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Considering Age Variation when Coining Drugs as High vs Low Hepatic Extraction Ratio

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Running Title Page

Running Title: Consequences of Age Variation in Hepatic Extraction Ratios

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Abbreviations

AAG, alpha-acid glycoprotein; B:P, Blood to plasma ratio; CLuint,H, Hepatic intrinsic clearance

of unbound drug; CL_{H,B}, Hepatic metabolic clearance; E_H, Hepatic extraction ratio; fu, Fraction

of drug in plasma unbound; fuB, Unbound drug in blood; MPPGL, Microsomal protein per gram

of liver; PopPK, Population pharmacokinetics; QH, Hepatic blood flow.

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Abstract

Hepatic extraction ratio (E_H) is commonly considered as 'an inherent attribute' of drug. It determines the main physiological and biological elements of the system (patient attributes) which are most significant in inter-individual variability of clearance. E_H consists of three agedependent parameters: fraction of unbound drug in blood (fu_B), hepatic intrinsic clearance of unbound drug (CLu_{int.H}) and hepatic blood flow (Q_H). When age-effects on these elements are not proportional a given drug may shift from so called "high extraction" status to "low extraction". To demonstrate the impact of age-related changes on fub, CLuint, and QH, EH of midazolam and two hypothetical drugs with 10-fold higher and 10-fold lower CLuint H than midazolam were investigated in paediatrics based on known ontogeny functions. EH was simulated using Simcyp V14. This was then complemented by a comprehensive literature survey to identify commonly applied covariates in paediatric population pharmacokinetic (PopPK) studies. Midazolam E_H decreased from 0.6 in adults to 0.02 at birth, making its clearance much more susceptible to changes in CLuint,H and fuB than adults and reducing impact of Q_H on clearance. The drug with 10-fold higher CLu_{int.H} was categorised as high extraction from 4 days old onwards, whilst the drug with 10-fold lower CLuint.H remained low extraction from birth to adulthood. Approximately 50% of collected PopPK studies (n=120) did not consider interaction between age and other covariates. Interaction between covariates and age should be considered as part of studies involving young paediatric patients. E_H cannot be considered as an inherent drug property without considering the effect of age.

Introduction

Hepatic metabolic clearance ($CL_{H,B}$) of intravenously administered drugs is determined by hepatic blood flow (Q_H) and their hepatic extraction ratio (E_H) according to the Equation 1:

$$CL_{HB} = E_H \times Q_H$$
 Equation 1

 E_H is calculated from fraction of drug unbound in blood (fu_B), hepatic intrinsic clearance of unbound drug ($CLu_{int,H}$) and Q_H according to Equation 2:

$$E_{_{H}} = \frac{fu_{_{B}} \times CLu_{_{int,H}}}{Q_{_{H}} + (fu_{_{B}} \times CLu_{_{int,H}})}$$
 Equation 2

Extraction ratio of the drug is generally classified as high (>0.7), intermediate (0.3-0.7) or low (<0.3) according to the fraction of drug removed during one pass through the liver. Commonly E_H of a drug is considered as an inherent attribute of the drug and presented with a fixed value. However, this classification does not consider that the parameters in Equation 2 are age-dependent and changes in these parameters will affect E_H. For example, a rise in fu_B, for low E_H drugs increases hepatic metabolic clearance, whereas for high extraction drugs this does not affect metabolic clearance. Unless the age-related physiological changes in fu_B, CLu_{int,H} and Q_H occur in parallel, it is expected that E_H of drugs varies with age. Therefore, a high extraction drug in adults will not necessarily remain a high extraction drug in neonates.

Age-varying E_H can potentially be used as a covariate in clearance models when analysing population pharmacokinetic (PopPK) studies. However, since applying extraction ratio directly in the model might not be straightforward this concept is considered in PopPK models through the interaction between covariate terms in the model. For example, age and body weight are commonly used as covariates in PopPK clearance models where body weight is also affected by age. The interaction between these two covariates should be considered in the model.

The primary aim of this study is to investigate relative differences in E_H with age using *in vivo* midazolam data and two hypothetical high and low extraction drugs through modelling and

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simulation techniques. We also use the concept of the age varying E_H to examine whether the interaction between covariate terms in modelling clearance have been considered in the PopPK studies.

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Materials and Methods

Literature Data collection

Data on midazolam systemic clearance in paediatrics from birth to 17 years were collected

from the literature. Literature search strategy and methodology for deconvolution of clearance

to arrive at CL_{H,B} from midazolam systemic clearance (using blood to plasma ratio (B:P)) and

Q_H based on cardiac output were explained previously (Salem et al., 2014).

Simulations

A drug with 10-fold higher and 10-fold lower CLuint,H than midazolam was designed by

multiplying and dividing the deconvoluted midazolam CLuint,H by 10 as proposed by (Salem et

al., 2014) to mimic a high and low extraction drug, respectively. Then, using the relevant

CLu_{int,H}, Q_H and fu_B in Equation 2, E_H was calculated.

A number of simulations in Simcyp Paediatric v14 were carried out for midazolam, a drug with

10-fold higher and 10-fold lower CLu_{int,H} than midazolam to show age related changes in the

magnitude of E_H. One hundred subjects were simulated consisting of equal proportion of males

and females and combination of age bands (1 day, 1 month, 2 years and 12 years as well as

adult). E_H was calculated using Equation 3 from the output data. Mean values of E_H at each

age band were plotted against age for each of the simulated drugs.

Calculation of hepatic extraction ratio

Hepatic extraction ratio was calculated from CL_H and Q_H for midazolam and the other two

hypothetical drugs assuming well-stirred model using Equation 3;

Hepatic Extraction Ratio = $\frac{CL_{H,B}}{O_{H,B}}$

Equation 3

Sensitivity analysis

Sensitivity analysis was carried out with a view to identify which component of the extraction

ratio (CLu_{int,H}, Q_H and fu_B) plays the most dominant role in variation of E_H from those of adult

values at any given age. The impact of age dependent Q_H was evaluated by fixing fu_B and $CLu_{int,H}$ (L/h/g of liver) to the adult values for all age ranges. This involved assumptions on lack of any ontogeny for abundance of the enzymes (pmol per mg of microsomal protein) and no age related changes in the level of microsomal protein per gram of liver (MPPGL). The values of $CLu_{int,H}$ (L/h/g of liver) were used to calculated the paediatric $CLu_{int,H}$ values per whole liver by applying age-related liver weight. E_H was plotted against age and patterns were compared. In another set, only fu_B values were fixed to adult values to demonstrate the sensitivity of E_H to age related changes in Q_H and $CLu_{int,H}$ (L/h) without impact of age-related changes in binding. E_H was calculated and plotted against age and compared with the original set of results (where all age-related parameters had been considered).

In order to separate the size related effects (i.e. liver mass and hepatic blood flow) from ontogeny-related factors on E_H, a graphical representation was devised to demonstrate the paediatric values of enzyme abundance relative to adults at given age (in this case CYP3A4) alongside relative values for liver volume, hepatic blood flow and MPPGL.

Population Pharmacokinetic studies (PopPK)

A comprehensive literature survey using Pubmed was carried out to identify commonly used covariates in paediatric PopPK studies for drugs after intravenous administration. No year or journal or language restriction applied to the search process. Collated publications were carefully checked for modelling covariates and the form of the covariates-clearance relationship in the reported model. Studies that considered the interaction between covariates and clearance were identified. Interaction between covariate terms was also considered if the presence of a covariate modifies the impact of another covariate in a multiplicative or exponential way. Where there are different clearance models for different paediatric age ranges, interaction with age is also considered. Interaction between covariate terms was not considered if only one covariate was considered in the final clearance model or if the covariates are in linear additive relationship to the clearance. Corresponding authors were contacted where modelling section was not clear.

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Results

Midazolam hepatic extraction ratio

Hepatic extraction ratio of midazolam, after deconvolution of clinical systemic clearance, increased with age. Figure 1 illustrates that midazolam is a low extraction drug until about the age of 10 months. However, in some individuals it remained low at the age of 9 years.

Figure 2 shows E_H increases with age for midazolam and two other hypothetical compounds. The degree of change in E_H with age depends on magnitude of CLu_{int,H} against the given enzyme. As shown in the figure, 10-fold reduction in CLu_{int,H} results in a drug with low hepatic extraction across the paediatric and adult age range whereas 10-fold increase in CLu_{int,H} shifts the drug from so called intermediate to high extraction status.

Sensitivity analysis

Figure 2 compares E_H when all age related components (CLu_{int,H}, Q_H and fu_B) considered (solid lines) with a scenario involving no age-related changes in fu_B (dashed lines). As shown in Figure 2 E_H is marginally lower in younger groups if age related fu_B is not considered. However, this might be different for drugs with higher protein binding.

When CLu_{int,H} (L/h/g of liver) and fu_B are fixed to adult values, the changes in E_H will be driven by age-related changes in Q_H and liver weight (Supplemental Figure 1). In this scenario, there are no significant differences between E_H values across paediatric age groups for three drugs since the low activity of CYP3A4 in younger age is not considered.

The rate of change with age for liver volume, hepatic blood flow and MPPGL as a fraction of adult values are shown in Figure 3. This figure shows the changes in underlying parameters of E_H. The changes in blood flow and liver volume relative to adults occurs almost in parallel to each other. Therefore, the discrepancy in Q_H and liver size alone cannot account for the observed differences in E_H instead changes in intrinsic activity to the level of enzyme abundance and to a lesser extent MPPGL are determinants of age-varying E_H. **Error! Reference source not found.** summarises the contributing parameters to E_H that are reported

in Figure 3. Needless to say, if relative values to adult for all these elements had a similar rate of change with age, no age-related differences would have been anticipated in E_H.

Analysis of covariates in population pharmacokinetic (PopPK) studies

A total of 120 PopPK studies were retrieved in paediatric age range (birth to 18 years) for intravenously administered drugs. The interaction between covariate terms was not considered in 50% of the studies (n=60). Table S1 in supplements summarises the most commonly used covariates in the analysed PopPK studies (Supplemental Table 1). Table S2 in supplements shows all the analysed PopPK studies with interaction between covariates (Supplemental Table 2).

Discussion and Conclusions

E_H of the drugs in this study increases with age due to rapid physiological changes in parameters determining E_H after birth including the ontogeny of enzyme abundance and to a lesser extent MPPGL. Although not relevant to the cases represented in this study, ontogeny of plasma proteins can play a significant role in age-related changes of E_H for highly bound, low extraction drugs. As shown previously by several authors, concentration of plasma proteins increases with age whereas unbound fraction of drugs in plasma and therefore in blood decreases with age (Sethi et al., 2015, Johnson et al., 2006 and McNamara and Alcorn, 2002). The ontogeny of plasma protein binding and enzyme abundance on any given compound depends on the extent of binding to particular protein and the importance of that enzymatic pathway to the overall elimination of drug. As a consequence, a drug that is coined a high, low or intermediate hepatic extraction compound in adults is not necessarily going to carry the same extraction category in paediatrics. For the particular case studies in this report, where the binding was not a major factor, and CYP3A4 was the main metabolising enzyme, we demonstrated the switch from high extraction or intermediate extraction to low extraction in neonates and younger children. However, the results from this study should be generalised to other drugs metabolised by other pathways with caution. CLuint. H value is an interplay between the enzyme abundance and kinetic parameters (V_{max} and K_m). The difference in enzyme abundance depending on the pathway and age can be masked or stressed by enzyme kinetic parameters and hence resulting in similar or different CLuint, H and EH values from what has been shown in this study. Changes in EH is not confined to age-varying parameters. Induction or inhibition of drug metabolising enzymes for a capacity limited drug and inhibition of drug metabolising enzyme for a flow limited drug can also change the extraction ratio of drugs. In addition changes to Q_H and fu_B due to hemodynamic changes occurring in clinical conditions and progression of disease may affect the extraction ratio of drugs. This consideration can be more important in preterm neonates due to prematurity of metabolic pathways, special populations such as elderly and pregnancy that can affect free fraction of the drug (f_u), enzyme activity and/or Q_H which ultimately can alter E_H .

Since the determinants of CL (covariates) change with age, it is not right to assume nointeraction between age and covariates. In some of the analysed PopPK studies here, the
interaction between covariate terms were not identified by their authors. The reason for lack
of such interactions can be the wide age range in some of these studies with limited number
of subjects at lower end of spectrum. In addition, several investigations only examined older
subsets of children, where the ontogeny of enzymes responsible for metabolism are likely to
be fully mature. These investigations are not likely to find that the addition of age into their
clearance models provide a better fit. Another reason for lack of interaction origins from
unbalanced blood sampling in the early life after birth compared with older children. Some
pharmacokinetic studies retrospectively analysed the available samples of drug concentration
in blood or plasma where the relevant covariates are not always available. Also some relevant
covariates such as f_u may not have been measured in new-borns. Only one PopPK study on
morphine concluded an independent clearance model is required for new-borns (Knibbe et al.,
2009).

The results in this study suggest that CLu_{int,H} is the most important parameter that affects E_H of drugs. Due to rapid physiological changes after birth and especially in neonatal period E_H of drugs can be significantly affected by changes in CLu_{int,H}.

Hepatic extraction also contributes to determination of oral bioavailability of drugs. Currently, there are contradicting evidence as to bioavailability of drugs is different between paediatrics and adults (Harper et al., 1988; Stratchunsky et al., 1991; Pinkerton et al., 1993; Hassan et al., 1994; Fujiwara et al., 1996; Anderson et al., 2002; Crill et al., 2006; Zane and Thakker, 2014). While in clinical practice bioavailability of drugs is assumed to be similar between paediatrics and adults, the current study supports bioavailability also can be an age-dependant parameter and reduce with age since E_H changes with age. Assuming a higher hepatic extraction and therefore lower bioavailability in neonates, oral clearance can be overestimated in this population and unnecessarily higher doses given to neonates. However, the clinical significance of this under and overestimations is not clear and requires further investigation.

In conclusion, a high extraction drug in adults is not necessarily a high extraction drug in paediatrics. Unless the age-related changes in factors determining E_H occur at the same rate, extraction ratio will be different between paediatrics and adults. More attention should be given to interaction terms of covariate during analysis of such data as the impact of certain physiological covariates might change with age. Further clarification of underlying mechanisms for metabolism (and bioavailability) of drugs should heavily rely on modelling and simulation techniques.

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Participated in research design: Salem, Abduljalil, Kamiyama, Rostami-Hodjegan

Performed data analysis: Salem, Abduljalil, Kamiyama

Wrote or contributed to the writing of the manuscript: Salem, Abduljalil

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Figure Legends

Figure 1 Hepatic extraction for intravenous midazolam calculated from reports of clinical studies in the literature using ontogeny functions in paediatric subjects and healthy adult volunteers (n=523).

Figure 2 Simulated hepatic extraction in Simcyp v14 shows changes with age for midazolam, a drug with 10-fold higher CLu_{int,H} and a drug with 10-fold lower CLu_{int,H}. A high or intermediate extraction drug in adults is not necessarily a high or intermediate exteraction drug in paediatric subjects. Dashed lines are the same E_H values with age when fu_B is remained unchanged (fu_B=0.05). Dotted lines show the limits for high (>0.7) and low (0.3>) extraction.

Figure 3 Age-related variations in parameters defining E_H shown as relative values to corresponding adult level of each parameter. Part (A) indicates the changes in liver size (Johnson et al., 2005), hepatic blood flow (Guyton., 1991) and MPPGL (Barter et al., 2008) which applies to all drugs. Part (B) demonstrates the relative values of serum albumin (Johnson et al., 2003, Johnson et al., 2006 and Sethi et al., 2015), CYP3A4 abundance (of relevance to current study) (Salem et al., 2014) alongside age-variation in serum alpha-acid glycoprotein (AAG) (Johnson et al., 2003 and Johnson et al., 2006) and CYP1A2 abundance (Salem et al., 2014). The impact of the parameters shown in Part (B) will depend on the relative importance of the protein binding to each protein and the role of the specific enzyme to overall elimination.

Tables

Table 1. Examples of age-related parameters defining E_{H} and prior knowledge on their age-dependency

Parameter	Definition	Age-dependency Model
Qн	Hepatic blood flow as a function of cardiac index	Guyton, 1991
MPPGL	mg of microsomal protein per gram of liver	Barter et al., 2008
CYP3A4 and CYP1A"	pmol of enzyme abundance	Salem et al., 2014
Liver volume	ml of liver	Johnson et al., 2005
Albumin and AAG	Plasma proteins concentration (g/L)	Johnson et al., 2006 and Sethi et al., 2015

Figure 1

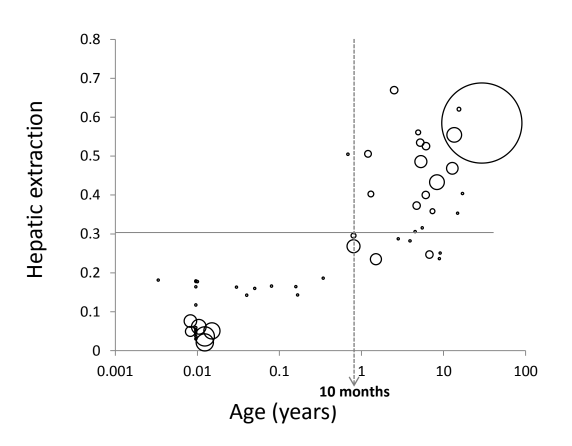


Figure 2

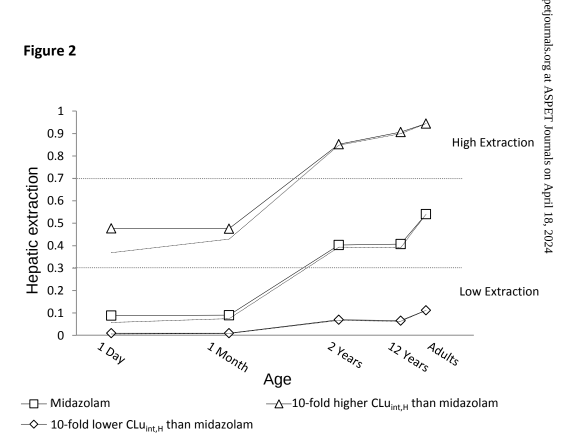
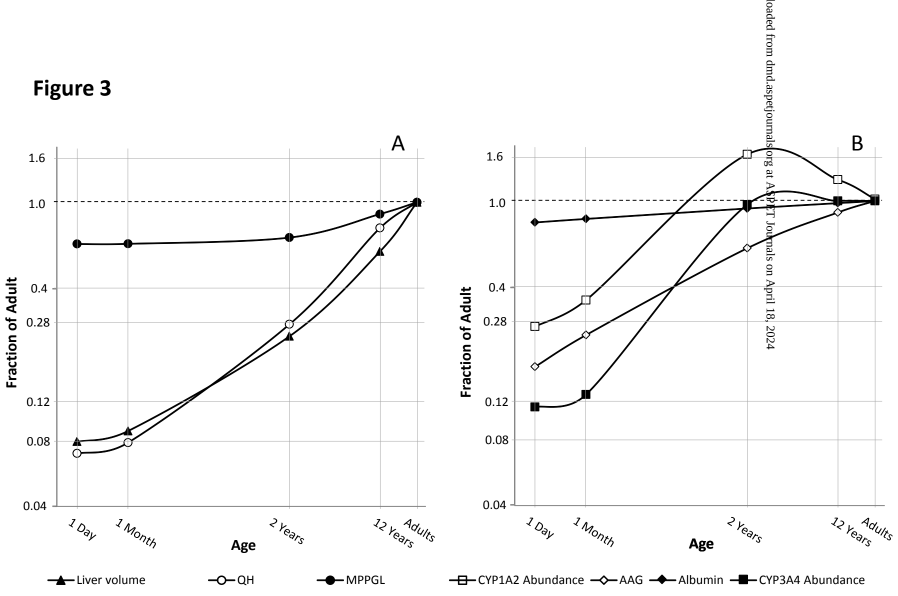


Figure 3



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Supplementary Material

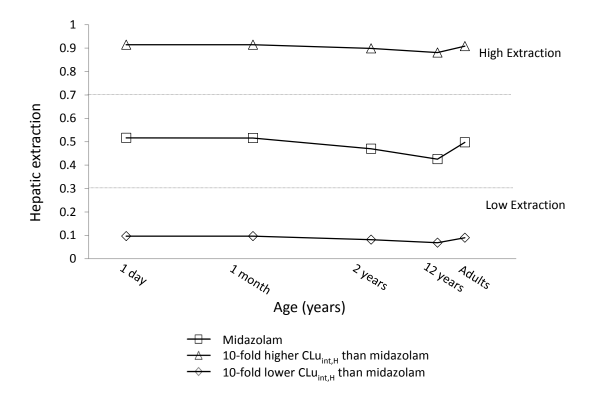


Figure S1 E_H changes with age for midazolam, a drug with 10-fold higher and lower CLu_{int}, are minimal when the enzyme abundance (pmol per MPPGL), MPPGL and fu_B are fixed to the adult values. The slight variation around 12 years relates to higher cardiac index (cardiac output to body surface area). Cardiac index values for 1 day, 1 month, 2 years, 12 years and 25 years are 2.33, 2.35, 2.50, 2.67, 3.80 and 2.5 L/min/m² (Johnson et al., 2006; Guyton., 1991).

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Table S1 Summary of covariates considered in analysing PopPK studies. However, not all of these covariates resulted in significant reduction of objective function and were included in the final model.

Covariate	Category	
Post-menstrual age		
Postnatal age	Age	
Post-conception age	, 190	
Starting age on breast milk		
Day of treatment		
Period of intubation before study entry	Time	
Post-transplant days		
Weight (birth weight or current weight)		
Weight (allometric scaling, linear and exponential model)		
Body mass index		
Body surface area	Size	
Fat free mass (Allometric scaling)		
Body length		
Head circumference		
Male and Female	Sex	
Race	Ethnicity	
Cratinine (Serum or clearance)		
Billirubin (unconjugated)		
Aminotransferase	Biomarker	
Alkaline phosphatase		
Globulin concentration		

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Platelet counts	
Glomerular filtration rate	-
Oxygen concentration	
Nephrectomy	
Systemic inflammatory response syndrome	
Critical illness	
Hepatic dysfunction	_
Health status	_
Apgar score at 1 minute and 5 minutes	Clinical Condition
Birth asphyxia	
Feed intolerance	-
Presence of respiratory distress syndrome	-
Patent ductus arteriosus	-
Respiratory support and form of support	_
Infection	-
Co-medication (phenobarbital, parental nutrition, diuretic,	
enzyme inducer, dexamethasone, nephrotoxic drug)	
Carboplatin therapy	Therapy
Surfactant therapy	
CYP2C19 inhibitors	
Route of delivery	
Prenatal smoking history,	_
Current multiple pregnancy	_ Maternal
Antepartum haemorrhage	Ivialemai
Chorioamnionitis	-
Preeclampsia	-
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Premature rupture of the membranes	
Genotypes (CYP2C19, 2C9, CYP2D6, OCT1)	Genotype
Study centre	Other
Inter-occasion variability	

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Table S2 Summary of the analyses PopPK studies with significant covariates. Where interaction between covariates was considered, the relevant covariates are identified.

ID	Compound	Population	Covariates	Covariate Interaction	Reference
1		PN=1-144 days	Weight, PMA, unconjugated bilirubin	Weight & PMA & unconjugated bilirubin	(Palmer et al., 2008)
2		27-42 weeks PCA	PCA, Weight	PCA & Weight	(Allegaert et al., 2004)
3		neonates (27-45 weeks PMA)	Weight, PCA	PCA & Weight	(Allegaert et al., 2011)
4		premature - 14 years	Weight (allometric),PCA	PCA & Weight	(Anderson et al., 2005)
5	Acataminanhan	6 months-2 years	Weight	No	(Prins et al., 2008)
6	Acetaminophen	Neonates to adolescents	PNA, Weight (allometric)	PNA and Weight	(Zuppa et al., 2011)
7		neonates	Weight, PMA	description of covariates interaction in the final model was not clear	(van Ganzewinkel et al., 2014)
8		1 day -34 years	Weight -dependent exponent model	No	(Wang et al., 2014)
9		4 days to 3 months.	age, Weight	age & Weight	(Anderson et al., 2000)

10		1 day to 6 months.	Weight & PCA	Weight & PCA	(Anderson et al., 2002)
11		< 18 years children	Weight	No	(Mohammed et al., 2012)
12	Actinomycin-D	children (1.6 to 20.3 years)	Weight	No	(Mondick et al., 2008)
13		< 21 years	Weight	No	(Hill et al., 2014)
14		preterm and term infants	Weight, PMA	Weight & PMA	(Sampson et al., 2014)
15	A a calaccia	0.8–19.9 children	Weight, CrCL	Weight & CrCL	(Zeng et al., 2009)
16	Acyclovir	0.25-17yr children	Weight, BSA, GFR, PCA,	Weight & BSA & GFR & PCA & Scr	(Tod et al., 2001)
17		gestational ages from 24 to 41 weeks	Weight, PMA	Weight & PMA	(Sherwin et al., 2009b)
18	Amikacin	gestational age 24–43 weeks; postnatal age 1– 30 days	Weight -dependent exponent & PNA	Weight & PNA	(De Cock et al., 2012)
19	Aminophylline	PNA=0-26 weeks	Weight, PNA	No	(Moore et al., 1989)
20	Bevacizumab	children	ВМІ	Weight & BMI	(Turner et al., 2014)

21	•	children	Woight	No	(Diestelhorst et al.,
21		Children	Weight	INO	2014)
22		0.45-16.7 years	Weight	No	(Nguyen et al., 2004)
23		10 days to 15 years	Weight (allometric)	No	(Paci et al., 2012)
24	Busulfan	94 children (0.4 - 18.8 years)	BSA, Weight	No	(Trame et al., 2011)
25		0.7-13.1 years children	Allometric Weight	No	(Veal et al., 2012)
26		0.2 to 23 years	Weight	No	(Zwaveling et al., 2008)
27		age 0.1-3.3 years old	Allometric Weight, Age	Weight and age	(Savic et al., 2013)
28		0.1 to 26 years	age, WT, BSA	No	(Bartelink et al., 2012)
29		premature neonates 27.6 weeks GSA	Weight, PNA	Weight & PNA	(Charles et al., 2008)
30	Caffeine	premature neonates	Weight, LGA, PNA and parental nutrition	GA & Weight &PNA	(Falcao et al., 1997)
31		premature neonates <32 GW	Weight, PNA	No	(Lee et al., 1997)

	_	premature neonates < 34			
32		GW	Weight	No	(Lee et al., 2002)
33		2 days-0.85 years	Weight, PMA	Weight & PMA	(Suyagh et al., 2013)
34	Canrenoate	2 days to 10 years children	Weight	No	(Suyagh et al., 2012)
35	carboplatin	2 m to 18 years	Weight, SeCr, nephrectomy	Weight & SeCr & nephrectomy	(Chatelut et al., 1996)
36	cefepime	Premature & infants <4 mon.	Weight, SeCr	Weight & SeCr	(Capparelli et al., 2005)
37	cefozopran	children	Weight	No	(Ikawa et al., 2009)
38	ceftazidime	10–45 years	Allometric scaling by FFM	No	(Bulitta et al., 2010)
39	ciprofloxacin	0.27-16.9 years	Weight, Age	Weight & Age	(Rajagopalan and Gastonguay, 2003)
40		6 - 16 years	Weight	No	(Schaefer et al., 1996)
41	Clindamycin	Premature Infants to Adolescents	Weight, PMA	Weight & PMA	(Gonzalez et al., 2014)
42	Clonidine	0-15 years	Weight	need full paper	(Potts et al., 2007)

	age range 1.30 – 9.37	GFR, gender	No	(McCune et al., 2009)
е	years	Or it, gondor		(Modano et al., 2000)
	children	BSA, posttransplant Days	BSA & posttransplant Days	(Zhou et al., 2013)
	0.26 47.5 40.00	Weight, Cholesterol, Hct,	Weight & Cholesterol & Hct	(Forto et al. 2007)
	0.36 - 17.5 years	Scr	& Scr	(Fanta et al., 2007)
cyclosporine		Weight, age, co-med	description of covariates	(0.1
	0.9 -20 years	(itraconazole &	interaction in the final model	(Schrauder et al.,
		tobramycine)	was not clear	2009)
cytosine	50 poodiatria patianta	ACE DOA	AOE 9 DCA	(Periclou and Avramis,
arabinoside	52 paediatric patients	AGE, BSA	AGE & BSA	1996)
daunorubicin	children	Weight	No	(Hempel et al., 2003)
dexmedetomidin	ah il duan	Majaht atudu santra	No	(Datron et al. 2000)
е	children	weight, study centre	NO	(Petroz et al., 2006)
Diclofenac	aged 1-14 years	Weight	No	(Standing et al., 2011)
digoxin	0.33 to 15 years children	SeCr	No	(el-Desoky et al., 2005)
	children older than 3	BSA, inter-occasion	N	(1/2) 1 1 00 (0)
doxorubicin	years	varaibility	No	(Kontny et al., 2013)
	cyclosporine cytosine arabinoside daunorubicin dexmedetomidin e Diclofenac	children 0.36 - 17.5 years cyclosporine 0.9 -20 years 52 paediatric patients arabinoside daunorubicin children dexmedetomidin e Diclofenac aged 1-14 years digoxin 0.33 to 15 years children children older than 3	cyclosporine Cyclosporine Cyclosporine Cyclosporine Cyclosporine Cytosine arabinoside daunorubicin Children Cytosine arabinoside Cytosine AGE, BSA AGE, BSA Weight Weight, study centre Weight, study centre Children Cytosine AGE, BSA AGE, BSA AGE, BSA BSA, inter-occasion Cytosine AGE, BSA AGE, BSA BSA, inter-occasion Children older than 3 BSA, inter-occasion	cyclosporine cytosine arabinoside cytosine arabinoside cyclosporine cytosine arabinoside cyclosporine cytosine arabinoside cyclosporine cytosine arabinoside cyclosporine cytosine arabinoside tobramycine) was not clear No dexmedetomidin No dexmedetomidin e children children Weight, study centre No Children SeCr No No children older than 3 BSA, inter-occasion No

	-			description of covariates	(Canadatua na at al
53	esomeprazole	0-17 years	Weight, age	interaction in the final model	(Sandstrom et al.,
				was not clear	2012)
54		0.53-13.21 years	Allometric WT, Age	Weight & Age	(Lin et al., 2012)
55	Etomidate	14 weeks to 16.7 years	Weight (alometry)	No	(Urien et al., 2011)
			Weight, postnatal age,		
56	fluconazole	PNA=0.14 to 12.6 weeks	gestational age at birth,	Weight & GA & PNA & Scr	(Wade et al., 2008)
			SeCr		
		3 mon 13 years			(Kumpulainen et al.,
57	flurbiprofen	children	Weight	No	2010)
58	Ganciclovir	PNA= 8-34 days	WT	No	(Acosta et al., 2007)
50		14 - 81 years cancer	CoinI	NI.	(Danasia at al. 4000)
59		patient	creatinine clearance	No	(Rosario et al., 1998)
	Gentamicin		WT, PNA	WT & PNA	(De Cock et al., 2014)
60		neonates	WT , PNA	WT & PNA	(Sherwin et al., 2009a)

	-	premature & term	WT, PNA, gestational		
61		neonates(PMA=24.2- 42.4wks)	age	WT, PNA & GA	(Fuchs et al., 2014)
62	hydrocortisone	critically ill neonates	WT, PMA	WT & PMA	(Vezina et al., 2014)
63	indomethacin	premature neonates	WT	No	(Al Za'abi et al., 2007)
64	indometriacin	PNA=1-77 days	WT, PNA	WT & APNA	(Smyth et al., 2004)
65	Irinotecan	children	WT, Age and bilirubin	WT & Age & bilirubin	(Thompson et al., 2008)
66	Ketamin	1.5-14 years children (and adult)	WT	No	(Herd et al., 2007)
67	Lamivudine	2 days to 18 years	WT (alometry), PMA (maturation)	WT & PMA	(Bouazza et al., 2011)
68	Levofloxacin	paediatric & adult	WT, Age	WT & Age	(Li et al., 2010)
69	levetiracetam	PNA=1 - 5 days	WT, PNA	WT & PNA	(Sharpe et al., 2012)
70	lorazepam	3 mon - <18 years	WT	No	(Chamberlain et al., 2012)

	victue on sin & Bisp		WT, GFR, carboplatin		
71	Melphalan	children	therapy	No	(Nath et al., 2007)
72	Meropenem	GA=29 - 42 weeks	CrCL, WT	No	(van den Anker et al., 2009)
73	Methotrexate	children	WT, Age	No	(Colom et al., 2009)
74	Metronidazole	32 pre-term	WT, PMA	WT & PMA	(Suyagh et al., 2011)
75	Micafungin	children and adult	WT, platelet counts	No	(Tabata et al., 2006)
76		paediatric intensive care unit	WT	No	(Bienert et al., 2013)
77		neonates	WT, GA	WT & GA	(Burtin et al., 1994)
78	Midazolam	1 month to 17 years children	critical illness, WT	critical illness & WT	(Ince et al., 2012)
79		Premature Infants to Adolescents	WT, WT-dependent exponent of allometric exponent function	No	(Ince et al., 2013)
80		6 months to < 16 years	Age	No	(Reed et al., 2001)

04	•	2 month to 17 years	WT-dependent exponent	NI-	(Ince et al., 2012)
81		children	function	No	
			WT (albumin was added,		
		paediatric intensive care	but removed from the		(5)
82		unit	final model not	No	(Bienert et al., 2013)
			significant)		
00	Montolyleast	C. OC vecre	WIT	Ne	(Ramakrishnan et al.,
83	Montelukast	6 - 86 years	WT	No	2005)
84		children	WT, OCT1 genotypes	WT & OCT1 genotypes	(Fukuda et al., 2013)
85	Morphine	preterm - < 3 years	WT, PNA	PNA (stratified)	(Knibbe et al., 2009)
86	Mycophenolic	0 - 3 year children	WT, PNA, creatinine,	WT & PNA WT & ciclosporin	(Bouwmeester et al.,
00		0 - 3 year cillidren	bilirubin,		2004)
87		children and young	WT, ciclosporin		(Zeng et al., 2010)
07		people	wi, didiosponin		(Zeng et al., 2010)
88	Myo-inositol	Infants born in 23-	WT (allometric), CrCL,	No	(Phelps et al., 2013)
00		29 weeks of gestation	gestational age		(1 Helps et al., 2013)
89	Netilmicin	2 to 28 days	cWT, PMA	cWT & PMA	(Sherwin et al., 2008)
					· ·

90	Omeprazole	critically ill children	WT (allometry)	No	(Solana et al., 2014)
91	Ondansetron	2–38 years	Wt, Age, Sex,	sex & age + wt	(de Alwis et al., 1998)
92	Palivizumab	Preterm infants to adult	age,WT, lung disease, race, antidrug antibody titer	WT, Age, race, lung disease and antidrug antibody titer	(Robbie et al., 2012)
93		0.025-16 years	WT, GA, Age, race, gender, 2C9 polymorphisms	WT, GA, Age, race, gender, 2C9 polymorphisms	(Knebel et al., 2011)
94	Pantoprazole	10 days to 16.4 years	WT, Age, hepatic dysfunction, CYP2C19 inhibitors, systemic inflammatory response syndrome	WT, Age, DDI , SIRS, Hepatic dysfunction	(Pettersen et al., 2009)
95	Phenobarbital	PNA= 0.0-206 days	WT (allometric)	No	(Marsot et al., 2014)
96	Phenytoin	neonates and infants	WT, PNA	WT & PNA	(Al Za'abi et al., 2006)
97		include children	WT	No	(Tanaka et al., 2013)

	victabolishi & Dispo	obese children and	NA/T	NI.	(Diepstraten et al.,
98		adolescents	WT	No	2012)
99		0.25-88 years	WT, age, sex, health status	WT & Age	(Eleveld et al., 2014)
100		3 - 11 years	WT	No	(Kataria et al., 1994)
101	Propofol	children	WT	No	(Knibbe et al., 2002)
102		pre-term to adults	WT	No	(Wang et al., 2012)
103		from 2.0 to 88 year	WT	No	(Schuttler and Ihmsen, 2000)
101		4 week to 40 year	WT Conding ourses	No (or yes WT & Cardiac	(Rigby-Jones et al.,
104		1 week to 12 year	WT, Cardiac surgery	Surgery)	2002)
105	Ranitidine	critically ill children (age: 0.042 - 15.51 years)	WT (allometry)	WT & Cardiac failure/surgery	(Hawwa et al., 2013)
106	Sildenafil	new-born (11-71 hr)	PNA	No	(Mukherjee et al., 2009)
107	Toonsline	1-14 years children	Age	No	(Sam et al., 2000)
108	Tacrolimus	children	WT, serum creatinine;	WT & SeCr	(Wallin et al., 2009)

109	Theophylline	premature neonates	WT, PNA	No	(Lee et al., 1996)
110		children	WT	No	(Hennig et al., 2008)
111	Tobramycin	0.1-18.8 years	WT	No	(Sherwin et al., 2014)
			WT, PNA	WT & PNA	(De Cock et al., 2014)
112	Tramadol	children (0-3 months)	WT, PCA	WT & PCA	(Allegaert et al., 2005)
	Trainage.	and adult (23-57 year)	, -	5 3	(r megaert et am, 2000)
113	Tranexamic acid	1.0 - 12 years	WT (alometry)	No	(Grassin-Delyle et al.,
113	Trancamic acid	1.0 - 12 years	w r (alometry)	NO	2013)
114	Valproic acid	1 to 17 years	Weight	No	(Williams et al., 2012)
115		neonates	current Weight, birth	current weight, birth weight,	(Zhao et al., 2013)
	Vancomycin	riconates	Weight, SeCr, PNA	SeCr & PNA	(Znao ot al., 2010)
116			Weight, PNA	Weight & PNA	(De Cock et al., 2014)
117		2 to <12 years	Weight, age, CYP2C19	Weight, age, 2C19	(Karlsson et al., 2009)
117		Z to <12 years	phenotypes, ALT	phenotype, ALT	(Nalissoli et al., 2009)
118	Voriconazole	2 years to adult	Weight, age, CYP2C19	Weight, age, CYP2C19	(Friborg et al. 2012)
110		2 years to adult	phenotypes	phenotypes	(Friberg et al., 2012)
119		2 to 11 years	Weight, 2C19 phenotype	Weight & 2C19 phenotype	(Walsh et al., 2004)

Considering Age Variation when Coining Drugs as High vs Low Hepatic Extraction Ratio. Farzaneh Salem, Khaled Abduljalil, Yoshi Kamiyama and Amin Rostami Hodjegan

Drug Metabolism & Disposition

Diag Wettoonsin & Disposition				
120	2–11 years children	Weight, 2C19 phenotypes, aminotransferase, alkaline phosphatase	Weight & 2C9 phenotype, alanine amino transferase, alkaline phosphatase	(Wahlby et al., 2004)

SeCr: serum creatinine

Farzaneh Salem, Khaled Abduljalil, Yoshi Kamiyama and Amin Rostami Hodjegan Drug Metabolism & Disposition

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