

1	Title:	A repository describing an aging population to inform physiologically based
2		pharmacokinetic models considering anatomical, physiological and biological age-
3		dependent changes.
4		
5	Authors:	Felix Stader <sup>1,2,3,\$</sup> , Marco Siccardi <sup>4</sup> , Manuel Battegay <sup>1,3</sup> , Hannah Kinvig <sup>4</sup> , Melissa A.
6		Penny <sup>2,3</sup> & Catia Marzolini <sup>1,3</sup>
7		
8	Affiliation:	1. Division of Infectious Diseases and Hospital Epidemiology
9		Departments of Medicine and Clinical Research
10		University Hospital Basel
11		Basel, Switzerland
12		
13		2. Infectious Disease Modelling Unit
14		Department of Epidemiology and Public Health
15		Swiss Tropical and Public Health Institute
16		Basel, Switzerland
17		
18		3. University of Basel
19		Basel, Switzerland
20		
21		4. Department of Molecular and Clinical Pharmacology
22		Institute of Translational Medicine
23		University of Liverpool
24		Liverpool, UK
25		
26		\$ corresponding author
27		E-mail: Felix.Stader@unibas.ch
28		Phone: +41 61 284 8738
29		
30	keywords:	Aging, elderly, PBPK, modelling, population database
31		
32	article type	Original Research Article

#### 33 **1** Abstract

34

#### 35 Background

Aging is characterized by anatomical, physiological and biological changes which can impact drug kinetics. Elderly are often excluded from clinical trials and knowledge about drug kinetics and drug-drug interaction (DDI) magnitudes are sparse. Physiologically based pharmacokinetic (PBPK) modelling can overcome this clinical limitation but detailed descriptions of the population characteristics are essential to adequately inform models.

41

# 42 Objective

The objective of this work was to develop and verify a population database for aging Caucasians
considering anatomical, physiological and biological system parameters required to inform a PBPK
model with included population variability.

46

## 47 Methods

48 A structured literature search was performed to analyze age-dependent changes of system parameters.

49 All collated data were carefully analyzed, and descriptive, mathematical equations were derived.

50

#### 51 Results

A total of 362 studies were found of which 318 studies were included in the analysis as they reported rich data for anthropometric parameters and specific organs (e.g. liver). Continuous functions could be derived for most system parameters describing a Caucasian population from 20 to 99 years with variability. Areas with sparse data have been identified like tissue composition, but knowledge gaps were filled with plausible, qualified assumptions. The developed population was implemented in Matlab<sup>®</sup> and estimated system parameters from 1,000 virtual individuals were in accordance to independent observed data showing the robustness of the developed population.

59

#### 60 Conclusion

The developed repository for aging subjects provides a singular specific source for key system parameters needed for PBPK modelling and can in turn be used to investigate drug kinetics and DDI magnitudes in elderly.

# 64 2 Key Points

The developed repository provides a singular, specific source of age-dependent anatomical, physiological and biological system parameters required to inform physiologically based pharmacokinetic (PBPK) models. The parameters and associated developed equations can be implemented into existing PBPK frameworks and can be used to overcome sparse clinical data in aging subjects older than 65 years to investigate age-dependent changes in drug kinetics and drug-drug interaction (DDI) magnitudes *in silico*. These parameterized and informed PBPK models for elderly can provide more rational frameworks for dose-adjustments to overcome DDIs.

#### 72 **3** Introduction

73 In recent years the number of elderly people worldwide has increased substantially [1]. An "elderly" is 74 defined as being above the age of 65 years [2], which is in line with the age of retirement in most Western 75 countries. Older individuals are prone to multi-morbidities and hence polypharmacy and in turn for drug-76 drug interactions (DDIs) [3-5], however there is no clear pharmacological or clinical definition of "elderly" 77 [6]. Often, elderly subjects are excluded from clinical trials resulting in a general lack of knowledge about 78 the efficacy, safety and kinetics of a drug at different ages [7]. There are certain age-dependent 79 anatomical, physiological and biochemical changes influencing drug kinetics including decreased kidney 80 weight [8], reduced renal blood flow [9], reduced glomerular filtration rate (GFR) [10] and reduction in 81 liver volume and blood flow [11-13]. For other parameters like enzyme and transporter abundance, or 82 the concentration of plasma binding proteins, data are limited, contradictory or missing. In addition, it is 83 difficult to investigate aging, because other environmental and behavioral factors like diseases, food and 84 smoking can have effects themselves or enhance the aging process [14].

85

86 Physiologically based pharmacokinetic modelling can help to overcome the lack of clinical data and to 87 understand drug absorption, distribution, metabolism and elimination at different ages. Furthermore, 88 PBPK models predict DDI magnitudes in aging individuals and support more rational identification of 89 dose adjustments to overcome DDIs. To develop a PBPK model, system data (where system refers to the population of interest – e.g. elderly) are required to inform the PBPK model. To generate reliable 90 91 predictions, a comprehensive description of system characteristics is essential to fully represent the 92 population of interest. To date only two databases have been published to inform PBPK models for 93 elderly, of which one does not distinguish between ethnicities [15] and the other does not consider 94 population variability and provides no descriptive functions of physiological and anatomical parameters 95 [16].

96

97 The objective of this work was to collate and analyze data from the literature with the view to create a 98 new comprehensive description of system characteristics for PBPK modelling and to address 99 shortcomings of previous databases. The work focuses on parameters to inform a PBPK model for aging 100 people that considers population variability, and to develop continuous functions describing 101 physiological parameters of interest between 20 and 99 years of age for a Caucasian population.

- 3 -

#### 102 **4 Methods**

# 103 4.1 Data source

104 A structured literature search was performed using the MEDLINE database for age-dependency of 105 anatomical, physiological and biological parameters required to inform a PBPK model for aging subjects. 106 Keywords used were "aging", "elderly" or "geriatric" plus the parameter of interest (Supplement S-Table 107 1 and S-Figure 1 for the investigated compartments of a PBPK model). No restrictions were applied 108 regarding the language or the publication year of the article. Abstracts were screened, and studies 109 included if the study population were Caucasians, at least age has been reported in addition to the 110 parameter of interest, and subjects were healthy or their disease / organ function was deemed unlikely 111 to affect the parameter of interest like the effect of chronic liver disease on brain blood flow [17]. Studies 112 performed with North Americans and Australians were considered if at least 80% of the study population were of European heritage. Studies including subjects over the age of 65 years should report at least a 113 114 mean age in age decades. The reference list of chosen articles was manually screened to identify further 115 references.

#### 116 4.2 Data analysis

Data analysis was performed in Matlab<sup>®</sup> 2015b. The parameter of interest was analyzed in age decades. Data were converted to consistent units and a normal distribution was assumed for each parameter to make published data comparable. If a study reported median, minimum and maximum, data were converted to the arithmetic mean and standard deviation according to Hozo *et al.* [18] and if the interquartile range was given, the conversion was done according to Wan *et al.* [19].

122

123 Collated data were separated into a development and verification dataset. Studies in the development 124 dataset were required to report age, sex, body height, body weight and the ethnicity in addition to the 125 parameter of interest as necessary covariates to describe correlations. Otherwise, studies with less 126 reported covariates were used in the verification dataset. If at least three different studies covering the 127 entire age range with at least one value in each age decade and all required covariates for the 128 development dataset were available for a parameter of interest, the data was randomly separated into 129 a development and a verification dataset. In the case of missing covariates like anthropometric parameters in the verification dataset or cardiac output for regional blood flow analysis, the covariates
have been estimated by the derived equations following the approach by Williams & Leggett [20]. The
body surface area was calculated according to DuBois & DuBois [21].

133

134 Weighted linear regression was performed to derive descriptive, continuous equations for the parameter 135 of interest from 20 to 99 years considering age, sex, anthropometric parameters, location of the study, 136 the publication year and methods of measurement as independent variables. Location was used as an 137 independent variable to investigate if studies conducted in Europe, North America and Australia can be 138 combined without bringing a bias into the data. Publication year has been used to investigate differences 139 in key parameters (e.g. body weight) over the last century and if different methods used at different 140 times have an impact. Data obtained by different methods have only been pooled when there was no 141 significant difference between methods.

142

143 Linear, polynomial and exponential functions were investigated during regression analysis. Covariates 144 with a p-value below 0.01 have been considered as significant. Visual and numerical regression 145 diagnostic were performed. The corrected Aikake's information criterion was used for numerical 146 diagnostics to select the best fitted function [22]. Variability for each parameter was calculated as the 147 weighted coefficient of variance (CV) of the development dataset for each individual mean and standard 148 deviation and it was visually investigated whether age has an impact on variability. The variability of a 149 parameter of interest is estimated by the variability of the covariates describing the parameter of interest 150 and if necessary additional random variability to fully capture the observed variability.

151

The derived equations for all parameters necessary to describe a white population have been implemented in Matlab<sup>®</sup> and 1,000 virtual men and women have been created and the estimated system parameters have been compared to the independent verification dataset. <u>Normal distribution with the</u> derived CV (Tab. 1) was used to describe variability of the parameter of interest. Furthermore, it was analyzed if the sum of organ weights and regional blood flows do not exceed body weight and cardiac output.

#### 158 **5 Results**

A total of 362 studies were found of which 318 studies were included in the analysis. Studies were 159 160 mostly excluded because age or ethnicity of the study population were insufficiently defined. Rich data 161 were found for anthropometric parameters, adipose, brain, heart, kidney and liver. Data for some 162 regional blood flows, such as to the bone, and in general composition of tissues were difficult to obtain 163 from the literature. Although including data for centenarians, most of the data were found for ages up to the mid-eighties identifying a general knowledge gap for the very old. Derived equations and the 164 165 population variability expressed as the CV can be found in Table 1. Detailed information on the number 166 of subjects in each age decade used in the development dataset (S-Table 2), the number of total studies 167 in the development and verification dataset, the methods used to measure the parameter of interest, the 168 study location and the references (S-Table 4) can be found for each investigated parameter in the 169 supplement.

# 170 5.1 Age and sex distribution

171 Data regarding age and sex distribution were taken from Eurostat [23] for all 28 member states of the 172 European Union and the Federal Office for Statistics of Switzerland [24] (Figure 1). The number of 173 subjects in each age decade was found to be uniform between 20 and 59 years. The number of subjects 174 declined from the age of 60 years, with only 2% of the Swiss population being above 90 years. A Weibull 175 distribution with  $\alpha = 1.55$  and  $\beta = 61.73$  best described the age distribution. The proportion of women 176 was found to be 50% of the population in Europe till the age of 69 years and increased to over 80% for 177 very old Swiss subjects above the age of 100 years. In all following equations, age is expressed in years 178 and sex is either 0 for men or 1 for women.

# 179 5.2 Body height and body weight

Anthropometric data of 106,698 Caucasians have been analyzed in the developmental dataset [24-70] and the derived equation has been verified with data from 14,096 subjects [71-86]. The mean body height of Caucasians aged 20 to 59 years was 178 cm for men and 166 cm for women with a genderindependent CV of 3.8%. Body height declined 2% per age decade from the age of 60 years (Figure 2). The difference between men and women was constant at all age ranges. Location was found to be a significant variable during regression, with lower height observed in Southern Europe, and an exclusion
of data reported from Portugal, Spain and Italy led to a non-significance of location.

187

The mean body weight of a Caucasian aged 20 to 49 years was 79.9 kg for men and 64.1 kg for women with a CV of 15.7% (Figure 2). Body weight increased in subjects in the 5<sup>th</sup> and 6<sup>th</sup> age decade about 4% and decreased afterwards about 10% in each age decade. In women, the decline started one age decade later than in men. In contrast to body height, location was not significant for body weight, but publication year was with a significant increase since 2000.

#### 193 **5.3 Liver**

# 194 **5.3.1** Liver weight

Liver is the major organ of metabolism. Liver weight was analyzed from over 3,000 subjects [29, 41, 51, 52, 55, 69, 72, 78, 87, 88] and was found to be on average 1.78 kg in men and 1.49 kg in women with a CV of 23.7% till the age of 65 years. Thereafter, liver weight decreased by 10 to 15% in women per age decade reaching 1.03 kg at the age of 100 years. The decrease in men was around 20% per age decade reaching 1.01 kg on average in 90 years old individuals (Figure 3).

200

# 201 5.3.2 Liver blood flow

Absolute total liver blood flow decreased by 60% between 60 and 90 years in men and women, but relative to cardiac output the changes were only significant between 90 and 100 years of age [13, 89]. The age-dependent changes in total liver blood flow might come from changes of the splanchnic blood flow [77, 89-94] explaining observed differences in the first pass effect between young and old subjects [95-97]. The hepatic arterial blood flow appears to be constant with age [20, 89, 98].

207

#### 208 5.3.3 In-vitro-in-vivo extrapolation factors

PBPK models are informed by *in-vitro-in-vivo* extrapolation meaning that for instance the *in vivo* clearance is extrapolated from measured *in vitro* data. Hepatic scaling factors like the hepatocellularity (HPGL) or microsomal proteins per gram liver (MPPGL) are needed [99]. Barter *et al.* reported agedependent equations for HPGL [100] and MPPGL [101] with the oldest individuals in the analysis being between the mid-seventies and the early eighties.

# 215 5.3.4 Hepatic enzyme activity

Studies concerning the age-dependency of hepatic CYP enzyme activity are sparse and contradictory. 216 217 The biggest challenge is the high variability in hepatic CYP enzyme abundance [102, 103] and the small 218 sample size generally used for analysis [104, 105]. In a recent large meta-analysis investigating hepatic 219 CYP abundance to inform PBPK models, age was only a significant covariate for CYP2C9 [103]. It is 220 worthwhile mentioning, that the different genotypes known for CYP2C9 increase the sample size needed 221 to identify age-dependency even further. A significant age-dependency was detected for CYP1A2, 222 CYP2D6 and CYP2E1 in a different study, but not for CYP2C9 [106]. In a third study, CYP1A2 activity 223 was reported to be independent of age [107]. Consistent between different studies, CYP3A4 activity is 224 reported to be independent of age [108-110].

225

Posalek *et al.* investigated drug clearances in elderly for probe substrates like caffeine (CYP1A2), warfarin (CYP2C9), phenytoin (CYP2C19), desipramine (CYP2D6) and midazolam (CYP3A4) and found a clearance decrease of 30 to 40% in 70 years old subjects compared to young individuals, which can be explained by the decline in liver volume and blood flow rather than hepatic CYP enzyme activity [111]. In addition inflammation affects CYP enzyme activity [112] making it difficult to analyze data from non-healthy elderly.

232

UGT enzyme activity is reported to be independent of age in the literature [106, 113-115]. Taken together, this lack of evidence and data to inform age dependency necessitates a more judicious approach to assume no age-dependent hepatic enzyme activity and thus assume the same values in aging subjects as in young individuals.

- 237
- 238 **5.3.5** Hepatic drug transporter activity

Recently, a compact meta-analysis about hepatic drug transporter abundance to inform a PBPK model was published and age was tested as a covariate in the analysis and was reported to be not significant for any hepatic drug transporter [116]. In a PBPK model, we are interested in activity rather than abundance because the activity of enzymes and drug transporters can explain the observed data. If the abundance of transporter does not change, there might still be an age-dependent difference in transport activity; however, these data are currently not available. Comparable to hepatic enzymes, it is therefore
recommended to use the same activity in elderly as in young subjects.

# 246 5.4 Kidney

#### 247 5.4.1 Kidney weight

The literature search yielded nine different studies with a total of 1,620 data points measuring kidney weight after autopsy [29, 41, 42, 51, 52, 55, 69, 78, 85] (Figure 4A). The average kidney weight in young males and females was 0.318 kg with a CV of 19.3% and 0.259 kg with a CV of 23.2%, respectively. The reduction in kidney weight increased with age starting from 5% at the age of 70 years to 15% at the age of 80 years to 25% up to the age of 100 years in both genders.

253

## 254 5.4.2 Kidney blood flow

Absolute kidney blood flow decreased by 5 to 10% per age decade till the age of 65 years and thereafter decreased by 25% per age decade (Figure 4B) [77, 90, 94, 117-125]. Kidney blood flow relative to cardiac output was 19.7% in young men and decreased to 11.9% at the age of 85 years. The decrease was 5 to 20% per age decade. In women, the average kidney blood flow relative to cardiac output was 16.5% and stayed constant till the age of 70 years. Thereafter, it decreased to 9.2% at the age of 85 years.

261

# 262 5.4.3 Glomerular filtration rate

263 Only studies using inulin or <sup>51</sup>Cr-EDTA as a biomarker for glomerular filtration rate have been considered in this work [117-123, 125-129]. Equations to estimate the glomerular filtration rate like Cockcroft-Gault 264 265 [10] and the modification of diet in renal disease [130] use serum creatinine, which is problematic 266 considering senile sarcopenia in aging subjects [131]. The average glomerular filtration rate was 267 between 130 - 140 mL/min in men aged between 20 and 50 years and around 120 mL/min in women 268 of the same age. In the 5<sup>th</sup> age decade, glomerular filtration rate declined in men to 115 mL/min, which 269 was like the value in women (112 mL/min). Afterwards, the decline in glomerular filtration rate was 270 roughly 10% per age decade independent of gender reaching 50% of the value of a young adult at the 271 age of 90 years (Figure 4C).

# 272 5.5 <u>Adipose</u>

#### 273 5.5.1 Adipose weight

Adipose weight is usually measured via X-ray absorptiometry and bioelectric impedance analysis. Data from 18 different studies from 12,323 subjects were available for the development dataset [25, 26, 36, 37, 41, 42, 45-48, 57, 59, 60, 62, 65, 68, 73, 132]. In young men, adipose weight was on average 17.8 kg with a CV of 24%. It increased by 5 to 10% per age decade to 22.9 kg at the age of 70 years. The CV increased to 28%. In young women, adipose weight was found to be 17.3 kg with a CV of 29%. Between 20 and 70 years, adipose weight increased to 25.2 kg with a CV of 37% in women and decreased again to 21.9 kg with a CV of 37% at the age of 85 years.

281

## 282 5.5.2 Adipose blood flow

Adipose blood flow increased from 5% in young to 9% in aged males and from 8% in young to 10% in aged females [133, 134].

285 5.6 Muscle

#### 286 5.6.1 Muscle weight

Data from 11 different studies with 5,542 participants were available to analyze muscle weight, which was measured by X-ray absorptiometry and bioelectrical impedance analysis [26, 41, 42, 45, 50, 64, 73, 79, 81, 83, 132]. The average muscle weight was 32.0 kg in men aged 20 to 65 years and 19.8 kg in women of the same age. Muscle weight decreased by 10% per age decade between 65 and 100 years. The CV was 11.8% and was similar for males and females.

292

#### 293 5.6.2 Muscle blood flow

294 Only sparse data concerning muscle blood flow have been found in the literature which do not cover all 295 age decades but suggesting 17.5% of cardiac output in men and 11.1% in women [135-138]. 296 5.7 Brain

#### 297 5.7.1 Brain weight

Brain weight was analyzed by using data from eight different studies with 2,425 participants [29, 41, 42,
51, 52, 55, 78, 139] and was found to be independent of age. The average brain weight was 1.39 kg in
males and 1.28 kg in females with a gender-independent CV of 9%.

301

### 302 5.7.2 Brain blood flow

The literature search yielded 12 different studies with 956 participants for brain blood flow [140-151]. brain blood flow relative to cardiac output was 11.8% in men and 15.6% in women below the age of 40 years and increased to 15.6% in men and 16.3% in women in the 4<sup>th</sup> age decade and was constant thereafter.

#### 307 5.8 Heart

#### 308 5.8.1 Heart weight

Heart weight was analyzed using data from 10 different studies measuring heart weight after autopsy [29, 41, 42, 53, 55, 61, 69, 78, 152, 153] and increased in both, males and females, from 0.325 kg and 0.241 kg at the age of 25 to 0.390 kg and 0.317 kg in the 9<sup>th</sup> age decade.

312

#### 313 5.8.2 Heart blood flow

Blood flow to the heart relative to cardiac output increased from 5.5% at the age of 25 years to 12% at the age of 85 years in men and from 4.3% at the age of 25 years to 11.3% at the age of 70 years in women [154-159].

317

#### 318 **5.8.3 Cardiac output**

Cardiac output is the volume of blood being ejected by the heart per minute. Data from 12 studies involving 645 subjects were used to analyze cardiac output [39, 63, 70, 74, 77, 84, 90, 94, 135, 138, 160, 161]. Cardiac output decreased from 352 L/h in 30 years old males and 312 L/h in young females between 5 and 10% every age decade to 258 L/h in aged males and 201 L/h in aged females (Figure 5). The CV was similar between both genders with a value of 21.1%. 324 **5.9 Blood** 

#### 325 5.9.1 Blood weight

Blood weight was analyzed from seven different studies with 382 male and 179 female participants [27, 30, 31, 44, 66, 75, 162]. In young males, blood weight was 5.8 kg with a CV of 10% and decreased to 5.0 kg at the age of 90 years (Figure 6). In young women, blood weight was lower with 3.8 kg, but stayed constant over different age decades. At the age of 70 years, female blood weight was still 3.7 kg. The CV was like in men.

331

## 332 **5.9.2** Hematocrit

Blood parameters that have been analyzed were hematocrit and the concentration of albumin and alphaacidic glycoprotein (Figure 6). Data of 1,752 subjects aged 21 till 90 years were available to analyze hematocrit [122, 142, 163-168]. Sex was the only significant covariate. Mean values were  $0.443 \pm 0.064$ for men and  $0.410 \pm 0.063$  for women.

337

### 338 5.9.3 Plasma binding protein concentration

Alpha-acidic glycoprotein showed no significant covariate when analyzing data of 472 subjects aged 24
to 90 years from five different studies [169-173]. The mean value was 0.798 g/L with a CV of 24.3%.

341

Regression analysis of albumin yielded age as a significant covariate [169, 174-181] with an overall CV of 7.9%. Albumin concentration declined about 1.5% in each age decade. Malnutrition and acute illnesses, occurring both often in the elderly, can have a significant impact on albumin concentration complication the analysis of age-dependent albumin concentration [172, 174, 179],. Therefore, only data from apparently healthy subjects have been used in the analysis.

# 347 5.10 Other organs

Other organs like spleen and pancreas are not described in detail here, but the descriptive equations to describe an aging Caucasian population can be found in Table 1 and more detailed information can be found in the Supplement (S-Table 2, 3 and 4).

# 351 5.11 <u>Tissue composition</u>

352 Tissue composition is an important parameter to predict the distribution of drugs into tissues in a PBPK model. Data regarding the composition of lipids and proteins of tissues are generally sparse in humans 353 354 and no age-dependency was found in the literature, but total body water, total extracellular water and 355 total body cell mass have been reported in aging subjects [26, 37, 65, 182-190]. Age-independent 356 fraction of tissue volumes [191] coupled with age-dynamic tissue volumes have been used to calculate 357 the vascular and interstitial space of tissues (representing the extracellular water) and the intracellular 358 space minus the intracellular water (representing the cell mass). Organ densities to convert organ weight 359 obtained from the derived functions to volumes have been used from the ICRP database [192, 193]. 360 The weighted mean of the organ density and the fraction of tissue compositions of investigated organs 361 was used for the remaining organ. The values of all tissues have been summed up and compared against the observed data (Figure 7). The prediction of total body water and total cell mass were well in 362 363 agreement with the observed data leading to the conclusion that the made assumptions were adequate 364 to inform a PBPK model.

# 365 5.12 Parameters affecting drug absorption

Physiological parameters having an impact on drug absorption are gastric pH, gastric emptying and
 small intestinal transit time, the surface area available for absorption, and intestinal enzyme and drug
 transporter abundance.

369

#### 370 5.12.1 Gastric pH

One study compared gastric pH in fasted and fed state between 24 young, healthy volunteers aged 21 371 372 to 35 years [194] and 79 subjects aged 65 to 83 years [195]. The study reported a significant age-373 dependent difference between the median pH in fasted state (interquartile range) with 1.72 (1.08 - 2.34) 374 in the young group and 1.28 (0.90 - 5.60) in the aged group. The variability appeared to be much greater 375 in older individuals, but the difference in sample size need to be kept in mind. Another study in young 376 subjects below the age of 65 years found a median fasted pH of 1.45 [196]. To conclude, it is doubtful 377 if there is an age-dependency of gastric pH in fasted state and more data need to be generated and 378 included in the meta-analysis to judge the age effect properly. Gastric pH in fed state was not 379 significantly different between young and elderly subjects [194, 195], but the decline of gastric pH from

fed to fasted state was exponential with a half-life of 1.8 hours (CV: 65%) in young and was linear with a half-life of 3.0 hours (CV: 80%) in aging subjects [195]. 8% of Caucasians are achlorhydric meaning they do not secret hydrochloric acid in the gastric juice [197] and thus having a gastric pH at fasted state of 7.1 [195]. In Japanese, the number of achlorhydric subjects increases with age [198], but this appears not to be the case in healthy aging Caucasians [195].

385

### 386 5.12.2 Gastric emptying time

387 Reports in the literature about gastric emptying time are contradictory. Some studies report a slower 388 gastric emptying time [199, 200] in aging subjects, some report no changes [201, 202] and some a faster 389 rate [203, 204]. A lot of influencing factors exist for gastric emptying time like gastric pH [205], particle 390 size [203] and food [202, 203, 206] making it difficult to analyze age-dependency. Furthermore, gastric 391 emptying has a circadian rhythm making a difference if the study is conducted in the morning or in the 392 evening [207]. Two studies have investigated gastric emptying time after fluid and food intake in young 393 controls and aging subjects [206, 208]. Both studies used the same marker, the same method and both 394 started in the morning. Gastric emptying time was different between fluids and food but did not show 395 any age-dependency, which was verified by the regression analysis. Therefore, it is recommended to 396 use the same gastric emptying time in aging subjects as in young individuals.

397

# 398 5.12.3 Small intestinal transit time

399 Small intestinal transit time appears to be independent of age and a fixed value can be used to inform400 a PBPK model [209, 210].

401

#### 402 5.12.4 Passive permeability

The mucosal area is reported to decline with age [211, 212], but enterocytes and villi appear to be unchanged [212]. Malnutrition, disease and drug intake could alter the mucosa and need to be carefully considered when investigating age-dependency. Passive permeability was reported to be impaired in aging subjects [211], but two studies investigating mannitol and lactulose, two carbohydrates which are passively absorbed, showed no difference in passive permeability between young controls and aging subjects after correcting the data for the age-dependent decline in glomerular filtration rate [213, 214].

- 409 It is therefore assumed that neither the surface area available for passive diffusion nor the rate of passive
- 410 diffusion differ in aging subjects compared to young individuals.
- 411

# 412 **5.12.5** Intestinal enzyme and drug transporter abundance

- 413 Data regarding intestinal enzyme and drug transporter abundance are generally sparse and therefore
- 414 age-dependency cannot be analyzed sufficiently.

#### 415 6 Discussion

The described population database for aging subjects summarizes anatomical, physiological and biological system parameters required to inform PBPK modelling. Descriptive, continuous functions for systems parameters from the age of 20 to 99 years have been derived and verified with observed data extracted from peer-reviewed literature. Population variability was considered for each parameter.

420

421 Two previous databases have been described in the literature for aging individuals. Thompson et al. 422 gathered extensive data from the literature, but the authors did not considered different ethnic groups 423 and combined data from Caucasians, Latin-Americans and Asians [15]. However, it is known that 424 ethnicity can have a significant impact on system parameters, for instance hepatic enzyme abundance, 425 and therefore on clearance prediction [215]. Schlender et al. published recently a database for elderly 426 individuals further processing the data from Thompson et al. for Caucasians only [16]. A limitation of this 427 study is that only values for organ weight and blood flow for each age decade were considered making 428 it difficult to extrapolate to other ages of interest. Furthermore, population variability of system 429 parameters was not considered by Schlender et al., which is an essential element for reasonable 430 predictions of drug kinetics using PBPK models [216].

431

432 One notable novelty of the presented repository for Caucasian subjects are the derived continuous 433 functions that allow prediction for a population from 20 to 99 years of age. The advantage of continuous 434 functions is the creation of only one population with one distinct value at a certain age. If two separated 435 populations would have been built with one representing young subjects from 20 to 65 years and one elderly individuals from 65 to 99 years, there would be two separated equations calculating system 436 437 parameters at the age of 65 which might lead to un-physiological steps. Another advantage for the 438 prediction of monoclonal antibody kinetics or long-term drug therapies could be to introduce time-varying 439 physiology [217] so that subjects age during the time of the simulation.

440

A few limitations need to be acknowledged. Data from individuals over the age of 85 are sparse (S-Table 2 in the Supplement) meaning the derived equations could be less robust and extrapolation to older ages might be difficult. However, data for centenarians have been included for some system parameters [78] and were adequately estimated by the derived functions. Clinical studies are usually not performed in the very old making it impossible to verify the described population by analyzing drug kinetics. It is therefore recommended to use the described repository with caution at older ages. This
holds particularly true for regional blood flows to adipose, heart, muscle and skin, because almost no
geriatric data are currently available in the literature.

449

Another area with sparse data, where more research is needed in the future, is tissue composition being important to predict the distribution into tissues accurately. It was shown that the assumptions used in this work are plausible for total body water and cell mass (Figure 7), however, exception for single tissues cannot be excluded and data for lipid composition in the elderly were generally not found in the literature.

455

The analysis of system parameters to inform a PBPK model for aging Caucasians was complicated by the fact that some studies combine age groups together, meaning individuals aged 65 to 100 years might have been included, but only a mean age is given. This can lead to a bias in the data and hinders the characterization of age-dependent changes. Reports that insufficiently described age should generally be excluded unless no other data are available. Furthermore, ethnicity, particular in European studies, is not always clearly defined and need to be assumed from the given study location.

462

463 Predictions of system parameters become more robust when model parameters are correlated with 464 each other and covariability can be described [218, 219]. To obtain such descriptive correlations, studies 465 need to report important covariates, which is unfortunately not always the case. Weighted regression 466 analysis has been used to correlate parameters and to receive a more robust aging population. Linear 467 regression can only describe linear relationships, however, using data transformation such as logarithm 468 might compensate. Using regression, it is easy to overfit and model the noise in the data rather than the 469 relationship between parameters. In this work, the corrected Akaike's information criterion was used to 470 select the best performing function among the tested ones, which in contrast to the coefficient of 471 determination exhibits no bias to higher parameterized models. Another limitation of regression analysis 472 is its sensitivity towards outliers. Visual inspection of the estimated mean and variability of each 473 parameter compared to observed data in this work, did show an adequate fit all investigated parameters 474 (Figure 2 to 7).

475

476

477 The evaluation of variability was further complicated by being unable to set boundaries for publication 478 year and study location. For a few parameters, for instance blood weight, data were only available from 479 specific regions (e.g. United States) and from the 1950s. Both, location and publication year have 480 therefore been used as independent variables during regression and their impact has been quantified 481 when sufficient data were available. Body height and body weight are key parameters to describe a 482 population adequately and data from 106,698 individuals were available. Location was found to have 483 an impact to body height, with lower height correlated with Southern Europe. Otherwise, location was 484 not a significant covariate for any variable and therefore combining data of studies conducted in Europe, 485 the United States and Australia appears not to bring a bias into the data. However, the derived equations should not be used to predict aging Africans or Asians as aging processes might be different. Publication 486 487 year had a significant impact on body weight showing the weight increase particularly in the last ten 488 years. Consequently, the developed population will require constant updates to include future potential 489 changes like body weight.

490

491 A challenge when studying older individuals is that the definition of elderly is not universal. The WHO 492 specifies elderly as being above the age of 65 years [2], which is in accordance with the age of retirement 493 in most Western countries, but a clear pharmacological or clinical age-cut off is missing [6]. For some 494 patient groups, like people infected with HIV, the age cut-off is even as early as 50 years [220]. We 495 compared organs parameters important for drug disposition for 50 and 70 years old men and women 496 with 30 years old subjects (Figure 8). There is a progressive decline in relevant system parameters, 497 such as adipose weight, liver and kidney blood flow, with age. However, it is challenging to conclude a 498 "pharmacological" or "clinical" age cut-off for elderly based on the age-dependent changes in anatomical 499 and physiological parameters, because it is unknown when those changes affect drug kinetics 500 significantly. No study has been undertaken to compare pharmacokinetics of a drug between different 501 age decades and correlate those data to age-dependent changes of organ parameters. Furthermore, 502 elderly subjects included in clinical trials can have diseases influencing the parameter of interest. It is 503 therefore a challenge to define "healthy" in terms of an aged person.

504

505 Despite the limitations, in this work it was possible to derive descriptive, continuous functions to generate 506 a virtual population from 20 to 99 years in accordance to observed, independent data. Elderly are a 507 growing vulnerable patient population with a high frequency of co-morbidities and in turn polypharmacy.

- 18 -

However, aging subjects are often excluded from clinical trials and knowledge concerning drug kinetics
and DDI magnitudes are scarce. The developed population database can be implemented into existing
PBPK frameworks and can in turn be used to predict drug kinetics and DDI magnitudes in aging subjects
overcoming the lack of clinical data and providing a rational framework for dose optimization to
overcome DDIs.

# 513 **7** *Conclusions*

The population database for aging subjects presented in this work can be implemented into existing PBPK frameworks and allows the prediction of drug kinetics and DDI magnitudes in elderly. It provides descriptive, continuous functions for anatomical and physiological parameters from 20 to 99 years necessary to inform PBPK models and provides a view of the current literature concerning metabolizing enzymes and drug transporters in aging individuals. Furthermore, population variability is considered for all system parameters providing a framework for realistic pharmacokinetic predictions.

520

# 521 **8 Funding.**

- 522 This study was supported by the Swiss National Foundation (Grant number 166204), the OPO
- 523 Foundation and the Isaac Dreyfus Foundation.

524

# 525 9 Conflict of Interest

- 526 Felix Stader, Marco Siccardi, Manuel Battegay, Hannah Kinvig, Melissa A. Penny, and Catia Marzolini
- 527 have no conflict of interest to declare.

528 10 References

529

- 530 1. United Nations. World Population Ageing. 2015 [cited 23/01/2018]; Available from: 531 http://www.un.org/en/development/desa/population/publications/pdf/ageing/WPA2015 Report. 532 pdf 2. World Health Organisation. Definition of an older or elderly person. 2010 [cited 23/02/2018]; 533 Available from: http://www.who.int/healthinfo/survey/ageingdefnolder/en/index.html 534 Maher RL, Hanlon J, Hajjar ER. Clinical consequences of polypharmacy in elderly. Expert 535 3.
- 5353.Maher RL, Hanlon J, Hajjar ER. Clinical consequences of polypharmacy in elderly. Expert536Opinion on Drug Safety. 2014;13(1):57-65.
- 537 4. Greene M, Steinman MA, McNicholl IR, Valcour V. Polypharmacy, drug–drug interactions, and
   538 potentially inappropriate medications in older adults with human immunodeficiency virus
   539 infection. Journal of the American Geriatrics Society. 2014;62(3):447-53.
- 5. Barnett K, Mercer SW, Norbury M, Watt G, Wyke S, Guthrie B. Epidemiology of multimorbidity 541 and implications for health care, research, and medical education: a cross-sectional study. The 542 Lancet. 2012;380(9836):37-43.
- 5436.Singh S, Bajorek B. Defining 'elderly'in clinical practice guidelines for pharmacotherapy.544Pharmacy Practice. 2014;12(4):489-98.
- 5457.Watts G. Why the exclusion of older people from clinical research must stop. BMJ: British546Medical Journal (Online). 2012;344.
- 5478.Dunnill M, Halley W. Some observations on the quantitative anatomy of the kidney. The Journal<br/>of Pathology. 1973;110(2):113-21.
- 5499.Hollenberg NK, Adams DF, Solomon HS, Rashid A, Abrams HL, Merrill JP. Senescence and<br/>the renal vasculature in normal man. Circulation Research. 1974;34(3):309-16.
- 10. Cockcroft DW, Gault H. Prediction of creatinine clearance from serum creatinine. Nephron.
   1976;16(1):31-41.
- 55311.Koff R, Garvey A, Burney S, Bell B. Absence of an age effect on sulfobromophthalein retention554in healthy men. Gastroenterology. 1973;65(2):300-2.

55512.Marchesini G, Bua V, Brunori A, Bianchi G, Pisi P, Fabbri A, et al. Galactose elimination capacity556and liver volume in aging man. Hepatology. 1988;8(5):1079-83.

- Wynne HA, Cope LH, Mutch E, Rawlins MD, Woodhouse KW, James OF. The effect of age upon liver volume and apparent liver blood flow in healthy man. Hepatology. 1989;9(2):297-301.
  Safar M. Ageing and its effects on the cardiovascular system. Drugs. 1990;39(1):1-8.
- Thompson CM, Johns DO, Sonawane B, Barton HA, Hattis D, Tardif R, et al. Database for
   physiologically based pharmacokinetic (PBPK) modeling: physiological data for healthy and
   health-impaired elderly. Journal of Toxicology and Environmental Health, Part B. 2009;12(1):1 24.
- Schlender J-F, Meyer M, Thelen K, Krauss M, Willmann S, Eissing T, et al. Development of a
  whole-body physiologically based pharmacokinetic approach to assess the pharmacokinetics
  of drugs in elderly individuals. Clinical Pharmacokinetics. 2016;55(12):1573-89.
- 567 17. O'Carroll R, Ebmeier K, Dougall N, Murray C, Goodwin G, Hayes P, et al. Regional cerebral
  568 blood flow and cognitive function in patients with chronic liver disease. The Lancet.
  569 1991;337(8752):1250-3.
- 57018.Hozo SP, Djulbegovic B, Hozo I. Estimating the mean and variance from the median, range,571and the size of a sample. BMC Medical Research Methodology. 2005;5(13):1-10.
- 57219.Wan X, Wang W, Liu J, Tong T. Estimating the sample mean and standard deviation from the<br/>sample size, median, range and/or interquartile range. BMC Medical Research Methodology.<br/>2014;14(1):135-59.
- 57520.Williams L, Leggett R. Reference values for resting blood flow to organs of man. Clinical Physics576and Physiological Measurement. 1989;10(3):187-217.
- 57721.DuBois D, DuBois EF. Fifth paper the measurement of the surface area of man. Archives of578Internal Medicine. 1915;15(5):868-81.
- 579 22. Burnham KP, Anderson DR. Model selection and multimodel inference: a practical information-580 theoretic approach: Springer Science & Business Media; 2003.
- 58123.European Union. Eurostat Population.2013 [cited 23/02/2017]; Available from:582<a href="http://ec.europa.eu/eurostat/data/database">http://ec.europa.eu/eurostat/data/database</a>
- 58324.Bundesamt für Statistik (Schweiz). Permanent resident population by age,sex and category of<br/>citizenship.584citizenship.2016[cited 01/05/2017];Availablefrom:585https://www.bfs.admin.ch/bfs/de/home/statistiken/bevoelkerung.assetdetail.299701.html

- 58625.Bedogni G, Pietrobelli A, Heymsfield SB, Borghi A, Manzieri AM, Morini P, et al. Is body mass587index a measure of adiposity in elderly women? Obesity Research. 2001;9(1):17-20.
- 588 26. Bosy-Westphal A, Mast M, Eichhorn C, Becker C, Kutzner D, Heller M, et al. Validation of air589 displacement plethysmography for estimation of body fat mass in healthy elderly subjects.
  590 European Journal of Nutrition. 2003;42(4):207-16.
- 591 27. Brown E, Hopper Jr J, Hodges Jr J, Bradley B, Wennesland R, Yamauchi H. Red cell, plasma,
  592 and blood volume in healthy women measured by radiochromium cell-labeling and hematocrit.
  593 Journal of Clinical Investigation. 1962;41(12):2182-90.
- 59428.Statistisches Bundesamt der Bundesrepublik Deutschland. Mikrozensus Fragen zur595Gesundheit. Körpermaße der Bevölkerung. 2013 [cited 31/05/2017]; Available from:596<a href="https://www.destatis.de/DE/Publikationen/Thematisch/Gesundheit/Gesundheitszustand/Koerp">https://www.destatis.de/DE/Publikationen/Thematisch/Gesundheit/Gesundheitszustand/Koerp</a>597ermasse5239003099004.pdf?blob=publicationFile
- 59829.Calloway N, Foley C, Lagerbloom P. Uncertainties in geriatric data. II. Organ size. Journal of<br/>the American Geriatrics Society. 1965;13(1):20-8.
- 60030.Chien S, Usami S, Simmons R, McAllister F, Gregersen M. Blood volume and age: repeated601measurements on normal men after 17 years. Journal of Applied Physiology. 1966;21(2):583-6028.
- 60331.Cohn JE, Shock NW. Blood volume studies in middle-aged and elderly males. American Journal604of Medical Sciences. 1949;217:388-91.
- 60532.Corish CA, Kennedy NP. Anthropometric measurements from a cross-sectional survey of Irish606free-living elderly subjects with smoothed centile curves. British Journal of Nutrition.6072003;89(1):137-45.
- Be Groot C, Perdigao A, Deurenberg P. Longitudinal changes in anthropometric characteristics
   of elderly Europeans. SENECA Investigators. European Journal of Clinical Nutrition.
   1996;50(Spl2):S9-15.
- 61134.Delarue J, Constans T, Malvy D, Pradignac A, Couet C, Lamisse F. Anthropometric values in<br/>an elderly French population. British Journal of Nutrition. 1994;71(2):295-302.
- 613 35. Dey DK, Rothenberg E, Sundh V, Bosaeus I, Steen B. Height and body weight in the elderly. I.
  614 A 25-year longitudinal study of a population aged 70 to 95 years. European Journal of Clinical
  615 Nutrition. 1999;53(12):905-14.
- 616 36. Dey D, Bosaeus I, Lissner L, Steen B. Body composition estimated by bioelectrical impedance
  617 in the Swedish elderly. Development of population-based prediction equation and reference
  618 values of fat-free mass and body fat for 70-and 75-y olds. European Journal of Clinical Nutrition.
  619 2003;57(8):909-16.
- 37. Dey DK, Bosaeus I, Lissner L, Steen B. Changes in body composition and its relation to muscle
  strength in 75-year-old men and women: a 5-year prospective follow-up study of the NORA
  cohort in Göteborg, Sweden. Nutrition. 2009;25(6):613-9.
- 62338.Eiben G, Dey D, Rothenberg E, Steen B, Björkelund C, Bengtsson C, et al. Obesity in 70-year-624old Swedes: secular changes over 30 years. International Journal of Obesity. 2005;29(7):810-6257.
- 626 39. Farinatti PT, Soares PP. Cardiac output and oxygen uptake relationship during physical effort
  627 in men and women over 60 years old. European Journal of Applied Physiology.
  628 2009;107(6):625-31.
- Gallagher D, Visser M, De Meersman RE, Sepúlveda D, Baumgartner RN, Pierson RN, et al.
  Appendicular skeletal muscle mass: effects of age, gender, and ethnicity. Journal of Applied
  Physiology. 1997;83(1):229-39.
- Gallagher D, Belmonte D, Deurenberg P, Wang Z, Krasnow N, Pi-Sunyer FX, et al. Organ-tissue
  mass measurement allows modeling of REE and metabolically active tissue mass. American
  Journal of Physiology-Endocrinology And Metabolism. 1998;275(2):E249-E58.
- Gallagher D, Allen A, Wang Z, Heymsfield SB, Krasnow N. Smaller organ tissue mass in the
  elderly fails to explain lower resting metabolic rate. Annals of the New York Academy of
  Sciences. 2000;904(1):449-55.
- 63843.Gavriilidou N, Pihlsgård M, Elmståhl S. Anthropometric reference data for elderly Swedes and639its disease-related pattern. European Journal of Clinical Nutrition. 2015;69(9):1066-75.
- 640 44. Gibson JG, Evans WA. Clinical studies of the blood volume. II. The relation of plasma and total blood volume to venous pressure, blood velocity rate, physical measurements, age and sex in ninety normal humans. Journal of Clinical Investigation. 1937;16(3):317-28.
- 64345.Gillette-Guyonnet S, Nourhashemi F, Lauque S, Grandjean H, Vellas B. Body composition and<br/>osteoporosis in elderly women. Gerontology. 2000;46(4):189-93.
- 645 46. Gnudi S, Sitta E, Fiumi N. Relationship between body composition and bone mineral density in
  646 women with and without osteoporosis: relative contribution of lean and fat mass. Journal of
  647 Bone and Mineral Metabolism. 2007;25(5):326-32.

- 47. Henche SA, Torres RR, Pellico LG. An evaluation of patterns of change in total and regional
  body fat mass in healthy Spanish subjects using dual-energy X-ray absorptiometry (DXA).
  European Journal of Clinical Nutrition. 2008;62(12):1440-8.
- 48. Horber FF, Gruber B, Thomi F, Jensen EX, Jaeger P. Effect of sex and age on bone mass, body composition and fuel metabolism in humans. Nutrition. 1997;13(6):524-34.
- 49. Launer LJ, Harris T. Weight, height and body mass index distributions in geographically and ethnically diverse samples of older persons. Age and Ageing. 1996;25(4):300-6.
- Legrand D, Adriaensen W, Vaes B, Matheï C, Wallemacq P, Degryse J. The relationship
  between grip strength and muscle mass (MM), inflammatory biomarkers and physical
  performance in community-dwelling very old persons. Archives of Gerontology and Geriatrics.
  2013;57(3):345-51.
- 659 51. Molina DK, DiMaio VJ. Normal organ weights in men: part II—the brain, lungs, liver, spleen, and 660 kidneys. The American Journal of Forensic Medicine and Pathology. 2012;33(4):368-72.
- 661 52. Molina DK, DiMaio VJ. Normal organ weights in women: Part II the brain, lungs, liver, spleen, 662 and kidneys. The American Journal of Forensic Medicine and Pathology. 2015;36(3):182-7.
- 663 53. Olivetti G, Giordano G, Corradi D, Melissari M, Lagrasta C, Gambert SR, et al. Gender
  664 differences and aging: effects on the human heart. Journal of the American College of
  665 Cardiology. 1995;26(4):1068-79.
- 666 54. Perissinotto E, Pisent C, Sergi G, Grigoletto F, Enzi G. Anthropometric measurements in the 667 elderly: age and gender differences. British Journal of Nutrition. 2002;87(2):177-86.
- 55. Puggaard L, Bjørnsbo KS, Kock K, Lüders K, Thobo-Carlsen B, Lammert O. Age-related
  decrease in energy expenditure at rest parallels reductions in mass of internal organs. American
  Journal of Human Biology. 2002;14(4):486-93.
- 671 56. Ravaglia G, Morini P, Forti P, Maioli F, Boschi F, Bernardi M, et al. Anthropometric
  672 characteristics of healthy Italian nonagenarians and centenarians. British Journal of Nutrition.
  673 1997;77(1):9-17.
- 674 57. Ravaglia G, Forti P, Maioli F, Boschi F, Cicognani A, Gasbarrini G. Measurement of body fat in
  675 healthy elderly men: a comparison of methods. The Journals of Gerontology Series A: Biological
  676 Sciences and Medical Sciences. 1999;54(2):M70-M6.
- 677 58. Rea I, Gillen S, Clarke E. Anthropometric measurements from a cross-sectional survey of
  678 community dwelling subjects aged over 90 years of age. European Journal of Clinical Nutrition.
  679 1997;51(2):102-6.
- 59. Santana H, Zoico E, Turcato E, Tosoni P, Bissoli L, Olivieri M, et al. Relation between body
  composition, fat distribution, and lung function in elderly men. The American Journal of Clinical
  Nutrition. 2001;73(4):827-31.
- 683 60. Schutz Y, Kyle U, Pichard C. Fat-free mass index and fat mass index percentiles in Caucasians 684 aged 18-98 y. International Journal of Obesity. 2002;26(7):953-60.
- 685 61. Smith HL. The relation of the weight of the heart to the weight of the body and of the weight of the heart to age. American Heart Journal. 1928;4(1):79-93.
- 687 62. Svendsen OL, Hassager C, Christiansen C. Age-and menopause-associated variations in body
  688 composition and fat distribution in healthy women as measured by dual-energy X-ray
  689 absorptiometry. Metabolism. 1995;44(3):369-73.
- 63. Tanner J. The construction of normal standards for cardiac output in man. Journal of Clinical Investigation. 1949;28(3):567-82.
- 692 64. Tichet J, Goxe D, Sallé A, Berrut G, Ritz P. Prevalence of sarcopenia in the French senior 693 population. The Journal of Nutrition Health and Aging. 2008;12(3):202-6.
- 694 65. Vache C, Rousset P, Gachon P, Gachon A, Morio B, Boulier A, et al. Bioelectrical impedance
  695 analysis measurements of total body water and extracellular water in healthy elderly subjects.
  696 International Journal of Obesity. 1998;22(6):537-43.
- 66. Wennesland R, Brown E, Hopper Jr J, Hodges Jr J, Guttentag O, Scott K, et al. Red cell, plasma and blood volume in healthy men measured by radiochromium (Cr51) cell tagging and hematocrit: influence of age, somatotype and habits of physical activity on the variance after regression of volumes to height and weight combined. Journal of Clinical Investigation. 1959;38(7):1065-77.
- Whimster WF, Macfarlane AJ. Normal lung weights in a white population. American Review of
   Respiratory Disease. 1974;110(4):478-83.
- 704 68. Ylihärsilä H, Kajantie E, Osmond C, Forsén T, Barker DJ, Eriksson JG. Body mass index during
  705 childhood and adult body composition in men and women aged 56–70 y. The American Journal
  706 of Clinical Nutrition. 2008;87(6):1769-75.
- 70769.de la Grandmaison GL, Clairand I, Durigon M. Organ weight in 684 adult autopsies: new tables708for a Caucasoid population. Forensic Science International. 2001;119(2):149-54.

- 709 70. Starr I, Donal J, Margolies A, Shaw R, Collins L, Gamble C. Studies of the heart and circulation
  710 in disease; estimations of basal cardiac output, metabolism, heart size, and blood pressure in
  711 235 subjects. Journal of Clinical Investigation. 1934;13(4):561-92.
- 712 71. Bartali B, Benvenuti E, Corsi AM, Bandinelli S, Russo CR, Di Iorio A, et al. Changes in anthropometric measures in men and women across the life-span: findings from the InCHIANTI study. Sozial-und Präventivmedizin. 2002;47(5):336-48.
- 715 72. Chouker A, Martignoni A, Dugas M, Eisenmenger W, Schauer R, Kaufmann I, et al. Estimation
  716 of liver size for liver transplantation: the impact of age and gender. Liver Transplantation.
  717 2004;10(5):678-85.
- 718 73. Clarys J, Martin A, Drinkwater D. Gross tissue weights in the human body by cadaver dissection.
   719 Human Biology. 1984;56(3):459-73.
- 720 74. Cournand A, Riley R, Breed E, Baldwin ED, Richards Jr D, Lester M, et al. Measurement of cardiac output in man using the technique of catheterization of the right auricle or ventricle.
  722 Journal of Clinical Investigation. 1945;24(1):106-16.
- 723 75. Davy KP, Seals DR. Total blood volume in healthy young and older men. Journal of Applied 724 Physiology. 1994;76(5):2059-62.
- 725 76. Gillette-Guyonnet S, Nourhashemi F, Andrieu S, Cantet C, Albarède JL, Vellas B, et al. Body
  726 composition in French women 75+ years of age: the EPIDOS study. Mechanisms of Ageing and
  727 Development. 2003;124(3):311-6.
- 728 77. Ho C, Beard J, Farrell P, Minson C, Kenney W. Age, fitness, and regional blood flow during 729 exercise in the heat. Journal of Applied Physiology. 1997;82(4):1126-35.
- 730 78. Ishii T, Sternby NH. Pathology of centenarians. I. The cardiovascular system and lungs. Journal of the American Geriatrics Society. 1978;26(3):108-15.
- 732 79. Janssen I, Heymsfield SB, Wang Z, Ross R. Skeletal muscle mass and distribution in 468 men and women aged 18–88 yr. Journal of Applied Physiology. 2000;89(1):81-8.
- 73480.Kasiske B, Umen A. The influence of age, sex, race, and body habitus on kidney weight in<br/>humans. Archives of Pathology & Laboratory Medicine. 1986;110(1):55-60.
- 81. Kemmler W, Teschler M, Goisser S, Bebenek M, von Stengel S, Bollheimer LC, et al.
  Prevalence of sarcopenia in Germany and the corresponding effect of osteoarthritis in females
  738 70 years and older living in the community: results of the FORMoSA study. Clinical Interventions
  rain Aging. 2015;10:1565-73.
- Kumar NT, Liestøl K, Løberg EM, Reims HM, Mæhlen J. Postmortem heart weight: relation to
  body size and effects of cardiovascular disease and cancer. Cardiovascular Pathology.
  2014;23(1):5-11.
- 74383.Masanes Toran F, Culla A, Navarro-Gonzalez M, Navarro-Lopez M, Sacanella E, Torres B, et744al. Prevalence of sarcopenia in healthy community-dwelling elderly in an urban area of745Barcelona (Spain). The Journal of Nutrition, Health & Aging. 2012;16(2):184-7.
- 746 84. Mezzani A, Grassi B, Giordano A, Corrà U, Colombo S, Giannuzzi P. Age-related prolongation
  747 of phase I of Vo2 on-kinetics in healthy humans. American Journal of Physiology.
  748 2010;299(3):R968-R76.
- 74985.Nyengaard J, Bendtsen T. Glomerular number and size in relation to age, kidney weight, and<br/>body surface in normal man. The Anatomical Record. 1992;232(2):194-201.
- 86. Sprogøe-Jakobsen S, Sprogøe-Jakobsen U. The weight of the normal spleen. Forensic Science
   International. 1997;88(3):215-23.
- 753 87. Thompson EN, Williams R. Effect of age on liver function with particular reference to bromsulphalein excretion. Gut. 1965;6(3):266-9.
- 75588.Meyer W, Peter B, Solth K. The weight of organs in the older age groups (70-92 years) and their756relation to age and body weight. Virchows Archiv für pathologische Anatomie und Physiologie757und für klinische Medizin. 1963;337:17-32.
- 75889.Carlisle K, Halliwell M, Read A, Wells P. Estimation of total hepatic blood flow by duplex759ultrasound. Gut. 1992;33(1):92-7.
- Minson CT, Wladkowski SL, Cardell AF, Pawelczyk JA, Kenney WL. Age alters the cardiovascular response to direct passive heating. Journal of Applied Physiology. 1998;84(4):1323-32.
- 91. Nakamura T, Moriyasu F, Ban N, Nishida O, Tamada T, Kawasaki T, et al. Quantitative
  measurement of abdominal arterial blood flow using image-directed Doppler ultrasonography:
  superior mesenteric, splenic, and common hepatic arterial blood flow in normal adults. Journal
  of Clinical Ultrasound. 1989;17(4):261-8.
- Vanis L, Gentilcore D, Lange K, Gilja OH, Rigda RS, Trahair LG, et al. Effects of variations in intragastric volume on blood pressure and splanchnic blood flow during intraduodenal glucose infusion in healthy older subjects. American Journal of Physiology. 2012;302(4):R391-R9.

- Zoli M, Iervese T, Abbati S, Bianchi G, Marchesini G, Pisi E. Portal blood velocity and flow in aging man. Gerontology. 1989;35(2-3):61-5.
- Part SL, Kenney WL. Effects of hormone replacement therapy on hemodynamic responses of postmenopausal women to passive heating. Journal of Applied Physiology. 2000;89(1):97-103.
- Robertson D, Waller D, Renwick A, George C. Age-related changes in the pharmacokinetics and pharmacodynamics of nifedipine. British Journal of Clinical Pharmacology. 1988;25(3):297-305.
- 778 96. Castleden CM, George CF. The effect of ageing on the hepatic clearance of propranolol. British
   779 Journal of Clinical Pharmacology. 1979;7(1):49-54.
- 780 97. Greenblatt DJ, Harmatz JS, Shapiro L, Engelhardt N, Gouthro TA, Shader RI. Sensitivity to 781 triazolam in the elderly. New England Journal of Medicine. 1991;324(24):1691-8.
- 782 98. Tygstrup N, Winkler K, Mellemgaard K, Andreassen M. Determination of the hepatic arterial blood flow and oxygen supply in man by clamping the hepatic artery during surgery. Journal of Clinical Investigation. 1962;41(3):447-54.
- Howgate E, Rowland Yeo K, Proctor N, Tucker G, Rostami-Hodjegan A. Prediction of in vivo drug clearance from in vitro data. I: impact of inter-individual variability. Xenobiotica. 2006;36(6):473-97.
- Barter ZE, Bayliss MK, Beaune PH, Boobis AR, Carlile DJ, Edwards RJ, et al. Scaling factors for the extrapolation of in vivo metabolic drug clearance from in vitro data: reaching a consensus on values of human micro-somal protein and hepatocellularity per gram of liver. Current Drug Metabolism. 2007;8(1):33-45.
- Barter ZE, Chowdry JE, Harlow JR, Snawder JE, Lipscomb JC, Rostami-Hodjegan A.
  Covariation of human microsomal protein per gram of liver with age: absence of influence of operator and sample storage may justify interlaboratory data pooling. Drug Metabolism and Disposition. 2008;36(12):2405-9.
- 796102.Rowland-Yeo K. Abundance of cytochrome P450 in human liver: a meta-analysis. British797Journal of Clinical Pharmacology. 2004;57(5):687-8.
- Achour B, Barber J, Rostami-Hodjegan A. Expression of hepatic drug-metabolizing cytochrome p450 enzymes and their intercorrelations: a meta-analysis. Drug Metabolism and Disposition. 2014;42(8):1349-56.
- 801 104. Shimada T, Yamazaki H, Mimura M, Inui Y, Guengerich FP. Interindividual variations in human liver cytochrome P-450 enzymes involved in the oxidation of drugs, carcinogens and toxic chemicals: studies with liver microsomes of 30 Japanese and 30 Caucasians. Journal of Pharmacology and Experimental Therapeutics. 1994;270(1):414-23.
- Achour B, Russell MR, Barber J, Rostami-Hodjegan A. Simultaneous quantification of the abundance of several cytochrome P450 and uridine 5'-diphospho-glucuronosyltransferase enzymes in human liver microsomes using multiplexed targeted proteomics. Drug Metabolism and Disposition. 2014;42(4):500-10.
- Parkinson A, Mudra D, Johnson C, Dwyer A, Carroll K. The effects of gender, age, ethnicity,
  and liver microsomes and inducibility in cultured human hepatocytes. Toxicology & Applied
  Pharmacology. 2004;199(3):193-209.
- 812 107. Simon T, Becquemont L, Hamon B, Nouyrigat E, Chodjania Y, Poirier J, et al. Variability of cytochrome P450 1A2 activity over time in young and elderly healthy volunteers. British Journal of Clinical Pharmacology. 2001;52(5):601-4.
- 815 108. Schwartz JB. Erythromycin breath test results in elderly, very elderly, and frail elderly persons.
  816 Clinical Pharmacology & Therapeutics. 2006;79(5):440-8.
- Hunt CM, Westerkam WR, Stave GM. Effect of age and gender on the activity of human hepatic
   CYP3A. Biochemical Pharmacology. 1992;44(2):275-83.
- Schmucker DL, Woodhouse KW, Wang RK, Wynne H, James OF, McManus M, et al. Effects
  of age and gender on in vitro properties of human liver microsomal monooxygenases. Clinical
  Pharmacology & Therapeutics. 1990;48(4):365-74.
- Polasek TM, Patel F, Jensen BP, Sorich MJ, Wiese MD, Doogue MP. Predicted metabolic drug
  clearance with increasing adult age. British Journal of Clinical Pharmacology. 2013;75(4):101928.
- 825112.Morgan E. Impact of infectious and inflammatory disease on cytochrome P450–mediated drug826metabolism and pharmacokinetics. Clinical Pharmacology & Therapeutics. 2009;85(4):434-8.
- 113. Court MH. Interindividual variability in hepatic drug glucuronidation: studies into the role of age,
   sex, enzyme inducers, and genetic polymorphism using the human liver bank as a model
   system. Drug Metabolism Reviews. 2010;42(1):209-24.

- Herd B, Wynne H, Wright P, James O, Woodhouse K. The effect of age on glucuronidation and sulphation of paracetamol by human liver fractions. British Journal of Clinical Pharmacology. 1991;32(6):768-70.
- Villesen HH, Banning A-M, Petersen RH, Weinelt S, Poulsen JB, Hansen SH, et al.
  Pharmacokinetics of morphine and oxycodone following intravenous administration in elderly
  patients. Therapeutics and Clinical Risk Management. 2007;3(5):961-7.
- Burt HJ, Riedmaier AE, Harwood MD, Crewe HK, Gill KL, Neuhoff S. Abundance of hepatic transporters in Caucasians: a meta-analysis. Drug Metabolism and Disposition.
  2016;44(10):1550-61.
- Bauer JH, Brooks CS, Burch RN. Renal function and hemodynamic studies in low-and normal renin essential hypertension. Archives of Internal Medicine. 1982;142(7):1317-23.
- 118. Davies DF, Shock NW. Age changes in glomerular filtration rate, effective renal plasma flow,
  and tubular excretory capacity in adult males. Journal of Clinical Investigation. 1950;29(5):496507.
- Fliser D, Zeier M, Nowack R, Ritz E. Renal functional reserve in healthy elderly subjects. Journal
  of the American Society of Nephrology. 1993;3(7):1371-7.
- Fliser D, Franek E, Joest M, Block S, Mutschler E, Ritz E. Renal function in the elderly: impact of hypertension and cardiac function. Kidney International. 1997;51(4):1196-204.
- B48 121. Ghose K, Burch A. Measurement of renal functions by double isotope techniques in elderly patients during tenoxicam therapy. Archives of Gerontology and Geriatrics. 1989;9(2):115-22.
- Boldring W, Chasis H, Ranges HA, Smith HW. Relations of effective renal blood flow and glomerular filtration to tubular excretory mass in normal man. Journal of Clinical Investigation.
  1940;19(5):739-50.
- 853123.McDonald RK, Solomon DH, Shock NW. Aging as a factor in the renal hemodynamic changes854induced by a standardized pyrogen. Journal of Clinical Investigation. 1951;30(5):457-62.
- 855 124. Miller JH, McDonald RK, Shock NW. The renal extraction of p-aminohippurate in the aged 856 individual. Journal of Gerontology. 1951;6(3):213-6.
- Fuiano G, Sund S, Mazza G, Rosa M, Caglioti A, Gallo G, et al. Renal hemodynamic response
  to maximal vasodilating stimulus in healthy older subjects. Kidney International.
  2001;59(3):1052-8.
- 860 126. Stevens LA, Coresh J, Greene T, Levey AS. Assessing kidney function measured and 861 estimated glomerular filtration rate. New England Journal of Medicine. 2006;354(23):2473-83.
- 127. Christensson A, Elmståhl S. Estimation of the age-dependent decline of glomerular filtration rate from formulas based on creatinine and cystatin C in the general elderly population. Nephron Clinical Practice. 2011;117(1):c40-c50.
- Van Den Noortgate NJ, Janssens WH, Delanghe JR, Afschrift MB, Lameire NH. Serum cystatin
   C concentration compared with other markers of glomerular filtration rate in the old old. Journal
   of the American Geriatrics Society. 2002;50(7):1278-82.
- BeSanto N, Anastasio P, Coppola S, Barba G, Jadanza A, Capasso G. Age-related changes in renal reserve and renal tubular function in healthy humans. Child Nephrology and Urology. 1991;11(1):33-40.
- 130. Levey AS, Bosch JP, Lewis JB, Greene T, Rogers N, Roth D. A more accurate method to
  estimate glomerular filtration rate from serum creatinine: a new prediction equation. Annals of
  Internal Medicine. 1999;130(6):461-70.
- 131. Musso CG, Oreopoulos DG. Aging and physiological changes of the kidneys including changes
   in glomerular filtration rate. Nephron Physiology. 2011;119(Suppl. 1):p1-p5.
- 876 132. Zoico E, Di Francesco V, Guralnik J, Mazzali G, Bortolani A, Guariento S, et al. Physical disability and muscular strength in relation to obesity and different body composition indexes in a sample of healthy elderly women. International Journal of Obesity. 2004;28(2):234-41.
- 133. Lesser GT, Deutsch S. Measurement of adipose tissue blood flow and perfusion in man by uptake of 85Kr. Journal of Applied Physiology. 1967;23(5):621-30.
- Andersson J, Karpe F, Sjöström L-G, Riklund K, Söderberg S, Olsson T. Association of adipose
  tissue blood flow with fat depot sizes and adipokines in women. International Journal of Obesity.
  2012;36(6):783-9.
- Proctor DN, Newcomer SC, Koch DW, Le KU, MacLean DA, Leuenberger UA. Leg blood flow
   during submaximal cycle ergometry is not reduced in healthy older normally active men. Journal
   of Applied Physiology. 2003;94(5):1859-69.
- 887136.Amery A, Bossaert H, Verstraete M. Muscle blood flow in normal and hypertensive subjects:888influence of age, exercise, and body position. American Heart Journal. 1969;78(2):211-6.
- 137. Johnson JM, Brengelmann GL, Rowell LB. Interactions between local and reflex influences on human forearm skin blood flow. Journal of Applied Physiology. 1976;41(6):826-31.

- 891138.Proctor DN, Koch DW, Newcomer SC, Le KU, Leuenberger UA. Impaired leg vasodilation during<br/>dynamic exercise in healthy older women. Journal of Applied Physiology. 2003;95(5):1963-70.
- Spann W, Dustmann H. Weight of the human brain and its dependence on age, body length,
  cause of death and occupation. Deutsche Zeitschrift fur die gesamte gerichtliche Medizin.
  1964;56(5):299-317.
- Bertsch K, Hagemann D, Hermes M, Walter C, Khan R, Naumann E. Resting cerebral blood
  flow, attention, and aging. Brain Research. 2009;1267:77-88.
- 898 141. Chen JJ, Rosas HD, Salat DH. Age-associated reductions in cerebral blood flow are 899 independent from regional atrophy. Neuroimage. 2011;55(2):468-78.
- Davis SM, Ackerman RH, Correia JA, Alpert NM, Chang J, Buonanno F, et al. Cerebral blood
   flow and cerebrovascular CO2 reactivity in stroke-age normal controls. Neurology.
   1983;33(4):391-.
- 903 143. Devous M, Stokely E, Chehabi H, Bonte F. Normal distribution of regional cerebral blood flow measured by dynamic single-photon emission tomography. Journal of Cerebral Blood Flow & Metabolism. 1986;6(1):95-104.
- 906144.Hagstadius S, Risberg J. Regional cerebral blood flow characteristics and variations with age907in resting normal subjects. Brain and Cognition. 1989;10(1):28-43.
- 908145.Leenders K, Perani D, Lammertsma A, Heather J, Buckingham P, Jones T, et al. Cerebral blood909flow, blood volume and oxygen utilization. Brain. 1990;113(1):27-47.
- 146. Lu H, Xu F, Rodrigue KM, Kennedy KM, Cheng Y, Flicker B, et al. Alterations in cerebral metabolic rate and blood supply across the adult lifespan. Cerebral Cortex. 2011;21(6):1426-34.
- Martin AJ, Friston KJ, Colebatch JG, Frackowiak RS. Decreases in regional cerebral blood flow
   with normal aging. Journal of Cerebral Blood Flow & Metabolism. 1991;11(4):684-9.
- 915 148. Parkes LM, Rashid W, Chard DT, Tofts PS. Normal cerebral perfusion measurements using
  916 arterial spin labeling: reproducibility, stability, and age and gender effects. Magnetic Resonance
  917 in Medicine. 2004;51(4):736-43.
- 918149.Scheinberg P, Blackburn I, Rich M, Saslaw M. Effects of aging on cerebral circulation and<br/>metabolism. AMA Archives of Neurology & Psychiatry. 1953;70(1):77-85.
- 920150.Shaw TG, Mortel KF, Meyer JS, Rogers RL, Hardenberg J, Cutaia MM. Cerebral blood flow921changes in benign aging and cerebrovascular disease. Neurology. 1984;34(7):855-.
- Shin W, Horowitz S, Ragin A, Chen Y, Walker M, Carroll TJ. Quantitative cerebral perfusion
  using dynamic susceptibility contrast MRI: evaluation of reproducibility and age-and genderdependence with fully automatic image postprocessing algorithm. Magnetic Resonance in
  Medicine. 2007;58(6):1232-41.
- Molina DK, DiMaio VJ. Normal organ weights in men: part I the heart. The American Journal of Forensic Medicine and Pathology. 2012;33(4):362-7.
- 928153.Molina DK, DiMaio VJ. Normal organ weights in women: part I the heart. The American Journal929of Forensic Medicine and Pathology. 2015;36(3):176-81.
- Baliga RR, Rosen SD, Camici PG, Kooner JS. Regional myocardial blood flow redistribution as
   a cause of postprandial angina pectoris. Circulation. 1998;97(12):1144-9.
- Bergmann SR, Herrero P, Markham J, Weinheimer CJ, Walsh MN. Noninvasive quantitation of myocardial blood flow in human subjects with oxygen-15-labeled water and positron emission tomography. Journal of the American College of Cardiology. 1989;14(3):639-52.
- 156. Chan SY, Brunken RC, Czernin J, Porenta G, Kuhle W, Krivokapich J, et al. Comparison of
  maximal myocardial blood flow during adenosine infusion with that of intravenous dipyridamole
  in normal men. Journal of the American College of Cardiology. 1992;20(4):979-85.
- 157. Duvernoy CS, Meyer C, Seifert-Klauss V, Dayanikli F, Matsunari I, Rattenhuber J, et al. Gender
  differences in myocardial blood flow dynamics: lipid profile and hemodynamic effects. Journal
  of the American College of Cardiology. 1999;33(2):463-70.
- 158. Leight L, Defazio V, Talmers FN, Regan TJ, Hellems HK. Coronary blood flow, myocardial oxygen consumption, and myocardial metabolism in normal and hyperthyroid human subjects.
  643 Circulation. 1956;14(1):90-9.
- 944159.Senneff MJ, Geltman EM, Bergmann SR. Noninvasive delineation of the effects of moderate945aging on myocardial perfusion. Journal of Nuclear Medicine. 1991;32(11):2037-42.
- 946160.Brandfonbrener M, Landowne M, Shock NW. Changes in cardiac output with age. Circulation.9471955;12(4):557-66.
- 948161.Lewis WH. Changes with age in the cardiac output in adult men. American Journal of949Physiology. 1938;121(2):517-27.
- 950 162. Smith RH. Normal blood volumes in men and women over sixty years of age as determined by
  951 a modified Cr51 method. The Journal of the American Society of Anesthesiologists.
  952 1958;19(6):752-6.

- 953 163. Smith WW, Wikler NS, Fox AC. Hemodynamic studies of patients with myocardial infarction.
   954 Circulation. 1954;9(3):352-62.
- 955164.Tietz NW, Shuey DF, Wekstein DR. Laboratory values in fit aging individuals sexagenarians956through centenarians. Clinical Chemistry. 1992;38(6):1167-85.
- 957165.Zauber NP, Zauber AG. Hematologic data of healthy very old people. Jama.9581987;257(16):2181-4.
- 166. Timiras ML, Brownstein H. Prevalence of anemia and correlation of hemoglobin with age in a geriatric screening clinic population. Journal of the American Geriatrics Society.
   1987;35(7):639-43.
- 962 167. Sklaroff D. Isotopic determination of blood volume in the normal aged. American Journal of
   963 Roentgenology. 1956;75:1082-3.
- 964168.Jernigan JA, Gudat JC, Blake JL, Bowen L, Lezotte DC. Reference values for blood findings in<br/>relatively fit elderly persons. Journal of the American Geriatrics Society. 1980;28(7):308-14.
- 966 169. Veering BT, Burm A, Souverijn J, Serree J, Spierdijk J. The effect of age on serum concentrations of albumin and alpha 1-acid glycoprotein. British Journal of Clinical Pharmacology. 1990;29(2):201-6.
- Routledge P, Stargel W, Kitchell B, Barchowsky A, Shand D. Sex-related differences in the plasma protein binding of lignocaine and diazepam. British Journal of Clinical Pharmacology. 1981;11(3):245-50.
- Paxton J, Briant R. Alpha 1-acid glycoprotein concentrations and propranolol binding in elderly
   patients with acute illness. British Journal of Clinical Pharmacology. 1984;18(5):806-10.
- 974 172. Denko CW, Gabriel P. Age and sex related levels of albumin, ceruloplasmin, alpha 1 antitrypsin, alpha 1 acid glycoprotein, and transferrin. Annals of Clinical & Laboratory Science. 1981;11(1):63-8.
- 977 173. Blain P, Mucklow J, Rawlins M, Roberts D, Routledge P, Shand D. Determinants of plasma alpha 1-acid glycoprotein (AAG) concentrations in health. British Journal of Clinical Pharmacology. 1985;20(5):500-2.
- 980 174. Campion EW, Delabry LO, Glynn RJ. The effect of age on serum albumin in healthy males:
   981 report from the Normative Aging Study. Journal of Gerontology. 1988;43(1):M18-M20.
- Fu A, Nair KS. Age effect on fibrinogen and albumin synthesis in humans. American Journal of
   Physiology-Endocrinology And Metabolism. 1998;275(6):E1023-E30.
- 984 176. Gardner M, Scott R. Age-and sex-related reference ranges for eight plasma constituents derived
   985 from randomly selected adults in a Scottish new town. Journal of Clinical Pathology.
   986 1980;33(4):380-5.
- 987 177. Garry PJ, Hunt WC, Van der Jagt DJ, Rhyne RL. Clinical chemistry reference intervals for 988 healthy elderly subjects. The American Journal of Clinical Nutrition. 1989;50(5):1219-30.
- 989 178. Gersovitz M, Munro HN, Udall J, Young VR. Albumin synthesis in young and elderly subjects using a new stable isotope methodology: response to level of protein intake. Metabolism. 1980;29(11):1075-86.
- 992179.Pickart L. Increased ratio of plasma free fatty acids to albumin during normal aging and in<br/>patients with coronary heart disease. Atherosclerosis. 1983;46(1):21-8.
- 180. Reed A, Cannon D, Winkelman J, Bhasin Y, Henry R, Pileggi V. Estimation of normal ranges
  995 from a controlled sample survey. I. Sexand age-related influence on the SMA 12/60 screening
  996 group of tests. Clinical Chemistry. 1972;18(1):57-66.
- 181. Wallace S, Whiting B. Factors affecting drug binding in plasma of elderly patients. British Journal
  of Clinical Pharmacology. 1976;3(2):327-30.
- 999182.Boddy K, King PC, Hume R, Weyers E. The relation of total body potassium to height, weight,<br/>and age in normal adults. Journal of Clinical Pathology. 1972;25(6):512-7.
- 1001 183. Bruce A, Andersson M, Arvidsson B, Isaksson B. Body composition. Prediction of normal body potassium, body water and body fat in adults on the basis of body height, body weight and age.
  1003 Scandinavian Journal of Clinical and Laboratory Investigation. 1980;40(5):461-73.
- 1004 184. Cornish B, Ward L, Thomas B, Jebb S, Elia M. Evaluation of multiple frequency bioelectrical impedance and Cole-Cole analysis for the assessment of body water volumes in healthy humans. European Journal of Clinical Nutrition. 1996;50(3):159-64.
- 1007 185. Fülöp T, Worum I, Csongor J, Foris G, Leövey A. Body composition in elderly people.
  1008 Gerontology. 1985;31(1):6-14.
- 1009186.Hume R, Weyers E. Relationship between total body water and surface area in normal and<br/>obese subjects. Journal of Clinical Pathology. 1971;24(3):234-8.
- 1011187.Schoeller DA. Changes in total body water with age. The American Journal of Clinical Nutrition.10121989;50(5):1176-81.

- 1013 188. St-Onge M-P, Wang Z, Horlick M, Wang J, Heymsfield SB. Dual-energy X-ray absorptiometry lean soft tissue hydration: independent contributions of intra-and extracellular water. American Journal of Physiology-Endocrinology and Metabolism. 2004;287(5):E842-E7.
- 1016 189. Steen B. Body composition and aging. Nutrition Reviews. 1988;46(2):45-51.
- 1017 190. Chumlea WC, Guo SS, Zeller CM, Reo NV, Baumgartner RN, Garry PJ, et al. Total body water 1018 reference values and prediction equations for adults. Kidney International. 2001;59(6):2250-8.
- 1019191.Gill KL, Gardner I, Li L, Jamei M. A bottom-up whole-body physiologically based1020pharmacokinetic model to mechanistically predict tissue distribution and the rate of1021subcutaneous absorption of therapeutic proteins. The AAPS Journal. 2016;18(1):156-70.
- 1022192.Snyder W, Cook M, Nasset E, Karhausen L, Howells GP, Tipton I. Report of the Task Group on1023Reference Man: Pergamon Press; 1975.
- 1024193.Valentin J. Basic anatomical and physiological data for use in radiological protection: reference<br/>values: ICRP Publication 89. Annals of the ICRP. 2002;32(3):1-277.
- 1026 194. Dressman JB, Berardi RR, Dermentzoglou LC, Russell TL, Schmaltz SP, Barnett JL, et al.
  1027 Upper gastrointestinal (GI) pH in young, healthy men and women. Pharmaceutical Research.
  1909;7(7):756-61.
- 1029 195. Russell TL, Berardi RR, Barnett JL, Dermentzoglou LC, Jarvenpaa KM, Schmaltz SP, et al.
  1030 Upper gastrointestinal pH in seventy-nine healthy, elderly, North American men and women.
  1031 Pharmaceutical Research. 1993;10(2):187-96.
- 1032 196. Fallingborg J, Christensen L, Ingeman-Nielsen M, Jacobsen B, Abildgaard K, Rasmussen H.
   1033 pH-profile and regional transit times of the normal gut measured by a radiotelemetry device.
   1034 Alimentary Pharmacology & Therapeutics. 1989;3(6):605-14.
- 1035197.Jamei M, Turner D, Yang J, Neuhoff S, Polak S, Rostami-Hodjegan A, et al. Population-based1036mechanistic prediction of oral drug absorption. The AAPS Journal. 2009;11(2):225-37.
- 1037198.Morihara M, Aoyagi N, Kaniwa N, Kojima S, Ogata H. Assessment of gastric acidity of Japanese1038subjects over the last 15 years. Biological and Pharmaceutical Bulletin. 2001;24(3):313-5.
- 1039199.Evans MA, Triggs EJ, Cheung M, Broe GA, Creasey H. Gastric emptying rate in the elderly:1040implications for drug therapy. Journal of the American Geriatrics Society. 1981;29(5):201-5.
- 1041200.Horowitz M, Maddern GJ, Chatterton BE, Collins PJ, Harding PE, Shearman DJ. Changes in<br/>gastric emptying rates with age. Clinical Science. 1984;67(2):213-8.
- 1043201.Gainsborough N, Maskrey V, Nelson M, Keating J, Sherwood R, Jackson S, et al. The<br/>association of age with gastric emptying. Age and Ageing. 1993;22(1):37-40.
- 1045202.Madsen JL. Effects of gender, age, and body mass index on gastrointestinal transit times.1046Digestive Diseases and Sciences. 1992;37(10):1548-53.
- 1047203.Graff J, Brinch K, Madsen JL. Gastrointestinal mean transit times in young and middle-aged1048healthy subjects. Clinical Physiology. 2001;21(2):253-9.
- 1049204.Kupfer R, Heppell M, Haggith J, Bateman D. Gastric emptying and small-bowel transit rate in<br/>the elderly. Journal of the American Geriatrics Society. 1985;33(5):340-3.
- 1051 205. Chaw C, Yazaki E, Evans D. The effect of pH change on the gastric emptying of liquids measured by electrical impedance tomography and pH-sensitive radiotelemetry capsule. International Journal of Pharmaceutics. 2001;227(1):167-75.
- 1054206.Moore JG, Tweedy C, Christian PE, Datz FL. Effect of age on gastric emptying of liquid-solid<br/>meals in man. Digestive Diseases and Sciences. 1983;28(4):340-4.
- 1056207.Goo R, Moore J, Greenberg E, Alazraki N. Circadian variation in gastric emptying of meals in<br/>humans. Gastroenterology. 1987;93(3):515-8.
- 1058 208. Henderson JM, Heymsfield SB, Horowitz J, Kutner MH. Measurement of liver and spleen volume by computed tomography. Assessment of reproducibility and changes found following a selective distal splenorenal shunt. Radiology. 1981;141(2):525-7.
- 1061209.Fischer M, Fadda HM. The effect of sex and age on small intestinal transit times in humans.1062Journal of Pharmaceutical Sciences. 2016;105(2):682-6.
- 1063210.Husebye E, Engedal K. The patterns of motility are maintained in the human small intestine<br/>throughout the process of aging. Scandinavian Journal of Gastroenterology. 1992;27(5):397-<br/>404.
- Bender AD. Effect of age on intestinal absorption: implications for drug absorption in the elderly.
   Journal of the American Geriatrics Society. 1968;16(12):1331-9.
- 1068212.Warren P, Pepperman M, Montgomery R. Age changes in small-intestinal mucosa. The Lancet.10691978;312(8094):849-50.
- 1070213.Saltzman JR, Kowdley KV, Perrone G, Russell RM. Changes in small-intestine permeability with<br/>aging. Journal of the American Geriatrics Society. 1995;43(2):160-4.
- 1072 214. Valentini L, Ramminger S, Haas V, Postrach E, Werich M, Fischer A, et al. Small intestinal 1073 permeability in older adults. Physiological Reports. 2014;2(4):1-10.

- 1074215.Barter ZE, Tucker GT, Rowland-Yeo K. Differences in cytochrome p450-mediated1075pharmacokinetics between chinese and caucasian populations predicted by mechanistic1076physiologically based pharmacokinetic modelling. Clinical Pharmacokinetics.10772013;52(12):1085-100.
- 1078<br/>1079216.Jamei M, Dickinson GL, Rostami-Hodjegan A. A framework for assessing inter-individual<br/>variability in pharmacokinetics using virtual human populations and integrating general<br/>knowledge of physical chemistry, biology, anatomy, physiology and genetics: a tale of 'bottom-<br/>up'vs 'top-down'recognition of covariates. Drug Metabolism and Pharmacokinetics.<br/>2009;24(1):53-75.
- 1083<br/>1084217.Abduljalil K, Jamei M, Rostami-Hodjegan A, Johnson TN. Changes in individual drug-<br/>independent system parameters during virtual paediatric pharmacokinetic trials: introducing<br/>time-varying physiology into a paediatric PBPK model. The AAPS Journal. 2014;16(3):568-76.
- Ludin H. Radiologic estimation of kidney weight. Acta Radiologica Diagnosis. 1967;6(6):561-74.
  McLachlan M, Wasserman P. Changes in sizes and distensibility of the aging kidney. The British Journal of Radiology. 1981;54(642):488-91.
- 1089220.Centers for Disease Control and Prevention. HIV among people aged 50 and over. 2018 [cited<br/>26/02/2018]; Available from: <a href="https://www.cdc.gov/hiv/group/age/olderamericans/index.html">https://www.cdc.gov/hiv/group/age/olderamericans/index.html</a>

1092

1093 **11 Figures** 



1094

Figure 1: Proportion of subjects (1A) and proportion of women (1B) per age decade. Data are from the28-member states of the European Union (black bars) and Switzerland (white bars)



1098

1099



1100

Figure 2: Body height (2A) and body weight (2B) per age decade in an aging population. The blue, red and black lines represent the predicted mean of virtual males, virtual females and from all virtual subjects, respectively. The dashed lines represent the 5 and 95% percentile of the predictions. Stars show observed data from the development and circles represent overserved data from the independent verification dataset. The size of the stars and circles indicates the size of the studied population





Figure 3: Liver weight (3A) and liver blood flow (3B) per age decade in an aging population. The blue, red and black lines represent the predicted mean of virtual males, virtual females and from all virtual subjects, respectively. The dashed lines represent the 5 and 95% percentile of the predictions. Stars show observed data from the development and circles represent observed data from the independent verification dataset. Black circles represent data from an undefined gender population. The size of the stars and circles indicates the size of the studied population

- 1114
- 1115
- 1116



1117

Figure 4: Kidney weight (4A), kidney blood flow (4B) and glomerular filtration rate (4C) per age decade in an aging population. The blue, red and black lines represent the predicted mean of virtual males, virtual females and from all virtual subjects, respectively. The dashed lines represent the 5 and 95% percentile of the predictions. Stars show observed data from the development and circles represent observed data from the independent verification dataset. Black circles represent data from an undefined gender population. The size of the stars and circles indicates the size of the studied population



Figure 5: Cardiac output per age decade in an aging population. The blue, red and black lines represent the predicted mean of virtual males, virtual females and from all virtual subjects, respectively. The dashed lines represent the 5 and 95% percentile of the predictions. Stars show observed data from the development and circles represent observed data from the independent verification dataset. The size of the stars and circles indicates the size of the studied population

# 1132



#### 1133

Figure 6: Blood weight (6A), hematocrit (6B), albumin (6C) and alpha-acid glycoprotein (6D) concentration per age decade in an aging population. The blue, red and black lines represent the predicted mean of virtual males, virtual females and from all virtual subjects, respectively. The dashed lines represent the 5 and 95% percentile of the predictions. Stars show observed data from the development and circles represent overserved data from the independent verification dataset. Black circles represent data from an undefined gender population. The size of the stars and circles indicates the size of the studied population



Figure 7: Total body water (7A) and total body cell mass (7B) per age decade in an aging population. The blue, red and black lines represent the predicted mean of virtual males, virtual females and from all virtual subjects, respectively. The dashed lines represent the 5 and 95% percentile of the predictions. Stars show observed data from the development and circles represent observed data from the independent verification dataset. The size of the stars and circles indicates the size of the studied population



- Figure 8: Comparison of a 50 and 70 years old man (8A and 8B) and women (8C and 8D) with a 30 years old subject, who was arbitrarily chosen to represent a young individual. Blood flow is relative to
- 1153 cardiac output and all values are relative to a 30 years old man and women, respectively

# 1154 **12 Tables**

1155

Table. 1: Descriptive equations and population variability for anatomical, physiological and biological
parameters necessary to inform a PBPK model. Virtual subjects from 20 to 99 years can be generated.
Blood flows are relative to cardiac output and the variability is only propagated from cardiac output. *m*indicates male and *f* female, when there was a gender-related difference in the CV

Parameter	Unit	Descriptive equation	CV [%]
Body height	cm	$-0.0039 \times Age^{2} + 0.238 \times Age - 12.5 \times Sex + 176$	3.8
Body weight	kg	$\begin{array}{l} -0.0039 \times Age^2 + 1.12 \times Body \ height + 0.611 \times \\ Age - 0.424 \times Sex - 137 \end{array}$	15.2
Lung weight	kg	$e^{(0.028 \times Body \ height + 0.0077 \times Age - 5.6)}$	0
Adipose weight	kg	$0.68 \times Body \ weight - 0.56 \times Body \ height + 6.1 \times Sex + 65$	29.6
Bone weight	kg	$e^{(0.024 \times Body \ height - 1.9)}$	13.2
Brain weight	kg	e <sup>-0.0075×Age+0.0078×Body</sup> height-0.97	9.0
Gonad weight	kg	$\begin{array}{l} -0.00034 \times Body \ weight - 0.00022 \times Age - 0.03 \times \\ Sex + 0.072 \end{array}$	34.8
Heart weight	kg	$0.34 \times BSA + 0.0018 \times Age - 0.36$	17.9 (m), 22.7 (f)
Kidney weight	kg	$-0.00038 \times Age - 0.056 \times Sex + 0.33$	19.3 (m), 23.2 (f)
Muscle weight	kg	$17.9 \times BSA - 0.0667 \times Age - 5.68 \times Sex - 1.22$	11.8
Skin weight	kg	$e^{(-0.0058 \times Age - 0.37 \times Sex + 1.13)}$	8.3
Thymus weight	kg	0.0221	44.8
Gut weight	kg	$3E^{-06} \times Body  height^{2.49}$	7.3
Spleen weight	kg	e <sup>1.13×BSA-3.93</sup>	51.7
Pancreas weight	kg	0.103	27.8
Liver weight	kg	$e^{(0.87 \times BSA - 0.0014 \times Age - 1.0)}$	23.7
Blood weight	kg	$e^{(0.067 \times BSA - 0.0025 \times Age - 0.38 \times Sex + 1.7)}$	10.4
Cardiac output (CO)	L/h	$159 \times BSA - 1.56 \times Age + 114$	21.1
Adipose blood flow	% of CO	$(0.044 + 0.027 \times Sex) \times Age + 2.4 \times Sex + 3.9$	
Bone blood flow	% of CO	5	
Brain blood flow	% of CO	$e^{-0.48 \times BSA + 0.04 \times Sex + 3.5}$	
Gonad blood flow	% of CO	$-0.03 \times Sex + 0.05$	
Heart blood flow	% of CO	$-0.72 \times Body \ height - 10 \times Sex + 134$	
Kidney blood flow	% of CO	$-8.7 \times BSA + 0.29 \times Body \ height - 0.081 \times Age - 13$	
Muscle blood flow	% of CO	$-6.4 \times Sex + 17.5$	
Skin blood flow	% of CO	5	
Thymus blood flow	% of CO	1.5	
Gut blood flow	% of CO	$2 \times Sex + 14$	

Parameter	Unit	Descriptive equation	CV [%]
Spleen blood flow	% of CO	3	
Pancreas blood flow	% of CO	1	
Liver blood flow	% of CO	$-0.108 \times Age + 1.04 \times Sex + 27.9$	
Albumin	g/L	$-0.0709 \times Age + 47.7$	7.9
GFR	mL/min	$e^{-0.0079 \times Age + 0.5 \times BSA + 4.2}$	14.7