01-13 15:01
Dissolution shape	1.0838799888	Parameter Identification-Parameter Identification-Value updated from 'Parameter Identification 1' on 2020-01-13 15:01
Use as suspension	Yes	

## Formulation: Erythromycin\_Weibull\_filmtab

Type: Weibull

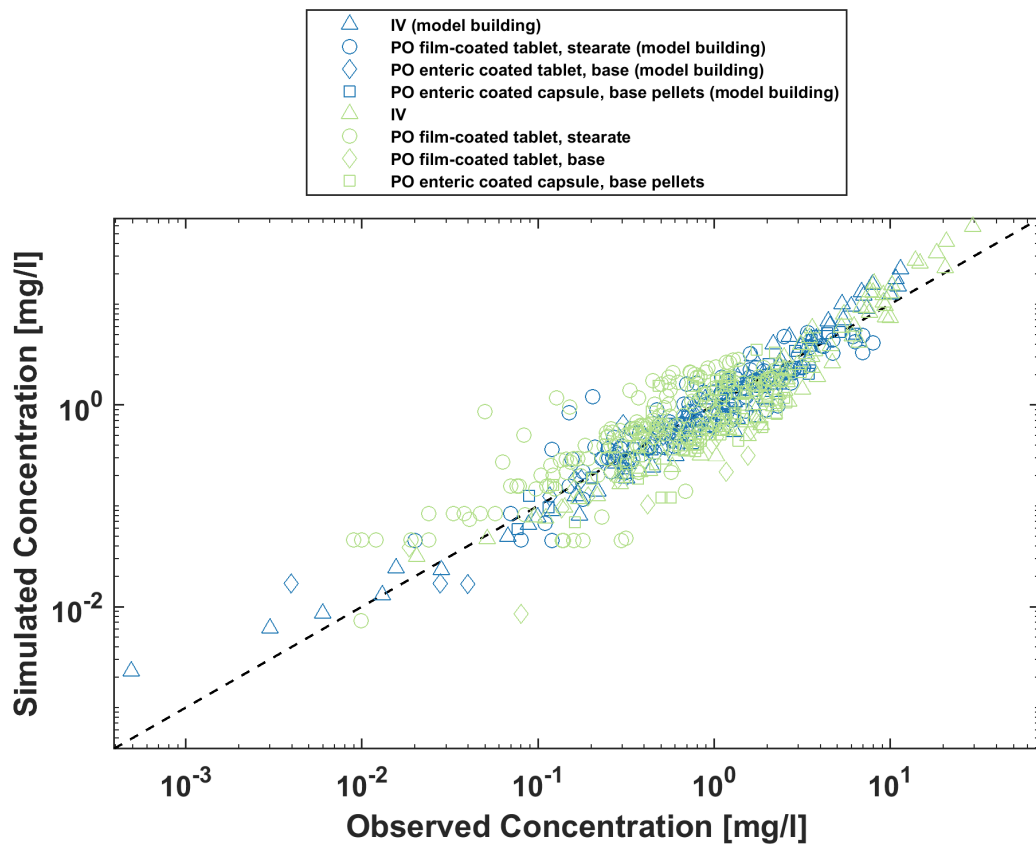
## Parameters

Name	Value	Value Origin
Dissolution time (50% dissolved)	1.7038947098 min	Parameter Identification-Parameter Identification-Value updated from 'Parameter Identification 1' on 2020-01-13 15:01
Lag time	0 min	Other-Assumption
Dissolution shape	1.0960212213	Parameter Identification-Parameter Identification-Value updated from 'Parameter Identification 1' on 2020-01-13 15:01
Use as suspension	Yes	

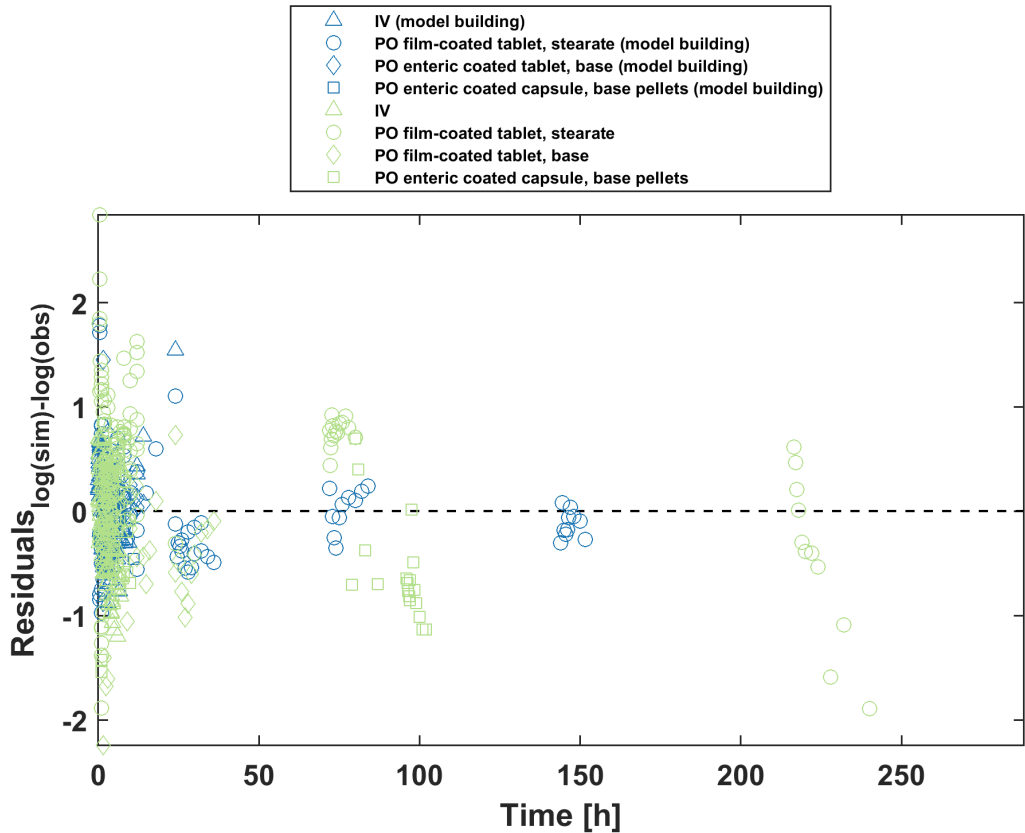
## 3.2 Diagnostics Plots

Below you find the goodness-of-fit visual diagnostic plots for the PBPK model performance of all data used presented in [Section 2.2.3](#).

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



Goodness of fit plot for concentration in plasma



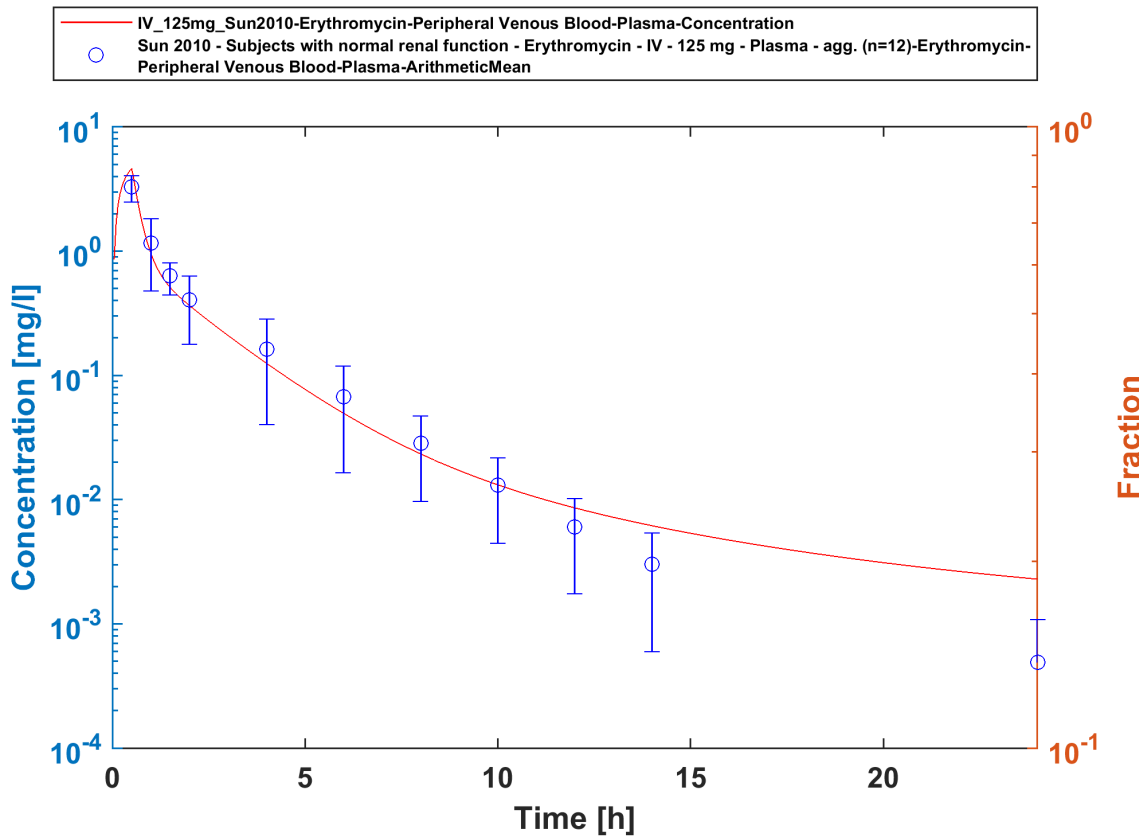
Goodness of fit plot for concentration in plasma

GMFE = 1.575079

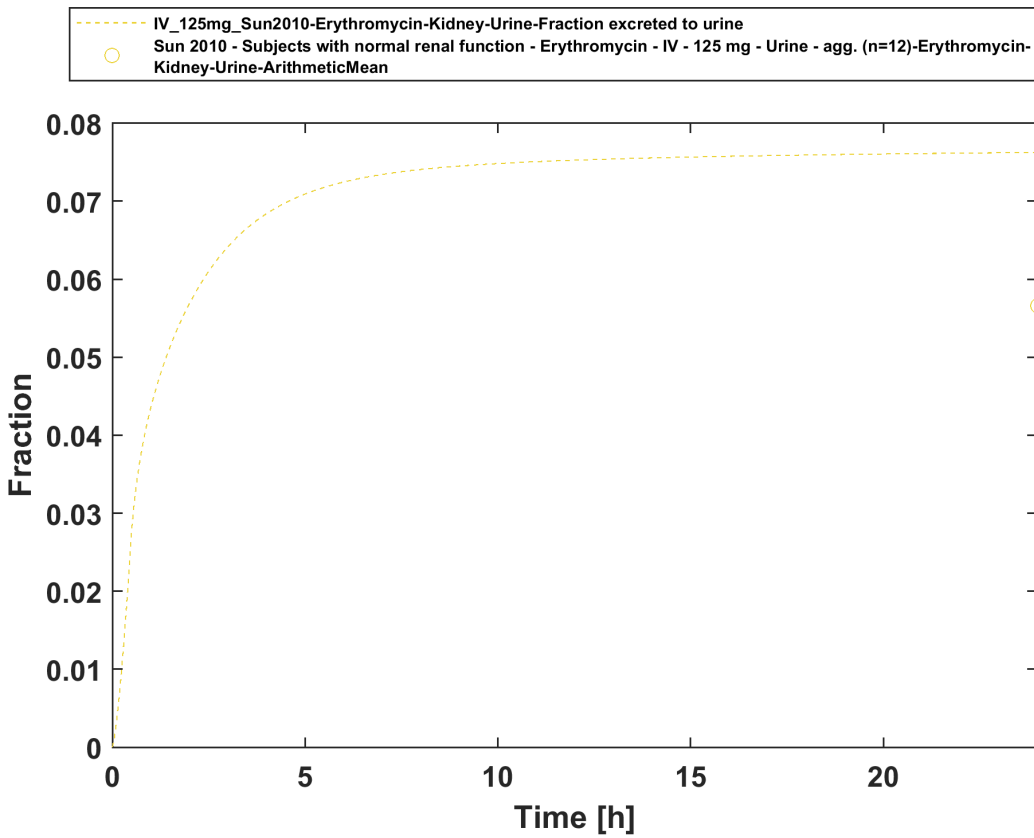
### 3.3 Concentration-Time Profiles

Simulated versus observed concentration-time profiles of all data listed in [Section 2.2.3](#) are presented below.

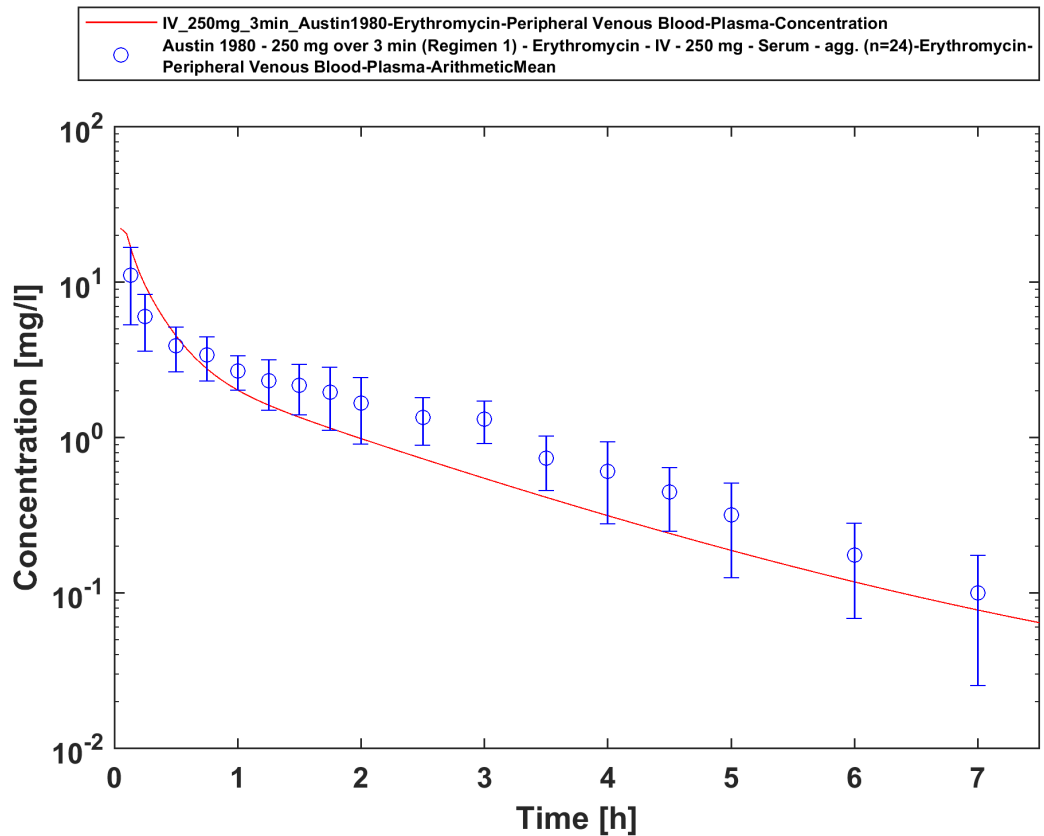
#### 3.3.1 Model Building



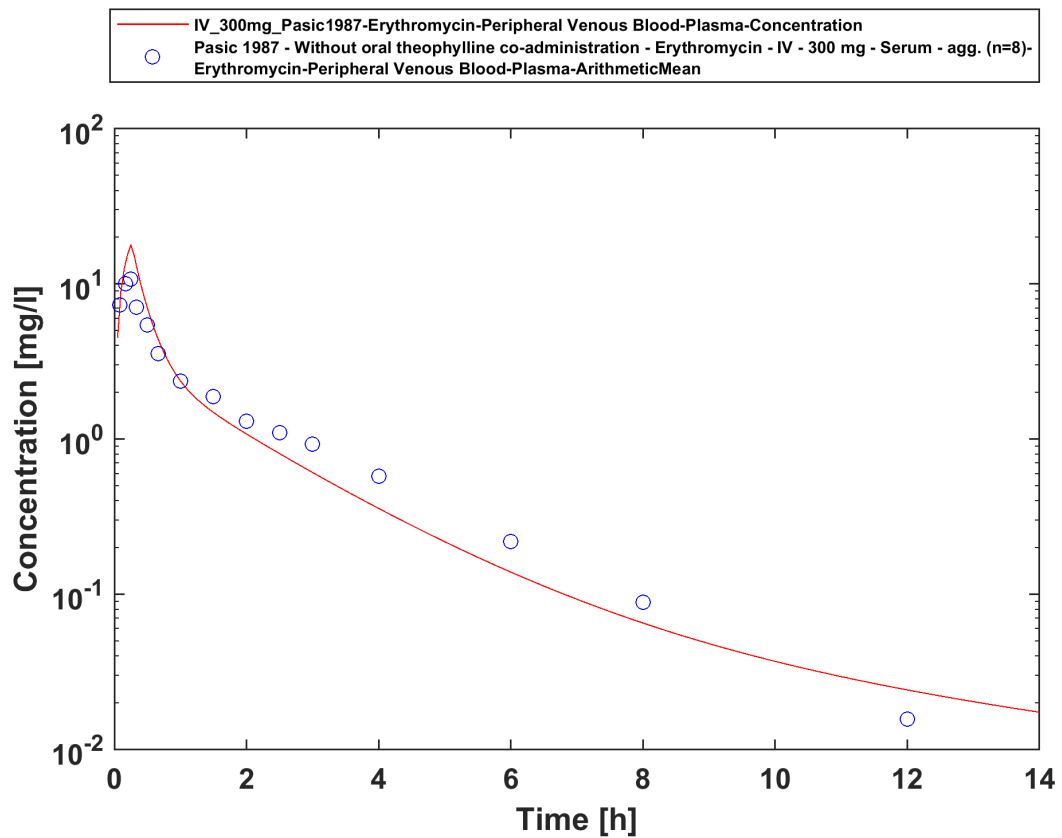
Time Profile Analysis



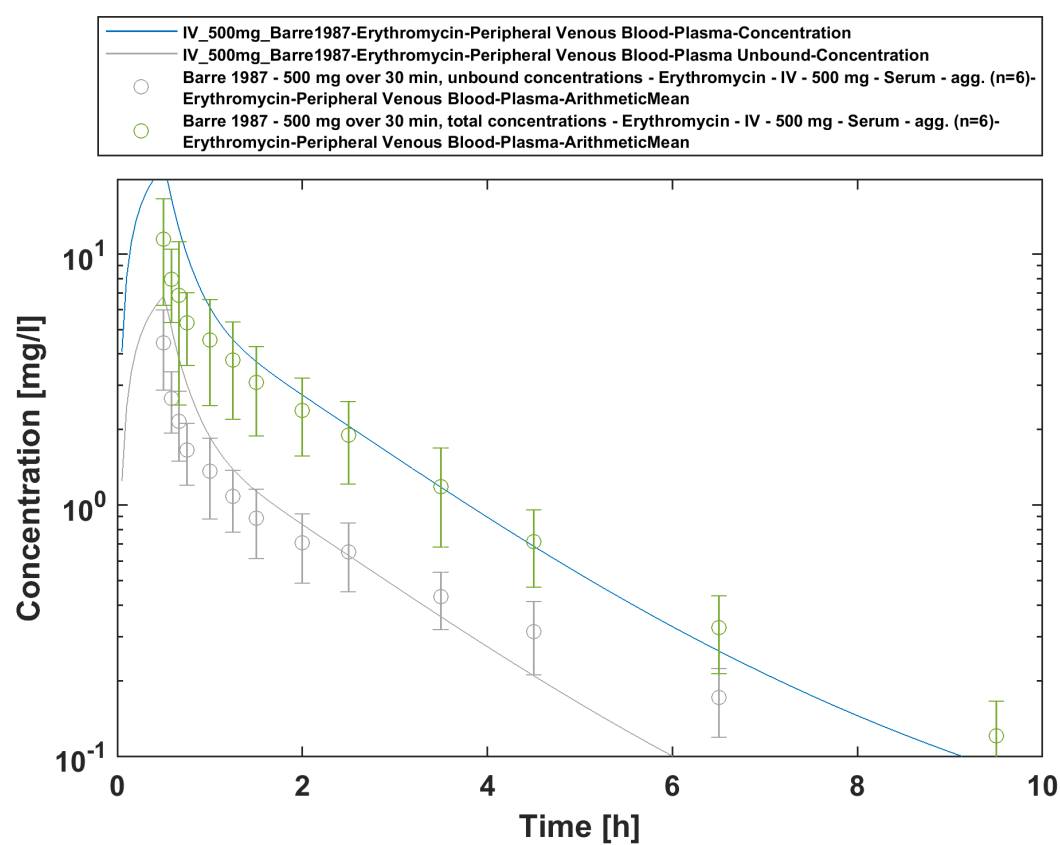
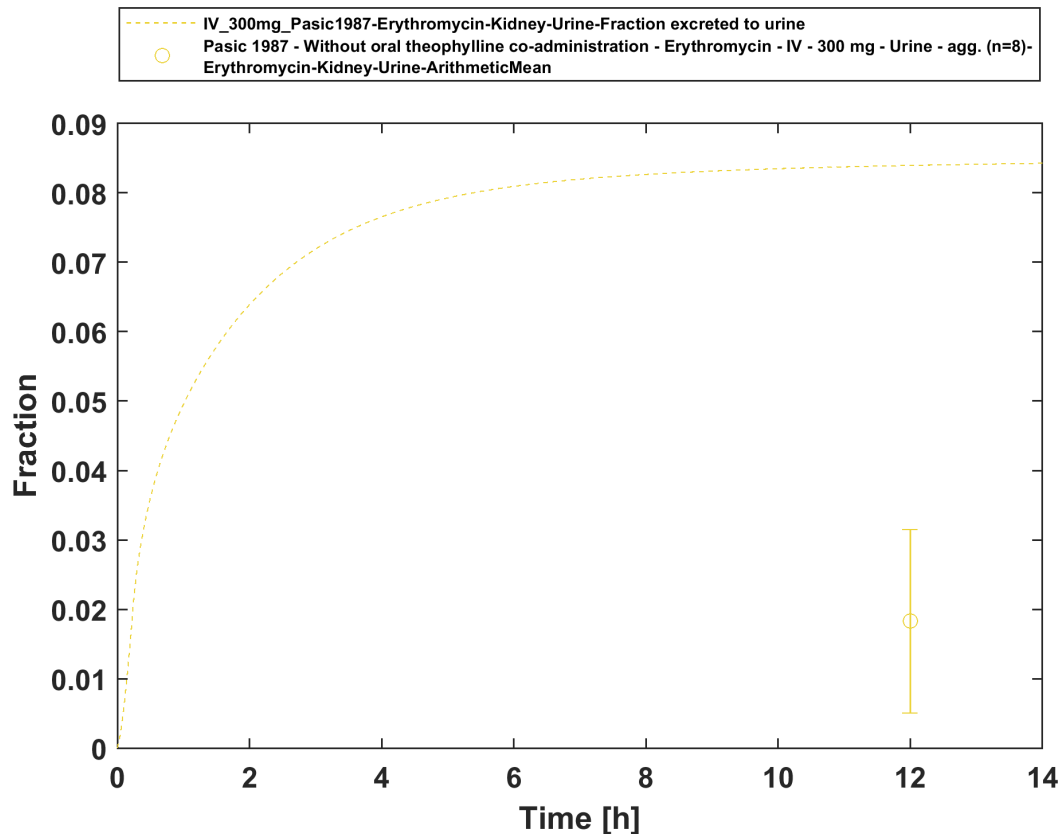
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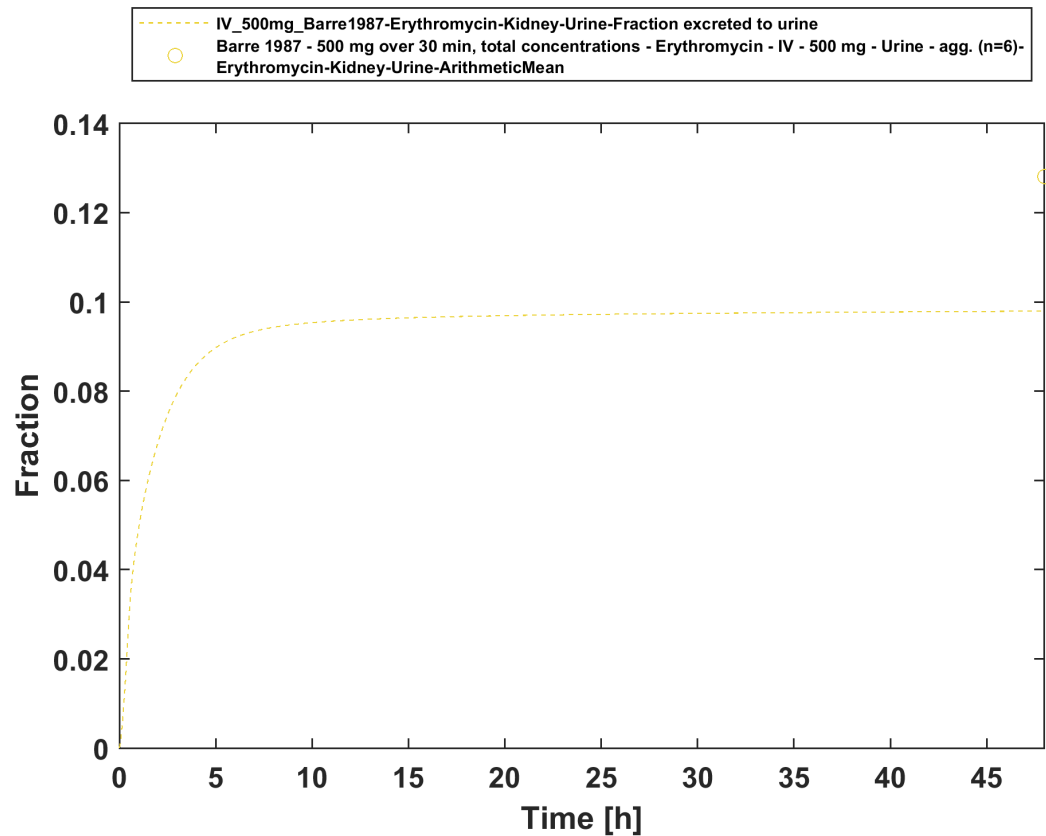


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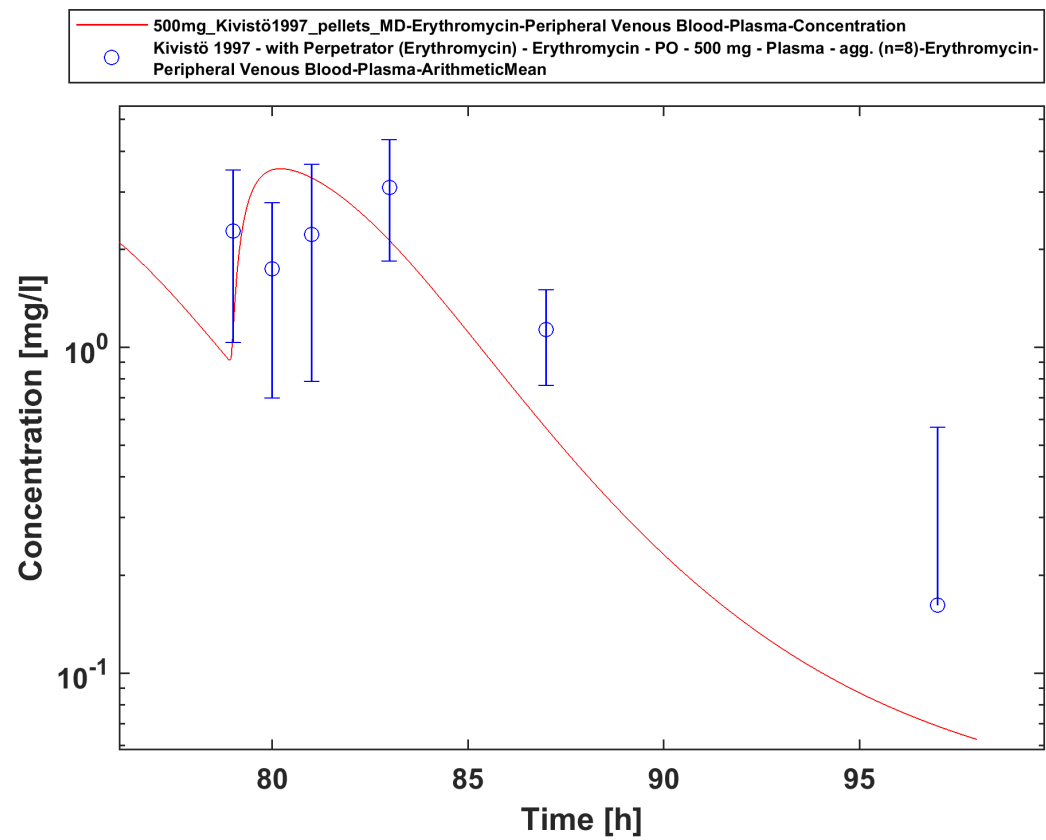


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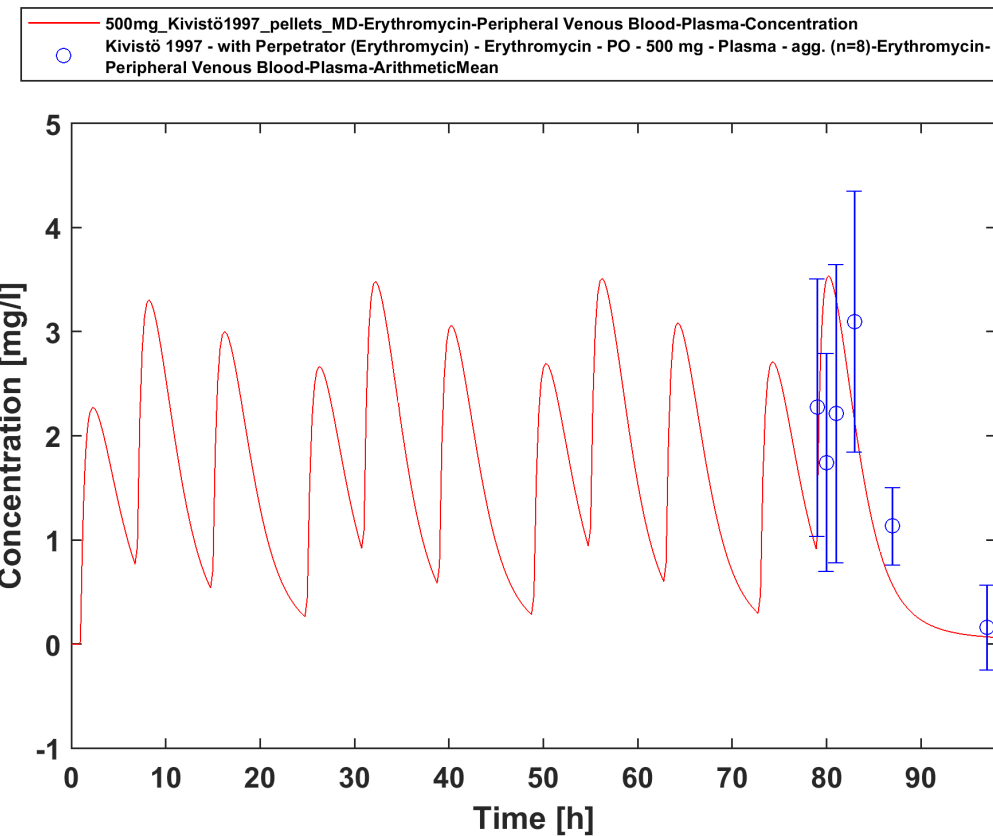




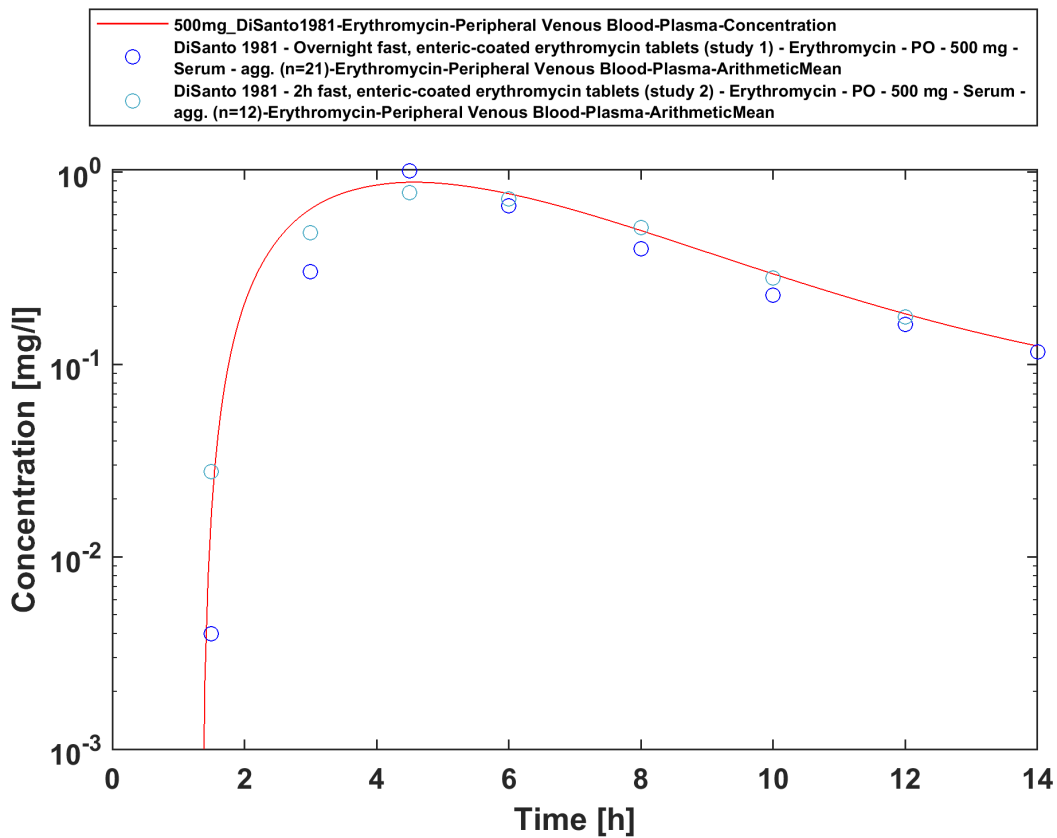
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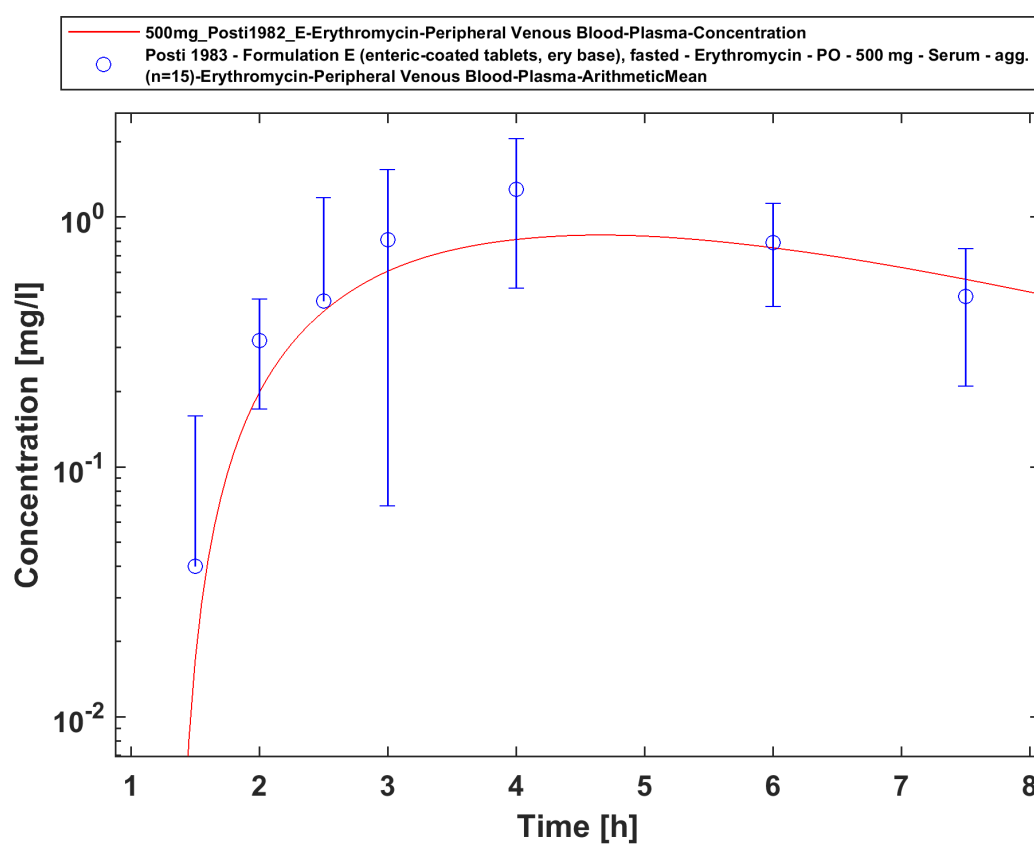
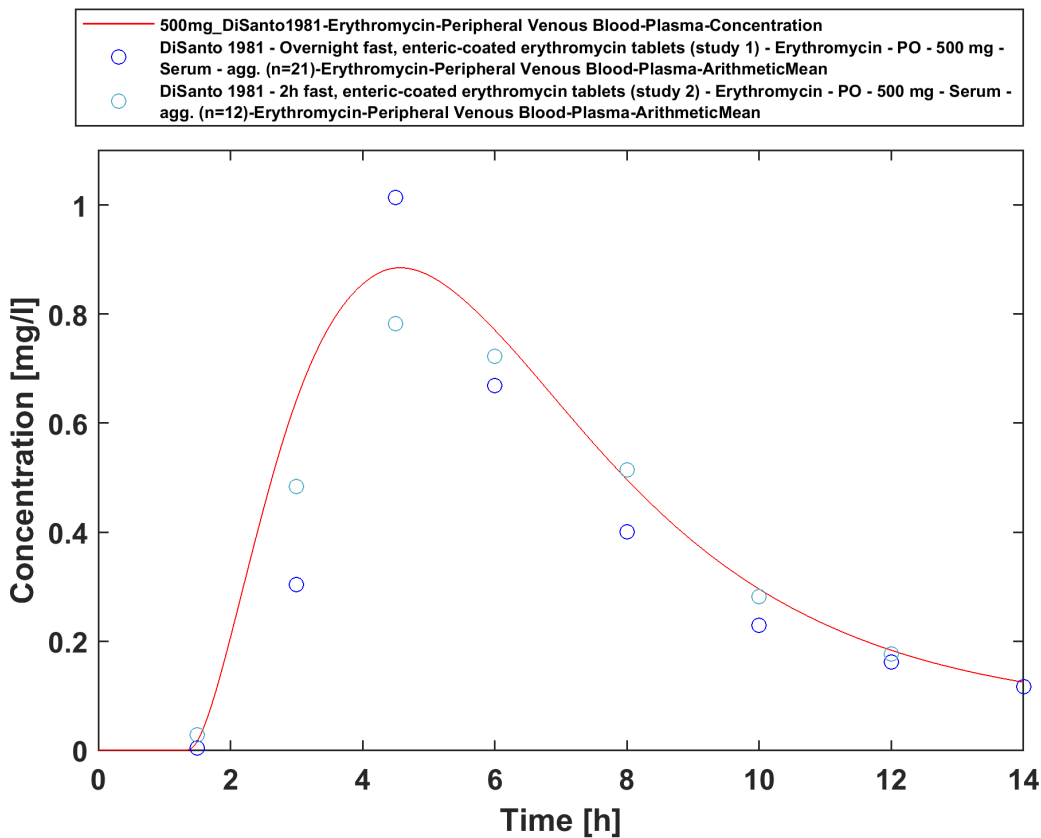
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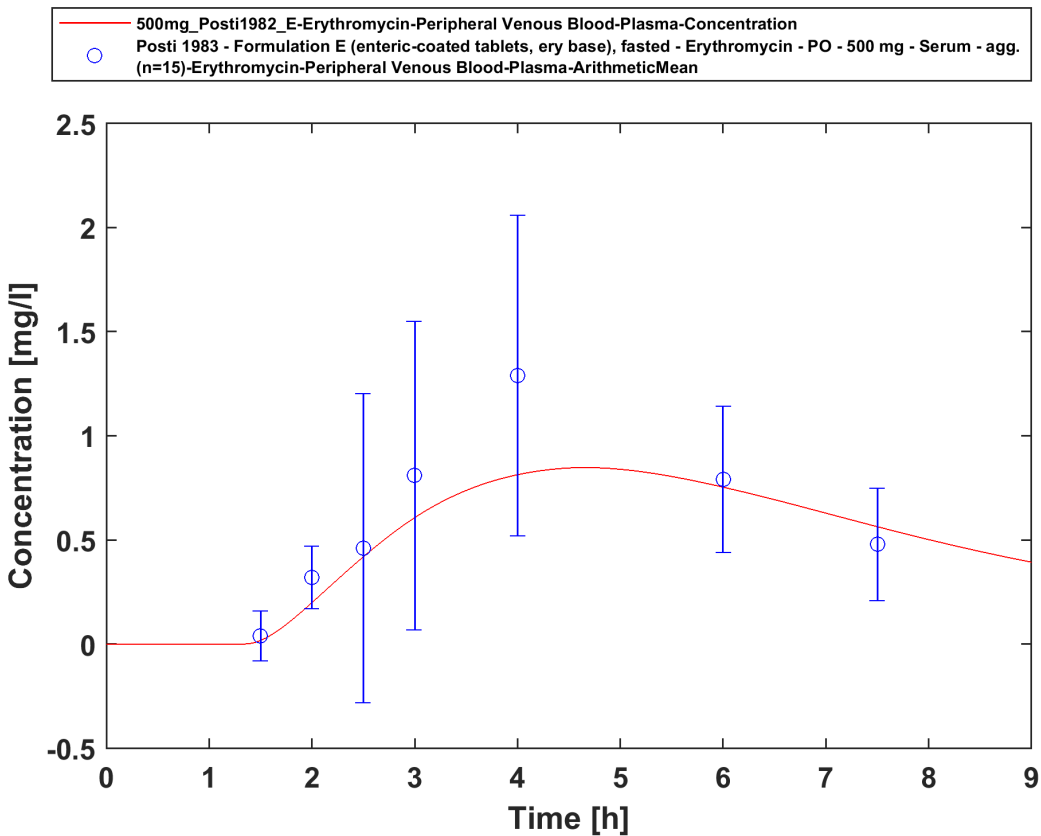


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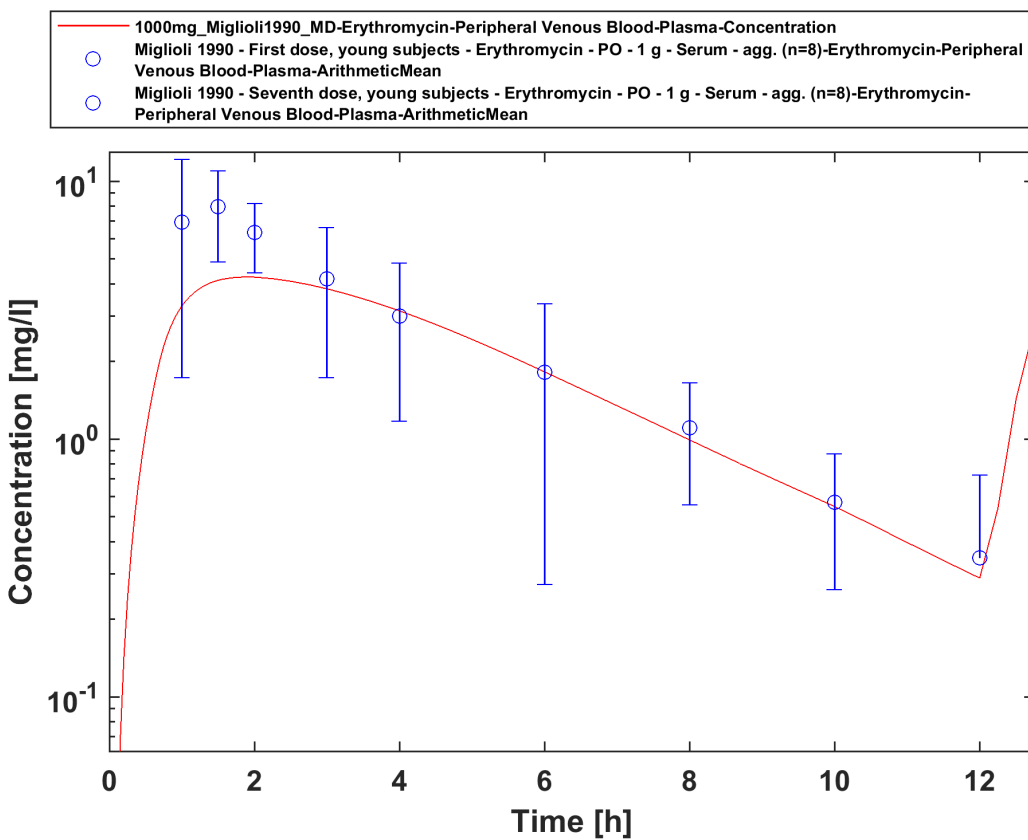


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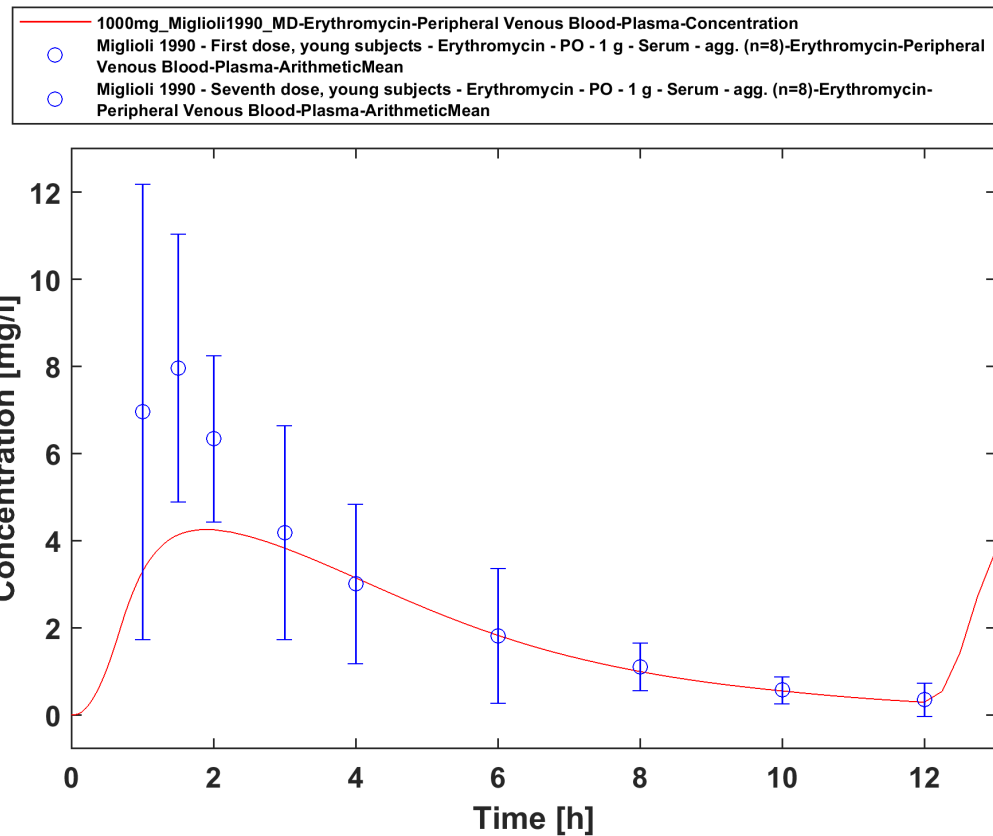




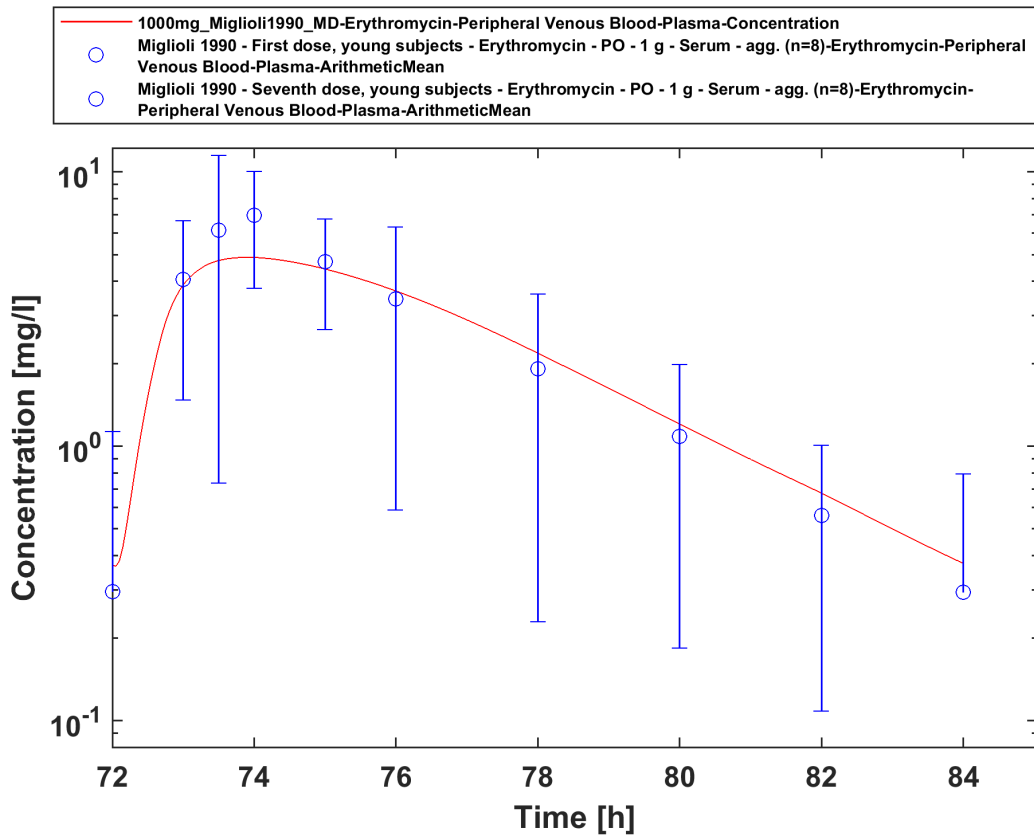
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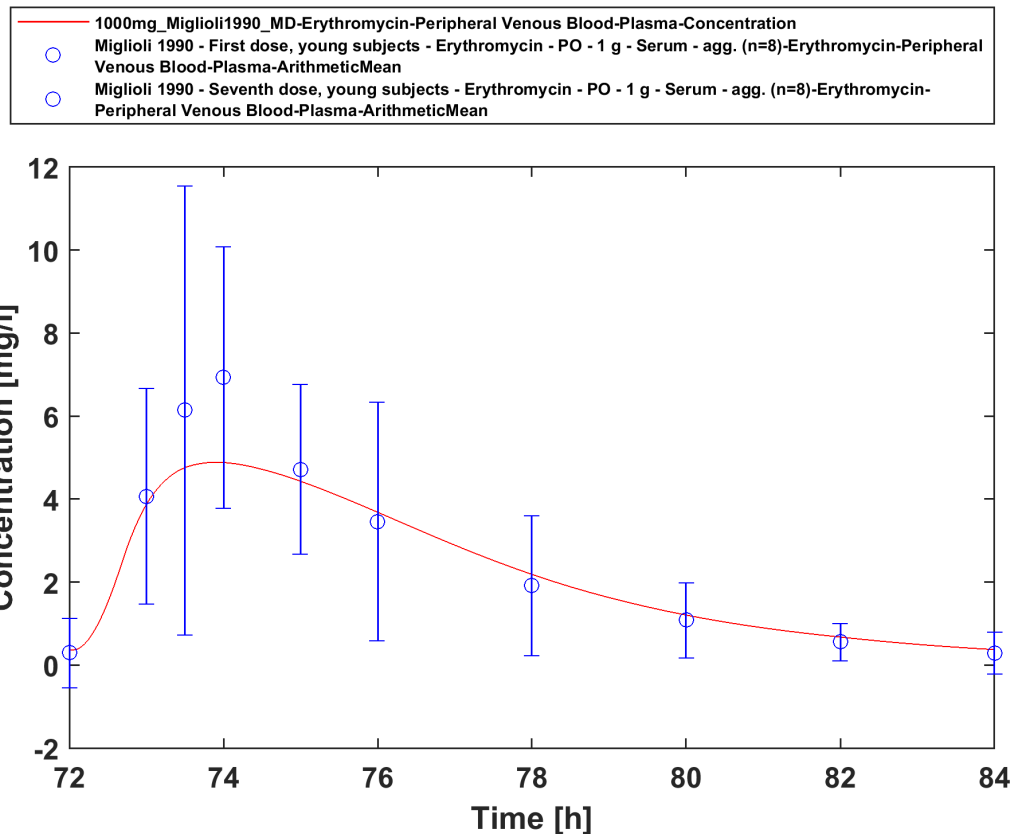
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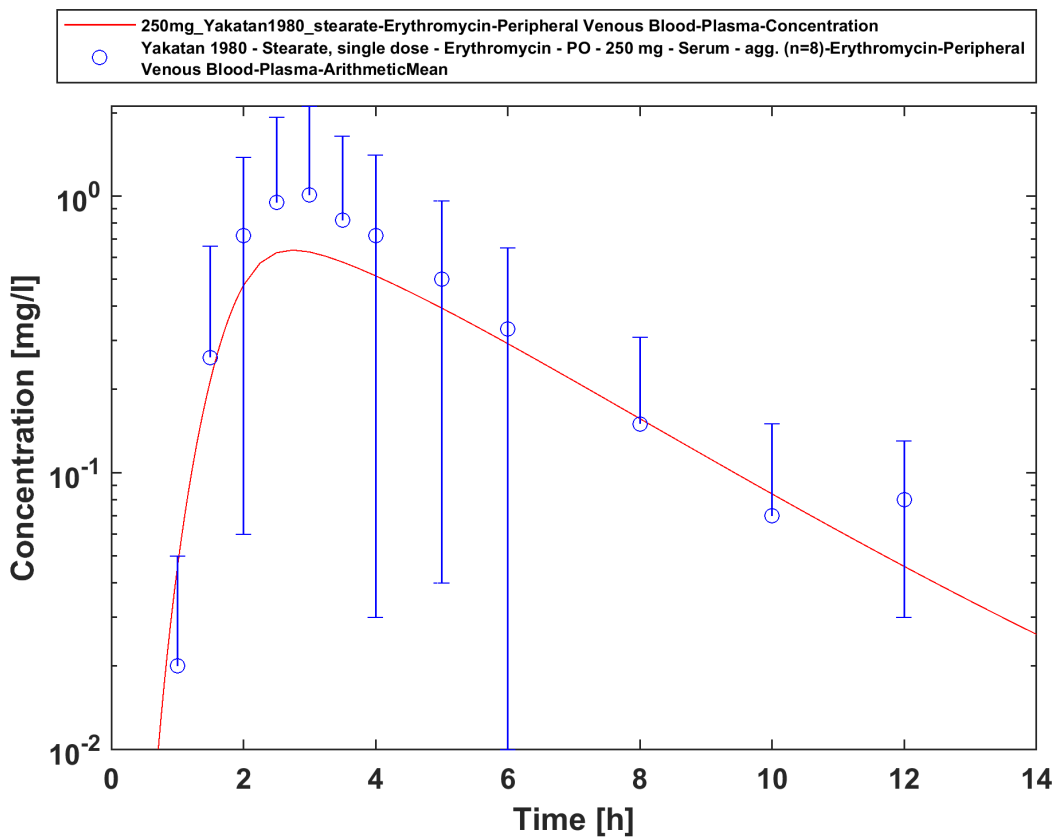
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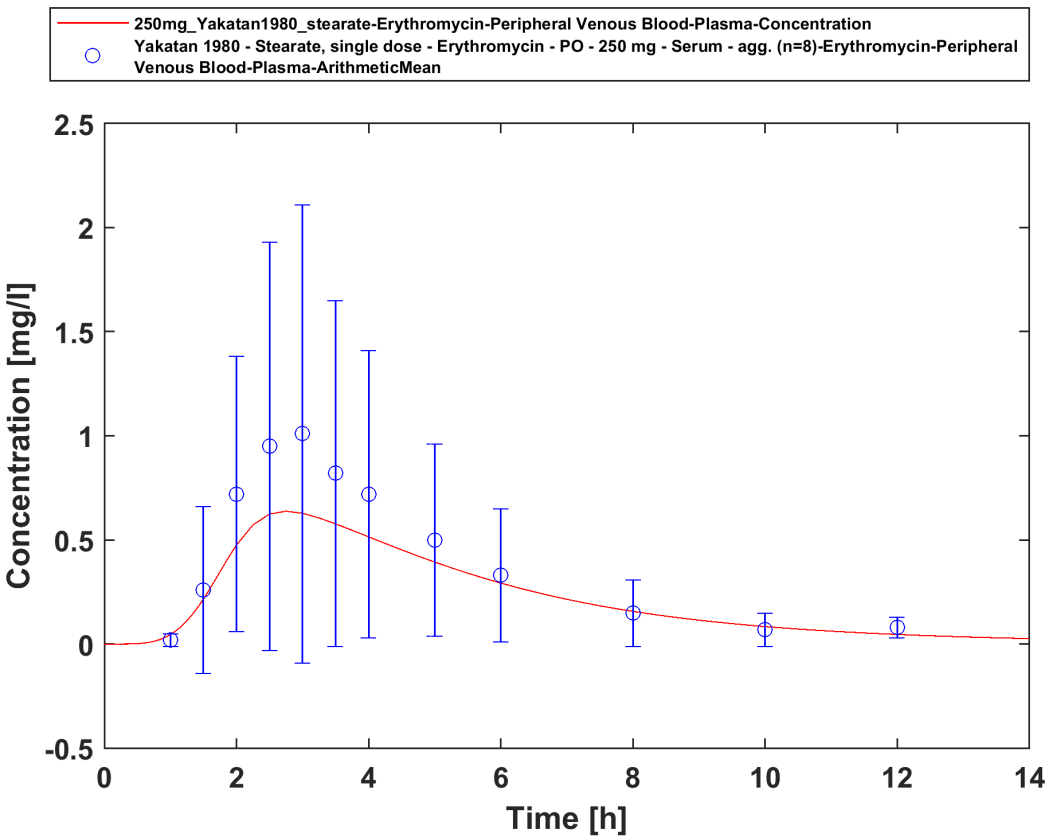
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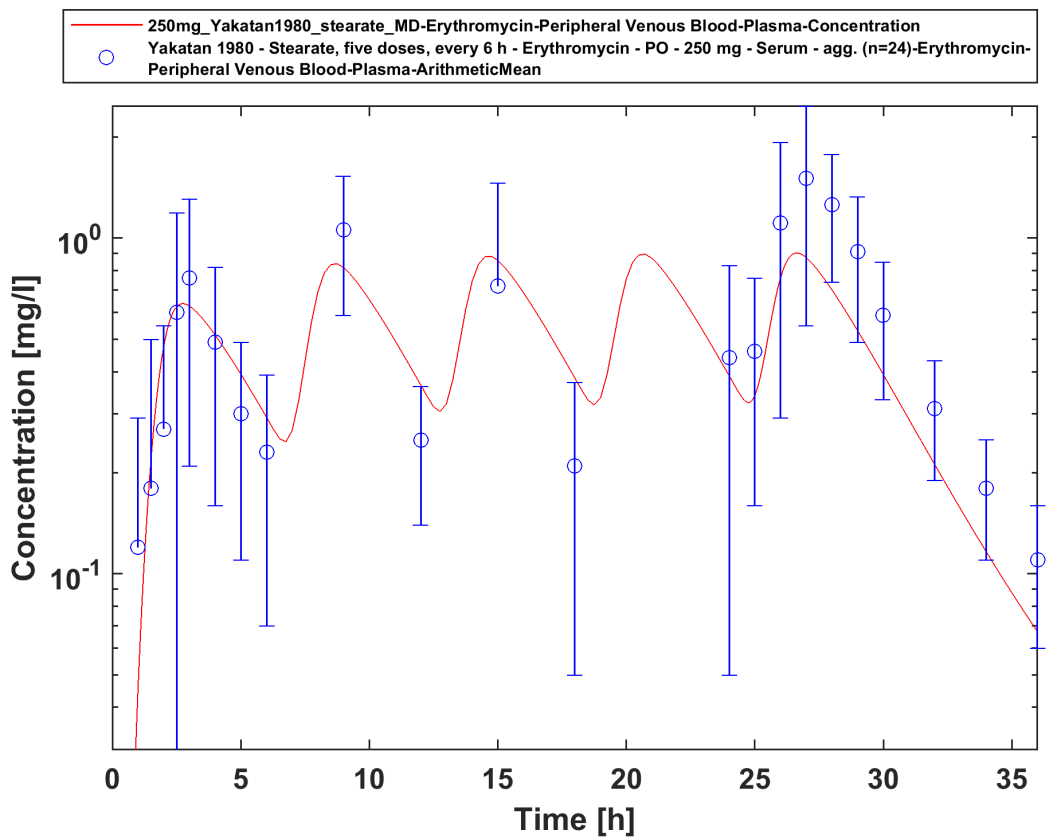
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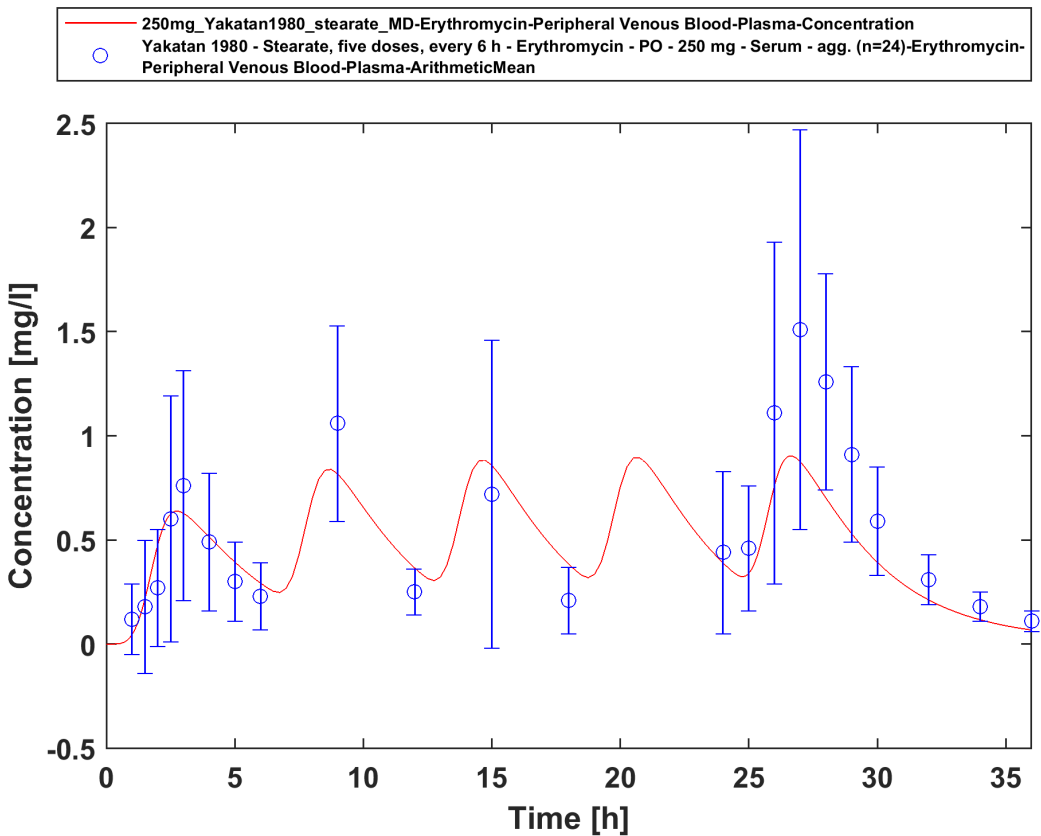
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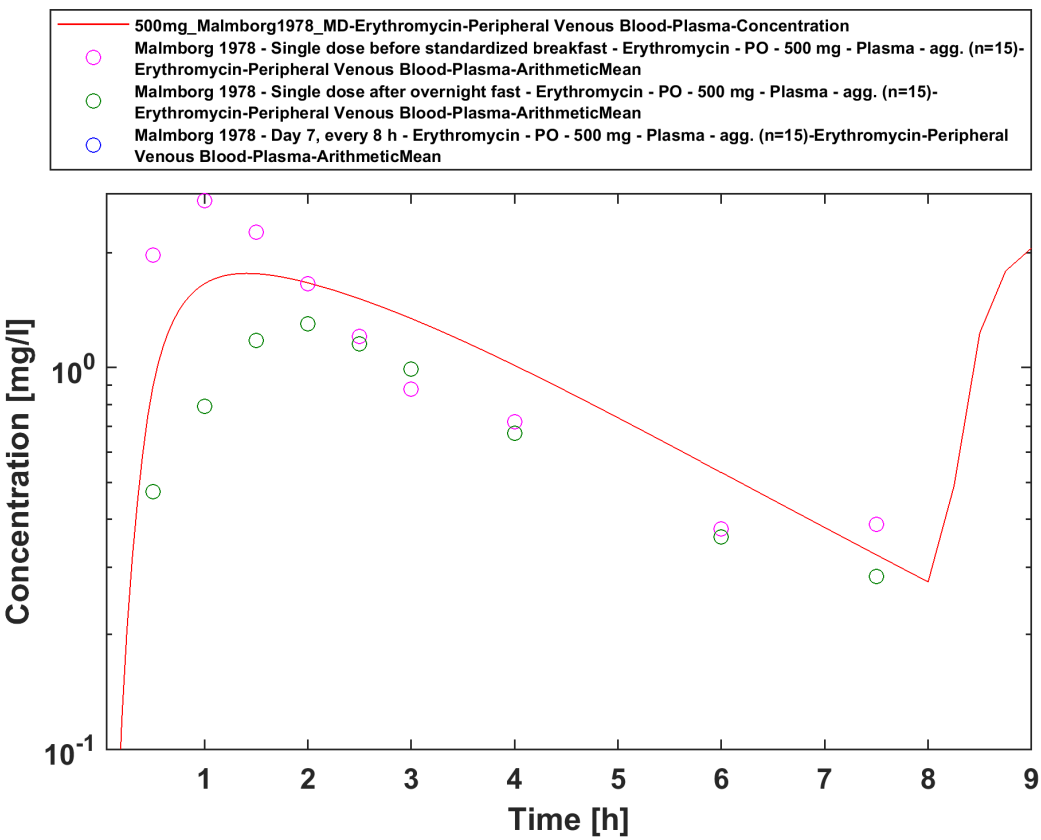
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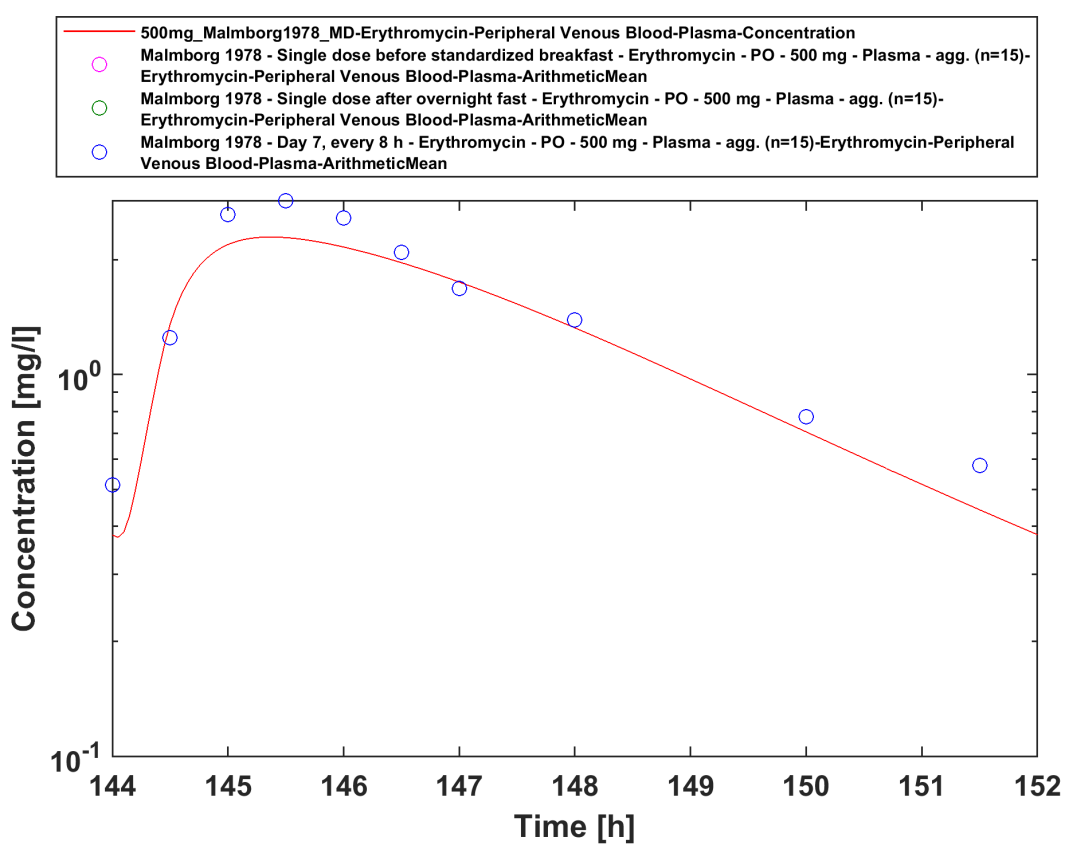
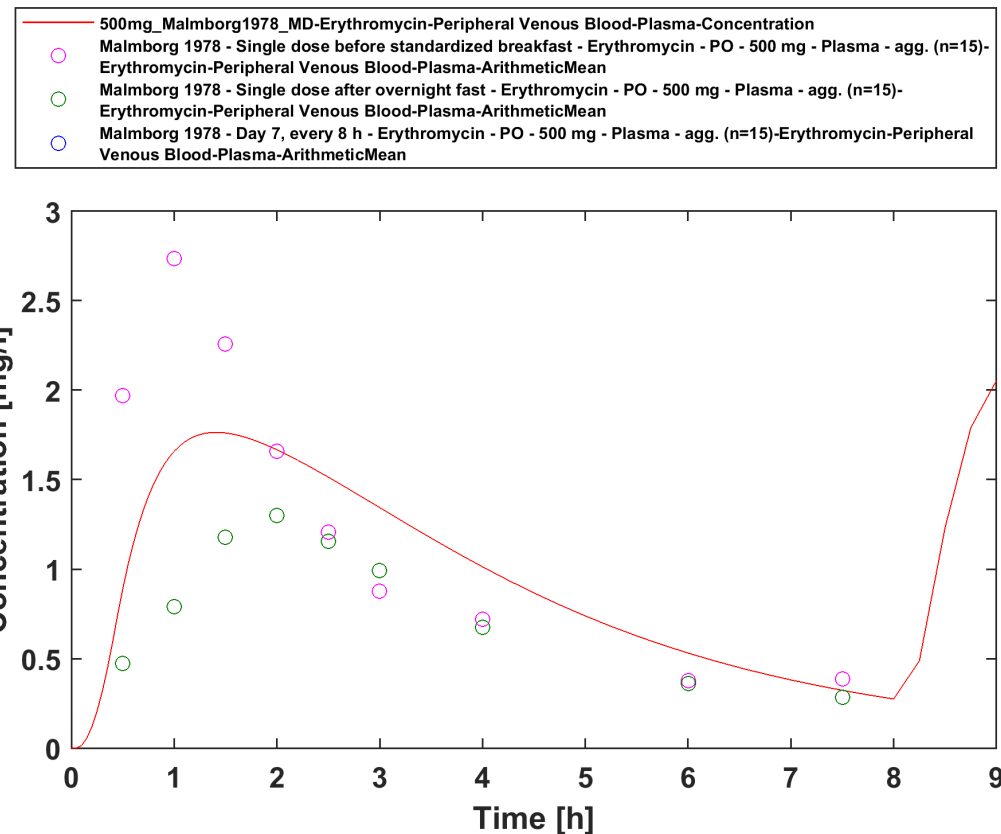
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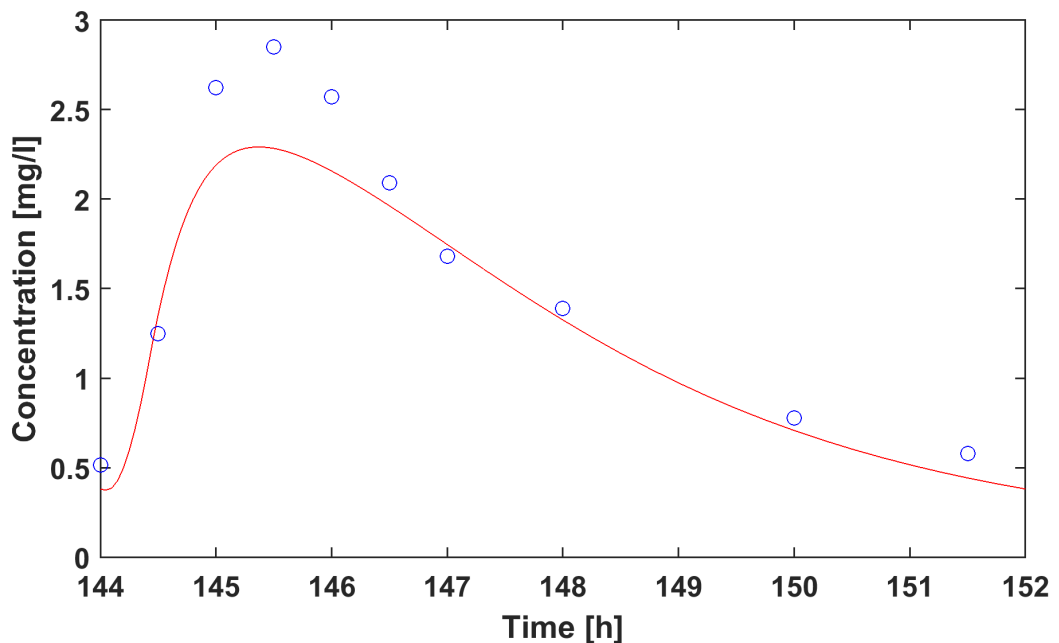
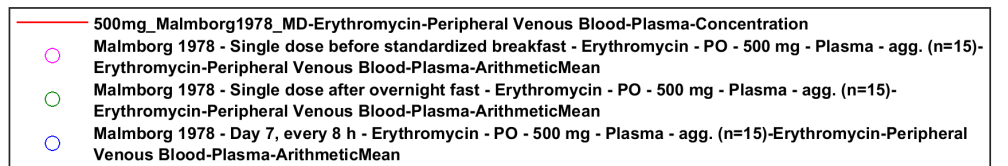


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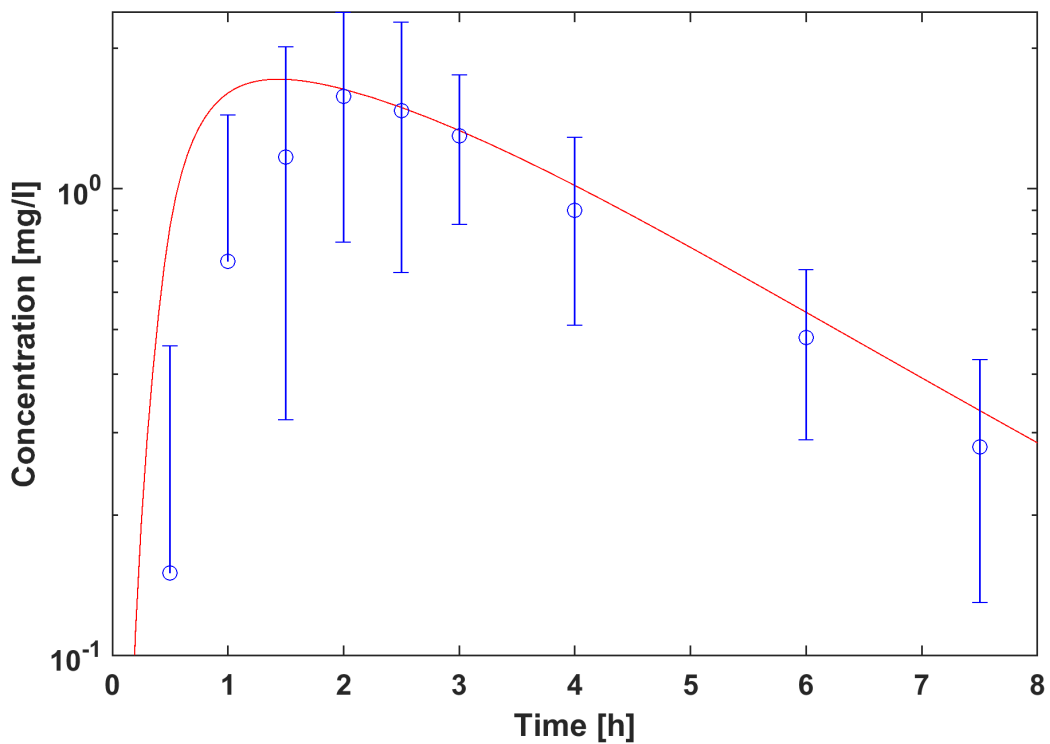
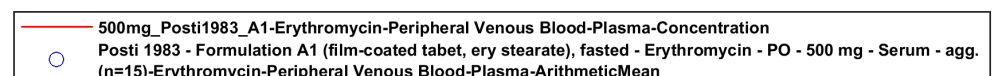


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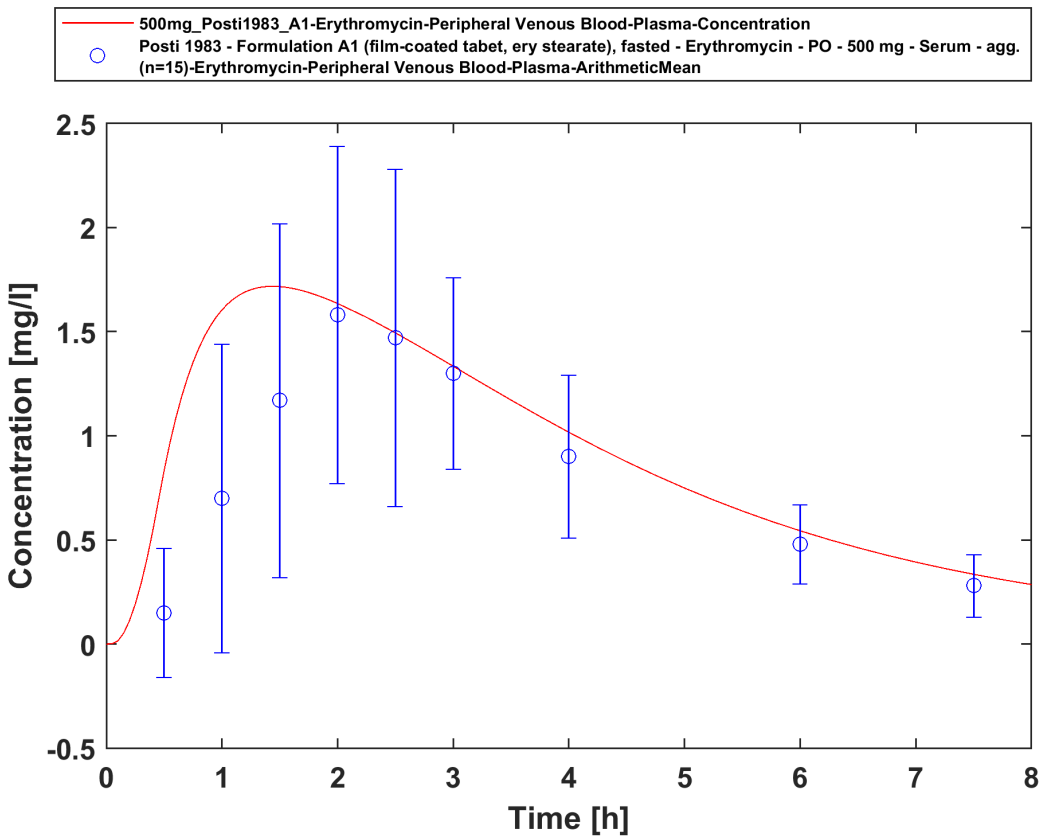




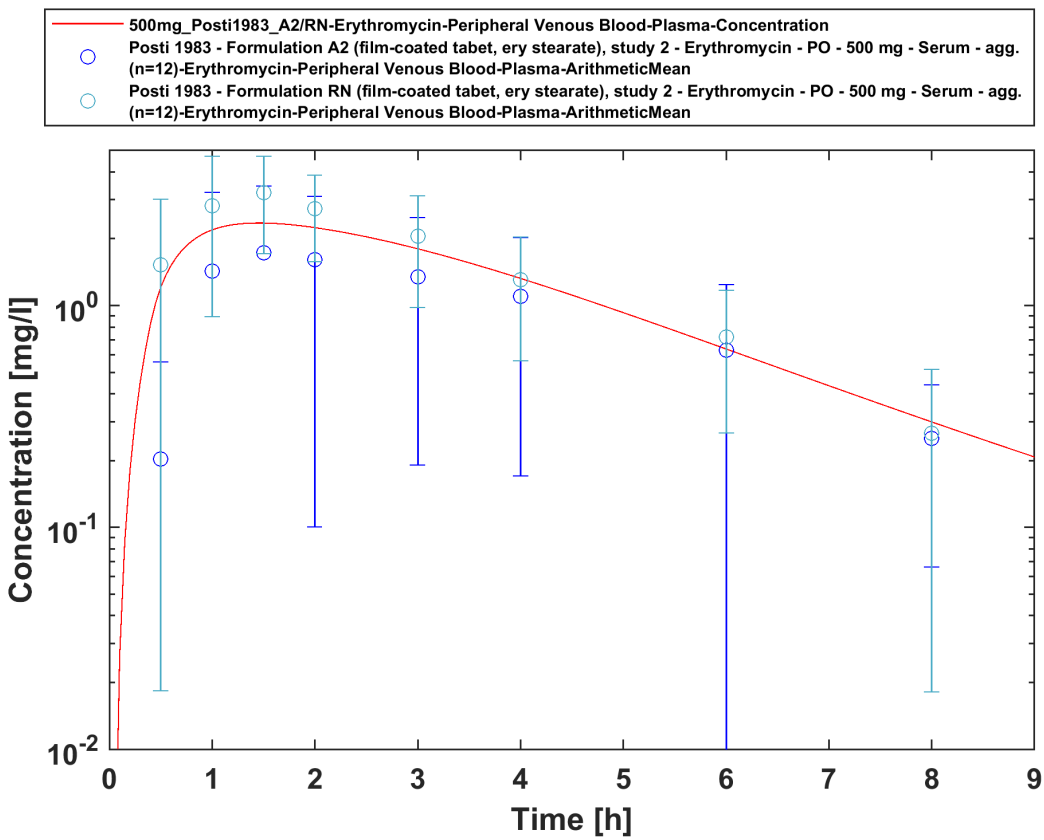
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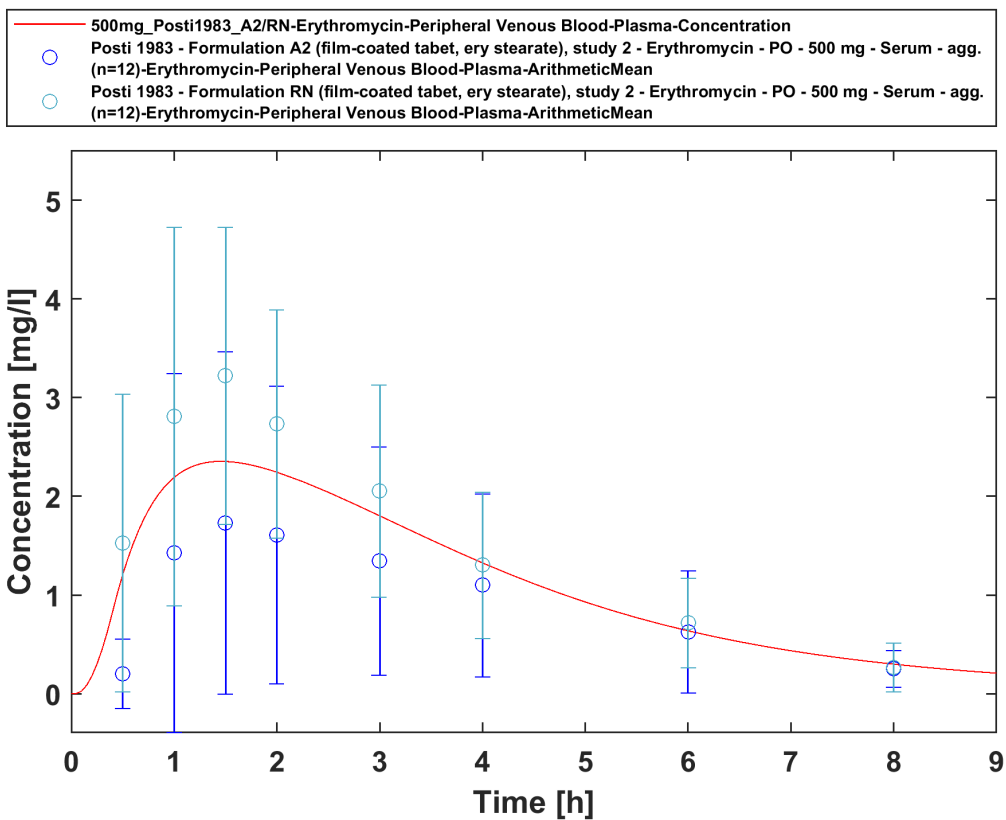
Time Profile Analysis



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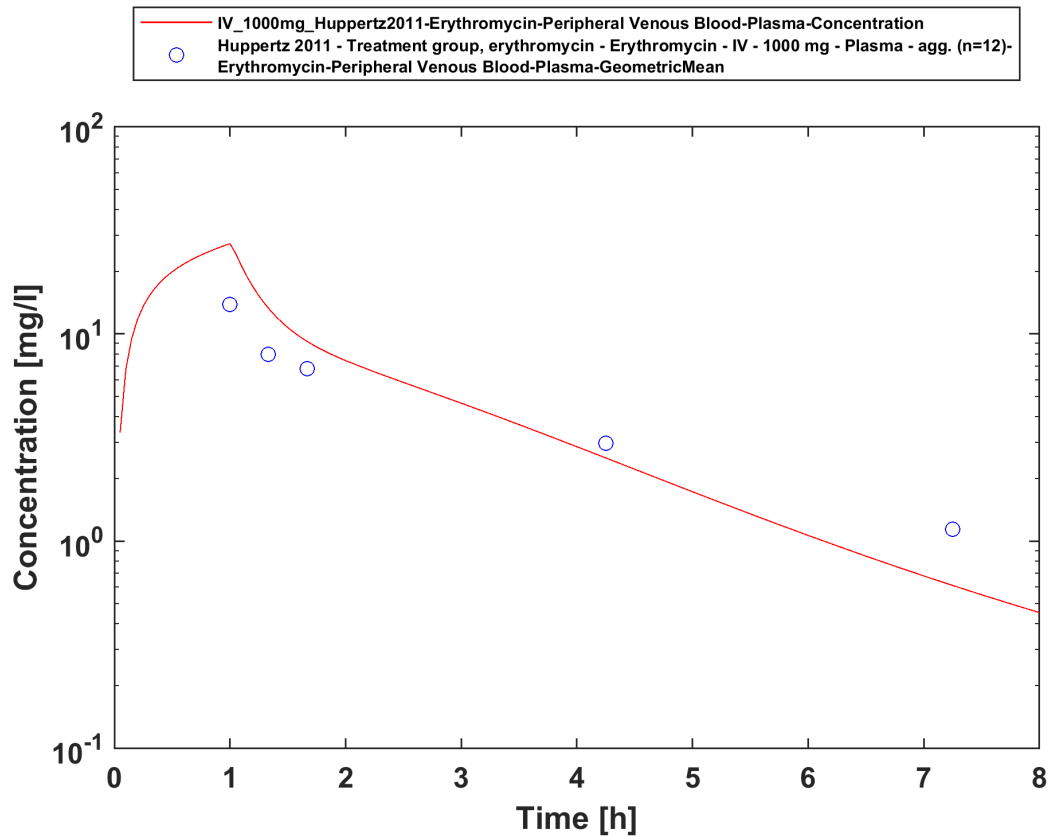


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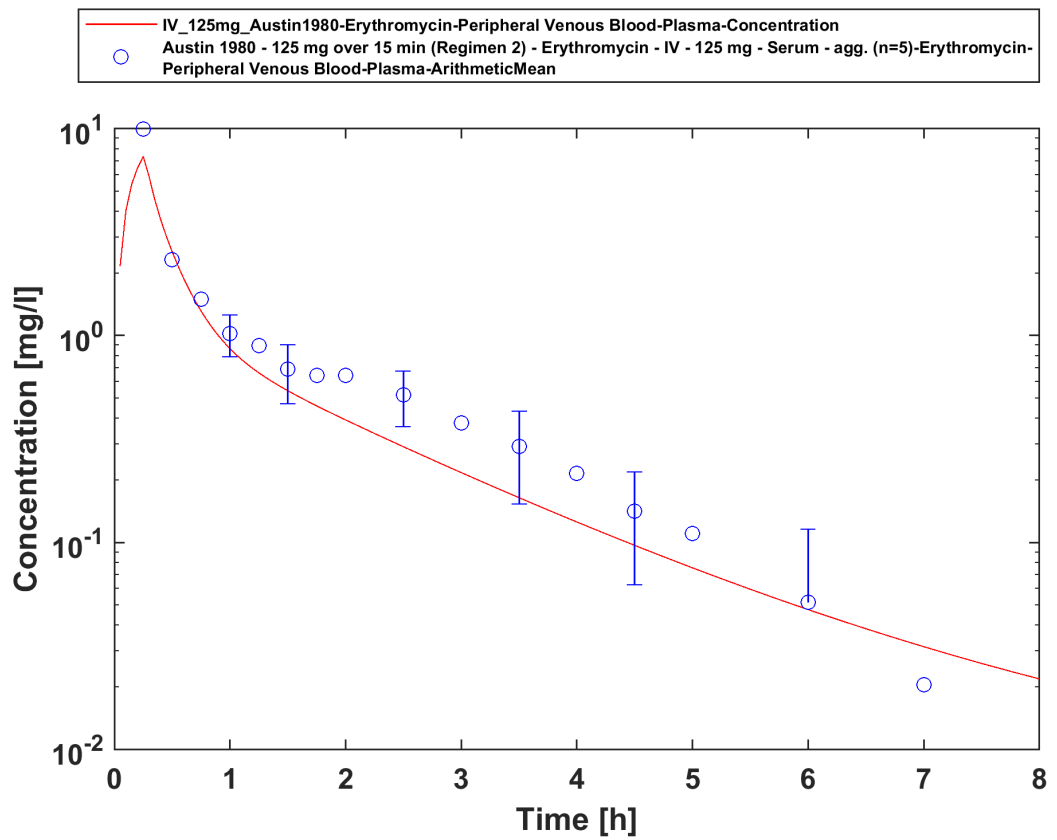


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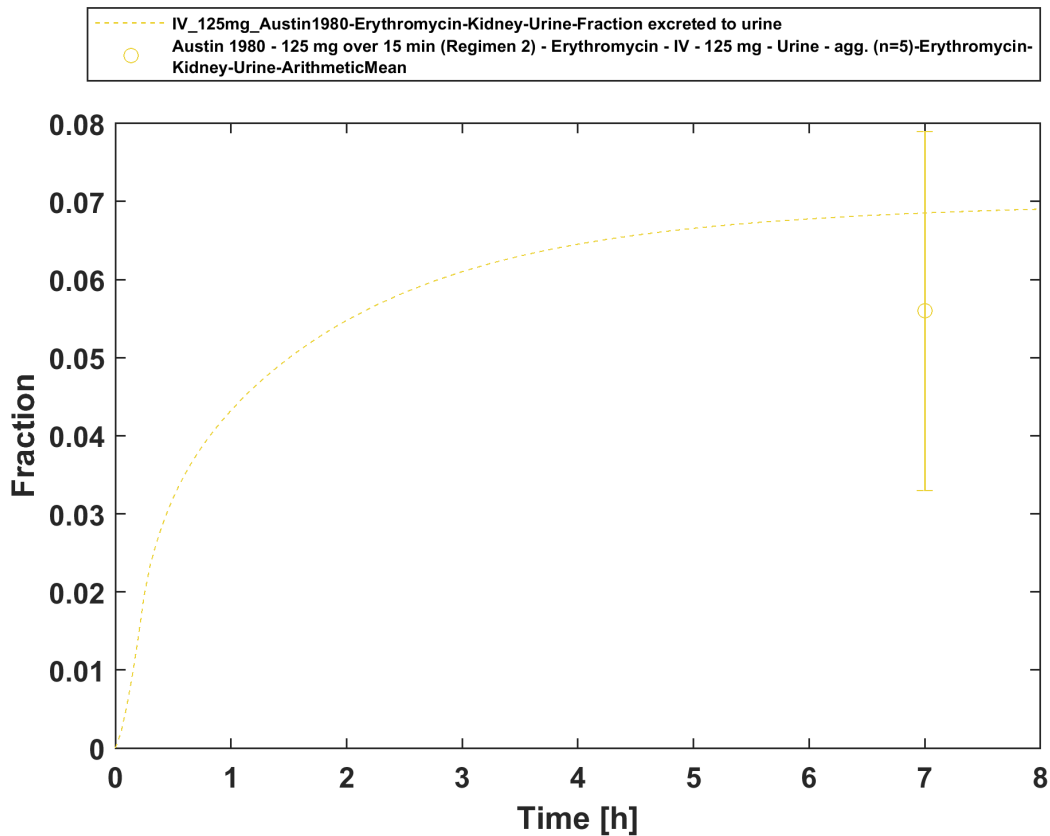
### 3.3.2 Model Verification



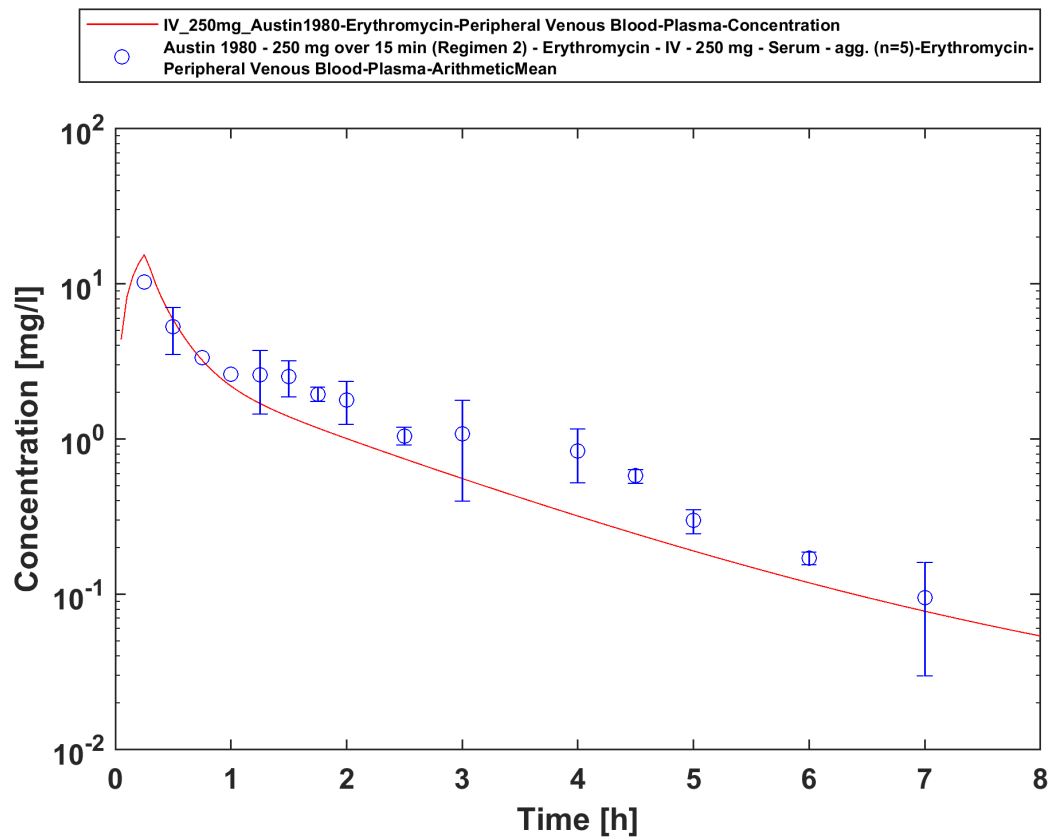
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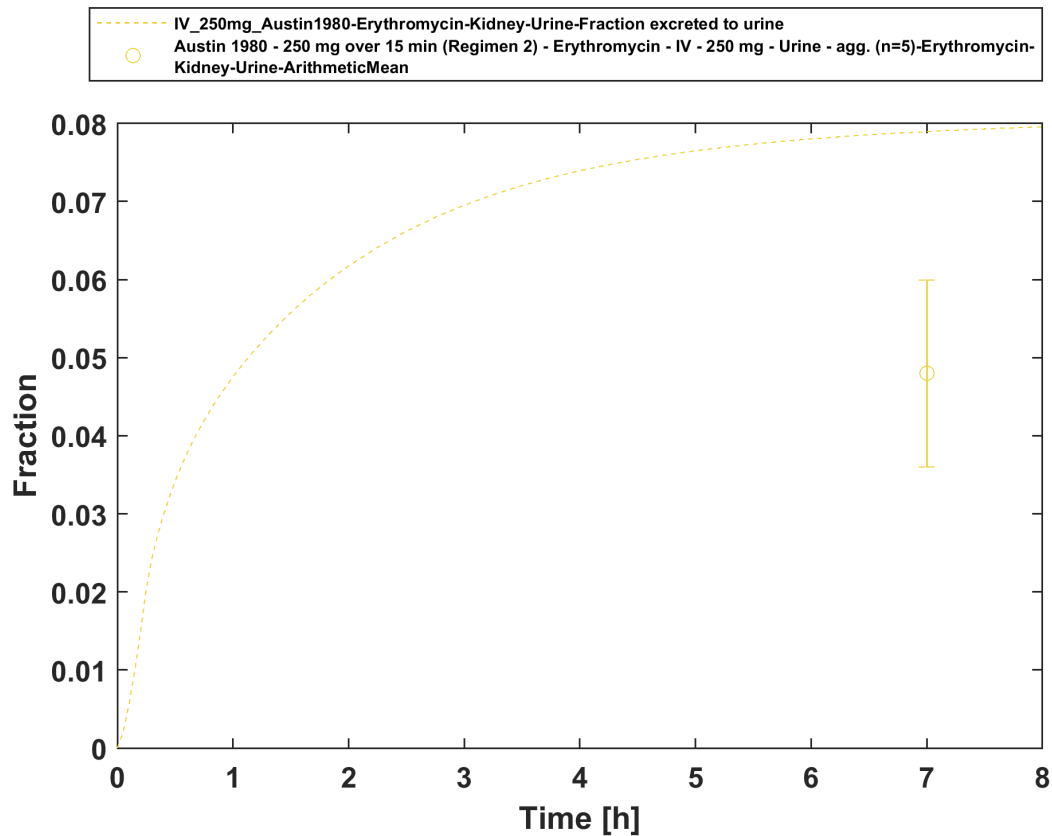
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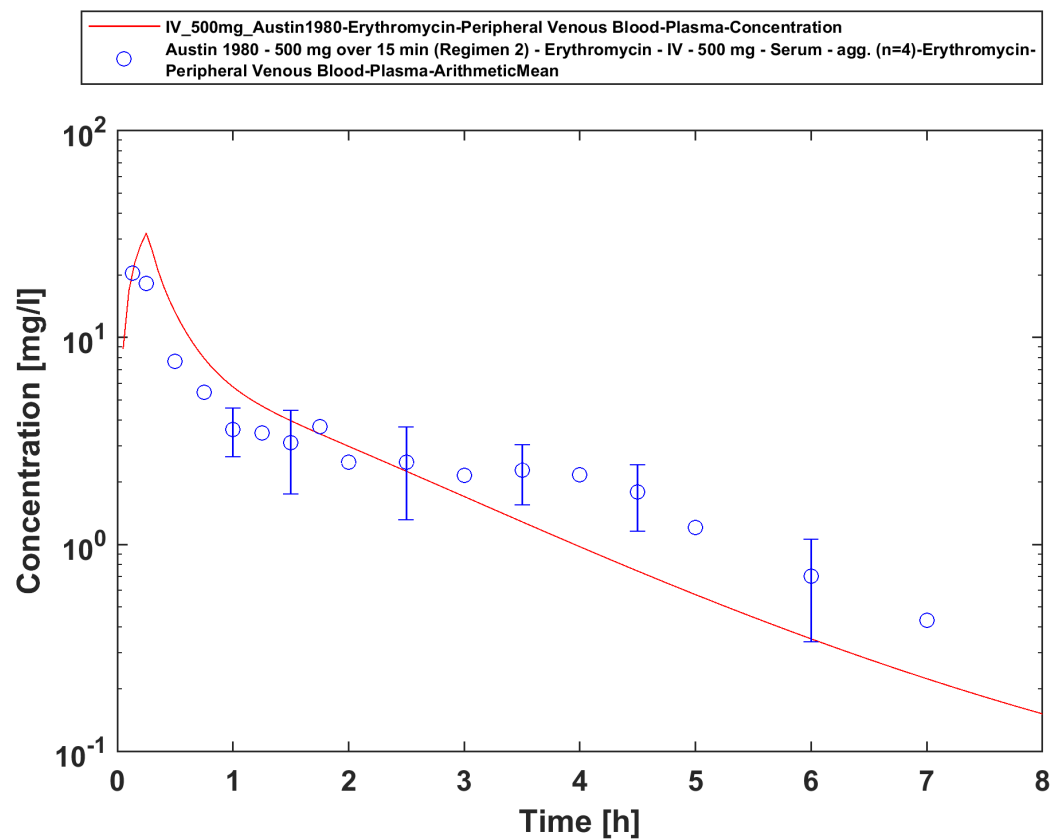
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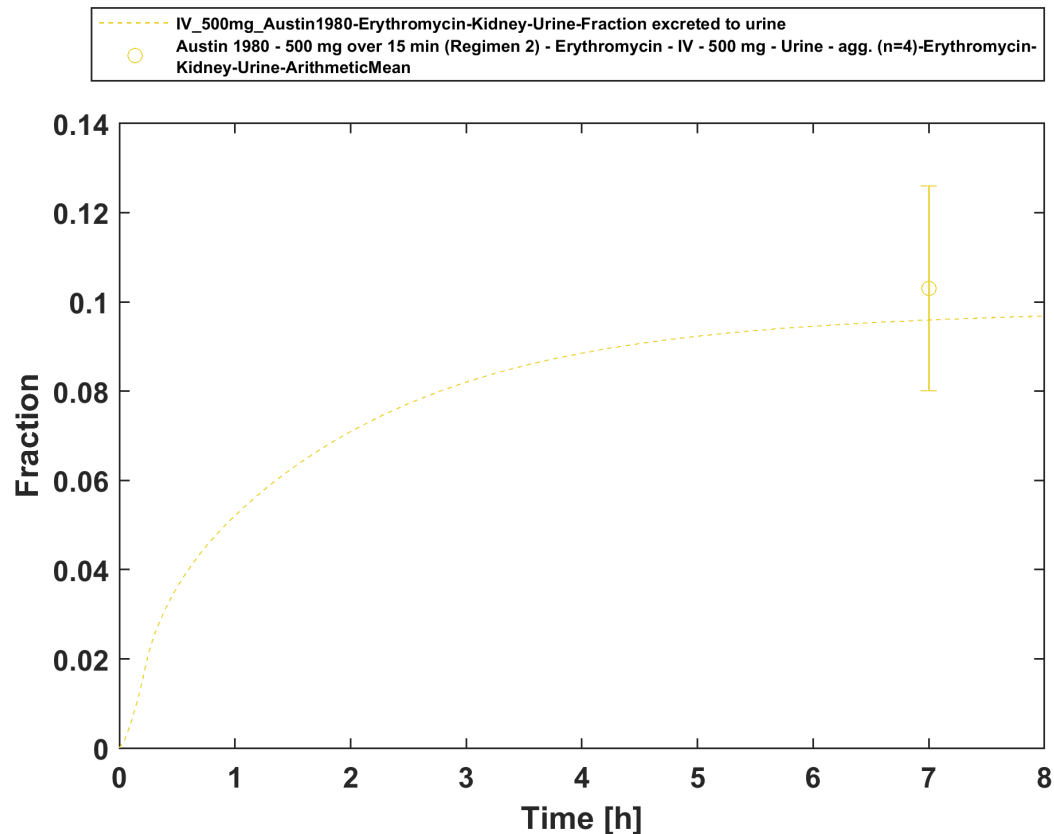
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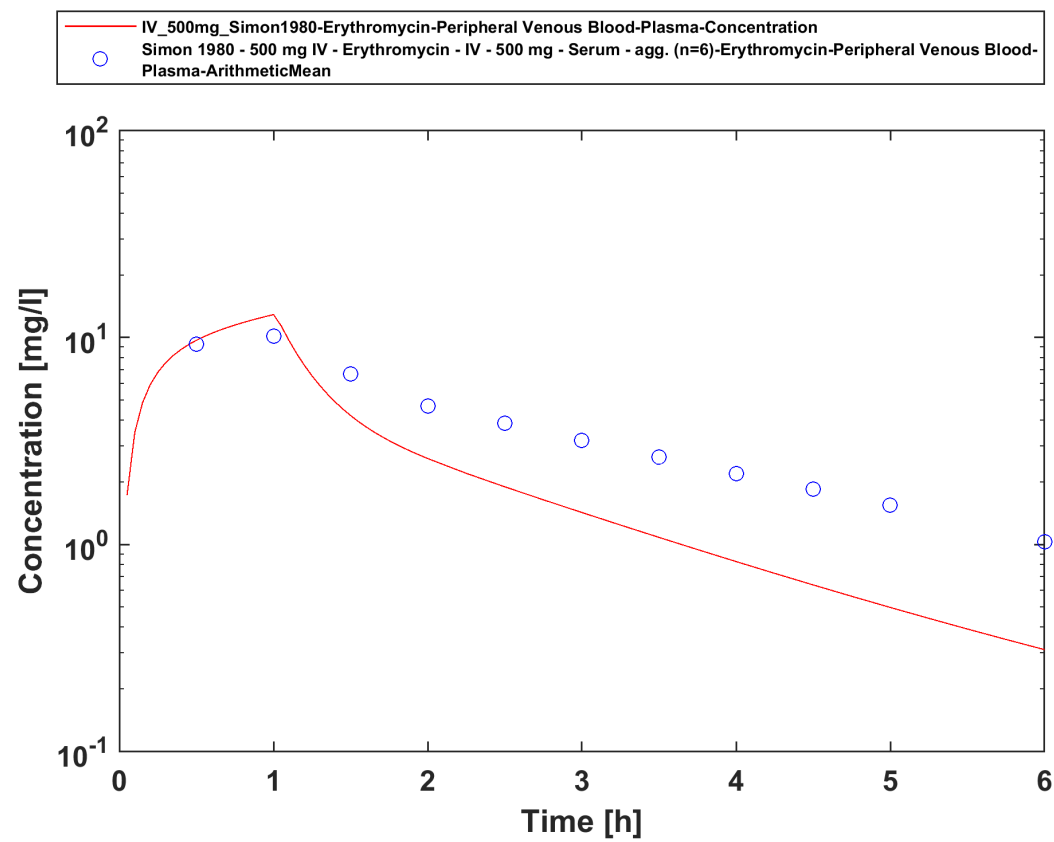
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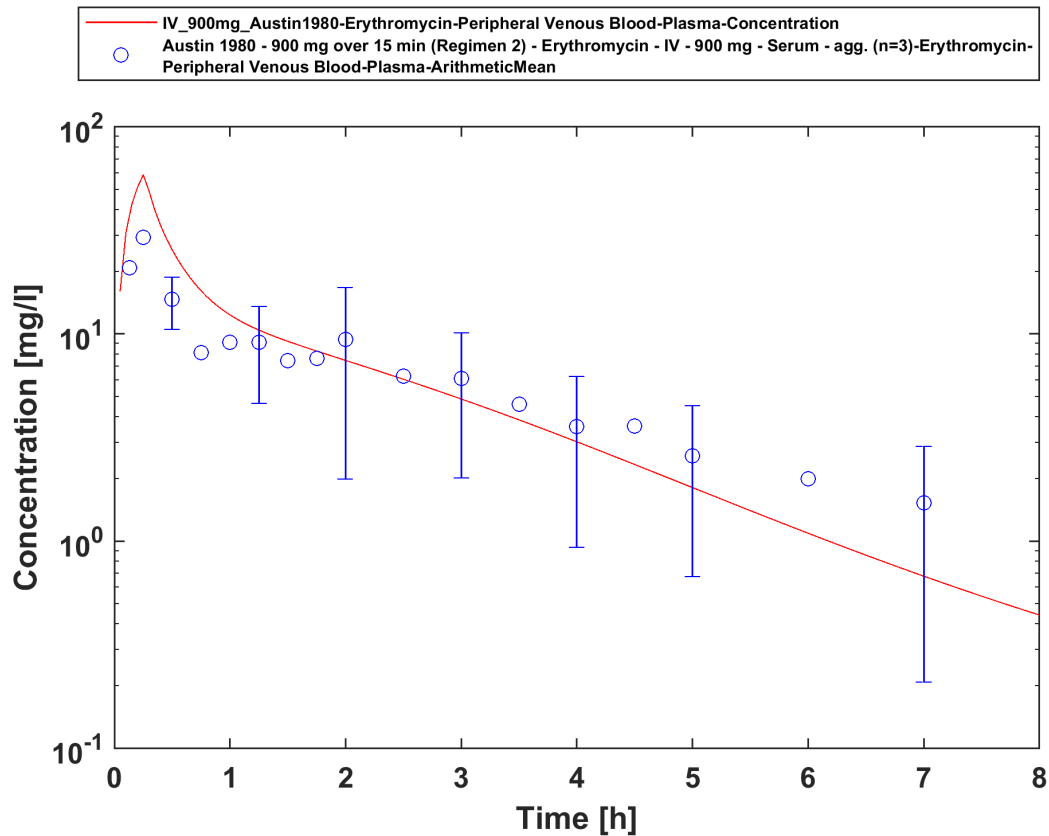
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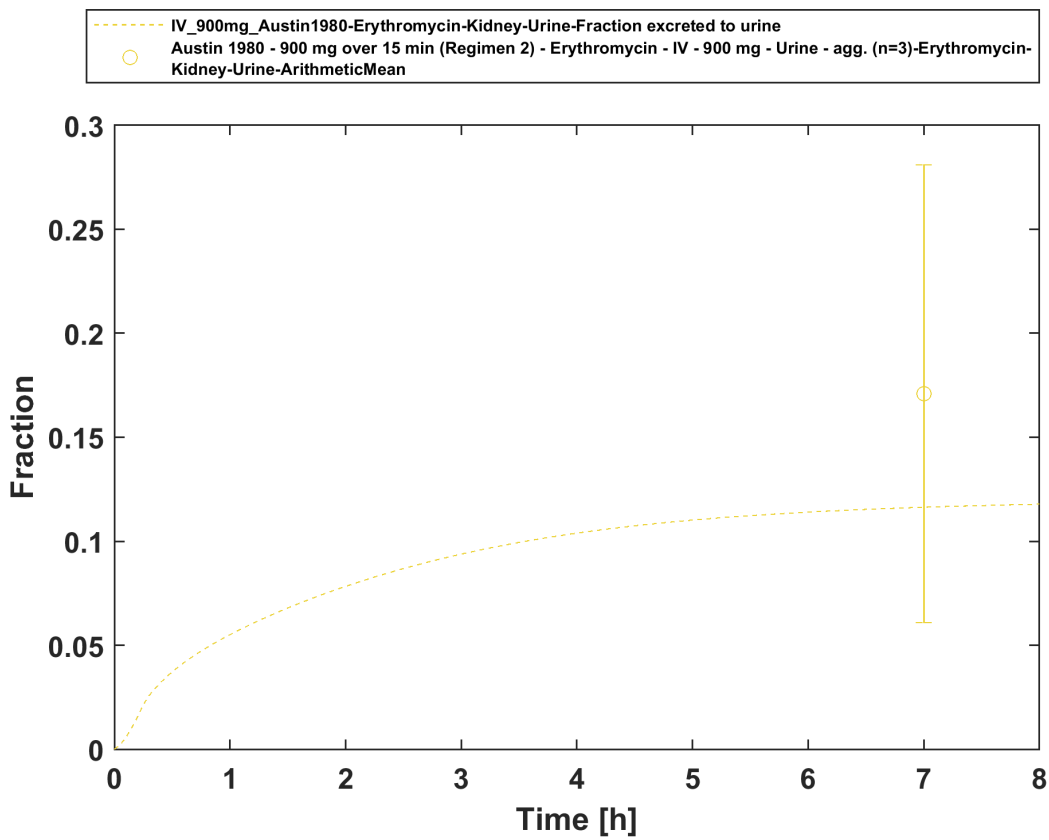
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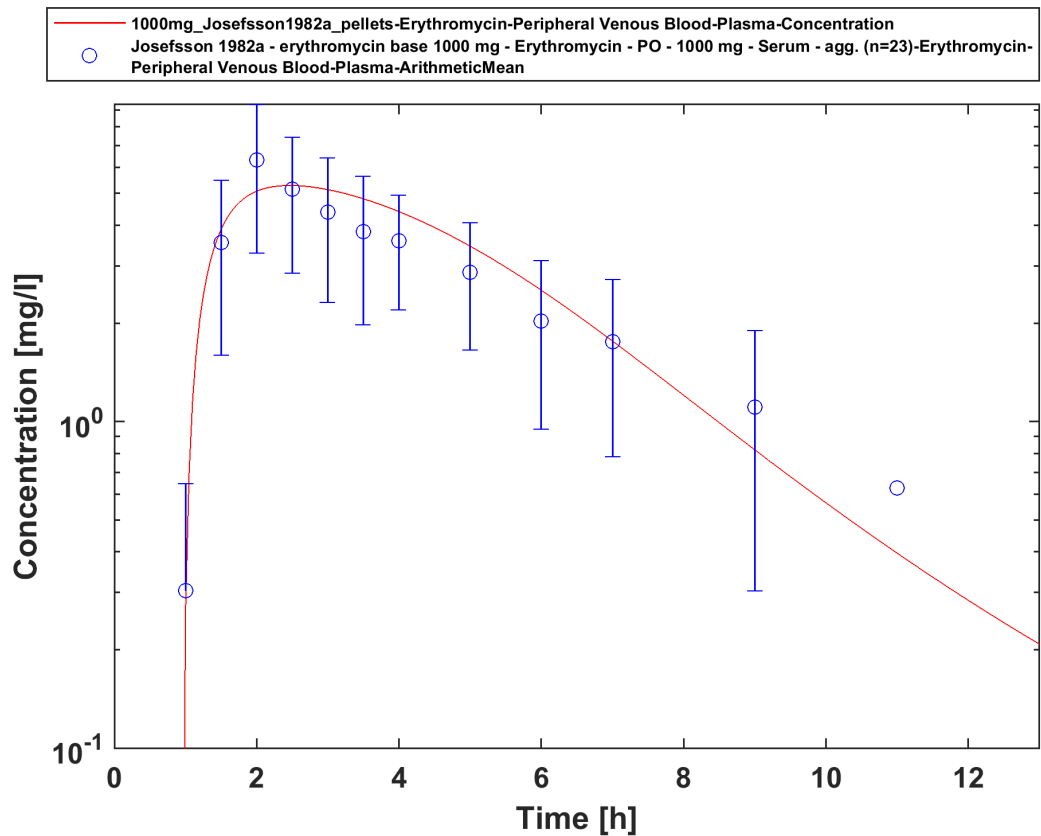
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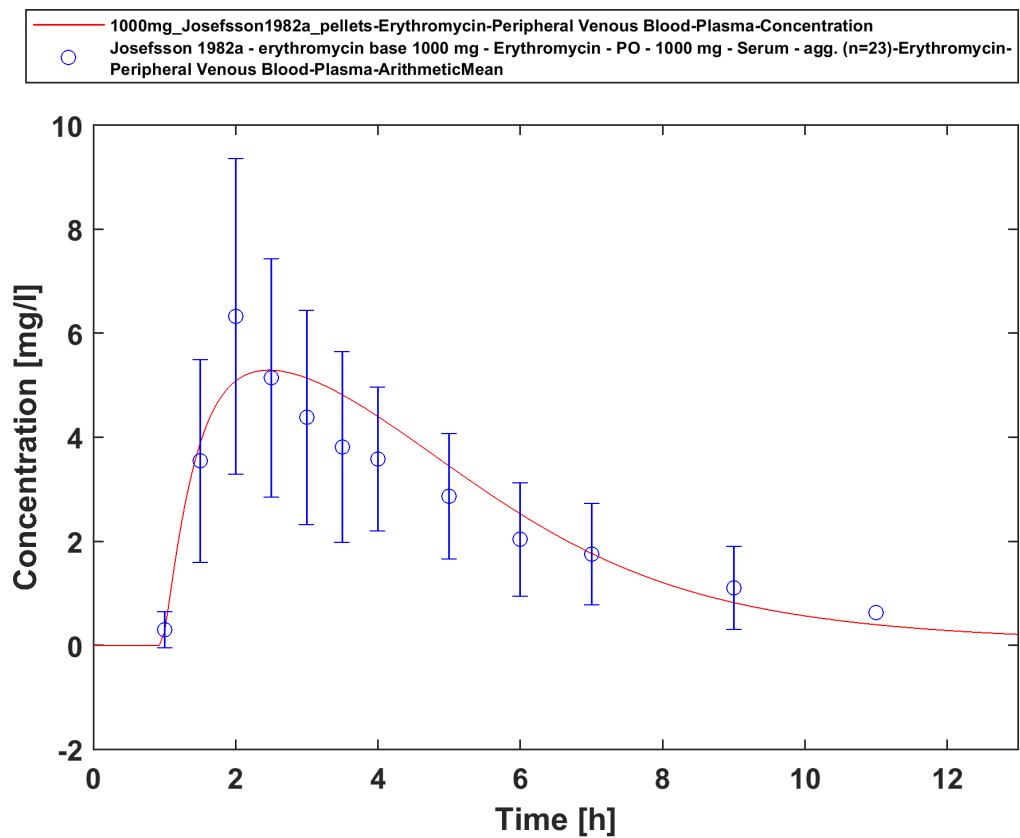
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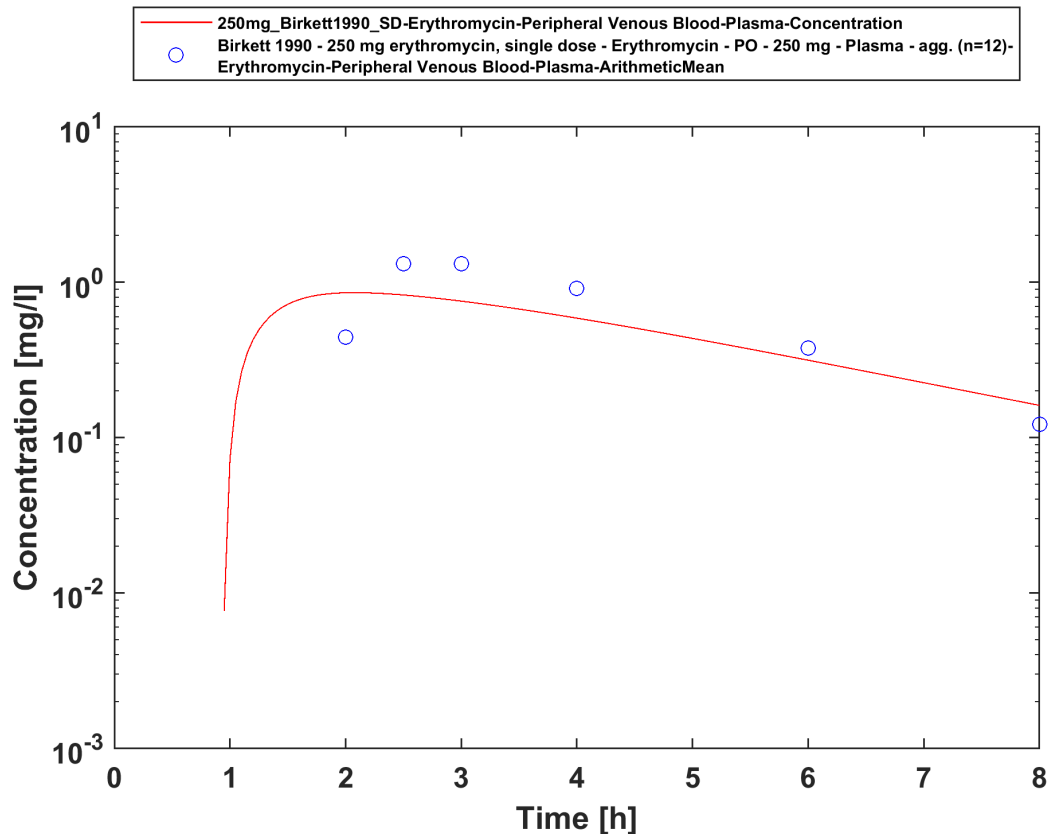
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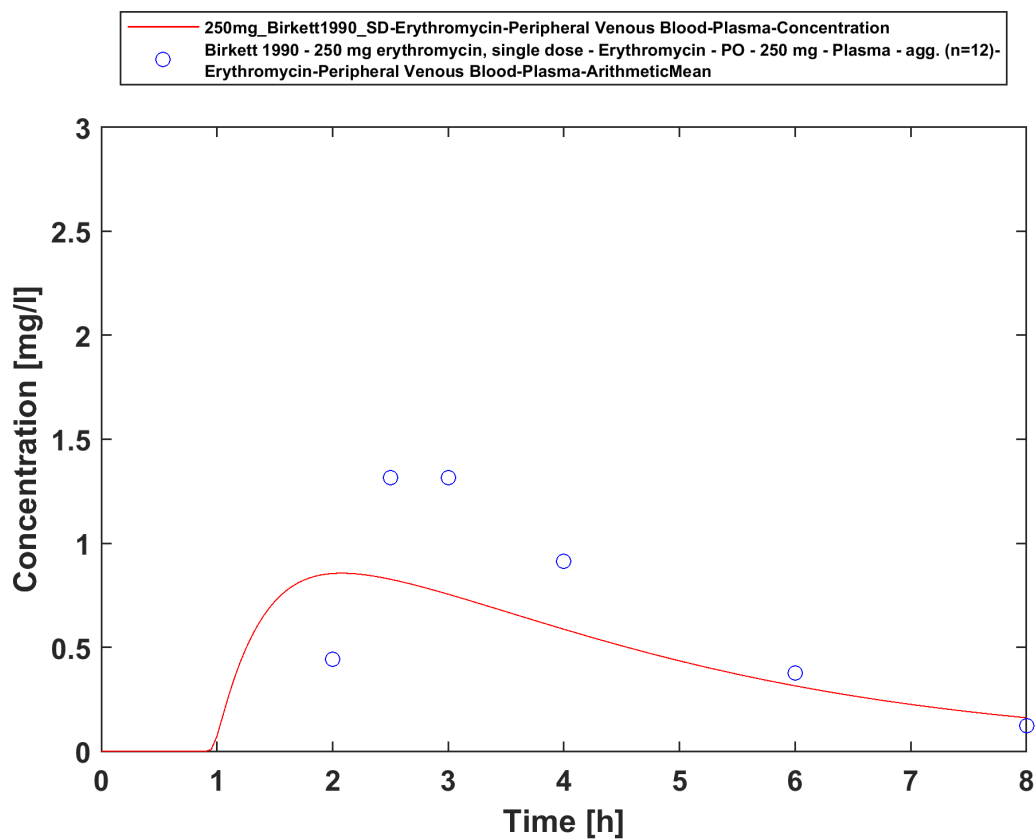
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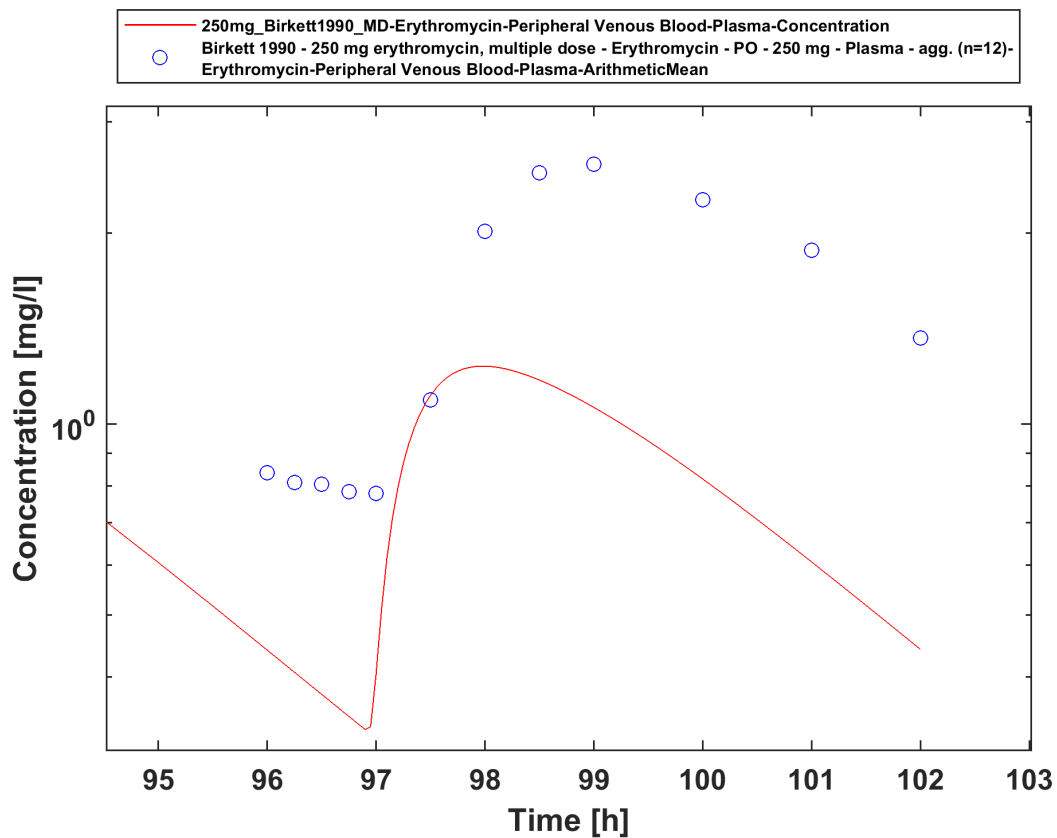
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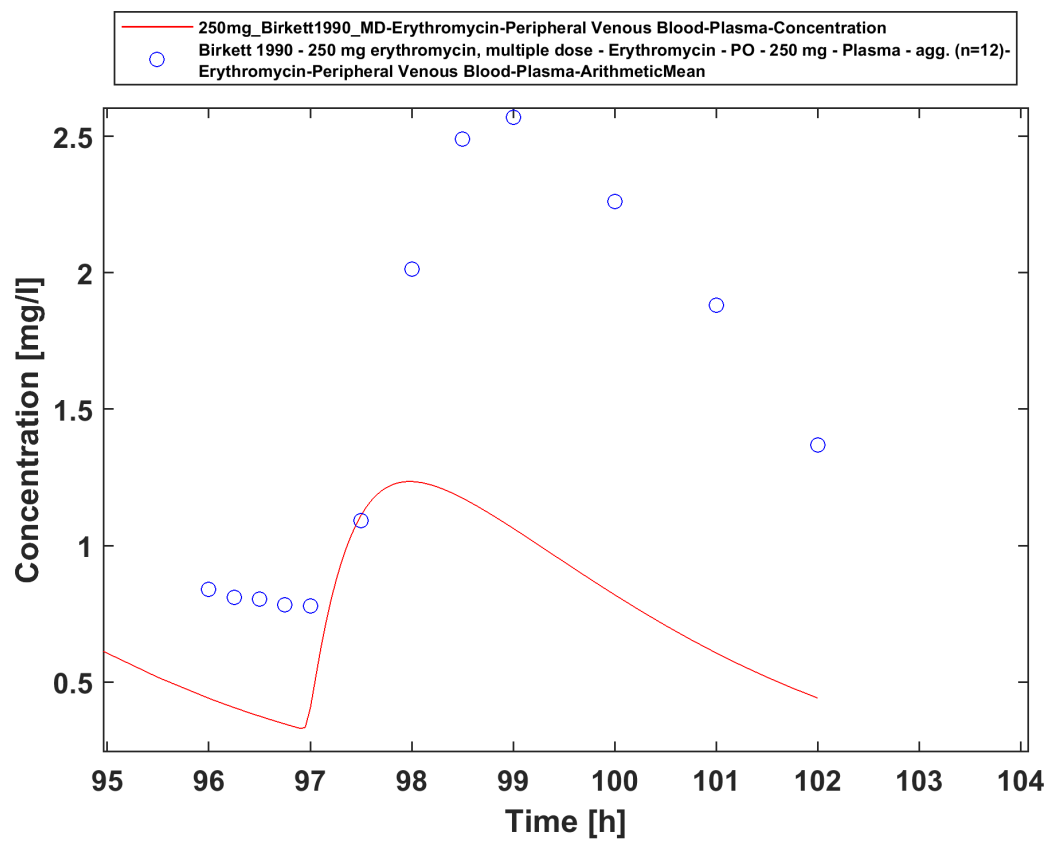
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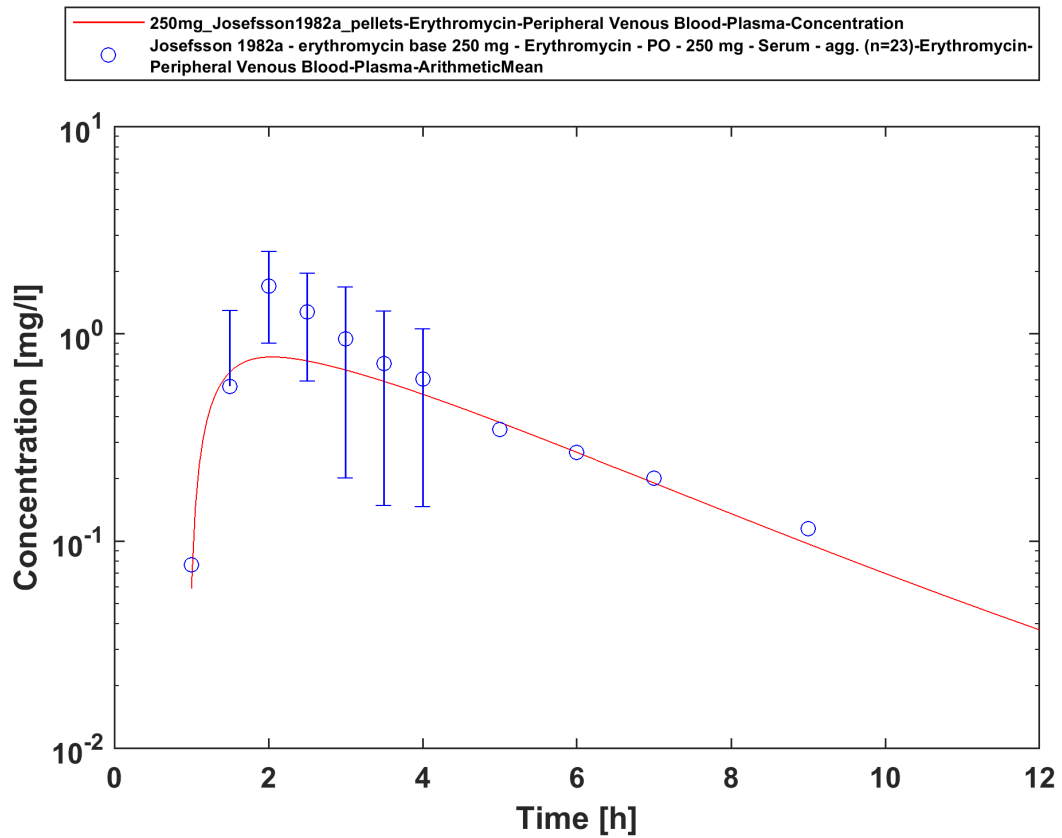
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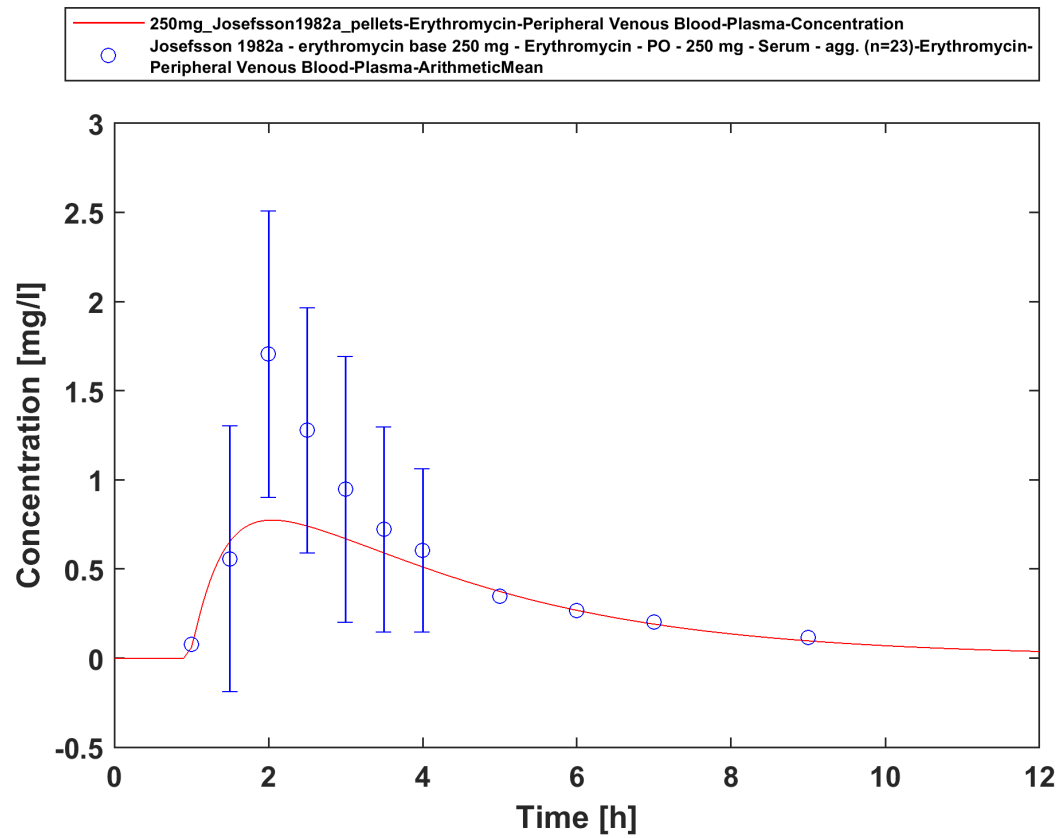
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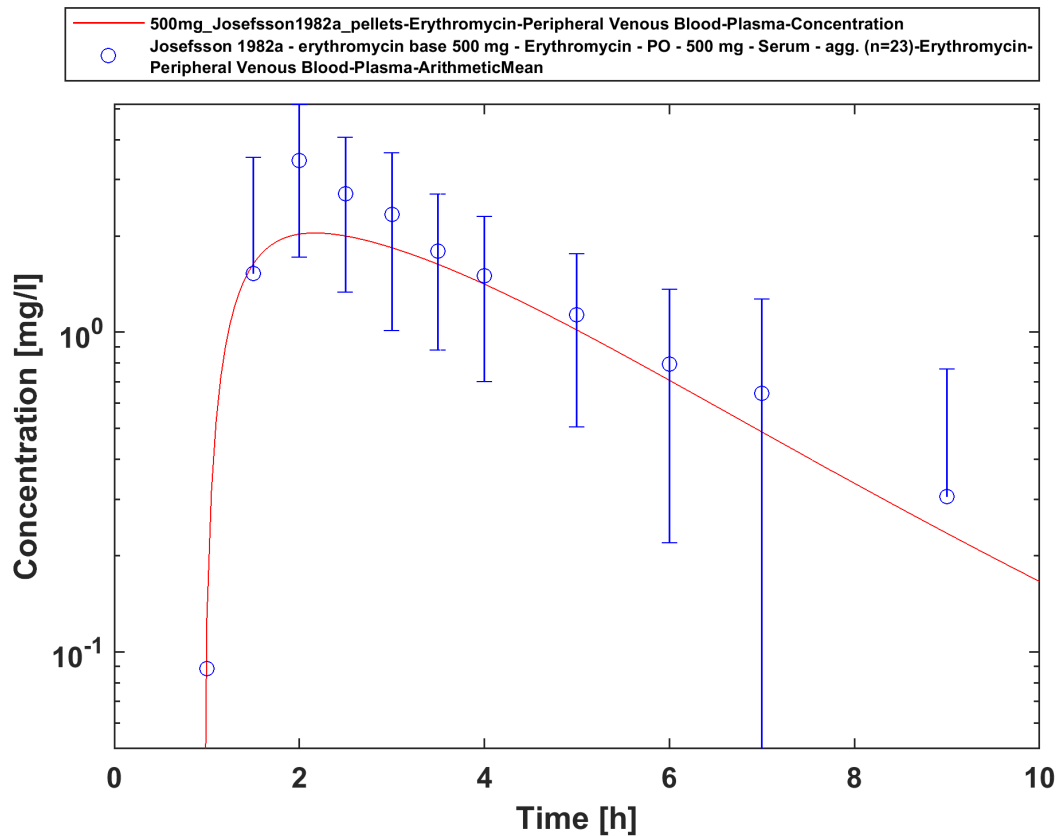
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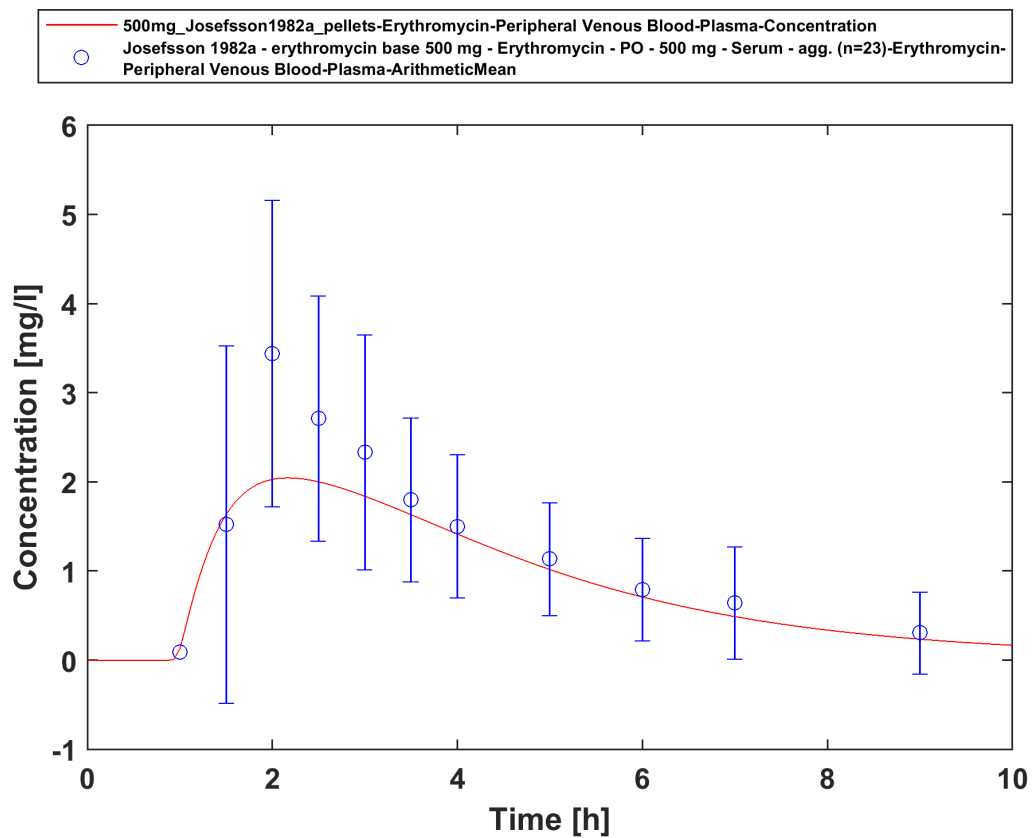
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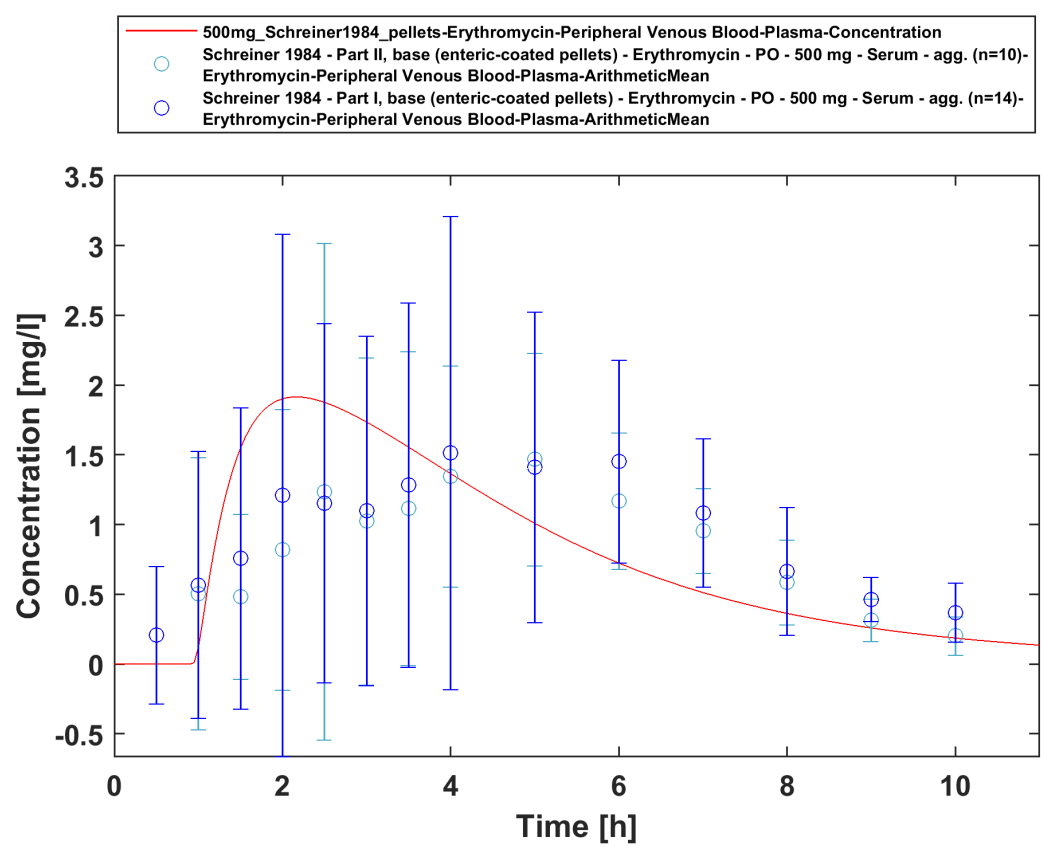
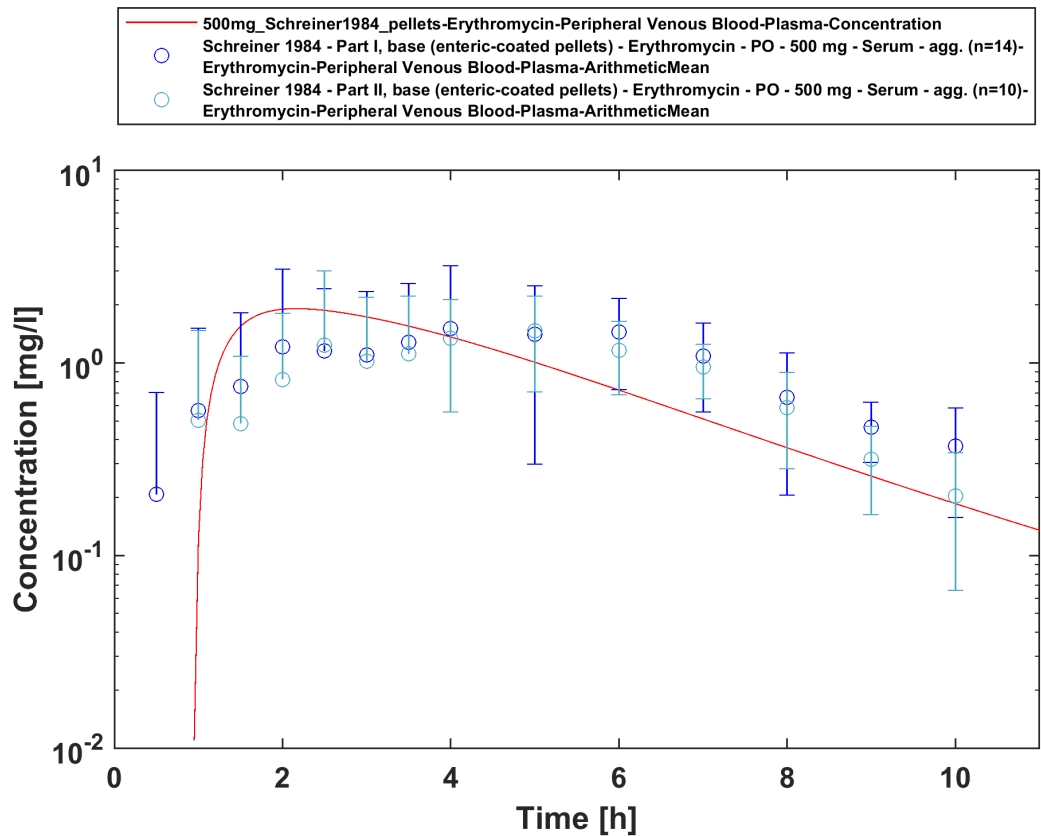
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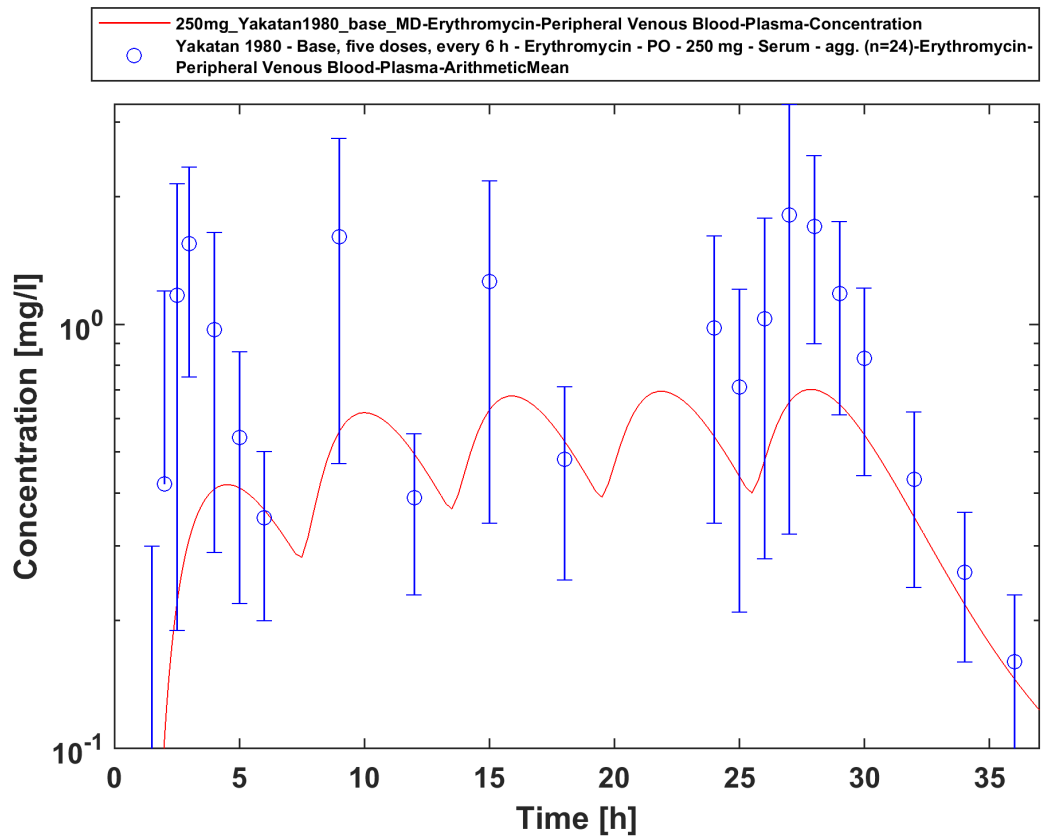


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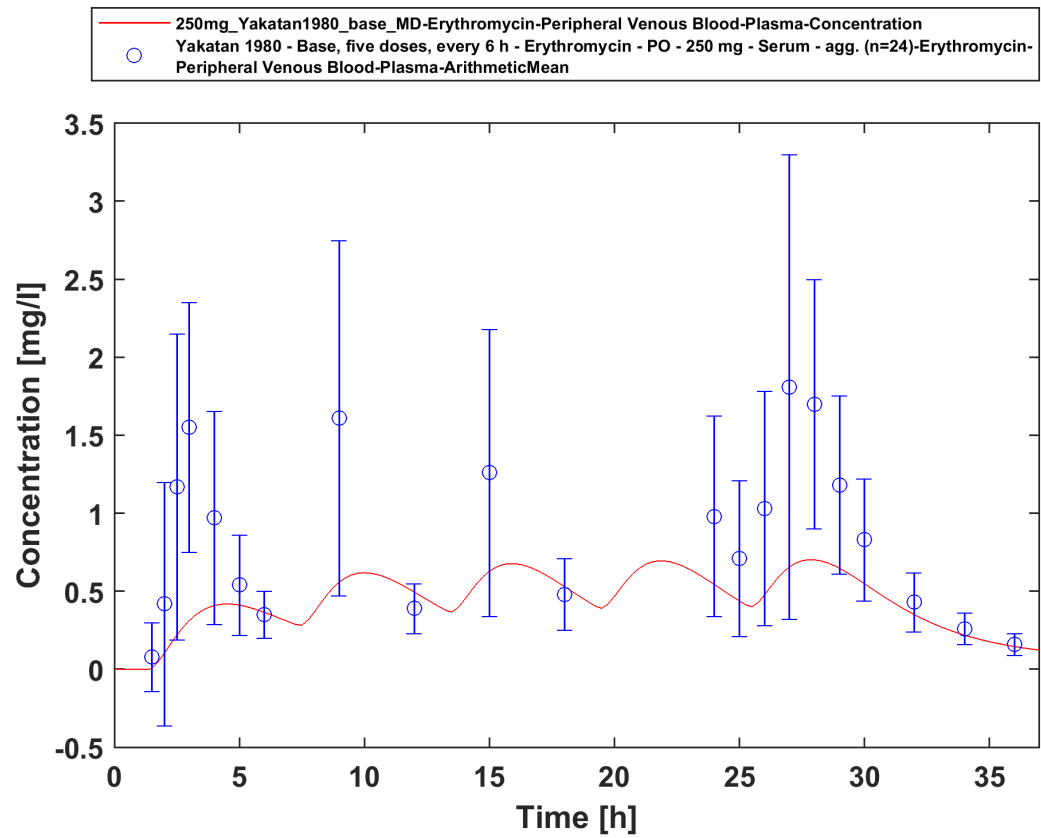


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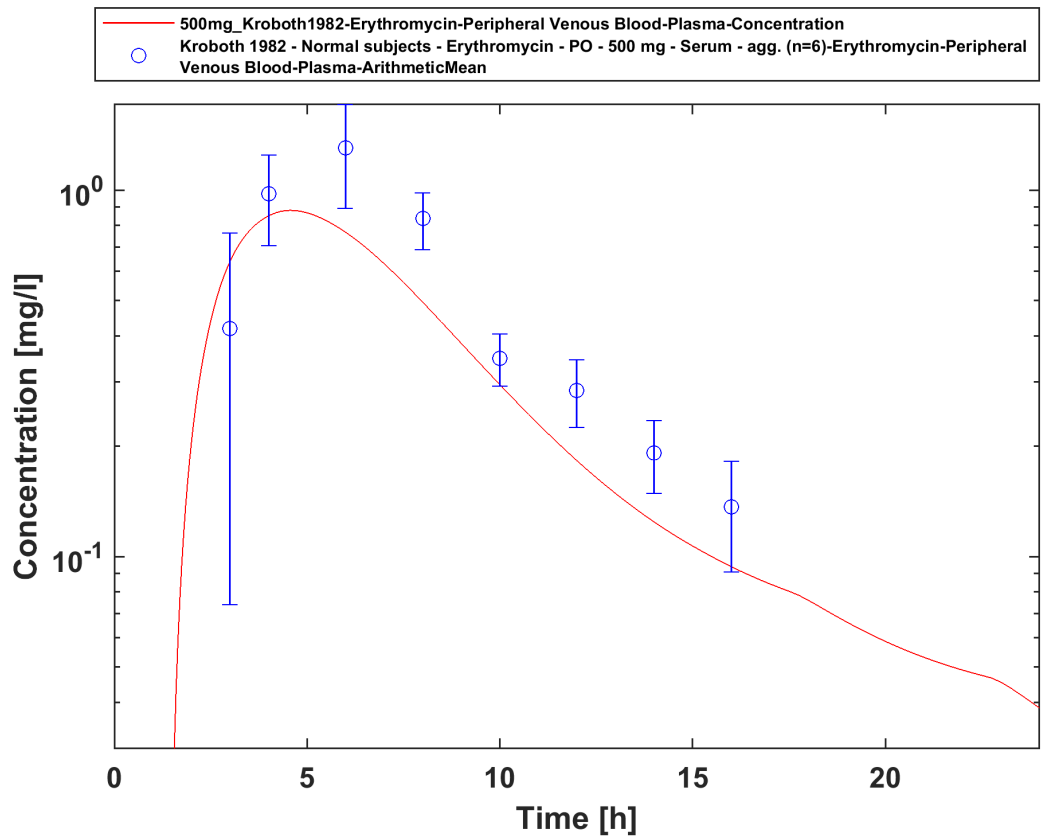




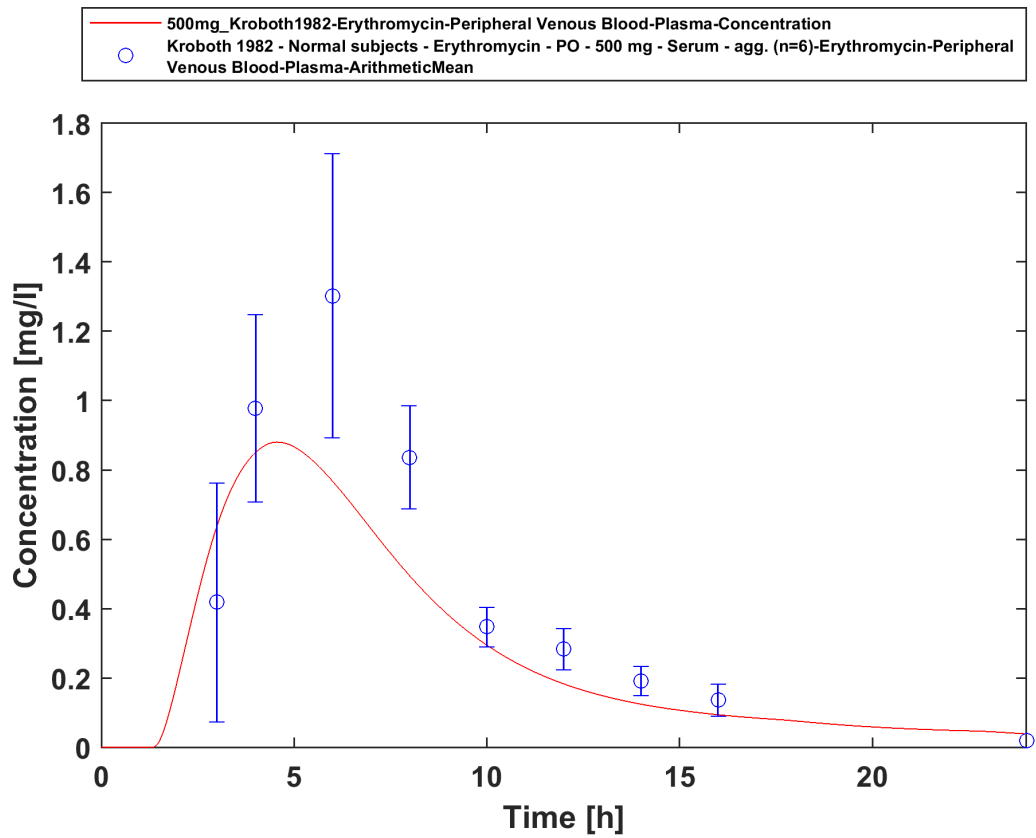
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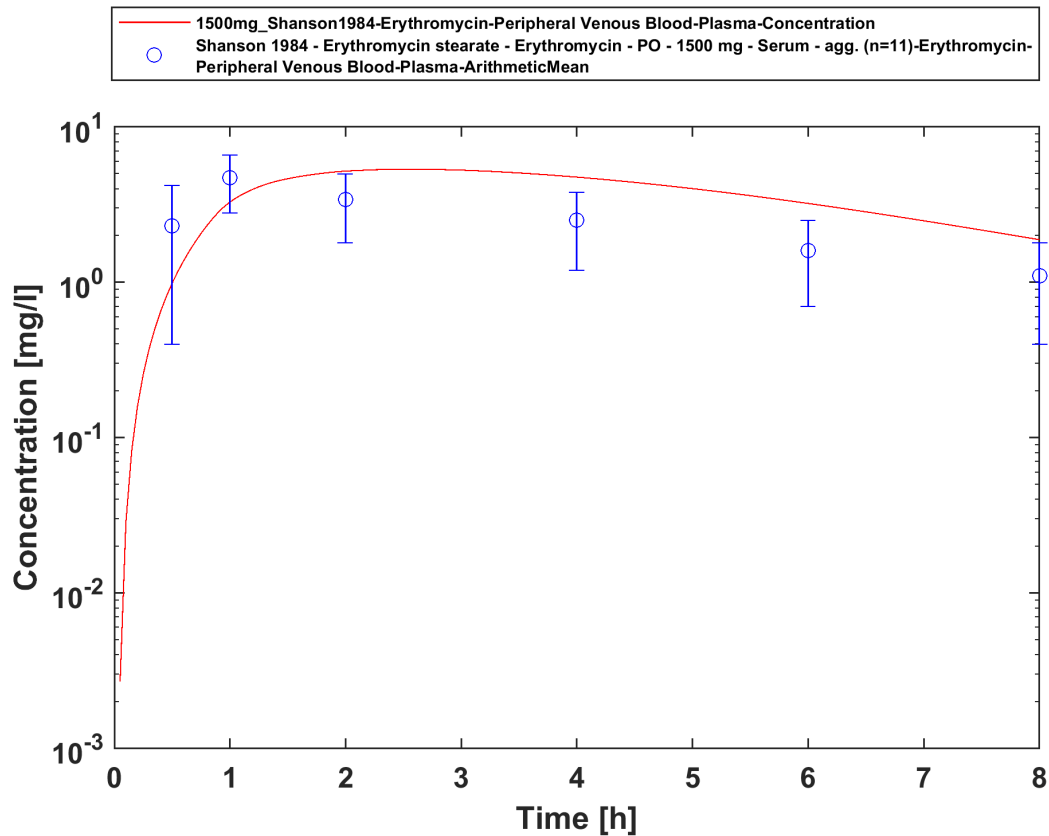
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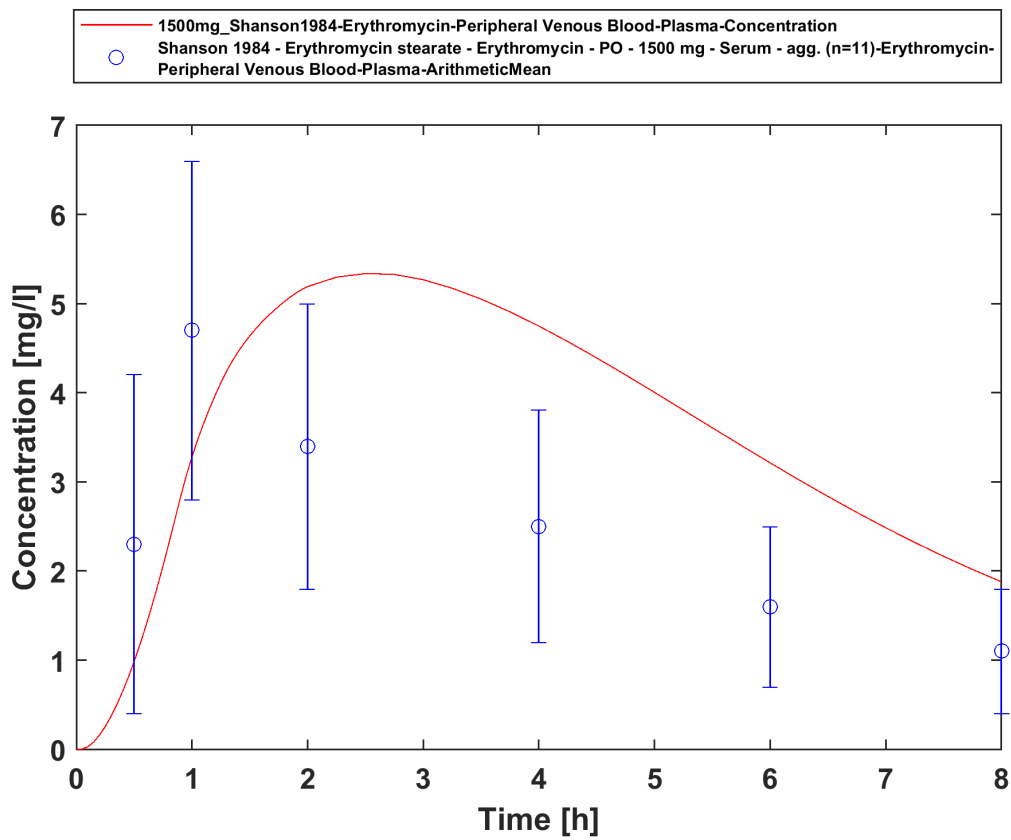
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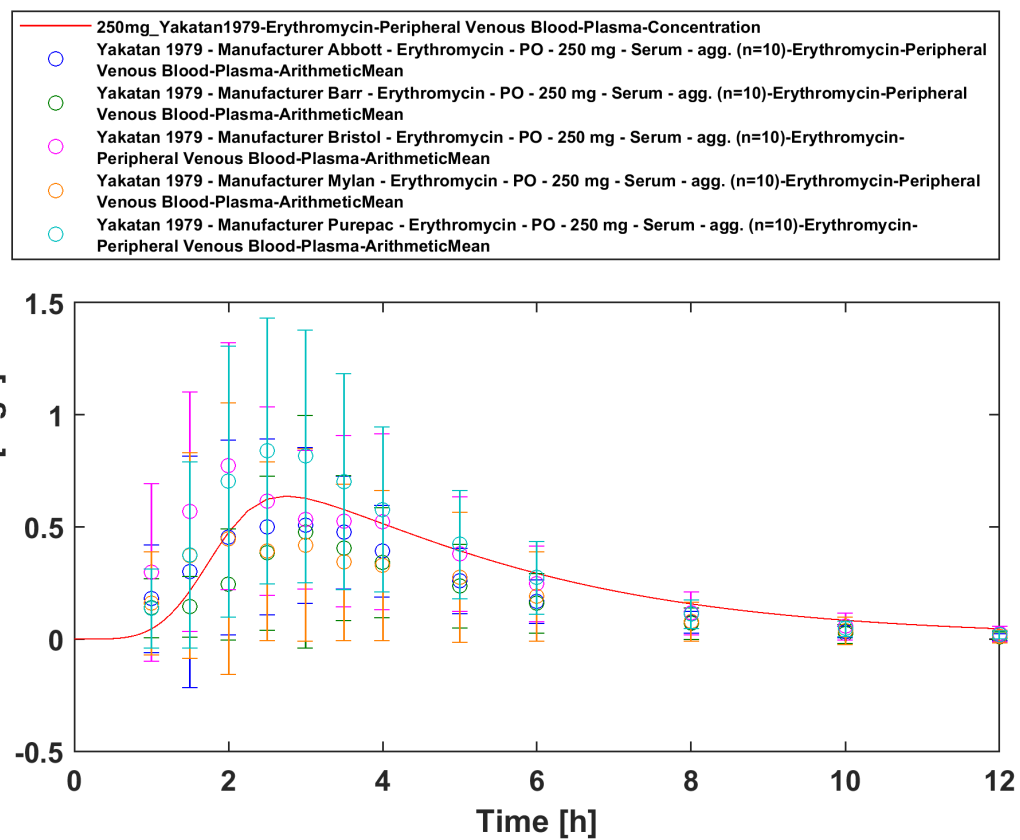
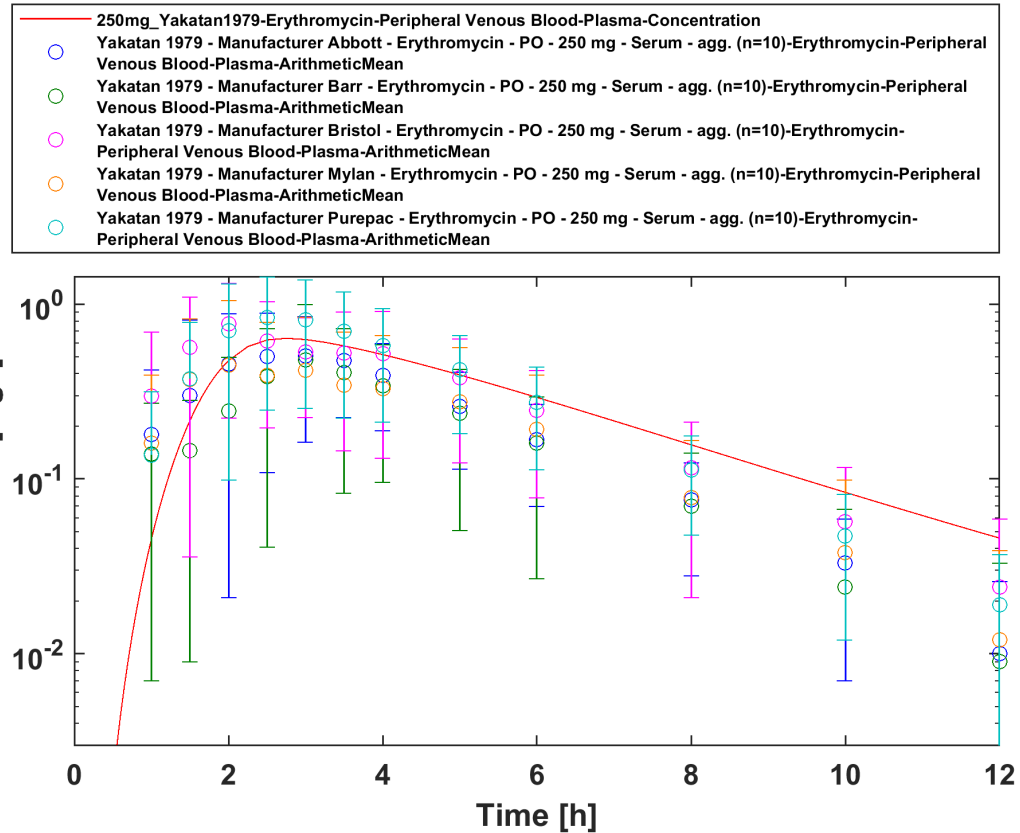
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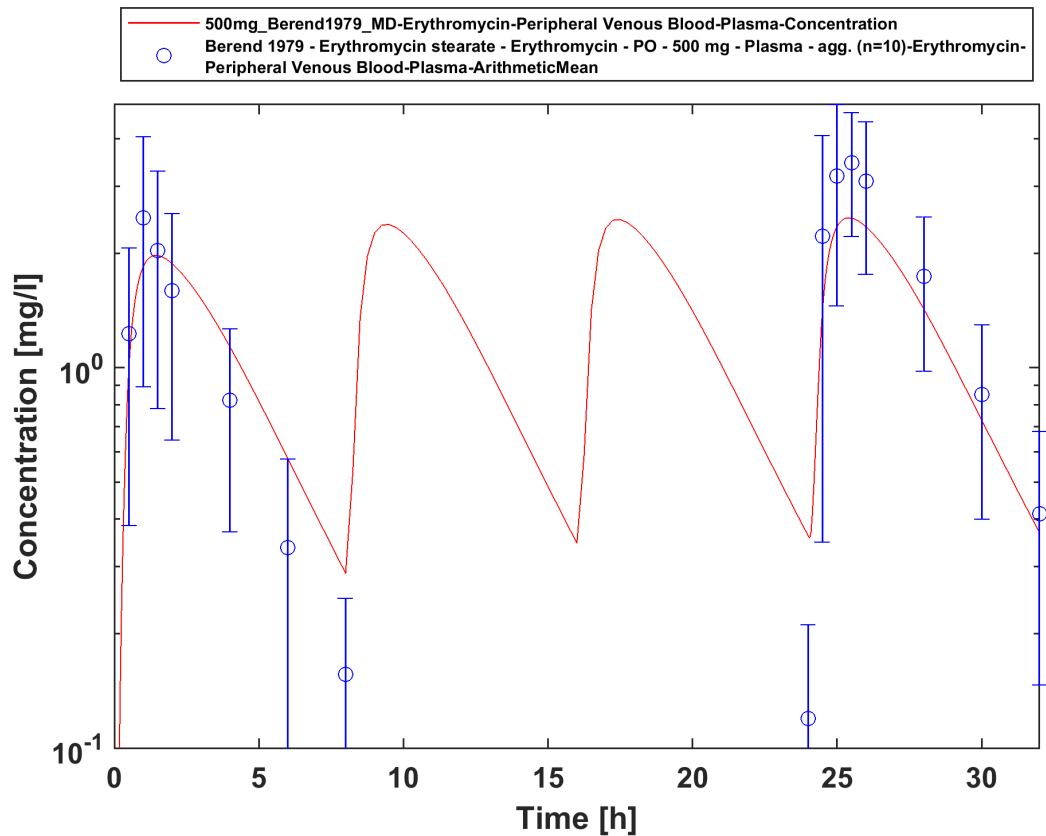


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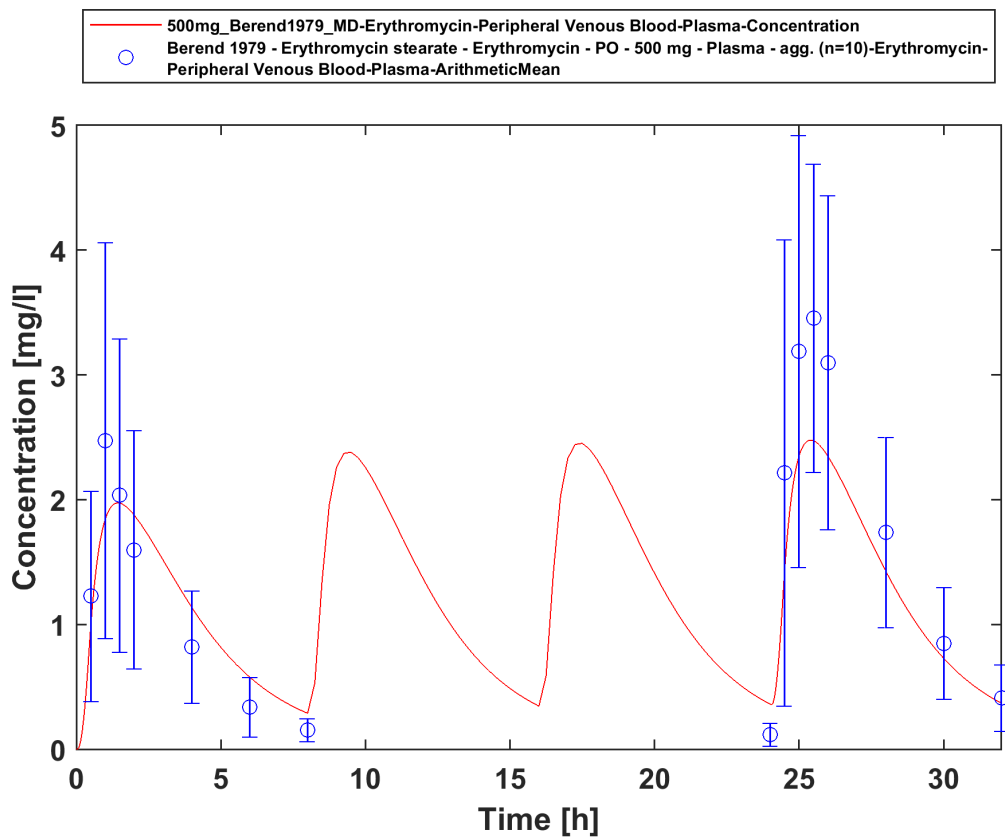


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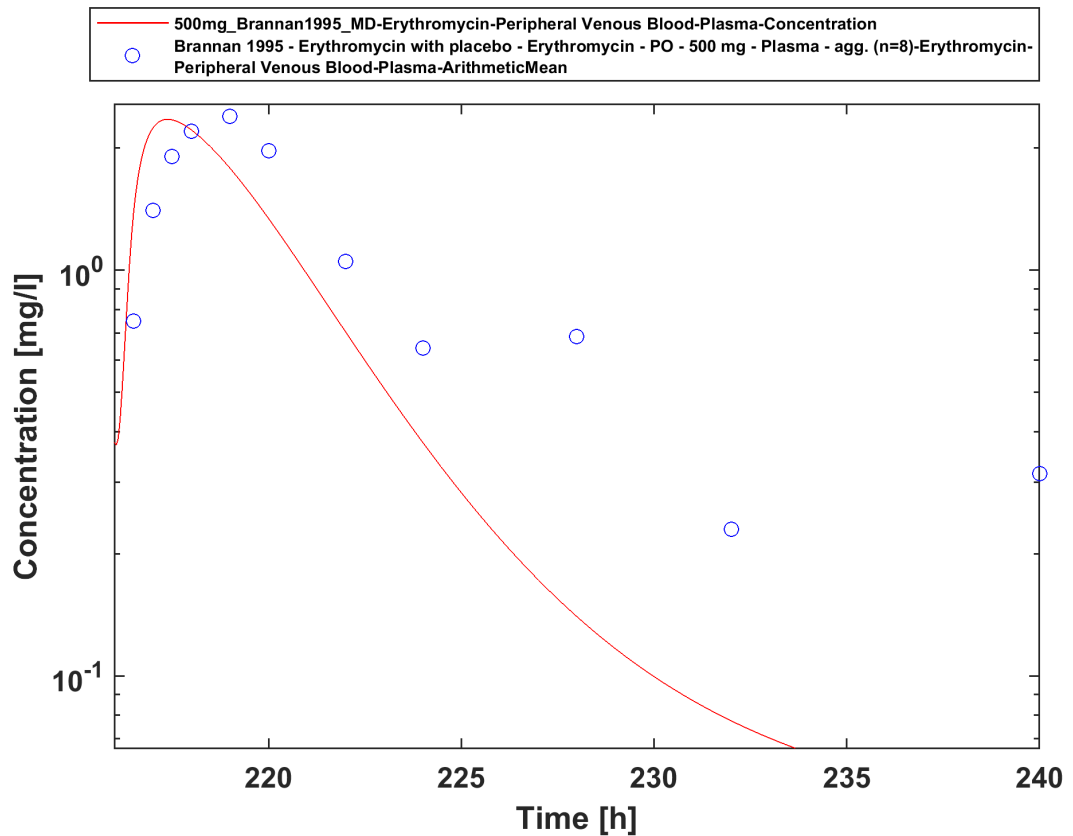




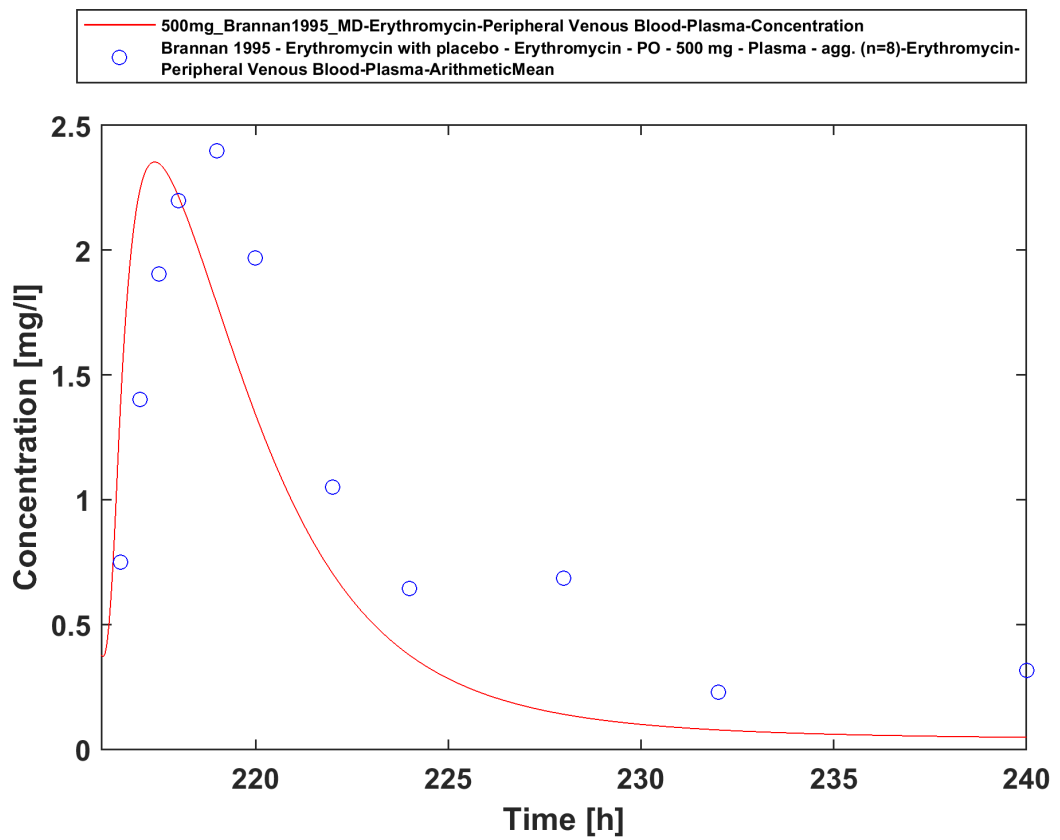
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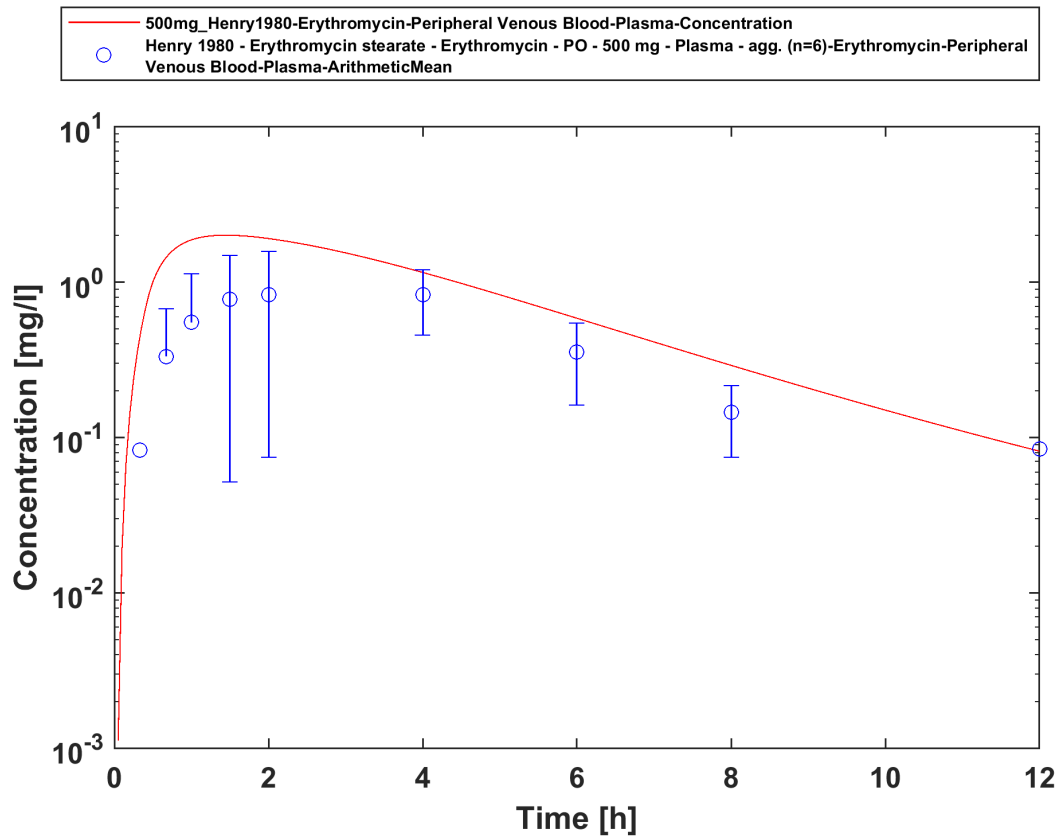
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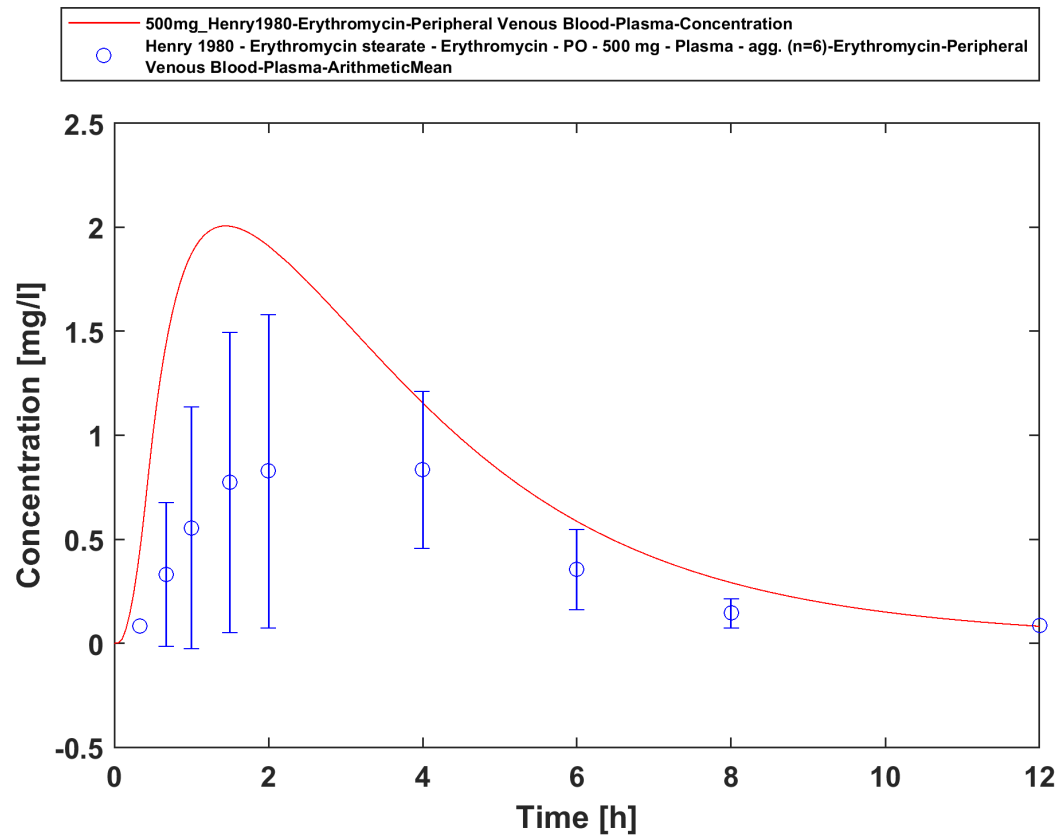
Time Profile Analysis



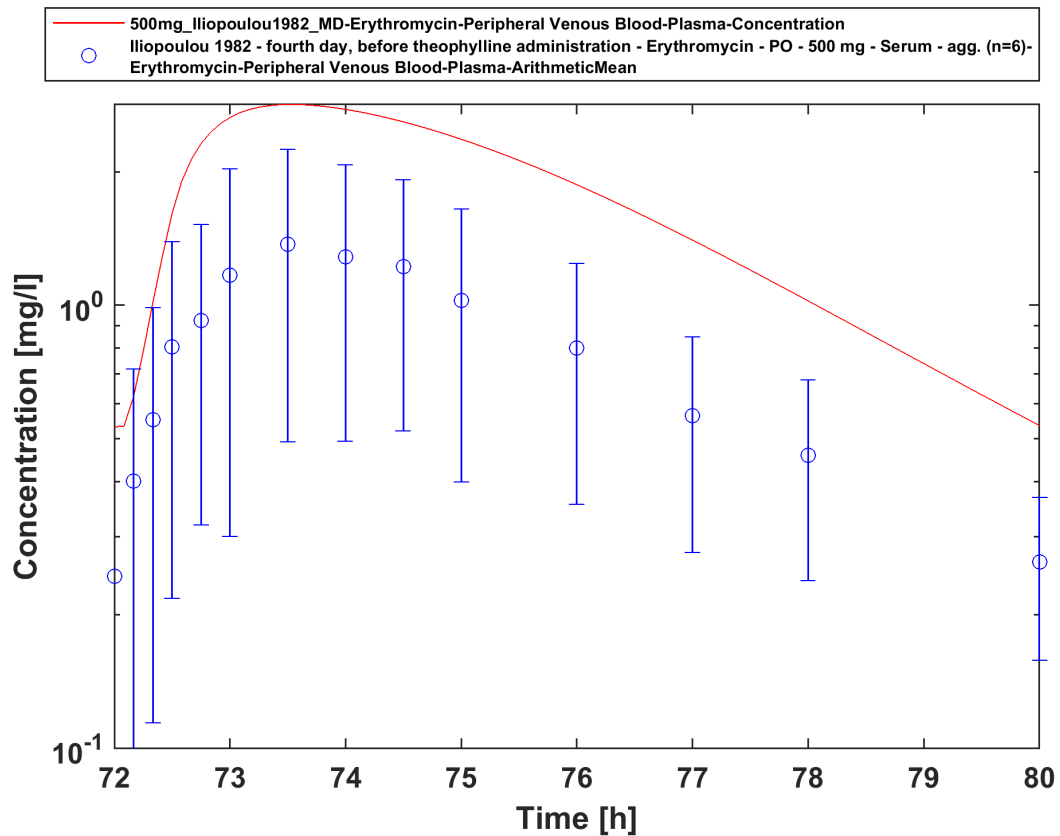
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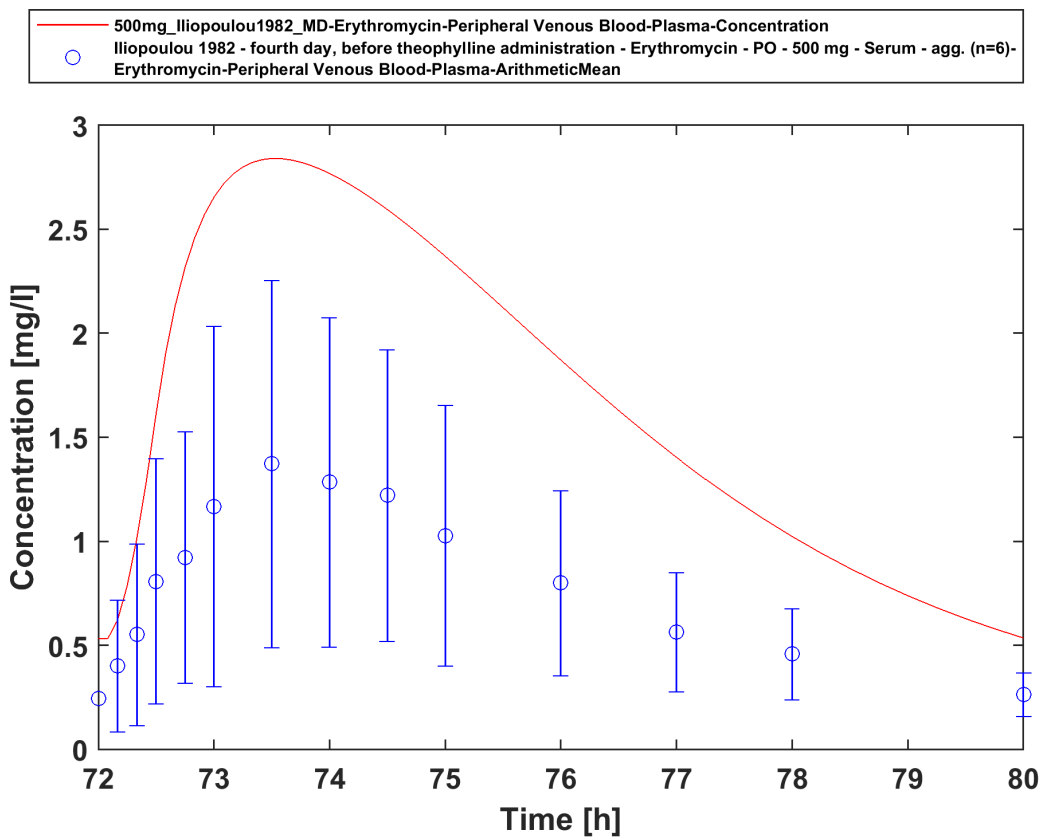
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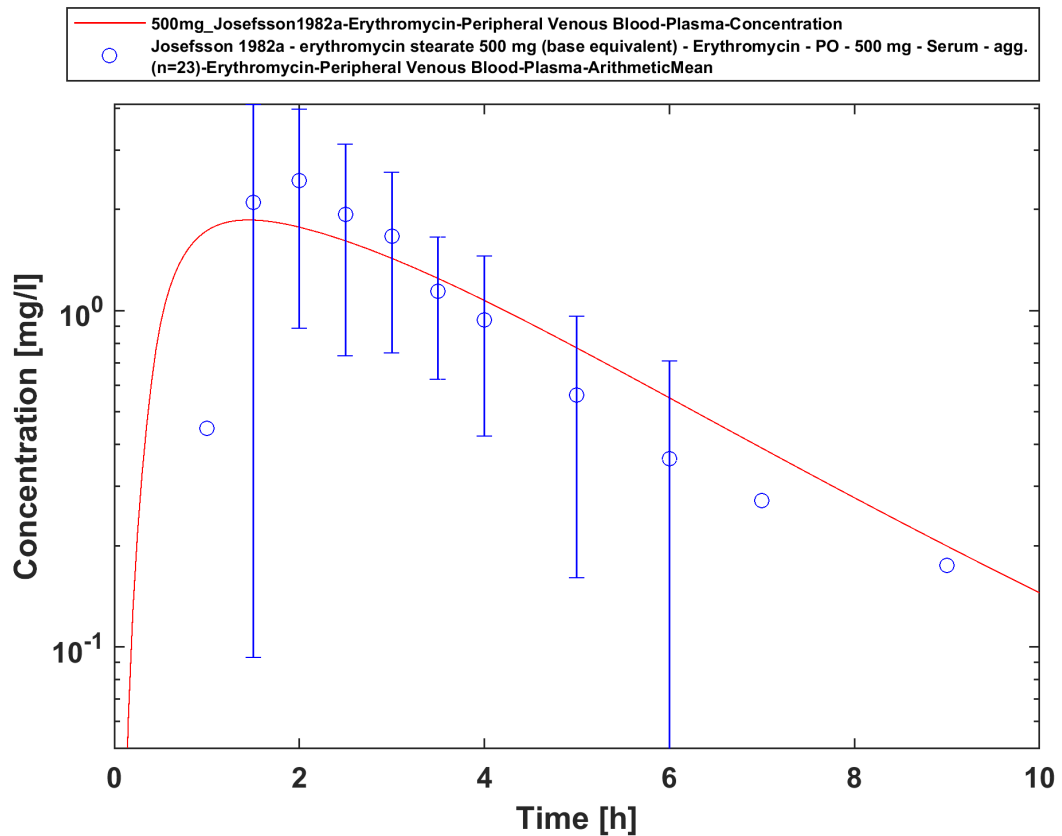
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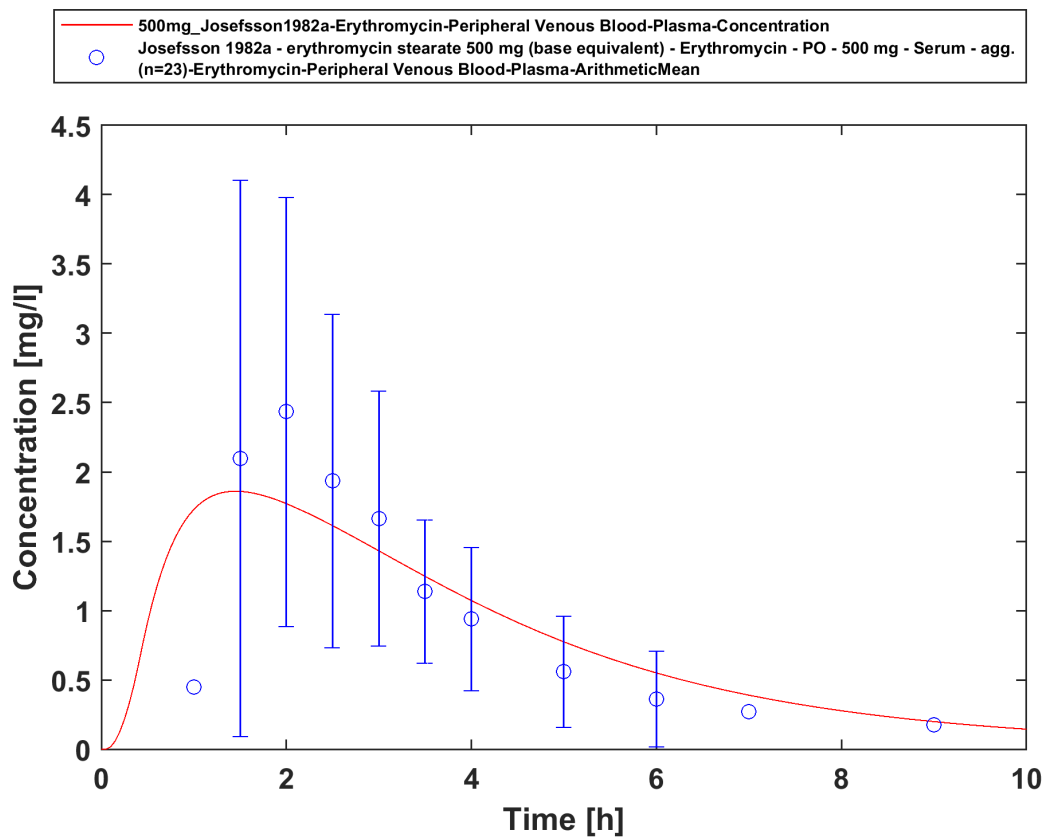
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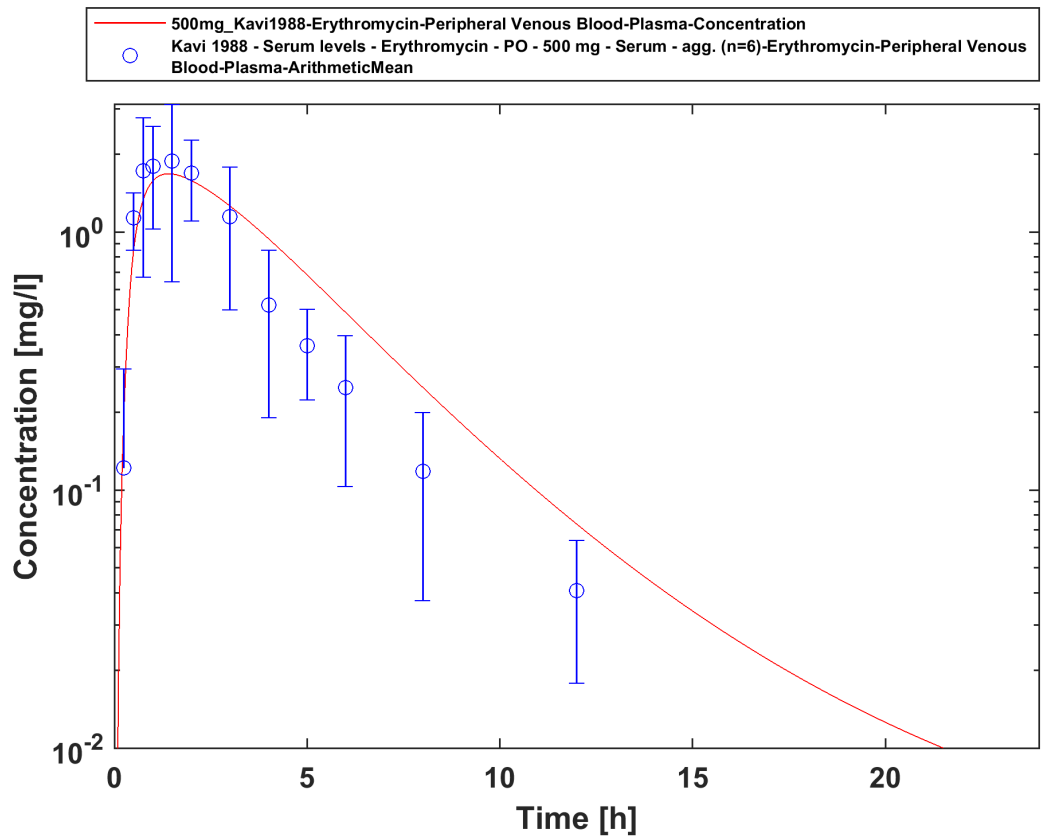
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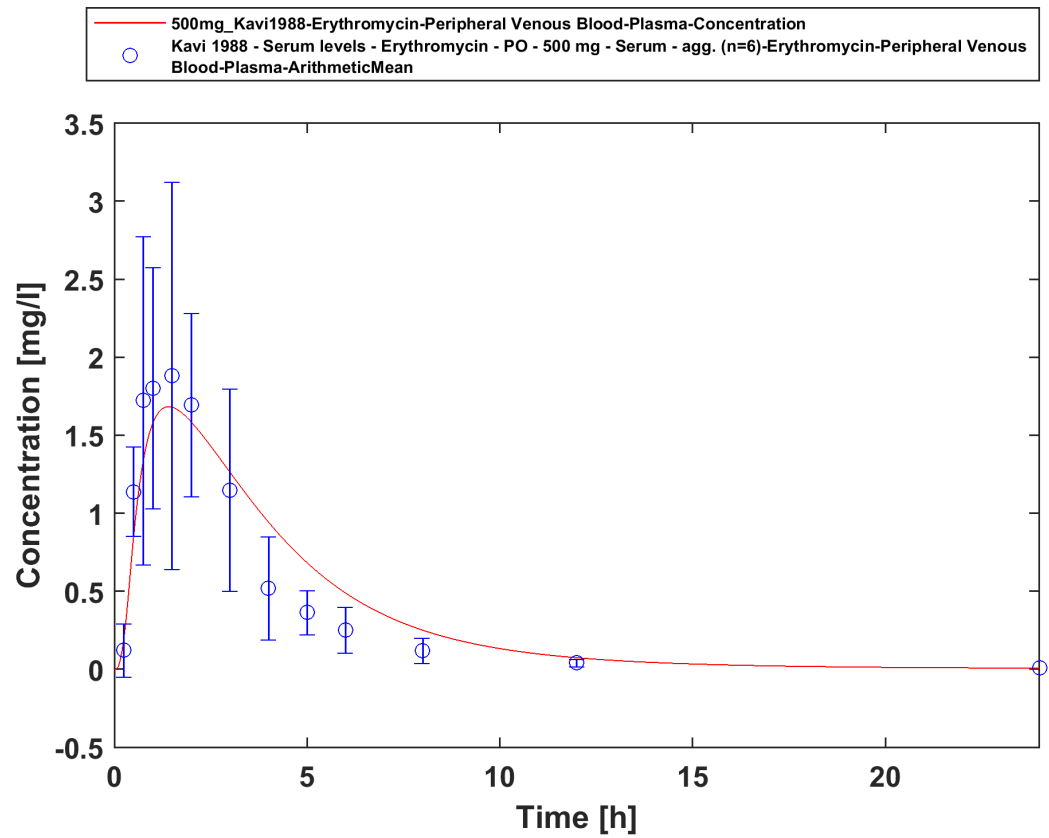
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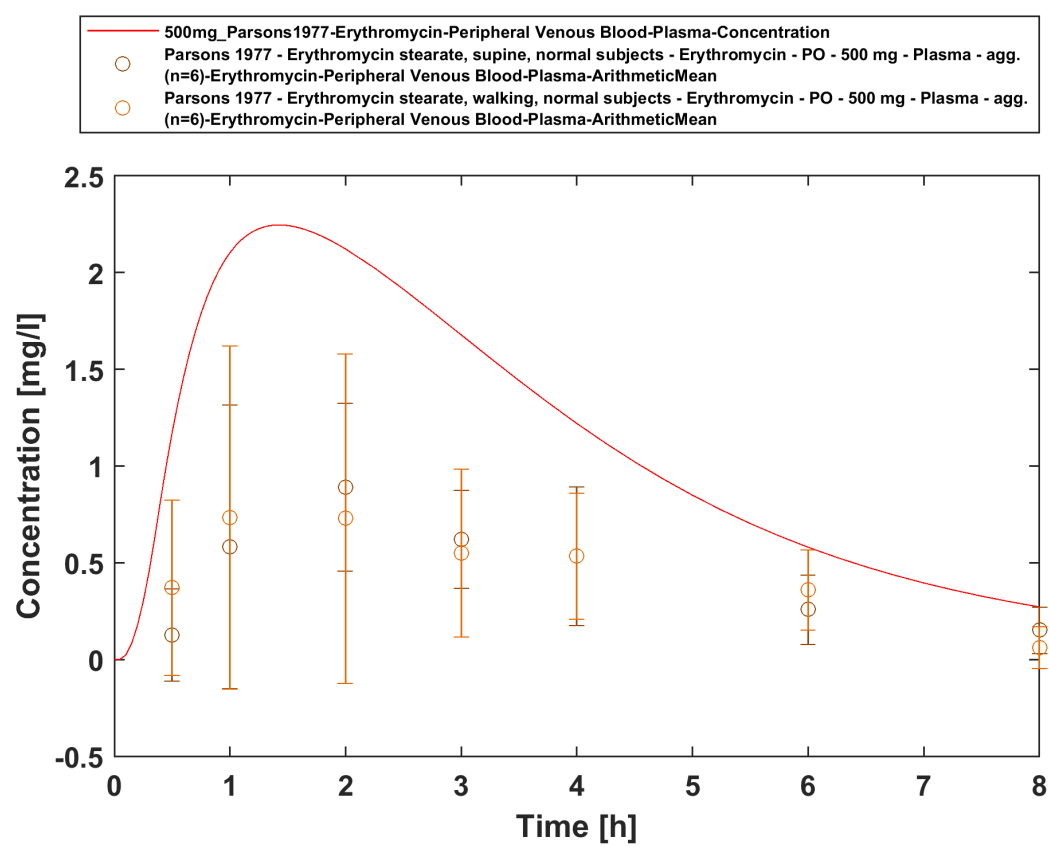
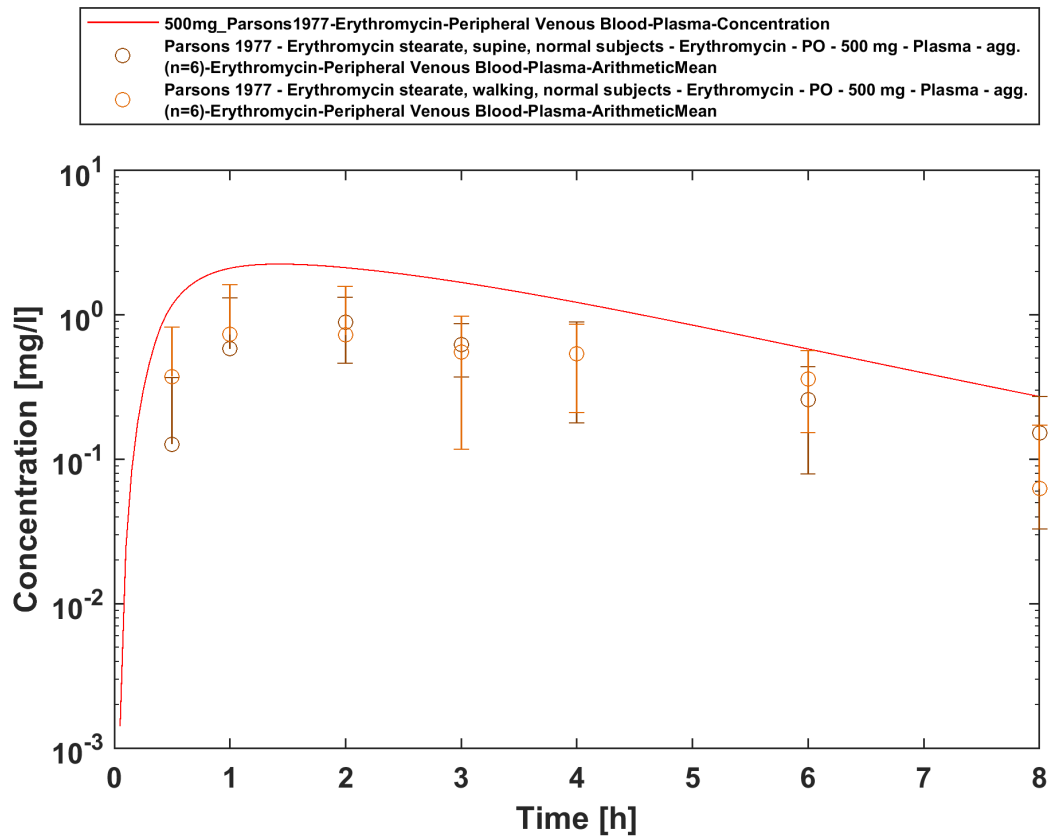
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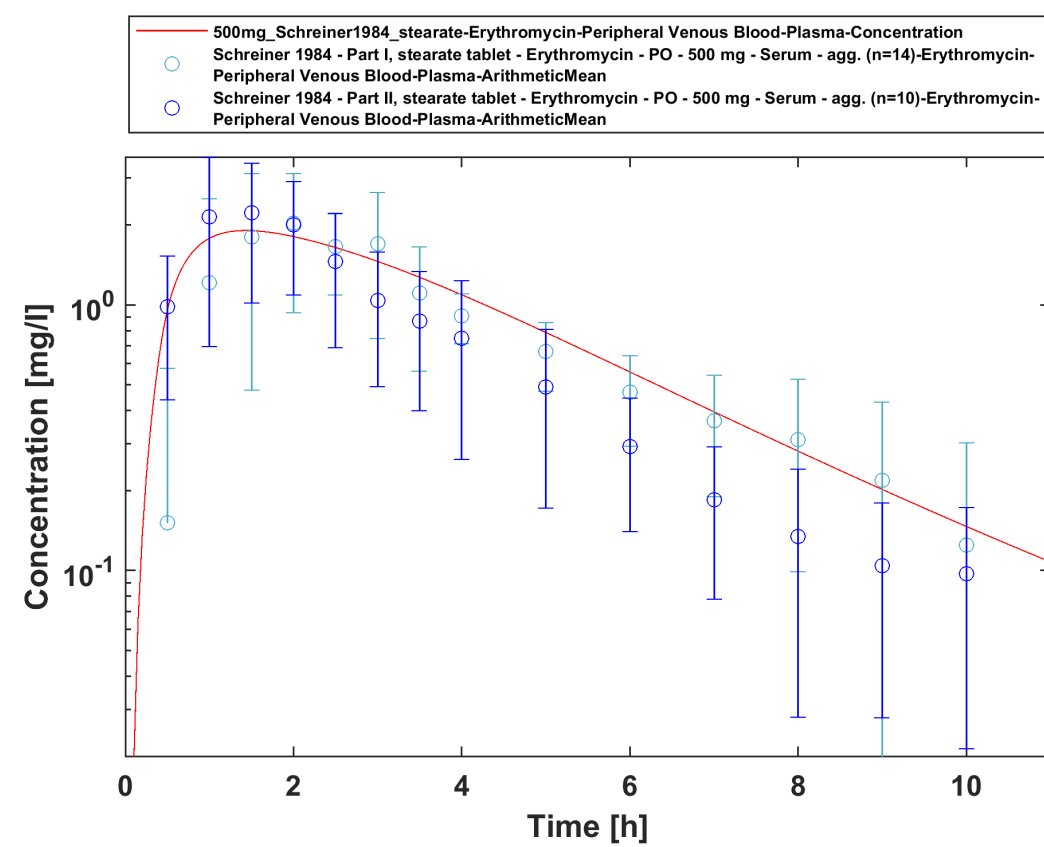
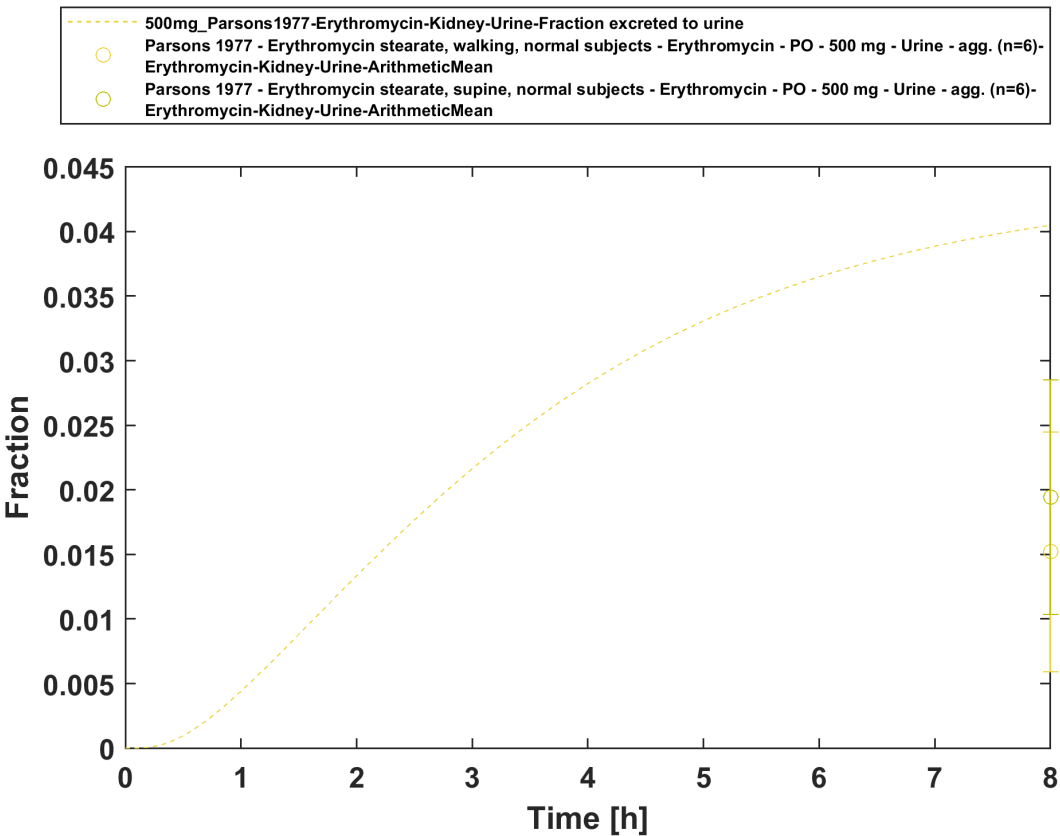


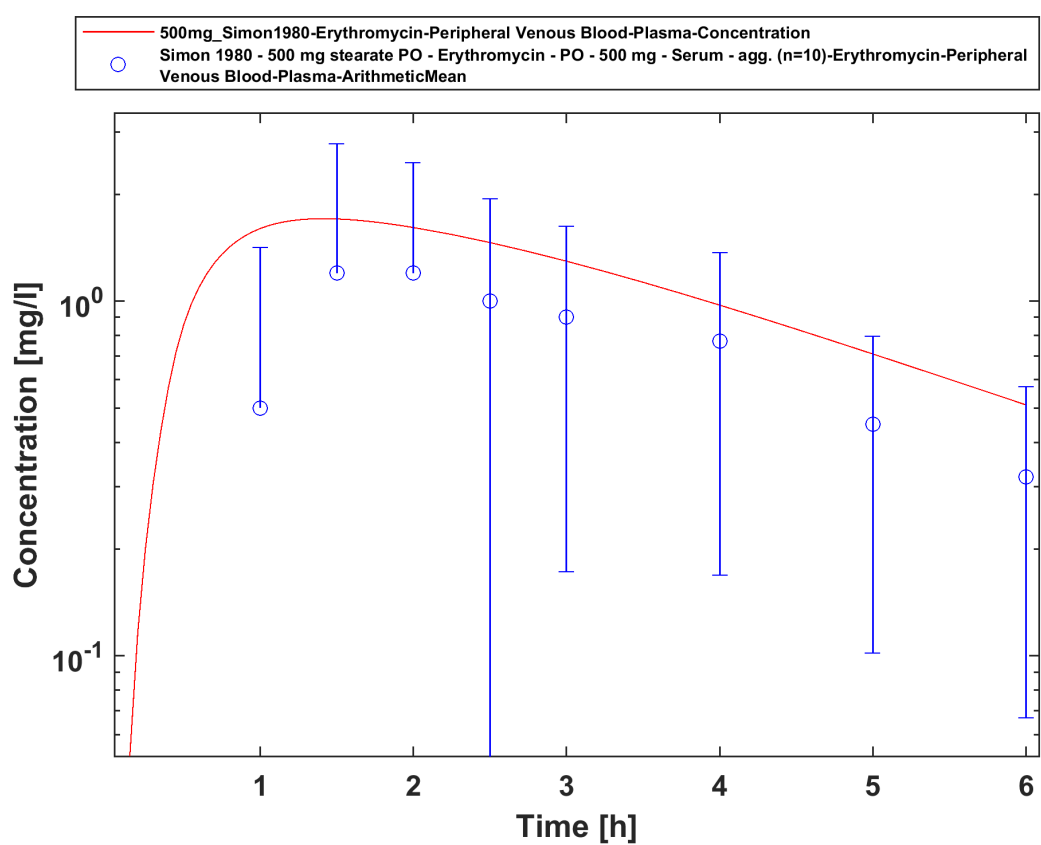
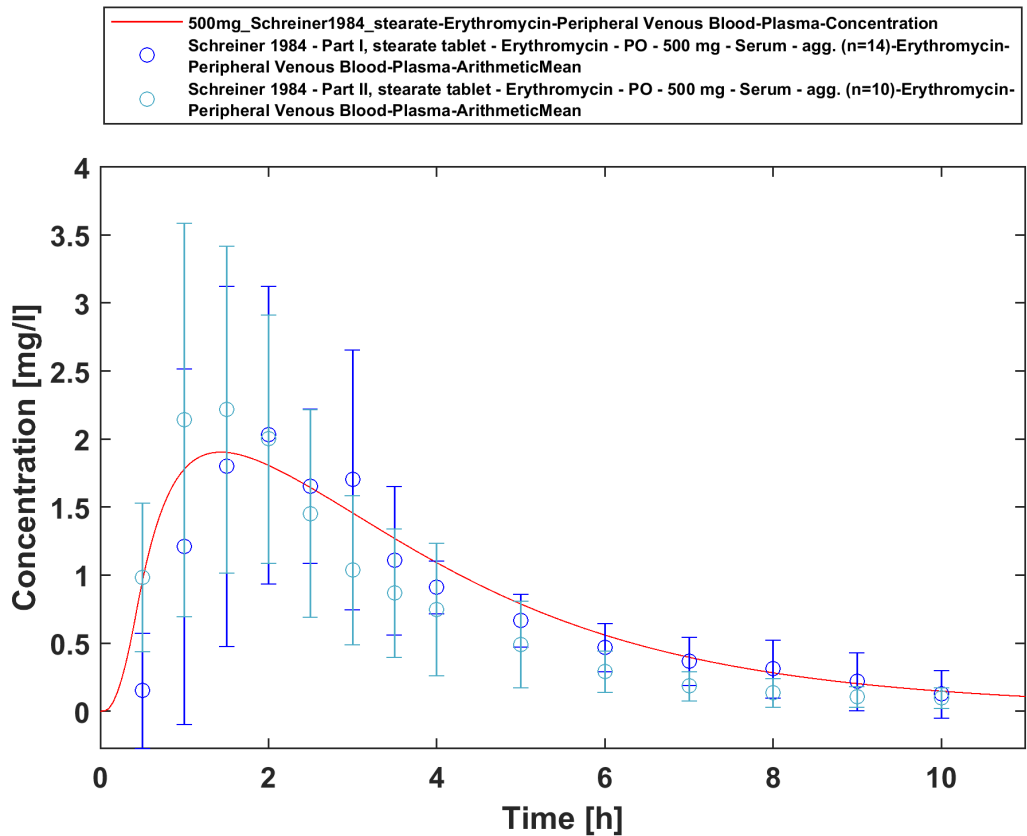
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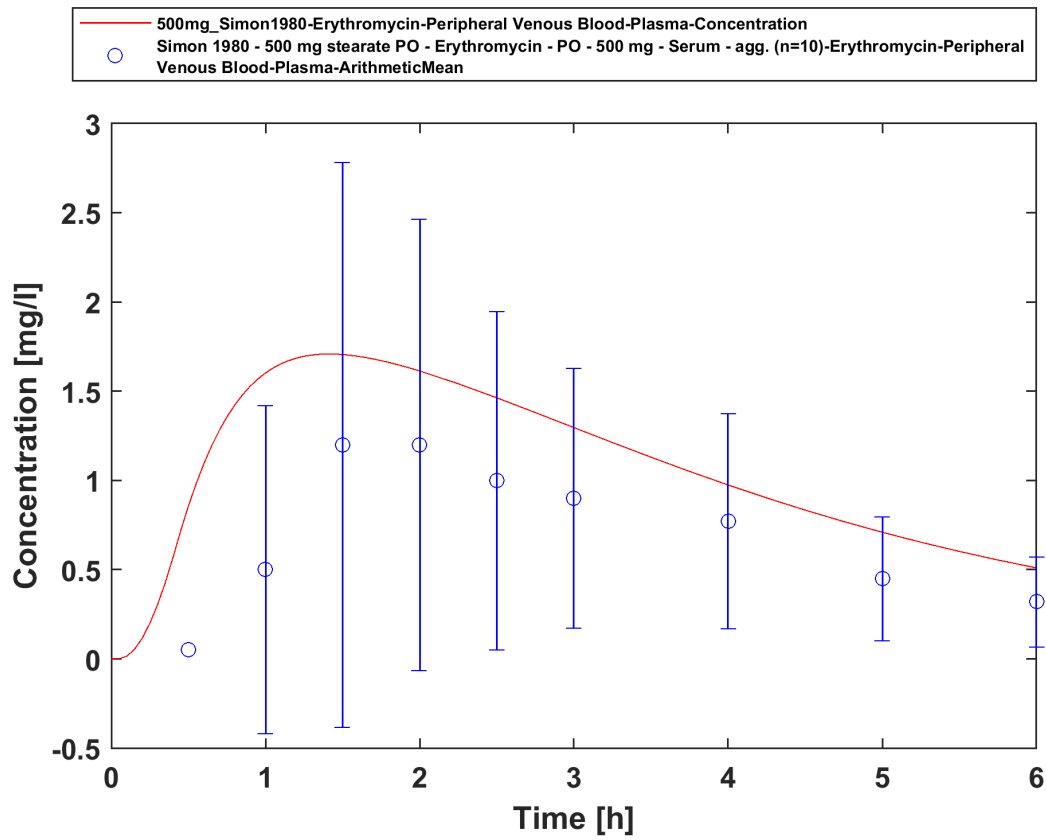


Time Profile Analysis 1









Time Profile Analysis 1

## 4 Conclusion

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The final erythromycin PBPK model applies metabolism by CYP3A4, glomerular filtration, and a dummy clearance technically implemented as hepatic plasma clearance accounting for additional clearance pathways, as well as mechanism-based inhibition of CYP3A4. Overall, the model adequately describes the oral pharmacokinetics of erythromycin in healthy adults receiving different single or multiple doses of several oral dosage forms and is considered verified for the use as a perpetrator drug in drug-drug interaction simulations.

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