A Multi-Individual Pharmacokinetic Model Framework for Interpreting Time Trends of Persistent Chemicals in Human Populations: Application to a Postban Situation

Roland Ritter, Martin Scheringer, Matthew MacLeod, Urs Schenker, and Konrad Hungerbühler

Safety and Environmental Technology Group, ETH Zurich, Zurich, Switzerland

BACKGROUND: Human milk and blood are monitored to detect time trends of persistent organic pollutants (POPs) in humans. It is current practice to use log-linear regression to fit time series of averaged cross-sectional biomonitoring data, here referred to as cross-sectional trend data (CSTD).

OBJECTIVE: The goals of our study are to clarify the interpretation of half-lives derived from fitting exponential functions to declining CSTD and to provide a method of estimating human elimination half-lives from CSTD collected in a postban situation.

METHODS: We developed a multi-individual pharmacokinetic model framework and present analytical solutions for a postban period. For this case, the framework quantitatively describes the relationships among the half-life for reduction of body burdens of POPs derived from CSTD, the half-life describing decline in daily intake, and the half-life of elimination from the human body.

RESULTS: The half-life derived from exponential fitting of CSTD collected under postban conditions describes the exposure trend and is independent of human elimination kinetics. We use a case study of DDT (dichlorodiphenyltrichloroethane) to show that CSTD can be combined with exposure data obtained from total diet studies to estimate elimination kinetics of POPs for humans under background exposure conditions.

CONCLUSIONS: CSTD provide quantitative information about trends in human exposure and can be combined with exposure studies to estimate elimination kinetics. The full utility of these data has not been exploited so far. An efficient and informative monitoring strategy for banned POPs in humans would coordinate sampling of consistent sets of CSTD from young adults with total diet studies.

KEY WORDS: biomonitoring, DDT, exposure science, modeling, persistent organic pollutants.

Environ Health Perspect 117:1280–1286 (2009). doi:10.1289/ehp.0900648 available via http://dx.doi.org/ [Online 1 May 2009]

The Stockholm Convention on Persistent Organic Pollutants (POPs) entered into force in 2004 and mandates restriction and eventual elimination of selected chemicals at an international level to protect human health and the environment [United Nations Environment Programme (UNEP) 2009]. Article 16 of the convention requires an effectiveness evaluation of measures taken to reduce emissions of POPs. To this end, a global monitoring plan is currently being developed with the goal of detecting trends in levels of POPs in air, human milk, and human blood over time (UNEP 2007). The prospective character of this global monitoring plan will make it possible to control variables such as age and, for females, parity of donor individuals.

So far, long-term trends of POPs in humans have often been evaluated retrospectively by collecting several sets of cross-sectional data (CSD), each describing levels in human tissue in different individuals at one point in time. Summary statistics such as arithmetic means and medians of CSD obtained at different times are used to assemble time series from which a trend can be derived. Here, we refer to time trends derived from CSD as “cross-sectional trend data” (CSTD). Declining trends in CSTD typically occur after use and production of a chemical has been restricted or banned, and are often fitted with exponential functions (Craan and Haines 1998; Erdogan et al. 2004; Jaraczewska et al. 2006; Minh et al. 2004; Noergrohati et al. 1992; Norén and Meironytė 2000; Smith 1999). Exponential fitting, that is, log-linear regression, is also a method proposed in the global monitoring plan (UNEP 2007).

Presently, no consistent nomenclature exists to describe this CSTD-based half-life or its corresponding rate constant. Noergrohati et al. (1992) used the expression “disappearance rate constant,” Norén and Meironytė (2000) used the term “decline half-time,” Sjödin et al. (2004) used “population half-life,” and Smith (1999) used “population level half-life” to distinguish it from the human elimination half-life. CSTD-derived half-lives have also been referred to as “human half-life” (Woodruff et al. 1994), “half-life in humans” (Al-Saleh et al. 2003; Ennaceur et al. 2007; Solomon and Weiss 2002; Wolff 1999), and “half-life in the human body” (Cocco et al. 2000).

Two factors have been reported in the literature as influencing the CSTD-based half-life. The first is the trend of exposure or intake. It is intuitively apparent that a declining trend in exposure will result in a declining trend in CSTD. Accordingly, CSTD studies of human blood or milk are used as markers of exposure and to track reductions in exposure (UNEP 2007; World Health Organization 2007). The second factor that has been reported to influence CSTD-based half-lives is the rate of elimination of a substance from the body by all possible pathways, which we refer to as the “human elimination half-life.” Currently, however, the influence of exposure trend and elimination kinetics on CSTD-derived trends is unclear, and a range of different interpretations exists in the literature. For instance, the CSTD-based half-life has been interpreted to reflect both elimination kinetics and intake trend (Sjödin et al. 2004), to be consistent with the half-life derived from sequential measurements in individuals (Wolff 1999), or to be confounded by ongoing exposure (Minh et al. 2004). Both a single-individual pharmacokinetic (PK) model (Ayyward and Hays 2002) and a multi-individual PK model (Lorber 2002) have been applied to describe the trend observed in CSTD. This variability in interpretations reflects the lack of a mechanistic description of the CSTD-based half-life and the lack of standard nomenclature.

In contrast to CSTD, which represent the population level, the role of the time trend of intake and of elimination kinetics is well established for single individuals. Concentration trends in individuals, which are referred to as “longitudinal data” (LD) (Te et al. 2003), have been mechanistically described by single-individual PK models of different complexity (van der Molen et al. 1996; Verner et al. 2008). In LD, the trend in body burden is determined by both the intake trend and elimination kinetics. As a consequence, background intake is a possible confounding factor in estimates of the human elimination half-life from LD when complete isolation from ongoing exposure is not feasible (Shirai and Kissel 1996).

Address correspondence to M. Scheringer, Safety and Environmental Technology Group, ETH Zurich, Wolfgang-Pauli-Str. 10, CH-8093 Zurich, Switzerland. Telephone: 41-44-632-3062. Fax: 41-44-632-1189. E-mail: scheringer@chem.ethz.ch

Supplemental Material is available online (doi:10.1289/ehp.0900648.S1 via http://dx.doi.org/).

We thank M. Aune for providing us with summary statistics of reported data.

This project was funded by the Swiss Federal Office for the Environment.

The authors declare they have no competing financial interests.

Received 2 February 2009; accepted 1 May 2009.
The present study has two major goals. The first is to clarify the interpretation of half-lives derived from CSTD fitted with an exponential function by using a transparent and mathematically explicit model of the relationships among a) CSTD-based half-life, b) time trend of intake, and c) the human elimination half-life, in a postban period. The second goal is to provide a new approach to estimating human elimination half-lives; this approach uses CSTD that represent the adult population under background exposure conditions in combination with exposure studies.

Methods

To derive our multi-individual PK framework, we first formulated a one-compartment PK model for a single individual with a time-dependent intake function, \( I(t) \), which is assumed to decline exponentially in a post-ban phase with a first-order rate constant \( k_{dec} \) (years\(^{-1}\)):

\[
I(t) = I_0 \times e^{-k_{dec} \times t}, \tag{1}
\]

where \( t \) (years) is time (starting at \( t_0 \), where \( t_0 = 0 \) in the following equations), and \( I_0 \) is the intake (nanograms per person per day) at \( t_0 \). This intake function is shown in Figure 1A.

Other intake functions can also be used if the model is solved numerically [see Supplemental Material (doi:10.1289/ehp.0900648.S1)].

In most cases, human exposure to POPs occurs mainly through dietary intake (Darnerud et al. 2006). In this context, we use the term “intake” as synonymous with background exposure. The one-compartment PK model for an individual born at \( t_0 \) is represented by the following first-order differential equation:

\[
dC(t) \over dt = -k_{elim} \times C(t) + U \times E_a \times I_0 \times e^{-k_{dec} \times t} \over bw \times f_{lipid}, \tag{2}
\]

where \( C(t) \) is the concentration (nanograms per gram lipid) in the body on a lipid-normalized basis (also referred to as body burden) as a function of time, \( k_{elim} \) (years\(^{-1}\)) is the first-order rate constant for elimination of contaminant from the human body, \( bw \) is body weight in kilograms, \( f_{lipid} \) (dimensionless) is the lipid fraction of the human body, and \( U \) is a unit conversion factor (days \times year\(^{-1}\) \times kg \times g\(^{-1}\)). \( E_a \) (dimensionless) describes the fraction of chemical absorbed (i.e., intake \( \times E_a \) = uptake) and is set at 0.9 (Moser and McLaughlin 2001). In some contexts, it is more common to use a half-life rather than a rate constant. The human elimination half-life follows from the rate constant as

\[
\frac{1}{k_{elim}} \text{ (years)} = \ln(2) / k_{elim}. \tag{3}
\]

Analogously, \( \frac{1}{k_{dec}} \text{ (years)} = \ln(2) / k_{dec} \) denotes the half-life describing the decline of intake. Integrating Equation 2 with boundary condition \( C(t_0) = 0 \) yields a function describing the body burden as a function of time for an individual born at \( t_0 \) for \( k_{dec} = k_{elim} \):

\[
C(t) = \frac{U \times E_a \times I_0}{(k_{elim} - k_{dec}) \times bw \times f_{lipid}} \times (e^{-k_{dec} \times t} - e^{-k_{elim} \times t}). \tag{3}
\]

For the derivation of this solution and the solution for the special case \( k_{dec} = k_{elim} \), see Supplemental Material (doi:10.1289/ehp.0900648.S1). This equation can be generalized to a family of equations for individuals born at \( t_{birth} \) by, first, substituting \( I_0 \) the intake at time \( t_0 \), by \( I_{birth} \) the intake at the time of birth of each individual:

\[
I_{birth} = I_0 \times e^{-k_{dec} \times t_{birth}}, \tag{4}
\]

and second, introducing a variable \( t^{pge} \) that describes the age of an individual born at \( t_{birth} \) and is given by \( t^{pge} = t - t_{birth} \). The function \( C(t) \) from Equation 3 is then generalized to a function in two variables, \( t_{birth} \) and \( t^{pge} \):

\[
C(t_{birth}, t^{pge}) = \frac{U \times E_a}{(k_{elim} - k_{dec}) \times bw \times f_{lipid}} \times I_{birth} \times (e^{-k_{dec} \times t^{pge}} - e^{-k_{elim} \times t^{pge}})
\]

\[
= \frac{U \times E_a}{(k_{elim} - k_{dec}) \times bw \times f_{lipid}} \times I_{birth} \times e^{k_{dec} \times \frac{t_{birth}}{t^{pge}}} \times (e^{-k_{dec} \times t^{pge}} - e^{-k_{elim} \times t^{pge}}). \tag{5}
\]

This general equation can be used to explain three types of concentration–time relationships. The first is the time course of LD for an individual that enters the population...
shown for early life years as dashed lines in Figures 1–3 should not be interpreted as estimates of actual concentration–time courses; these parts of the curves are plotted for illustrative purposes only.

The second relationship derived from Equation 5 is the concentration time course observed in CSTD. CSTD represent a time trend in data taken from groups of individuals at different times. Each data point in the time trend represents the arithmetic mean or median of measurements in several individuals; that is, each of the data points in a set of CSTD is representative of the cohort sampled in the underlying cross-sectional study. To be comparable, all CSD sets that are used to build the CSTD should represent a population with the same or at least similar characteristics such as age, body weight, and parity. We refer to empirical CSTD that have been derived from similar CSD sets as “consistent CSTD.” We define \( t^{\text{ref}} \) as the constant “characteristic age” that is representative of the cohorts measured in all CSD sets included in a set of consistent CSTD. In our framework, each set of CSTD is modeled as an average individual representing a specific birth cohort born at \( t^{\text{birth}} \) and measured at age \( t^{\text{ref}} \). With constant \( t^{\text{ref}} \) we can derive from Equation 5 an explicit expression for CSTD as function of \( t^{\text{birth}} \):

\[
C^{\text{CSTD}}(t^{\text{birth}}) = \frac{U \times E_0}{(k_{\text{elim}} - k_{\text{dec}}) \times \text{bw} \times \text{f lipid}} \times I_0 \times e^{-k_{\text{dec}} \times \text{bw} \times \text{f lipid} \times t} \times (1 - e^{-(k_{\text{elim}} - k_{\text{dec}}) \times \text{bw} \times \text{f lipid} \times t^{\text{ref}}})
\]

with \( K_{\text{CSTD}} \) as a constant independent of \( t^{\text{ref}} \). Equation 7 is illustrated in Figure 1B.

Third and finally, Equation 5 can be used to derive an explicit relationship between concentration and age in CSTD at a specified time. If we recall that \( t = t^{\text{birth}} + t^{\text{ref}} \) and, thus, \( t^{\text{birth}} = t - t^{\text{ref}} \), replacing \( t^{\text{birth}} \) by \( t - t^{\text{ref}} \) in Equation 5 yields

\[
C(t, t^{\text{ref}}) = \frac{U \times E_0}{(k_{\text{elim}} - k_{\text{dec}}) \times \text{bw} \times \text{f lipid}} \times I_0 \times e^{-k_{\text{dec}} \times \text{bw} \times \text{f lipid} \times t} \times (1 - e^{-(k_{\text{elim}} - k_{\text{dec}}) \times \text{bw} \times \text{f lipid} \times t^{\text{ref}}})
\]

In a set of CSD, the time \( t \) at which the CSD were measured is constant, here denoted by \( t_{\text{meas}} \). Equation 8 can then be simplified to an expression for concentration as function of age:

\[
C^{\text{CSD}}(t^{\text{ref}}) = \frac{U \times E_0}{(k_{\text{elim}} - k_{\text{dec}}) \times \text{bw} \times \text{f lipid}} \times I_0 \times e^{-k_{\text{dec}} \times \text{bw} \times \text{f lipid} \times t^{\text{ref}}} \times (1 - e^{-(k_{\text{elim}} - k_{\text{dec}}) \times \text{bw} \times \text{f lipid} \times t_{\text{meas}}})
\]

with \( K_{\text{CSD}} \) as a constant independent of \( t^{\text{ref}} \). Equation 9 is illustrated in Figure 1C.

**Results**

**Properties of modeled CSTD.** Equation 7 provides two new insights into the exponential decay function that has been fitted to CSTD in numerous publications. First, Equation 7 demonstrates that in a postban situation the half-life derived from exponential fitting of CSTD is equal to the half-life of decline in intake, \( t^{\text{elim}} \), and is completely independent of human elimination kinetics. Second, the initial value of the exponential function, \( K_{\text{CSTD}} \), is now mechanistically characterized, which reveals the influence of specific PK parameters on CSTD. These two insights provide the basis for a new methodology to estimate a human elimination half-life, \( t^{\text{elim}} \), that is representative of the general adult population under background exposure conditions.
Figure 2A illustrates the influence of changes in $k_{elim}$ and $p_{D}$ on modeled CSTD. A change in $k_{elim}$ causes a parallel shift of the logarithmic CSTD function. As can be deduced from Equation 7, other parameters such as $I_{0}$, $b_{w}$, $E_{a}$, and $f_{lipid}$ will also change the value for $k_{CSTD}$ and thus cause a parallel shift of the CSTD function. However, as stated above, the slope of the CSTD function is solely determined by the rate constant of the intake trend, $k_{dec}$. Figure 2A also shows that chemicals with slower elimination show larger differences in concentrations measured in individuals of different ages that may be included in CSD. Hence, the slower the elimination kinetics of a chemical, the more important it is to include only empirical CSD with consistent age structure in the time trend analysis.

The relationship of body burden and age is explicitly described by Equation 9, which is illustrated in Figure 2B. As shown in the Supplemental Material (doi:10.1289/ehp.0900648.S1), the relationship between concentration and age is only linear if $k_{dec} = k_{dec}$; otherwise, the relationship is described by either a convex or concave curve, as has been reported in earlier modeling studies (Alcock et al. 2000; Moser and McLachlan 2002).

**Model application: a case study of DDT (dichlorodiphenyltrichloroethane).** Besides providing an analytical model to clarify the interpretation of the CSTD-based half-life, the second goal of this study is to employ the model to obtain information on human elimination kinetics from CSTD. In particular, we estimated human elimination half-lives of $p,p'$-DDT and $p,p'$-dichlorodiphenyl dichloroethylene ($p,p'$-DDE) by applying Equation 7 to CSTD from Sweden (Glynn et al. 2007). We present a five-step procedure, graphically illustrated in Figure 3, that has been implemented in an MS Excel spreadsheet (Microsoft Corporation, Redmond, WA, USA) that is available for download (ETH Zurich, Safety and Environmental Technology Group 2009).

- **Step 1:** Collect and assemble CSTD, including only individuals whose life history (or at least the major part of it) falls in the postban period [i.e., after 1970 for DDT in Sweden (Hayes 1969)].
- **Step 2:** Estimate $k_{dec}$ from the CSTD by log-linear regression.
- **Step 3:** Use at least one empirical intake estimate, such as data from a total diet study, in combination with the slope $k_{dec}$ from step 2 to describe the time trend of the intake as in Equation 1, again on a log-linear scale.
- **Step 4:** Extrapolate the function $I(t)$ from step 3 to the daily intake at $t_0$, $I_0$.
- **Step 5:** Estimate the human elimination rate constant, $k_{elim}$, by fitting Equation 7 to the empirical CSTD. Because $k_{dec}$ and $I_0$ have been estimated from the slope of the CSTD and the total diet study in steps 2 and 4, $k_{elim}$ is the only fitting parameter in Equation 7.

In principle, $k_{dec}$ could be derived from total diet studies rather than from CSTD. However, comprehensive sets of CSTD will become available from biomonitoring programs such as the National Health and Nutrition Examination Survey (NHANES) [Centers for Disease Control and Prevention (CDC) 2009] and the effectiveness evaluation of the Stockholm Convention, which will provide a consistent basis for estimating $k_{dec}$ of various chemicals in many parts of the world. In addition, at least a few, ideally several, total diet studies for the same population groups are needed to provide information on $I_0$; if these studies also yield an estimate of $k_{dec}$, this can be used to cross-check the CSTD-based estimate.

Figure 3 illustrates the steps of this procedure with DDT and DDE data from Sweden. Step 1 yields $I_{0}$ values of 6.3 years and 8.8 years for $p,p'$-DDT and $p,p'$-DDE, respectively. We took the intake estimates needed for steps 3 and 4 (open and solid circles in Figure 3A) from two Swedish market-based studies from 1999 (Danneur et al. 2006) and 2005 (Ankarberg et al. 2006). Because only the sum of DDTs ($\Sigma$DDT) and $p,p'$-DDE were reported explicitly, we estimated $p,p'$-DDT intakes as 25% of $p,p'$-DDE intakes according to findings from two dietary-intake studies from Canada (Rawn et al. 2004) and Greenland from the same time (Deutch et al. 2006). To assess the plausibility of the intake trends for $p,p'$-DDE and $p,p'$-DDT extrapolated in steps 3 and 4 (solid and dashed lines in Figure 3A), we collected additional empirical intake estimates that were available only for

![Figure 3](image-url)

**Figure 3.** Illustration of the five-step procedure for estimating human elimination half-lives: results for $p,p'$-DDE and $p,p'$-DDT are based on empirical data from Sweden (Ankarberg et al. 2006; Danneur et al. 2006; Glynn et al. 2007). (A) Modeled and empirical daily intakes used in steps 3 and 4 of the procedure. (B) Modeled and empirical CSTD used for steps 1, 2, and 5 of the procedure. Each circle represents the average of one set of CSD. See “Model application: a case study of DDT (dichlorodiphenyltrichloroethane)” for a detailed description of the five-step procedure. For additional information, see Supplemental Material, Tables S1 and S2 (doi:10.1289/ehp.0900648.S1).
\[ \Sigma \text{DDT (Vaz 1995). To compare them with the modeled intake curves for } p,p'-\text{DDE and} \ p,p'-\text{DDT (dotted line in Figure 3A). Modeled and empirical intake trends of } \Sigma \text{DDT are in good agreement. Modeled intake is slightly below the empirical } \Sigma \text{DDT data possibly because modeled } \Sigma \text{DDT includes only} \ p,p'-\text{DDE and} \ p,p'-\text{DDT, whereas empirical } \Sigma \text{DDT also includes dichlorodiphenyldichloro-} \ethane (\text{DDD}) \text{ and} \ \alpha,\beta'-\text{isomers.}

\]

Using the five-step procedure, we obtained estimates of the human elimination half-lives, \( t_{1/2} \), of 6.2 years for \( p,p'-\text{DDE} \) and 2.2 years for \( p,p'-\text{DDT} \). Our estimate for \( p,p'-\text{DDE} \) is slightly lower than estimates from sequential measurements in individuals (i.e., LD). Wolff et al. (2000) obtained a median elimination half-life of 8.6 years. For \( p,p'-\text{DDT} \) Morgan and Roan (1972) derived an elimination half-life of 1.25 years from measurements in three individuals who had received high doses of 5–20 mg/day. This compares well with our estimate of 2.2 years for individuals exposed to background levels. A slightly faster elimination in the highly dosed individuals is expected because of the concentration dependency of the elimination half-life (Leung et al. 2007). We conducted a second case study using data from the United Kingdom (see Supplemental Material, Table S3 [doi:10.1289/ehp.0900648.S1]). Elimination half-lives of 7.6 years and 2.1 years for \( p,p'-\text{DDE} \) and \( p,p'-\text{DDT} \), respectively, are in very good agreement with the estimates from the Swedish data.

From the most general equation of our PK framework, Equation 5, we have also derived an explicit equation for the body burden measured in CSD as a function of age. This equation, Equation 5, predicts body burdens of chemicals with relatively short elimination half-lives to be less dependent on age. This is consistent with empirical data for DDT, DDE, and different polychlorinated biphenyls (PCBs) (Greve and Vanzoonen 1990; Thomas et al. 2006). For a more detailed description of the properties of Equation 9, see Supplemental Material, Figure S1 and Equation S26 (doi:10.1289/ehp.0900648.S1).

**Discussion**

Our analysis provides a transparent interpretation of the half-lives derived from exponential fitting of CSTD. We show that in the postban phase these half-lives reflect a mechanistically defined quantity—the decline of intake by the population—and are independent of human elimination kinetics. To identify CSTD-based half-life estimates from a postban phase unambiguously, we suggest they be referred to as “population exposure half-life.”

In cases where individuals included in the CSTD have spent a significant fraction of their lifetime in a preban period, declining trends may be observed in the CSTD that can also be reasonably fitted with an exponential function (Craan and Haines 1998; Norén and Meironyté 2000). However, in such cases the half-life derived from the empirical fit does not reflect a mechanistically clearly defined quantity. Rather, it reflects influences of the intake trend, which in this case also includes preban intake, the elimination half-life, and the characteristic age of the sampled population (see Supplemental Material, Figure S2 [doi:10.1289/ehp.0900648.S1]). Under any exposure conditions, CSTD-derived half-lives are specific to the characteristics of the sampled population and the exposure trend, and not individual humans. Therefore, terms including “human” should be avoided when referring to half-lives derived from CSTD.

For a postban period, our analysis clarifies the relationships among a) CSTD-based half-life, \( k \) (time trend of intake, and c) the human elimination half-life. In this case, the trend in CSTD directly reflects the trend in exposure; therefore, exponential fitting of CSTD implies the assumption that intake is declining exponentially. For DDT, this is supported by total diet studies presented in Figure 3 and elsewhere (Morisawa et al. 2002). Empirical evidence shows that the assumption of first-order declining intake is also reasonable for other chemicals such as dioxins and PCBs (De Mul et al. 2008). Furthermore, exponential fitting implicitly assumes that the sets of CSD used to derive the CSTD have been obtained from similar groups of individuals. This implicit assumption allows us to model CSTD using a one-compartment PK model that represents individuals with the same average characteristics but born in different years. As a consequence, the solutions to the PK model in Figure 1A do not provide information about interindividual differences, nor do they account for complete lifetime exposure profiles.

Physiologic factors are implemented as constant values in the model, that is, as independent of age. This simplification makes it possible to solve the model analytically; it is supported by two findings. First, background intake and body weight are generally assumed to be proportional in adults (i.e., intake is expressed in units of nanograms per kilogram of body weight per day). Under this assumption, a one-compartment model parameterized for average adults, as presented in Equation 2, is equivalent to a one-compartment model parameterized with age-dependent body weight and background intake proportional to body weight [see Supplemental Material (doi:10.1289/ehp.0900648.S1)]. Second, large changes in daily intake, as they occur in early life years (e.g., due to breast-feeding), also have large effects on the body burden (Moser and Mclachlan 2002). However, these concentration peaks level out no later than 20 years of age (Kreuzer et al. 1997; Verner et al. 2008). In other words, predictions of concentrations in average adults as they are represented in CSTD are not strongly influenced by differences in body burdens occurring during early life years due to different breast-feeding regimes (Lober 2002). If required, the conceptual framework developed here could also be set up with age-dependent physiologic parameters and more general intake functions. In such cases, numerical integration of the model will be necessary.

**Scope of the model framework.** Our modeling framework can also be applied to cases that are different from the DDT example. First, the intake function, \( I(t) \), can represent any intake pathway, not only food. In addition, when the model is solved numerically, \( I(t) \) can have any shape [see Supplemental Material, Figure S2 (doi:10.1289/ehp.0900648.S1)]. This means that the model framework is not limited to postban cases. Second, for chemicals stored in parts of the body other than lipids, \( bw \times f_{\text{liquid}} \) in Equation 2 can be replaced by a description of the actual storage medium, such as the volume of distribution for perfluorinated chemicals. Finally, application of the framework is not limited to persistent chemicals with slow elimination. However, for chemicals that are eliminated quickly, \( k_{\text{lim}} \) can be more readily derived from LD collected from single individuals (Koch and Angerer 2007). Our modeling framework is therefore most powerful when applied to chemicals with slow elimination kinetics such as DDT for two reasons. First, LD studies aiming to estimate \( k_{\text{lim}} \) of slowly eliminating chemicals are difficult to perform because subjects must be tracked over several years, and fasting to eliminate background exposure is not feasible. Second, as demonstrated in Figure 2A, CSTD of slowly eliminating chemicals can only be described by a multi-individual PK model as presented here.

**Model application and performance.** For many persistent chemicals, information about human elimination half-lives is scarce (Wolff 1999), and available data exhibit a high degree of uncertainty. For instance, a review of PCB elimination half-lives found reported values ranging from < 1 year to virtually infinity even within a specific congener group (Shirai and Kiesel 1996). Different methods have been applied to estimate human elimination half-lives. A common method is to use sequential measurements in individuals who experienced high exposure for a limited time period (e.g., occupational exposure, accident). High-exposure cases are chosen in LD studies to mitigate the problem of ongoing background exposure (Grandjean et al. 2008). However, half-lives derived from such cases may be not representative for the general population because the rate of elimination...
has been shown to depend, to some extent, on the absolute level of body burden (Leung et al. 2002). An alternative approach to the problem of confounding through ongoing exposure is to account for it in the PK model used to estimate the elimination half-life. Such an expression has been presented for a constant intake (steady-state situation) (Shirai and Kissel 1996). However, because intake often changes over time, integration of the actual intake trend (rather than assuming steady state) would be desirable, but information about intake of measured individuals is rarely available. Here we have shown how time trend information present in CSTD can be combined with information from total diet studies to determine the intake trend and the whole body elimination half-life for the general adult population. To our knowledge, our method represents the first approach that uses CSTD to estimate human elimination half-lives. The further advantage is that CSTD are available in many countries and represent a broad empirical basis, whereas LD-inferred half-lives are often based on very few (often only two) measurement points (Phillips 1989) with a limited number of subjects available.

In two DDT case studies, we employed the model to estimate human elimination half-lives for $p,p'$-DDE and $p,p'$-DDT using intake estimates and CSTD from Sweden and the United Kingdom as empirical data. Results obtained from both CSTD sets are consistent with each other and with estimates from LD.

Data quality and availability. The CSTD from Sweden presented in Figure 3 represent only primiparous women with a median age of 28.8 years from the same region (Uppsala County). Hence, they exhibit a relatively high level of consistency. This is because the recurrent measurements had been planned prospectively aiming to establish time trends. In contrast, most studies in the literature that present CSTD-based half-lives have been assembled retrospectively. In many cases only summary statistics are reported, which makes it difficult to compare different sets of CSTD. As a result, confounding effects from different age structures and other differences between the studies may occur. In particular, an inconsistent age structure would introduce large uncertainties in CSTD-derived half-lives of slowly eliminated chemicals like POPs. Therefore, in the future, cross-sectional studies designed to evaluate time trends of POPs in populations should be conducted in such a way that they yield consistent sets of CSTD. Measuring levels in blood and human milk every year in sufficiently large groups of young adults that meet clearly defined physiologic and lifestyle criteria is a particularly effective monitoring strategy. Preferably, cohorts of young adults should be monitored because they meet post-natal conditions first. Total diet studies, which are already performed in many countries on a regular basis, should be coordinated with these biomonitoring programs to ensure optimal use of resources.

Conclusions

The present study demonstrates that CSTD used in combination with exposure data can provide quantitative information about human exposure trends and elimination kinetics that has not been exploited so far. We have demonstrated that, under postnatal conditions, the CSTD-based half-life is equivalent to the population exposure half-life and that the problem of scarcity and uncertainty in estimates of human elimination kinetics of POPs can be reduced by using CSTD in combination with exposure data to derive estimates that represent the adult population under background exposure conditions. Accurate estimates of exposure trends and human elimination kinetics are only possible when the model is applied to consistent CSTD; the acquisition of consistent daily intake data of the monitoring programs. Programs such as NHANES and the forthcoming effectiveness evaluation of the Stockholm Convention offer opportunities to apply the conceptual framework presented here to improve the interpretation of biomonitoring data.

References

Alcock RE, Sweetman AJ, Juan CY, Jones KC. 2000. A generic model of human lifetime exposure to persistent organic pollutants: application and PCB-101. Environ Pollut 110(2):253–265.

Al-Saleh I, Shinwari N, Basile P, El-Douh I, Al-Zahrani M, Al-Shanshouri M, et al. 2003. DDT and its metabolites in breast milk from two regions in Saudi Arabia. J Occup Environ Med 45(2):197–203.

Ankargren E, Törnqvist A, Darnerud PO, Aune M, Pettersson-Grawe K, Nordqvist Y, et al. 2006. Dietary Intake Estimations of Persistent Organic Pollutants (Dioxin, PCB, PBDE, Chlorinated Paraffins, Phenolic Compounds) Based on Swedish Market Basket Data and Levels of Mercury in Fish. Contract 215/0506. Uppsala, Sweden: Swedish Environmental Protection Agency.

Ayward LL, Hays SM. 2002. Temporal trends in human total body burden: decreases over three decades and implications for exposure levels. J Expo Anal Environ Epidemiol 12(5):319–328.

CDC Centers for Disease Control and Prevention. 2009. National Health and Nutrition Examination Survey Homepage. Available: http://www.cdc.gov/nchs/nhanes.htm [accessed 31 March 2009].

Corpet P, Kazerouni N, Zehm SH. 2000. Cancer mortality and environmental exposure to DDE in the United States. Environ Health Perspect 108:1–4.

Craan AG, Haines DA. 1998. Twenty-five years of surveillance for contaminants in human breast milk. Arch Environ Contam Toxicol 35(2):172–179.

Darnerud PO, Atuma S, Aune M, Petersson-Grawe K, Nordqvist Y, et al. 2006. Dietary Intake Estimations of Persistent Organic Pollutants (Dioxin, PCB, PBDE, Chlorinated Paraffins, Phenolic Compounds) Based on Swedish Market Basket Data and Levels of Mercury in Fish. Contract 215/0506. Uppsala, Sweden: Swedish Environmental Protection Agency.

Deutche B, Dyerberg J, Pedersen HS, Asmund G, Moller P, Hansen JC. 2006. Dietary composition and contaminants in north Greenland, in the 1970s and 2004. Sci Total Environ 370(2–3):372–381.

Einarson DR, Gaudoura N, Drsir MR. 2007. Organochlorine pesticide residues in human milk of mothers living in northern Tunisia. Bull Environ Contam Toxicol 78(5):325–329.

Erdogul D, Covaci A, Kurtul N, Schepens P. 2007. Levels of organochlorinated persistent pollutants in human milk from Kahramanmaras region, Turkey. Environ Int 30(3):659–666.

ETH Zurich, Safety and Environmental Technology Group. 2009. CSTD Halflife Tool. Available: http://sust-chem.ethz. ch/downloads [accessed 25 June 2009].

Glynn A, Aune M, Lignell S, Ankargren E, Darnerud PO. 2007. Polychlorinated Dibenzo-p-dioxins (PCDDs) and Dibenzofurans (PCDFs). Polychlorinated Biphenyls (PCBs). Chlorinated Peptidol and Brominated Fire Retardants in Mother’s Milk from Primiparae Women in Uppsala County, Sweden—Levels and Trends 1996–2006. Available: http://www.imm.ki.se/Datavard/Rapporter/ Sakraptorsrund6.pdf [accessed 22 January 2009].

Graeve PA, Vanzoopen P. 1990. Organochlorine pesticides and PCBs in tissues from Dutch citizens (1968–1988). Int J Environ Anal Chem 38(2):265–277.

Hays WJ. 1989. Sweden bans DDT. Arch Environ Health 186(6):872.

Jaraczewska K, Lulek J, Covaci A, Voorspoels S, Kaluba-Skotarczak A, Drees K, et al. 2006. Distribution of polychlorinated biphenyls, polybrominated diphenyl ethers in human umbilical cord serum, maternal serum and milk from Wielkopolska region, Poland. Sci Total Environ 372(1–2):41–51.

Koch HM, Angerer J. 2007. Diisoo-nonylmonophthalate (DIMP) metabolites in human urine after a single oral dose of deuterium-labelled DIMP. Int J Hyg Environ Health 210(1):9–19.

Kreufer PE, Csanady GA, Baur C, Kessler W, Papke O, Greim H, et al. 1997. 2,3,7,8-TetraChlorodibenzo-p-dioxin (TCDD) and congeners in infants. A toxicokinetic model of human lifetime body burden by TCDL with special emphasis on its uptake by nutrition. Arch Toxicol 71(6):385–400.

Leung HW, Kerger BD, Paustenbach DJ, Ryan JJ, Masuda Y. 2007. Concentration and age-dependent elimination kinetics of polychlorinated dibenzo-furans in Yucheng and Yusho patients. Toxicol Ind Health 23(3):493–501.

Lorber M. 2002. A pharmacoKinematic model for estimating exposure of Americans to dioxin-like compounds in the past, present, and future. Sci Total Environ 288(1):215–39.

Mirin NH, Somey A, Miin TB, Kuniiue T, Iwata H, Watanabe M, et al. 2004. Persistent organochlorine residues in human breast milk from Hanoi and Hochiminh City, Vietnam: contamination, accumulation kinetics and risk assessment for infants. Environ Pollut 129(3):431–441.

Morgan DP, Rean CC. 1972. Long-term DDT storage from human body fat. Nature 238(S811):221–222.

Morisawa S, Kato A, Yoneda M, Shima Y. 2002. The dynamic performances of DDTs in the environment and Japanese exposure to them: a historical perspective after the ban. Risk Anal 22(3):245–263.

Moser GA, McLachlan MS. 2001. The influence of dietary concentration on the absorption and excretion of persistent lipophilic organic pollutants in the human intestinal tract. Chemosphere 45(3):201–211.

Moser GA, McLachlan MS. 2002. Modeling digestive tract absorption and desorption of lipophilic organic contaminants in humans. Environ Sci Technol 36(19):3318–3325.

Noegahr S, Sardjito, Untung R, Wijaya W. 1992. Impact of DDT spraying on the residue levels in soil, chicken, fish, pond water, carp, and human-milk samples from malaria infested villages in central Java. Toxicol Environ Chem 34(2–4):237–251.

Norén K, Meironyté D. 2000. Certain organochlorine and organobromine contaminants in Swedish human milk in perspective of past 20–30 years. Chemosphere 40(2):1111–1123.

Phillips DL. 1989. Propagation of total diet studies. Estimated half-life estimates based on 2 measurements. Arch Environ Contam Toxicol 18(4):508–514.

Rawn DFF, Cao XL, Doutcet J, Davies DJ, Sun WF, Dabekw DA, et al. 2004. Canadian Total Diet Study in 1998: pesticide levels in foods from Whitehorse, Yukon, Canada, and corresponding dietary intake estimates. Food Addit Contam 21(3):232–250.

Shirai JH, Kissel JC. 1996. Uncertainty in estimated half-lives.
Organohalogen chemicals in human blood from the United Kingdom. Environ Pollut 141(1):30–41.

UNEP (United Nations Environment Programme). 2009. Stockholm Convention on Persistent Organic Pollutants. Available: http://www.pops.int (accessed 2 April 2009).

UNEP (United Nations Environment Programme). 2007. Guidance on the Global Monitoring Plan for Persistent Organic Pollutants. Available: http://www.pops.int/documents/meetings/cop_3/meetingdocs/inf14/GMP%20Guidance%20CD/Guidance.pdf (accessed 2 April 2009).

van der Molen GW, Kooijman S, Slob W. 1996. A generic toxicokinetic model for persistent lipophilic compounds in humans: an application to TCDD. Fundam Appl Toxicol 31(1):83–94.

Vaz R. 1995. Average Swedish dietary intakes of organochlorine contaminants via foods of animal origin and their relation to levels in human milk, 1975–90. Food Addit Contam 12(4):543–558.

Verner MA, Charbonneau M, Lopez-Carrillo L, Haddad S. 2008. Physiologically based pharmacokinetic modeling of persistent organic pollutants for lifetime exposure assessment: a new tool in breast cancer epidemiologic studies. Environ Health Perspect 116:886–892.

Wolff MS. 1999. Letter to the Editor. Arch Environ Contam Toxicol 36(4):504.

Wolff MS, Zeleniuch-Jacquotte A, Dubin N, Toniolo P. 2000. Risk of breast cancer and organochlorine exposure. Cancer Epidemiol Biomark Prev 9(3):271–277.

Woodruff T, Wolff MS, Davis DL, Hayward D. 1994. Organochlorine exposure estimation in the study of cancer etiology. Environ Res 65(1):132–144.

World Health Organization. 2007. Fourth WHO-Coordinated Survey of Human Milk for Persistent Organic Pollutants. Available: http://www.who.int/foodsafety/chem/PDPprotocol.pdf (accessed 30 January 2009).