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### Coupled mother-child model for bioaccumulation of POPs in nursing infants

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### 1 Environmental Pollution 2008, 156, 90-98

# 2 Coupled Mother-Child Model for Bioaccumulation of POPs in Nursing

# 3 Infants

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36	Capsule: This paper addresses a model for accumulation of organic compounds by
37	mother and breast-fed infant, applicable for exposure assessment within larger
38	frameworks.
39	
40	
41	Abstract
42	
43	Bioaccumulation of persistent organic pollutants (POPs) leads to high levels in human
44	milk and high doses of POPs for nursing infants. This is currently not considered in
45	chemical risk assessment. A coupled model for bioaccumulation of organic chemicals in
46	breastfeeding mother and nursing infant was developed and tested for a series of organic
47	compounds. The bioaccumulation factors (BAF) in mother, breast milk and child were
48	predicted to vary with log Kow and, for volatile compounds, with Kaw and concentration
49	in air. The concentrations of POPs in the infant body increase the first half year to about
50	factor 3 above mother and decline thereafter to lower levels. The predicted results are
51	close to empirical data and to an empirical regression. The new mother-child model is
52	compact due to its easy structure and the analytical matrix solution. It could be added to
53	existing exposure and risk assessment systems, such as EUSES.
54	
55	
56	
57	
58	Keywords: Accumulation; Breast milk; Human exposure; Infant; Model; POP

#### **1** Introduction 59

60

61	Persistent organic pollutants (POPs) are "chemicals that remain intact in the environment
62	for long periods, become widely distributed geographically, accumulate in the fatty tissue
63	of living organisms, and are toxic to humans and wildlife" (UNEP 2007). POPs, such as
64	polychlorinated dibenzodioxins and -furans (PCDD/F), polychlorinated biphenyls (PCB)
65	and chloroorganic pesticides, have been detected in human milk samples all over the
66	world (Rogan et al. 1986, Schecter et al. 1996, Filser et al. 1997, Raab et al. 2007, Shen
67	et al. 2007, Wittsiepe et al. 2007, Tanabe and Kunisue 2007). This raised considerable
68	concern about adverse health effects on nursing infants (Harrison 2001, CEHAPE 2004,
69	US EPA 2006).
70	

71 The uptake of POPs, such as PCDD/F, by adults is mainly via food ingestion (Travis and 72 Hattemeyer-Frey 1991). The uptake by nursing infants via breast milk has been reported 73 to be higher than by adults via diet, for some POPs at levels above the acceptable daily 74 intake (Dahl et al. 1995, Kreuzer et al. 1997, Schade and Heinzow 1998, BGVV 2000, 75 Tanabe and Kunisue 2007). POPs may accumulate for a longer period in the body of the 76 mother and then be transferred to the nursing infant via mother's milk. Travis et al. (1988) 77 developed empirical relations for the accumulation of chemicals in human adipose tissue 78 and human milk. The regressions are based on 12 (tissue) or 6 (milk) organic chemicals 79 with a log  $K_{OW}$  between 1.32 and 6.50 (tissue) or 5.16 to 6.50 (milk). The bioaccumulation factors  $B_f$  (tissue) and  $B_m$  (milk) were defined as 80

82 Br = Concentration of organic in adipose tissue (mg/kg lipid)  
83 Br = Concentration of organic in adipose tissue (mg/kg lipid)  
84 Average daily intake of organic (mg/d)  
85 Br = Concentration of organic in breast milk (mg/kg lipid)  
86 Average daily intake of organic (mg/d)  
91 Travis et al. (1988) related these bioaccumulation factors to the log 
$$K_{OW}$$
  
92 substances.  
93  
94 Br = 2.0 x 10<sup>-4</sup> Kow<sup>1.05</sup>  $\left[\frac{d}{kg}\right]$  (n=12, r=0.98)  
95 Br = 9.8 x 10<sup>-5</sup> Kow<sup>1.14</sup>  $\left[\frac{d}{kg}\right]$  (n=6, r=0.97)

of the

96

Besides this empirical approach, several mathematical model approaches exist to predict 97 98 human tissue concentrations after uptake, e.g. the models prepared by Kreuzer et al. 99 (1997) or Filser et al. (1997) and Maruyama et al. (2003) for PCDD/F. Accumulation in 100 the food chain with subsequent accumulation in humans was addressed by Czub and 101 McLachlan (2004a,b). To summarize, compound-specific models, comprehensive 102 numerical models and also easy empirical models for the prediction of the accumulation 103 of POPs in humans are available. 104 105 However, what lacks is a model predicting accumulation of POPs or other compounds in

106 breastfeeding mother and nursing infant after uptake of chemicals via diet or other

107 relevant sources by mother, which is compact enough to be combined with other models

and estimation routines, e.g., for chemical safety assessment tools such as EUSES (EC1091996).

110

111	"Traditional risk assessment approaches and environmental health policies have focused
112	mainly on adults and adult exposure patterns, utilizing data from adult humans or adult
113	animals" (CEHAPE 2004). Indeed, current chemical risk assessment in the EU (EC 2003)
114	considers only grown-ups (70 kg bodyweight). An additional focus on children and in
115	particular nursing infants, which are one trophic level higher and are eventually also more
116	sensitive to chemicals, requires a compact exposure estimation method that can run with
117	a minimum data set.
118	
119	This paper addresses the development, parameterization, sensitivity analysis, validation

120 and application of a coupled model for accumulation of organic compounds by nursing

121 mother and child. The coupled differential equations were solved analytically. The model

122 was tested with 2,3,7,8-TCDD and compared to empirical data for 11 other compounds

123 collected by Travis et al. (1988).

124 2 Methods

125

#### 126 **2.1 Model Development**

Figure 1 gives an overview of the system considered by the model. The human body is 127 128 considered as a flux-through system. The input of chemical occurs via diet (mother) or 129 milk (child) and inhalation (both). Inside the body, phase equilibrium is assumed. The 130 compound is eliminated from the body by exhalation and excretion (both together are 131 named "outflux"), by metabolism and, in case of the nursing mother, with breast milk. 132 133 <Figure 1> 134 135 Mother before birth of the child 136 The input of chemical into the mother is independent of the concentration in her body,  $C_H$ , while the output is proportionally related to it. This yields a linear differential 137 138 equation for the mass balance of the form 139  $\frac{dm}{dt} = I - k \times m$ 140 (1) 141 where m [mg] is the mass of chemical in the human body,  $I \text{ [mg d}^{-1}\text{]}$  is the sum of daily 142 uptake of chemical and  $k [d^{-1}]$  is the loss rate constant. 143 144

145 The input *I* can be derived from measurements or exposure assessments. The loss rate

146 constant *k* is calculated from the flux of chemical out of the body.

The human body is considered as composed of the phases lipids and water. Lipids were assumed to dissolve the chemical similar to octanol. The phase equilibrium between concentration in human body,  $C_H$  [mg kg<sup>-1</sup>], and concentration in water,  $C_W$  [mg L<sup>-1</sup>], is

152 
$$K_{HW} = \frac{C_H}{C_W} = W_H + \frac{L_H}{\rho_L} \times K_{OW} \qquad \left[\frac{L}{kg}\right]$$
(2)

153

where  $K_{HW}$  is the partition coefficient human body to water [L kg<sup>-1</sup>],  $W_H$  is the water content [L kg<sup>-1</sup>] and  $L_H$  is the lipid content of the human body [kg kg<sup>-1</sup>],  $\rho_L$  is the density of lipids [kg L<sup>-1</sup>] and  $K_{OW}$  [L L<sup>-1</sup>] is the partition coefficient between octanol and water.

158 The change of chemical mass in time due to outflux of chemical from the body  $dm_F/dt$ 159 [mg d<sup>-1</sup>] is the sum of outflux with water, lipid and air

160

161 
$$\frac{dm_F}{dt} = F_W \times C_{F,W} + F_L \times C_{F,L} + F_A \times C_{F,A} = F \times C_F \qquad \left\lfloor \frac{mg}{d} \right\rfloor$$
(3)

162

163 where  $F_W$  is the outflux of water [L d<sup>-1</sup>],  $F_L$  is the outflux of lipids [kg d<sup>-1</sup>] (with feces) 164 and  $F_A$  is the outflux of air [L d<sup>-1</sup>] (exhalation).  $C_{F,W}$  [mg L<sup>-1</sup>],  $C_{F,L}$  [mg kg<sup>-1</sup>] and  $C_{F,A}$  [mg 165 L<sup>-1</sup>] are the concentrations in the water, lipid and gas fraction of the outflux;  $C_F$  [mg kg<sup>-1</sup>] 166 is the weighted average concentration in the outflux. The total material outflux F [kg d<sup>-1</sup>] 167 is the sum of the outfluxes of water, lipids and air,

169 
$$F = F_W \times \rho_W + F_L + F_A \times \rho_A \qquad \left[\frac{kg}{d}\right] \tag{4}$$

173 
$$F \times C_F = F \times f_W \times C_{F,W} + F \times f_L \times K_{OW} \times C_{F,W} + F \times f_A \times K_{AW} \times C_{F,W}$$
(5)

175 where  $K_{AW}$  is the partition coefficient [L L<sup>-1</sup>] between air and water (also known as

176 dimensionless Henry's Law constant), and f are the flux fractions [L/d : kg/d] of water W,

- 177 lipids L and air A of the total flux F,

179 
$$f_W = \frac{F_W}{F}, f_L = \frac{F_L/\rho_L}{F} \text{ and } f_A = \frac{F_A}{F} \qquad \left[\frac{L}{kg}\right]$$
 (6)

- 181 The average concentration of chemical in the outflux,  $C_F$ , is then

183 
$$C_F = f_W \times C_{F,W} + f_L \times K_{OW} \times C_{F,W} + f_A \times K_{AW} \times C_{F,W}$$
(7)

185 Note that for phase equilibrium,  $C_{F,W}$  (concentration in aqueous phase of outflux) equals 186  $C_W$  (concentration in aqueous phase of human body), and thus we derive 

188 
$$K_{FW} = \frac{C_F}{C_W} = f_W + f_L \times K_{OW} + f_A \times K_{AW} \qquad \left[\frac{L}{kg}\right]$$
(8)

190 where  $K_{FW}$  [L kg<sup>-1</sup>] is the partition coefficient between outflux [kg d<sup>-1</sup>] and water [L d<sup>-1</sup>]. 191 Then, the partition coefficient between human body and outflux,  $K_{HF}$  [kg/kg], is 192

193 
$$K_{HF} = \frac{C_H}{C_F} = \frac{K_{HW}}{K_{FW}} \qquad \qquad \left[\frac{kg}{kg}\right]$$
(9)

194

195 where  $C_H$  and  $C_F$  are the concentrations [mg kg<sup>-1</sup>] in human body and outflux in phase 196 equilibrium. It follows for the loss rate constant k [d<sup>-1</sup>] in eq. 1, which is the sum of the 197 losses by outflux and by metabolism or degradation with first-order  $k_{deg}$  [d<sup>-1</sup>]

198

199 
$$k = \frac{F}{M_H \times K_{HF}} + k_{deg} \qquad \left[\frac{1}{d}\right]$$
(10)

200

where  $M_H$  [kg] is the bodyweight. The analytical solution of equation (1) for the chemical mass *m* [mg] in human body at time *t* is

203

204 
$$m(t) = m_0 \times e^{-kt} + \frac{I}{k}(1 - e^{-kt})$$
 (11)

205

206 which gives in steady-state  $(t \rightarrow \infty)$ 

207

$$208 \qquad m(\infty) = \frac{I}{k} \tag{12}$$

210 Concentrations  $C_H$  [mg/kg] in the human body were derived from  $C_H=m/M_H$ , assuming a 211 constant bodyweight  $M_H$ 

212

213 
$$C_{H}(\infty) = \frac{m(\infty)}{M_{H}} = \frac{I}{k \times M_{H}}$$
  $\left[\frac{mg}{kg}\right]$  (13)

214

This solution was used to calculate the concentration of chemical in the woman before birth of the child (and before pregnancy, the bodyweight is constant at 60 kg).

217

### 218 Nursing mother with child

219 In this scenario, the mother gives birth to a child and nurses the infant. Equations for the

220 mother were modified, and new equations for breast milk and nursing child were

221 introduced.

222

223 **Mother.** Nursing changes the outflux from the mother. Milk consists in the model of 224 water and lipids. The flux of milk  $F_M$  [kg d<sup>-1</sup>] was added to the outflux F in equation (4) 225

226 
$$F = F_W \times \rho_W + F_L + F_A \times \rho_A + W_M \times \rho_W \times F_M + L_M \times F_M \qquad \left[\frac{kg}{d}\right] \qquad (14)$$

227

where  $W_M$  [L kg<sup>-1</sup>] is the water content and  $L_M$  [kg kg<sup>-1</sup>] is the lipid content of human milk. Fractions of outflux [L kg<sup>-1</sup>]  $f_W$ ,  $f_L$  and  $f_A$  were recalculated for the case of nursing.

231 
$$f_W = \frac{F_W + W_M \times F_M}{F}, f_L = \frac{F_L + L_M \times F_M}{F \times \rho_L} \text{ and } f_A = \frac{F_A}{F} \qquad \left[\frac{L}{kg}\right]$$
(15)

The other equations (eqs. 1,2, 8-13) were applied without changes, but the new values of F and f were entered.

235

236 Milk. With breast milk, chemical is lost from the mother and transferred to the baby

237 (Schecter et al. 1996). To calculate the concentration of chemical in milk, phase

equilibrium between milk and mother was assumed. The concentration in milk  $C_M$  [mg

239 kg<sup>-1</sup>] is

240

241 
$$C_M = K_{MH} \times C_H$$
  $\left\lfloor \frac{mg}{kg} \right\rfloor$  (16)

242

243 where  $K_{MH}$  [kg kg<sup>-1</sup>] is the partition coefficient between milk and human. The partition 244 coefficient milk to water  $K_{MW}$  [L kg<sup>-1</sup>] is 245

246  $K_{MW} = \frac{C_M}{C_W} = W_M + \frac{L_M}{\rho_L} \times K_{OW}$   $\left[\frac{L}{kg}\right]$  (17)

247

The partition coefficient between milk and human body  $K_{MH}$  [kg/kg] is then

250 
$$K_{MH} = \frac{C_M}{C_H} = \frac{K_{MW}}{K_{HW}}$$
  $\left[\frac{kg}{kg}\right]$  (18)

252 Child. The breast-fed infant can take up chemicals by breast milk and by inhalation.

Breathing is external input to the child,  $I_c = F_A \times C_A$ , where  $F_A$  [here: m<sup>3</sup> d<sup>-1</sup>] is the flux of inhaled air and  $C_A$  [mg m<sup>-3</sup>] is the concentration of chemical in air. Loss of chemical occurs via outflux and by metabolic elimination with first-order rate constant  $k_{deg}$  [d<sup>-1</sup>]. The mass balance for the child is

257

258 
$$\frac{dm_c}{dt} = I_c + C_M \times F_M - C_F \times F_C - k_{deg} \times m_C$$
(19)

259

where  $C_M$  [mg kg<sup>-1</sup>] denotes the concentration in breast milk,  $F_M$  [kg d<sup>-1</sup>] is the flux of milk from mother to child,  $C_F$  [mg kg<sup>-1</sup>] is the concentration in the outflux of the child and  $F_C$  [kg d<sup>-1</sup>] is the outflux from the child.

263

264 Using the partition coefficients, the equation can be rewritten to

$$266 \qquad \frac{dm_C}{dt} = I_C + K_{MH} \frac{F_M}{M_H} \times m_H - \frac{F_C}{K_{CF} \times M_C} \times m_C - k_{deg} \times m_C \qquad \left[\frac{mg}{d}\right] \tag{20}$$

267

where  $m_H$  [mg] is the chemical mass in mother (human H),  $m_C$  [mg] is the chemical mass in the child,  $K_{CF}$  [kg kg<sup>-1</sup>] is the partition coefficient between child and outflux and  $M_C$ [kg] is the body mass of the child. The phase equilibrium between child body (index C) and water (index W) is

273 
$$K_{CW} = \frac{C_C}{C_W} = W_C + \frac{L_C}{\rho_L} \times K_{OW} \qquad \left[\frac{L}{kg}\right]$$
(21)

where  $K_{CW}$  [L kg<sup>-1</sup>] is the partition coefficient child body to water, *C* is the equilibrium concentration in child, index C [mg kg<sup>-1</sup>], or water, index W [mg L<sup>-1</sup>],  $W_C$  [L kg<sup>-1</sup>] is the water content and  $L_C$  [kg kg<sup>-1</sup>] is the lipid content of the child body. The initial concentration in the child  $C_C(0)$  [mg kg<sup>-1</sup>] was calculated from phase equilibrium to mother

280

$$281 C_C(0) = \frac{K_{CW}}{K_{HW}} \times C_H (22)$$

282

The outflux  $F_C$  [kg d<sup>-1</sup>] from the child was summed up, as was done for the outflux from the mother:

285

$$286 F_C = F_W \times \rho_W + F_L + F_A \times \rho_A (23)$$

287

where indeces W, L and A indicate water, lipid and air. Again, the flux fractions were used to calculate the phase equilibrium between outflux and water,  $K_{FW}$ :

291 
$$K_{FW} = \frac{C_F}{C_W} = f_W + f_L \times K_{OW} + f_A \times K_{AW} \qquad \left\lfloor \frac{L}{kg} \right\rfloor$$
(24)

292

293 The partition coefficient between child body and outflux,  $K_{CF}$  [kg/kg], is

295 
$$K_{CF} = \frac{C_C}{C_F} = \frac{K_{CW}}{K_{FW}} \qquad \left[\frac{kg}{kg}\right]$$
(25)

296

#### 297 2.2 Matrix Solution

The differential equations of mother and child are coupled and were treated as a linear
2×2 matrix system of the form

300

$$301 \qquad \frac{dm_1}{dt} = a_{11}m_1 + a_{12}m_2 + I_1 \tag{26}$$

$$302 \qquad \frac{dm_2}{dt} = a_{21}m_1 + a_{22}m_2 + I_2 \tag{27}$$

303

Matrix element 1 is the mother. The matrix constant  $a_{11}$  [d<sup>-1</sup>] is the sum of all loss processes from the mother and is identical with the negative loss rate *k* (eq. 10). The matrix constant  $a_{12}$  [d<sup>-1</sup>] is what mother receives from the child and is zero (therefore, the equation for chemical mass in mother can be solved independently of that for the child, eqs. 1 and 11). Input  $I_I$  [mg d<sup>-1</sup>] is the sum of all input to mother.

309

Matrix element 2 is the nursed child. The matrix constant  $a_{21}$  [d<sup>-1</sup>] describes the transfer via milk from mother to child,  $a_{21}=K_{MH}\times F_M/M_H$ . The matrix constant  $a_{22}$  [d<sup>-1</sup>] describes all losses of chemical from the child,  $a_{22} = -F_C/(K_{CF}\times M_C)-k_{deg}$ . Input  $I_2$  includes all chemical input independent from the mother, i.e. via inhalation,  $I_2=F_A\times C_A$ . A standard

314 solution for this system of differential equations exists for the case of constant rates and

315 inputs (Nazaroff and Alvarez-Cohen 2001). Concentrations [mg/kg] were derived by

dividing the chemical mass [mg] by the bodyweight [kg].

317

### 318 **2.3 Parameterization of the Model**

319 Input data (Table 1) was selected from several sources, preferably from existing models

320 (Kreuzer et al. 1997, Czub and McLachlan 2004b), in order to allow a comparison of the

321 results. The application of the steady-state solution (eqs. 12, 13) for the mother before the

birth of her child avoids the need to chose an appropriate initial mass  $m_0$  for the first

323 generation. The 95%-steady-state is reached for latest t=18 years for all chemicals with

324 the default parameterization.

325

326 The total daily uptake  $I \text{ [mg d}^{-1}\text{]}$  was calculated as the sum of uptake via diet  $i_D \text{ [mg/d]}$ 327 and inhalation of air:

328

329 
$$I = i_D + F_A \times C_A \qquad \left[\frac{mg}{d}\right]$$
 (28)

330

331 For the breast-fed baby,  $i_D$  is 0.

332

Outflux of lipids was assumed to be 10% of lipids in the diet. With 70 g d<sup>-1</sup> as average lipid ingestion, 0.007 kg d<sup>-1</sup> outflux of lipids results. For the baby, 0.0045 kg d<sup>-1</sup> (1/10 of influx of lipids with milk) was used. Table 1 lists the input data chosen as default for the model and used in the following simulations.

338	To calculate concentrations in the body of the child during	the simulation period,	the
339	respective bodyweight was used, to account for growth effe	ects. The bodyweight o	of the
340	child with age (in years) was approximated by a second-ord	ler polynom fitted to g	rowth
341	data for girls in Germany (Hesse et al. 1997) (eq. 29)		
342			
343	$bw = -0.053 \times age^2 + 3.76 \times age + 3.54$	(n=36, R <sup>2</sup> =0.98)	(29)
344			
345	<table 1=""></table>		
346			

#### 347 **3 Results**

348

#### 349 **3.1 Example Simulation TCDD**

350 To illustrate the general behaviour of the model, an example simulation with 2,3,7,8-351 tetrachlordibenzo-p-dioxin (TCDD) was performed. TCDD is a highly toxic, persistent 352 lipophilic (log  $K_{OW}$  6.76) and semivolatile ( $K_{AW}$  0.0015) compound (Rippen 1991). The concentration of TCDD in air was set to 4 fg m<sup>-3</sup> (background concentration in Southern 353 Germany, McLachlan 1992). Ingestion of TCDD by the mother with diet was 25  $pg d^{-1}$ 354 355 (Kreuzer et al. 1997). Figure 2 shows the simulated concentration of TCDD in lipids for 356 mother and child over a three-years period. The starting concentration of the mother [3.6 ng kg<sup>-1</sup> lipid] is the steady-state concentration (eq. 13). For t>0, the matrix solution was 357 358 applied. For t>0, the concentration of TCDD in mother decreases exponentially and falls 359 to 63% of the initial concentration after 1/2 year and to 42% after 1 year of nursing. The initial concentration in the infant [3.6 ng kg<sup>-1</sup> lipid] is in equilibrium to mother. It steeply 360 increases to 12.3 ng kg<sup>-1</sup> lipids after  $\frac{1}{2}$  year. Hereafter, it falls, due to depletion of the 361 mother's body burden and growth dilution, to 8.8 ng kg<sup>-1</sup> lipids after 1 year and to 1.73 ng 362 kg<sup>-1</sup> lipid after 3 years (of course, 3 years nursing is rare). The concentration in lipids of 363 364 milk is identical to that in lipids of the mother body and was not plotted. During the 365 period of nursing, the loss of TCDD from mother with milk is higher than the daily 366 intake, which is the reason for the depletion of TCDD from the body of the mother. Figure 3 shows the ratio of the TCDD-dose taken up by the infant (per kg bw) divided by 367 the dose taken up by the mother (25 pg  $d^{-1} = 0.42$  pg kg<sup>-1</sup> bw  $d^{-1}$ ). The ratio is initially 110 368 369 and falls later to 45 (t=1/2a), 22 (t=1a) and 4.5 (t=3a). The dose ratio is much higher than

370	the concentration ratio (Figure 2). Uptake of TCDD with air is neither for mother
371	(inhalation 11 m3 per day, uptake 44 fg TCDD per day) nor infant (inhalation 4.5 m3 per
372	day, uptake 18 fg TCDD per day) of relevance. The maximum concentration ratio child
373	to mother is reached after t= $1/2a$ . The concentration in the child is maximally 3.4 times
374	that in mother before birth and falls to 2.5times (t=1a) and to 0.48times (t=3a), due to
375	rapid elimination and growth dilution. The calculated elimination half-time (ln 2 divide
376	by rate constant k) of TCDD from the body is 4.6 years for the mother before birth, 0.6
377	years for the nursing mother and only 0.34 years for the infant.
378	These simulation results can be confronted to empirical data (Kreuzer et al. 1997, Filser
379	et al. 1997). Measured concentrations of TCDD in lipids of adipose tissue and blood for
380	adults in Germany early 1990ies range from <0.1 ng kg <sup>-1</sup> lipid to 16 ng kg <sup>-1</sup> lipids, with
381	an average background level of 3 ng kg <sup>-1</sup> lipids (Filser et al. 1997). Concentrations in
382	breast milk vary between 1 and 3.9 ng kg <sup>-1</sup> lipids, decreasing during the period of
383	nursing, with an average of about 2 ng kg <sup>-1</sup> lipids. TCDD-concentrations in stillborn
384	range from of 1.3-2.1 ng/kg lipids. Concentrations of TCDD in lipids of adipose tissue,
385	faeces and blood of infants did not differ much and ranged from $<0.2$ to 7.3 ng kg <sup>-1</sup> lipids.
386	TCDD levels in adipose tissue of 20 breast-fed infants aged between 0 and 44 weeks
387	ranged from 0.16 to 4.1 ng kg <sup>-1</sup> tissue and were higher than that of non-breast-fed
388	children (0.16-0.76 ng kg <sup>-1</sup> lipids) (Kreuzer et al. 1997). Predicted half-life of TCDD in
389	infants was short (0.42 years), and increased to about 10 years for adults between 40 and
390	60 years of age. These results are throughout close to the outcome of the simulations with
391	the mother-child model, without any conflicting results.
392	<figure 2=""> <figure 3=""></figure></figure>

**<Figure 2> <Figure 3>** 

### 393 **4 Discussion**

394

### 395 4.1 Comparison of Regression and Mother-Child Model

396 The regression of Travis et al. (1988) uses only one physico-chemical parameter, the 397  $K_{OW}$ , while the mother-child model requires  $K_{OW}$  and  $K_{AW}$ . The bioaccumulation factors 398 (BAF), related to concentration in lipids, derived from model and regression were 399 compared. The concentration in air was set to 0. The steady-state BAF of mother at birth 400 of the child (t=0) and the BAF milk after t= $\frac{1}{2}$  year were plotted in Figure 4. The BAF for 401 mother and milk are practically identical, except for very hydrophilic compounds (the 402 relation to lipids gives artificially higher concentrations for milk if compounds do not 403 partition into lipids). The model BAF differ more than two orders of amount with low  $(10^{-9})$  or high (0.1)  $K_{AW}$  except for high log  $K_{OW}$ , because volatile compounds are rapidly 404 405 lost from the body via exhalation. Within its regression range (log  $K_{OW}$  1.32 to 6.50), the 406 regression gives similar results as the model with high  $K_{AW}$ , probably because the less 407 lipophilic compounds in the training set of the regression were all solvents with high  $K_{AW}$ 408 (Table 2). With increasing lipophilicity, the BAF predicted by the mother-child model reach a plateau (mother at t=0: BAF is 143 [d kg<sup>-1</sup> lipid], milk at t=0.5 years: BAF is 90 409 [d kg<sup>-1</sup> lipid]), while the BAF derived by the regression increase unlimited with  $K_{OW}$ . This 410 411 is unrealistic, except for short time-periods, as the loss of super-lipophilic compounds via 412 milk would be several orders of amount higher than the daily intake. The daily intake (1 mg  $d^{-1}$ ) is balanced at a BAF milk (4.5% lipids) of 22 [d kg<sup>-1</sup> lipid]. The regression gives 413 a BAF=22 [d kg<sup>-1</sup> lipid] with log  $K_{OW}$ =4.7, but higher BAF for all log  $K_{OW}$  above that 414 415 value. Contrary, the steady-state  $(t=\infty)$  BAF milk predicted by the mother-child model for

416	compounds with log $K_{OW} > 4.7$ is constant at 19 [d kg <sup>-1</sup> lipid]. In the initial period of
417	nursing, the BAF milk is above steady-state, therefore, mother is depleted from POPs by
418	nursing (Fig. 2).
419	
420	<figure 4=""></figure>
421	
422	4.2 Uptake via inhalation compared to uptake via food
423	
424	The impact of exhalation on BAF of hydrophilic to medium lipophilic compounds (log
425	$K_{OW}$ < 5) is evident from Figure 4: fugitive compounds with high $K_{AW}$ show much lower
426	bioaccumulation, due to this process. On the other hand, the $K_{AW}$ may also impact the
427	uptake by inhalation. Basically, this uptake is calculated from the product of
428	concentration in air and inhalation (eq. 28). Under certain conditions, such as ubiquitous
429	background distribution of persistent compounds, we may assume that the concentrations
430	in diet and air are near phase equilibrium. Using the formalism of section 2.1, the
431	equilibrium ratio $K_{DA}$ [m <sup>3</sup> kg <sup>-1</sup> ] between concentration in diet $C_D$ [mg kg <sup>-1</sup> ] and in air $C_A$
432	$[mg L^{-1}]$ is
433	

434 
$$\frac{C_D}{C_A} = K_{DA} = \frac{W_D + L_D \times K_{OW}}{K_{AW}} \qquad \left\lfloor \frac{L}{kg} \right\rfloor$$
(30)

436 where  $W_D$  is the water content [kg kg<sup>-1</sup>] and  $L_D$  is the lipid content [kg kg<sup>-1</sup>] of the diet. 437

438 The relation between the input data  $i_D$  (uptake of chemical with diet, mg d<sup>-1</sup>) and  $C_D$  is

441

442 where  $F_D$  is the daily dietary consumption [kg d<sup>-1</sup>]. Thus, the equilibrium concentration in 443 air  $C_{A,eq}$  [mg L<sup>-1</sup>] is

444

445 
$$C_{A,eq} = \frac{i_D}{K_{DA} \times F_D}$$
(32)

446

447 The dose via inhalation  $i_A$  [mg d<sup>-1</sup>] is subsequently

448

450

451 where  $F_A$  is the inhalation of air [mother 11 m<sup>3</sup> d<sup>-1</sup> and child 4.5 m<sup>3</sup> d<sup>-1</sup>].

452

453 A typical diet of an adult Danish female  $(F_D)$  contains 60 g lipids and 2 L water, hereof

454 1.4 L drinking water. Using these numbers, the ratio of uptake via air to uptake via diet,

455 assuming phase equilibrium between air and food (including water), was calculated.

456

Figure 5 shows that the relevance of inhalation as uptake pathway for chemicals into the human body depends much on the value of the partition coefficient air to water  $K_{AW}$ . For non-volatile compounds (low  $K_{AW}$ , 10<sup>-6</sup> L L<sup>-1</sup>), inhalation is not relevant at all. With very

460 low  $K_{AW}$  (10<sup>-9</sup> L L<sup>-1</sup>), the ratio of uptake with inhalation versus uptake with diet is never

461	above 1 : 100 000 (not shown). On the other hand, inhalation is the dominant way of
462	entry into the body for volatile compounds (high $K_{AW}$ , 0.1 L L <sup>-1</sup> ) with up to log $K_{OW}$ 4.
463	With moderate $K_{AW}$ (10 <sup>-3</sup> L L <sup>-1</sup> in Fig. 5), the relative importance of inhalation for the
464	body burden decreases, but it is still higher than uptake by diet for the less lipophilic
465	compounds (log $K_{OW} \le 2$ ). For lipophilic compounds (log $K_{OW} > 5$ ), which have the
466	highest bioaccumulation, uptake by inhalation is generally not of much relevance.
467	Compared to the mother, uptake via inhalation has similar (hydrophilic compounds) or
468	lower importance (lipophilic compounds) for the child.
469	
470	Note that these calculations were done for the rare case of near-equilibrium conditions. In
471	real life, many individuals live in urban centers, while the agricultural production is in
472	remote rural areas. It may be expected that the air pollution is higher in the cities, in
473	particular when additional indoor sources of pollutants are present. Furthermore,
474	lipophilic compounds may be strongly adsorbed to particles, which are inhaled
475	simultaneously with air. Thus, these conclusions are surely not of general validity, and

the relative importance of inhalation for the uptake of pollutants may be higher in real life

477 than expected from the calculations displayed in Fig. 5.

**<Figure 5>** 

### **4.3 Validation Against Empirical Data**

482 To derive their regressions for bioaccumulation in adipose tissue and breast milk, Travis
483 et al. (1988) collected twelve bioaccumulation factors (BAF) for human adipose tissue

484	and six BAF for breast milk from literature and pharmacokinetic models. The model was
485	tested against these data. Additionally, BAF for TCDD were calculated from data in
486	Kreuzer et al. (1997). The concentrations are related to lipid content. Log $K_{OW}$ -values
487	given in the original reference (Travis et al. 1988) were used, except for TCDD (Rippen
488	1991) (Table 2). One compound, pentachlorophenol, had to be excluded from the analysis
489	because it is not a neutral compound but a weak acid (Rippen 1990). The uptake of weak
490	electrolytes into living cells follows principles which are not covered by the model
491	(Trapp 2004).
492	
493	<table 2=""></table>
494	
495	In order to reproduce the experimental conditions under which empirical BAF were
496	derived, concentration in air was set to zero. Figure 6 shows the measured BAF for

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497 human adipose tissue of the 12 organic compounds, the results from the regression by 498 Travis et al. (1988) and the model outcome for mother before birth at steady-state. The 499 measured BAF range from 0.013 (TCE) to 724 (DDE) and are lowest for the volatile 500 compounds with low log  $K_{OW}$ . Naturally, the regression predicts this range, and its results are generally less than factor 5 from the measurements, except for TCDD (over-predicted 501 502 factor 21), which is out of the regression range. The model simulations, too, are close to 503 the measured data. The results differ maximally factor 7 (dieldrin). The averaged ratio 504 between predicted and measured BAF is 1.54 for the regression (3.15, including TCDD) 505 and 2.0 for the model (including TCDD). 506

507	Figure 7 shows the measured BAF for human milk of seven organic compounds. The
508	measured BAF range from 43 (dieldrin) to 1660 (PCB). The regression results are quite
509	close to the measured BAF, except for TCDD. To derive the BAF, averaged values from
510	milk samples in the period between birth and up to 18 month after birth have been used
511	(Rogan et al. 1986). Therefore, the measured BAF were compared to the model result at
512	birth (t=0) and for t=1 year. The calculated BAF milk are higher for t=0 and do not vary
513	much, as all compounds are lipophilic with log $K_{OW}$ >4.7. The predicted BAF are
514	somewhat too low, except for dieldrin and TCDD. The largest deviation is seen for PCB,
515	which is not a single compound but a mix of 209 congeners. The averaged ratio between
516	regression result and measured BAF is 2.02 (1.09 without TCDD). The ratio between
517	model prediction and measurements is 0.99 (0.87 without TCDD) for t=0 and 0.42 (0.37
518	without TCDD) for t=1 year.
519	

### 520 **<Figure 6> <Figure 7>**

521

#### 522 **4.4 Comment on Nursing**

The question is often raised whether nursing may have an adverse impact on the health of the child (BgVV 2000). While the high dose of POPs (here: TCDD) that the infant receives with breast milk suggests so, the moderate increase of infant body concentration gives less reasons to be concerned. There is evidence that after a few life-years, the difference between breast-fed and formula-fed infants in their body-burden with POPs, such as TCDD, vanishes (Kreuzer et al. 1997). If the mother nurses more than one child without longer periods in between, the model predicts lower body-burdens for the later

530 children, i.e., for the second child after one year nursing, the body concentration is about 531 the same as in the mother, if she never had breast-fed. Empirical studies confirm that the 532 first born child is at higher risk to be exposed to POPs that have accumulated in mother 533 and are transferred via mother milk (Tanabe and Kunisue 2007), and that levels of POPs 534 decrease during lactation (Harris et al. 2001). An argument pro nursing may also be that 535 the mother reduces her POP pool (Schecter et al. 1996). Metabolism in the body of the 536 mother reduces the dose transferred to the nursing infant. With metabolism half-times 537 below 14 days, the model predicts that the dose the nursing infant receives is always 538 below the dose for the mother. According to the model, the mother has a "filter effect" for 539 less lipophilic and volatile compounds: for those, the dose for the infant via breast milk is 540 lower than the dose mother takes up (per kg bodyweight) (Figure 4).

541

#### 542 **4.5 Limitations and Application Range of the New Model**

543 The new mother-child model is, due to the underlying equations for phase equilibrium, 544 not valid for inorganic (Wuenschmann et al. 2008) and electrolytic organic compounds 545 (Trapp 2004, Trapp and Horobin 2005). The assumption of phase equilibrium within the 546 body for neutral lipophilic organic compounds is supported by the results of Kreuzer et 547 al. (1997), who found comparable levels of TCDD in lipids of adipose tissue, feces, blood, liver, breast milk and new-borns. Deviations from equilibrium could in particular 548 549 occur for compounds with rapid metabolism. However, for those the model predicts low 550 transfer into infants anyhow, thus, a "false alarm" due to over-prediction would not occur,

551 if accurate metabolism rate constants are at hand.

552

553	The new mother-child model is more complex than the regression of Travis et al. (1988),
554	but still the structure is relatively easy, and the analytical solution of the differential
555	equations keeps the calculations compact and robust. The model requires five chemical
556	input parameters ( $i_D$ , $K_{OW}$ , $C_A$ , $K_{AW}$ and $k_{deg}$ ). It may be more troublesome to acquire these
557	data, but the differences in the accumulation behavior of persistent and reactive
558	compounds can be considered, and uptake via diet and inhalation can be calculated
559	simultaneously or separately. Therefore, results from diet studies can be used as input
560	data, and bioaccumulation factors as defined by Travis et al. (1988) can be calculated,
561	using $i_D = 1 \text{ mg d}^{-1}$ and $C_A = 0$ . The regression necessarily will fail if uptake from air plays
562	a major role.

Another advantage of the deterministic approach, compared to empirical relations, is that

the relevant processes behind the BAF can be identified. The variation of physiological

566 parameters (for the human body) allows to determine the influence of age, diet,

567 bodyweight, growth, metabolism etc. Furthermore, the regression violates the mass

balance for more lipophilic compounds with high  $\log K_{OW}$  and gives unrealistically high

569 BAF, as was shown before.

570

571 In comparison to more sophisticated models for bioaccumulation (Kreuzer et al. 1997,

572 Molen et al. 1996, Maruyama et al. 2003), the new mother-child model is more compact

and more variable (i.e., it does not require the measurement of any chemical-specific

574 data, besides the minimum data set, and it needs no calibration steps). Compared to the

575 human bioaccumulation model ACC (Czub and McLachlan 2004b), which calculates the

576 body concentration of a single human over the whole life-time, the mother-child model is 577 less complex and more flexible, due to the analytical solution. If, for the purpose of risk 578 assessment, only the dose for the infant is required, the differential equation system is 579 decoupled, and the solution for the breast-feeding mother alone can be solved (eq. 11). 580 581 The development of the new mother-child model was driven by the need to predict the 582 exposure of nursing children within the framework of chemical risk assessment and/or 583 risk assessment of polluted sites. Model systems for these purposes exist (EC 1996, 584 Rikken et al. 2001, Kulhanek et al. 2004) but none of them considers nursing infants (in 585 fact, children are not considered at all in most of them). The new model could be added 586 with small effort to existing exposure assessment tools, in order to fill this gap. 587

#### 588 Model availability

The new mother-child model is available as unprotected excel-spreadsheet version fromthe first author. Please mail to stt@er.dtu.dk.

591

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Parameter	Symbol	Value	Unit	Reference
Mother				
Age	t	25	а	Kreuzer et al. (1997)
Body mass	M <sub>H</sub>	60	kg	Maruyama et al. (2003)
Body water fraction	W	0.71	L/kg	Czub and McLachlan (2004b)
Body lipid fraction	L	0.284	kg/kg	Deurenberg et al. (1991)
Outflux of water	$F_W$	1.24	$Ld^{-1}$	Maruyama et al. (2003)
Outflux of lipid	$F_L$	0.007	$kg d^{-1}$ m <sup>3</sup> d <sup>-1</sup>	10% of lipids in diet
In/exhalation of air	F <sub>A</sub>	11	$m^3 d^{-1}$	Layton (1993)
Breast milk data				
Milk flux	F <sub>M</sub>	1	kg d⁻¹	Kreuzer et al. (1997)
Milk water content	$W_M$	0.87	L kg <sup>-1</sup>	Czub and McLachlan (2004b)
Milk lipid content	L <sub>M</sub>	0.045	kg kg⁻¹	Kreuzer et al. (1997)
Child				
Age	t	0 - 3	а	
Body mass	$M_b$	3.5 - 7.25	kg	Hesse et al. (1997)
Body water fraction	W	0.71	L/kg	Czub and McLachlan (2004b)
Body lipid fraction	L	0.233	kg/kg L d <sup>-1</sup>	Deurenberg et al. (1991)
Outflux of water	$F_W$	0.87	L d <sup>-1</sup>	water content of 1 kg milk
Outflux of lipid	$F_L$	0.0045	$kg d^{-1}$ m <sup>3</sup> d <sup>-1</sup>	10% of influx
Outflux of air	F <sub>A</sub>	4.5	$m^3 d^{-1}$	Layton (1993)
Other data				
Density of water	$\rho_W$	1	kg L⁻¹	
Density of lipids	$ ho_{L}$	0.82	kg L <sup>-1</sup>	
Density of air	ρΑ	1.3×10 <sup>-3</sup>	kg L <sup>-1</sup>	

# **Table 1.** Default input data for the mother-child model.

Abbreviation	Compound	log K <sub>ow</sub> <sup>a</sup>	K <sub>AW</sub> <sup>b</sup>
Benzene	benzene	2.13	0.23
DDE	1,1-bis(4-chlorphenyl)-2,2-dichlorethen	5.83	0.05
DDT	1,1-bis(4-chlorphenyl)-2,2,2-trichlorethan	5.76	0.0016
DCM	dichlormethane	1.32	0.087
Dieldrin	dieldrin	5.16	0.0044
HE	heptachlor epoxide	5.40	0.01
HCB	hexachlorbenzene	5.45	0.028
PCE	perchlorethene	2.53	0.54
PCB	polychlorinated biphenyls	6.50	0.001 <sup>c</sup>
TCE	trichlorethene	2.33	0.35
MC	methylchloroform	2.47	0.715
TCDD	2,3,7,8-tetrachlordibenzo-p-dioxin	6.76 <sup>b</sup>	0.0015

743	Table 2. Names and	physico-chemical data of the compounds in Travis et al. (198	8).
110		singside diterinear adda of the desing stands in fractis et al. (198	0).

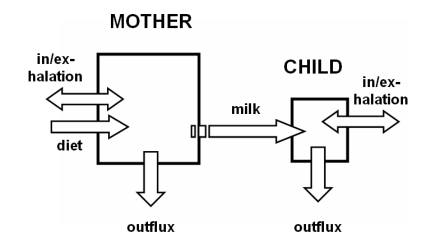
745 a) Travis et al. (1988) if not given otherwise; b) Rippen (1991-2007) if not given otherwise; c) estimate; PCB is a mix of 209 compounds.

746 747 748 749 750 751 752	Stefan Trapp, Li Ma and Charlotte N. Legind Figures to Coupled Mother-Child Model for Bioaccumulation of POPs in Nursing Infants
753	Figure Captions
754	
755	Figure 1. System overview
756 757 758 759 760	<b>Figure 2.</b> Concentrations in nursing mother and child (ng kg <sup>-1</sup> lipids) after uptake of 25 pg TCDD per day with diet by the mother.
761 762 763	<b>Figure 3.</b> Ratio of the TCDD-dose taken up by the nursing infant (per kg bw) to the dose taken up by the mother (25 pg $d^{-1} = 0.42$ pg kg <sup>-1</sup> bw $d^{-1}$ ).
764 765 766 767	<b>Figure 4.</b> Calculated bioaccumulation factor (BAF) mother (t=0) and milk (t=0.5 a) with varying log $K_{OW}$ for low $K_{AW}$ ( $K_{AW} = 10^{-9}$ L L <sup>-1</sup> ) and high $K_{AW}$ ( $K_{AW} = 0.1$ L L <sup>-1</sup> ) compared to the result derived with the regression of Travis et al. (1988).
768 769 770 771 772 773	<b>Figure 5.</b> Calculated ratio of uptake via inhalation to uptake via diet for the assumption of phase equilibrium for mother and child (t=0.5 a) with varying log $K_{OW}$ for high $K_{AW}$ ( $K_{AW} = 0.1 \text{ L L}^{-1}$ ), moderate $K_{AW}$ ( $K_{AW} = 0.001 \text{ L L}^{-1}$ ) and low $K_{AW}$ ( $K_{AW} = 10^{-6} \text{ L L}^{-1}$ ). Dotted line shows ratio 1:1.
774	<b>Figure 6.</b> Bioaccumulation factors (related to lipid content) for human adipose tissue for

ose tissue for 12 neutral organic compounds collected from literature (Lit) compared to the regression 775 776 by Travis et al. (1988) and the model outcome for mother before birth at steady-state.

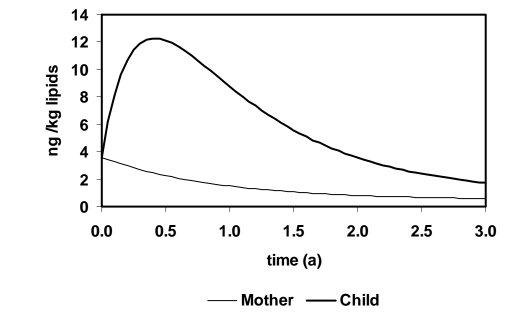
777

778 Figure 7. Bioaccumulation factors (related to lipid content) for human milk for 7 neutral organic compounds collected from literature (Lit) compared to the regression by Travis et 779 al. (1988) and the model outcome for t=0 (model t=0, at birth) and t=1 year (model t=1). 780







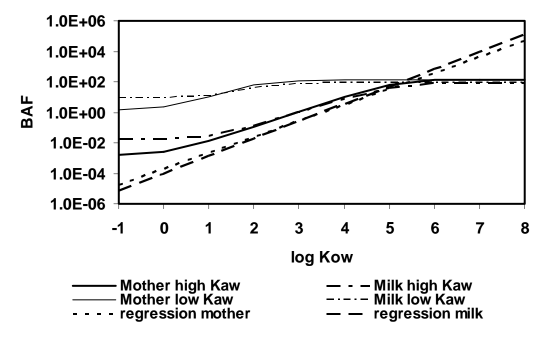


**Figure 2** 

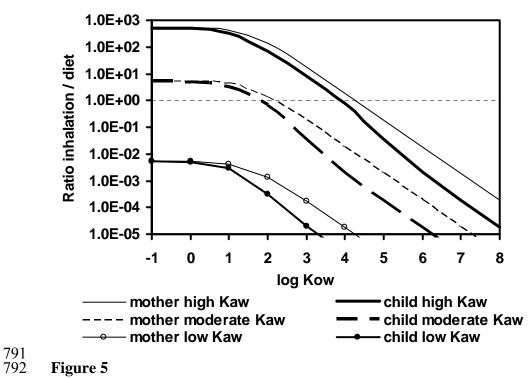


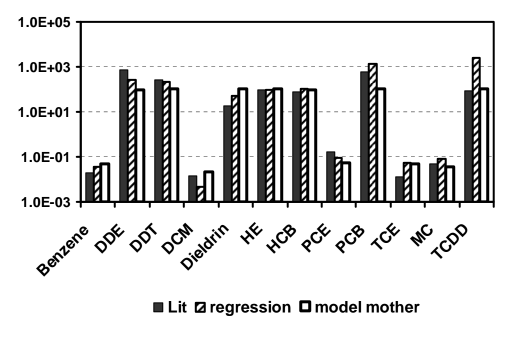


**Figure 3** 



**Figure 4** 





**Figure 6** 

