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Karlsson, Maja Viktoria, Carter, Laura Jayne, Agatz, Annika orcid.org/0000-0003-3228-8822 et al. (1 more author) (2017) Novel Approach for Characterizing pH-Dependent Uptake of Ionizable Chemicals in Aquatic Organisms. Environmental science & technology. pp. 6965-6971. ISSN: 1520-5851

https://doi.org/10.1021/acs.est.7b01265

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A novel approach for characterising pH-dependent uptake of ionisable chemicals in aquatic organisms

Journal:	Environmental Science & Technology	
Manuscript ID	es-2017-012652.R2	
Manuscript Type:	Article	
Date Submitted by the Author:	24-May-2017	
Complete List of Authors:	Karlsson, Maja; University of York, Environment Department Carter, Laura; University of York, Environment Department Agatz, Annika; University of York, Environment Department Boxall, Alistair; University of York, Environment Department	

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- 1 A novel approach for characterising pH-dependent uptake of ionisable chemicals in
- 2 aquatic organisms
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Abstract

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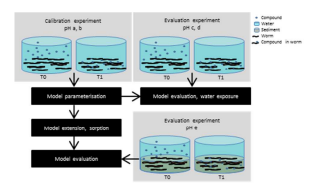
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Here, we present and evaluate a combined experimental and modelling approach for characterising the uptake of ionisable chemicals from water and sediments into aquatic organisms under different pH conditions. We illustrate and evaluate the approach for two pharmaceuticals (diclofenac and fluoxetine) and one personal care product ingredient (triclosan) for the oligochaete Lumbriculus variegatus. Initially, experimental data on the uptake of the three chemicals at two pH values were fitted using a toxicokinetic model to derive uptake and depuration constants for the neutral and ionised species of each molecule. The derived constants were then used to predict uptake from water and sediment for other pH conditions. Evaluation of predictions against corresponding experimental data showed good predictions of uptake for all test chemicals from water for different pH conditions and reasonable predictions of uptake of fluoxetine and diclofenac from a sediment. Predictions demonstrated that the level of uptake of the study chemicals, across pH ranges in European streams, could differ by up to a factor of 3035. Overall, the approach could be extremely useful for assessing internal exposure of aquatic organisms across landscapes with differing pH. This could help support better characterisation of the risks of ionisable chemicals in the aquatic environment.

Keywords

26 Lumbriculus variegatus; toxicokinetic modelling; pH; sorption; ionisable chemicals



TOC Art

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Introduction

A wide range of pharmaceuticals and ingredients used in personal care products has been
detected in natural environments across the globe. 1-5 The presence of these chemicals in the
environment has prompted concerns over potential toxic effects in non-target organisms. For
a chemical to elicit an effect in an organism, it must usually be first taken up from the
ambient environment. Understanding the internal exposure of a chemical can provide
valuable insights to inform our understanding of the effects of chemicals in organisms. The
information can also help in extrapolating from effects in standard laboratory studies to
effects across different exposure scenarios. $^{6,\ 7}$ For example, for active pharmaceutical
ingredients (APIs), it has been suggested that by understanding the internal concentrations
in organisms in the natural environment and the presence/absence of the target receptors
and pathways for the API, it may be possible to predict potential ecological effects of
pharmaceuticals based on preclinical and clinical pharmacological data that are produced in
the drug development process.8
It has been estimated that between 85 and 95 % of APIs are ionisable 9 and therefore there
is the potential that the behaviour of these chemicals in the environment may be affected by
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(Gammarus pulex and Notonecta glauca) could be based on a chemicals pH-corrected liposome-water partition coefficient (Log D_{lip-water}). Nichols, et al. ¹⁵ modelled the uptake of the weak base, diphenhydramine, into fish plasma using a model that accounts of acidification at the gill surface. The model assumed that the undissociated form of the molecule diffuses freely across the branchial epithelium. The membrane transport of the cation is estimated in a relation to the neutral form using a term varying from 0 to 1. Fu, et al. 16 proposed regression equations that estimate bioconcentration factors of acids and bases based on the octanol-water partition coefficient (K_{ow}) and the logarithmic acid dissociation constant (pKa) of a molecule. These previous studies have focused on the situation where the external and internal concentrations of the ionisable chemical are in equilibrium and have typically taken an overly simplistic approach to dealing with the ionised form of the molecules. By using a more mechanistic approach that considers the rate of uptake of the ionised and neutral species of a molecule, it may be possible to better estimate the internal exposure of an organism over time for varying pH conditions typically found in the natural environment. This could be invaluable for assessing the degree of risk of these chemicals. Here, we present a new combined experimental and modelling approach for characterising the uptake of ionisable chemicals, such as APIs, in aquatic invertebrates over time for different pH conditions. A schematic of the model underlying the approach is shown in Figure 1. The approach characterises the uptake of an ionisable chemical into an organism over time based on the fraction of the ionised and non-ionised species of a molecule in water

concentration of the compound in the sediment and the sediment sorption coefficient (K_d). This is based on the assumption that uptake is occurring only from the pore water. We illustrate and evaluate the approach for two ionisable APIs and one ionisable personal care product ingredient and the oligochaete *Lumbriculus variegatus* for water-only exposures at a range of pH values and using previously published data on uptake of the chemicals from one sediment type.

Methods

91 Underlying model

The kinetic model used is based on a first order one compartment toxicokinetic model that is used to describe the internal concentrations within an organism over time (dC_{int}/dt) [mass/volume] based on exposure medium concentrations and uptake and depuration rates (Eq. 1). Here we extend this toxicokinetic model to account for differences in uptake of the neutral and ionised forms over time (Eq. 2).

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$$dC_{int} / dt = k_{in} * C_w - k_{out} * C_{int}$$
 Equation 1

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$$dC_{int} / dt = (k_{in-ion} * (f_{ion} * C_w) + k_{in-neut} * (f_{neut} * C_w)) - k_{out} * C_{int}$$
 Equation 2

Where: C_w [mol/mL] is the concentration of the ionisable chemical in the water; C_{int} [mol/g] is the concentration of the chemical in the organism; $k_{in\text{-}ion}$ [d⁻¹] is the uptake rate constant of the ionic species of the chemical; $k_{in\text{-}neut}$ [d⁻¹] is the uptake rate constant for the neutral form of the chemical; f_{ion} [-] is the fraction of the chemical in the ionised form at the test pH; f_{neut} [-] is the fraction of the neutral form of the chemical at the test pH; and k_{out} [d⁻¹] is the depuration rate constant. The depuration rate constant is not altered in comparison to the original model because the pH within the organism is assumed to be independent of the

external pH and thus fractioning of the chemical into the ionised and neutral form is constant within the organism and the depuration rate constant is independent of pH alterations once derived. This assumption had to be made because, to our knowledge, no information is available on the internal pH of *L. variegatus* and the circumstances on if, when and to what extent this pH changes over the life span or is independent of external pH.

The uptake and depuration rates needed to parameterise the model are obtained from uptake and depuration experiments on the organism of interest. The tests need to be performed at a minimum of two pH values within the naturally occurring environmental pH ranges. One of the pH values needs to be chosen so that the test chemical is either fully or not ionised to allow the parameterisation of k_{in_ion} and k_{in_neut} in a two-point calibration. The Henderson Hasselbach Equation is used to estimate the fraction of dissociation of the study chemicals at each of the test pH values. By fitting Equation 2 to experimental data for both pH conditions, k_{in_neut} , k_{in_ion} and k_{out} can be derived. These rate constants can then be used, in conjunction with ionisation predictions from the Henderson Hasselbach Equation, in the toxicokinetic equation to predict uptake from water for other pH conditions.

If the organism of interest is a sediment-dwelling organism then concentrations in the organism can also be estimated according to concentrations in pore water of a known pH based on the sediment-water distribution coefficient K_d (L/Kg) of the chemical for the sediment of interest which can be obtained from batch sorption studies based on the OECD 106 Batch Equilibrium Method. Concentrations of test chemicals in sediment over time are used, alongside the sorption coefficient, to estimate concentrations in the sediment pore water over time using Equation 3.

$$C_{pw} = C_{sed} / ((K_d * (\%sed / \%water) * bulk density) + 1)$$

Equation 3

Where $C_{\it pw}$ and $C_{\it sed}$ are the concentrations of the chemical in pore water [mol/mL] and				
sediment [mol/g wwt], respectively, and %sed and %water are calculated based on the				
moisture content of the sediment.				
Illustration and evaluation of the approach for diclofenac, fluoxetine and triclosan				
To illustrate and test the approach, we performed studies into the uptake and depuration of				
diclofenac (a non-steroidal anti-inflammatory compound), fluoxetine (an antidepressant) and				
triclosan (an antimicrobial compound). Studies were done using water at four pH values -				
two of these being used to derive uptake and depuration constants for the neutral and ionic				
species of each molecule and two being used to test the predictive power of the approach				
for other test conditions. A previous dataset on uptake of the study chemicals from sediment				
was used to evaluate whether the method can be extended to predict uptake from the				
sediment compartment.				
Test chemicals				
Experimental studies were done using ¹⁴ C-labelled versions of the test chemicals, ranging in				
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5.5, 7 and 8.5 based on an approach recommended by the USEPA ¹⁹. The pH was not buffered or manipulated in the APW (pH 7.4) treatment. For the SRW treatments, a NaH₂PO₄ was used and the pH was maintained for the duration of the uptake and depuration phase through the addition of either 0.1 M HCl or NaOH.

Prior to the studies, the *L. variegatus* were acclimatized to the test conditions for 18 h. For

Prior to the studies, the *L. variegatus* were acclimatized to the test conditions for 18 h. For the uptake studies, animals were then exposed in groups of 10 animals, contained in 40 ml of either APW or the SRW at the different pH values, to between 3 - 12 nmol Γ^1 of test chemical for 3, 6, 12, 24 or 48 h. All test concentrations were below toxicological thresholds. For the depuration studies, groups of animals were exposed to the test chemical for 48 h after which time they were transferred to either APW or pH-adjusted SRW for 3, 6, 12, 24 or 48 h. Three replicates per time point and pH treatment were used. The study temperature was 20 ± 2 °C and the beakers were kept in the dark throughout the test to minimize potential photodegradation of the test chemical. Control beakers containing SRW and radiolabelled test chemical were used to monitor sorption to the jars. At the end of the exposure, samples of the test media were taken for chemical analysis. Exposed worms were rinsed with distilled water, blotted dry on tissue paper, weighed and then analysed.

Chemical analysis

Concentrations of the study chemicals in test media and worm extracts were determined using Liquid Scintillation Counting (LSC) using a Beckman LS 6500 LSC counter (Beckman Coulter Inc., Fullerton, USA). For the analysis of test media, 1 ml of sample was taken and placed into a 20 ml scintillation vial and 10 ml Ecoscint A scintillation cocktail (National Diagnostics) was added. For the analysis of worm samples, animals were placed in 20 ml scintillation vials, 2 ml of tissue solubilizer (Soluene®-350, Perkin Elmer, Waltham, Massachusetts) was then added and the vials were left for 24 h to allow the worm tissue to dissolve completely. Prior to scintillation counting, 10 ml of Hionic Fluor scintillation cocktail (Perkin Elmer) was added to the vials.

Samples were counted three times for 5 min. Counts were corrected for background activity by using blank controls. Counting efficiency and colour quenching were corrected using the external standard ratio method. A mass balance was performed to account for all radioactivities present in the experiments by summing the mass of chemical contained in the organism and in the test media for each treatment and timepoint.

Parameterisation and testing of the model against experimental data

For the water-only studies, for each chemical tested, there was one experiment that was conducted at a pH where the molecule was almost fully dissociated (diclofenac, pH 8.5; fluoxetine, pH 5.5) or non-dissociated (triclosan, pH 5.5). These experiments were used to fit the depuration rate constant and the uptake rate constant (k_{in_ion} for diclofenac and fluoxetine and k_{in_neut} for triclosan) for the chemical to measured internal concentrations by fixing the fraction of ionisation to either 0 or 1. The experiment with the most deviating pH (i.e., the highest variation in the fractioning of ionisation) was then used to fit the 2nd uptake rate constant keeping the prior fitted uptake rate constant and the depuration rate constant fixed and adjusting the fraction of ionisation and neutralisation. Measured pH values were used in these calculations. Modelling was conducted in OpenModel V 2.4.2. (http://openmodel.info/) using the Runge-Kulta (4th Order) ordinary differential equation method (with Monte Carlo simulations to obtain the 95% confidence interval and the Nash–Sutcliffe Efficiency calculation for goodness of fit indication where a value ≥0 shows an acceptable fit/prediction and a value <0 indicates an unacceptable fit) using the full data set and also using the minimised design method described in Carter et al. ²⁰

These fitted rate constants and the fractions of ionisation, derived from the pH measurements, in the other two experiments were then used to predict the internal concentration over time in the uptake studies performed at the other pH values of 7.0 (SRW) and 7.4 (APW) using both the full and minimised methods.

To evaluate the approach for chemical uptake from sediment, we used data from a previously reported study into the uptake and depuration of diclofenac, fluoxetine and

triclosan into *L. variegatus* from a sediment obtained from Buttercrambe in Yorkshire ²¹ (See Supporting Information Section 1 for more details). The pH of the sediment was 7.67 and the bulk density of the test sediment was estimated to be 1.16 g/mL using the approach proposed by Avnimelech, et al. ²². The sorption coefficient between water and the sediment was obtained using a batch sorption test and were found to be 4.2, 422.5 and 241.2 for diclofenac, fluoxetine and triclosan respectively. A full description of the methods used to calculate the sediment sorption coefficients is provided in the Supporting Information (Section 2). The sorption data were used to estimate concentrations of the study chemicals in pore water over the duration of the study (Eq. 3) and internal concentrations in the organisms were then estimated from the pore water concentration using the same approach as used in the water only studies (Eq. 2). Experimental observations were then compared to the predictions.

Results and discussion

Uptake and depuration experiments in Lumbriculus variegatus

The pH in the APW treatments ranged from 7.5-8.3. For the SRW, the pH of the pH 5.5 and 7 treatments remained within ± 0.3 pH units of the nominal value. For the pH 8.5 treatment, measured pH decreased by up to 1.1 pH unit during the experiment. Radioactivity in the media in the chemical controls, containing test chemical and aqueous media only, was stable for the duration of the study indicating that there was no sorption to the vessels. Losses of activity from the water phase in the beakers with organisms could be explained by uptake into the study organisms. Mass balance calculations showed recoveries of greater than 89% of the applied radioactivity in the different treatments (Supporting Information, Section 3, Table S1). No mortality was observed either in the treatments or in the controls during the uptake or depuration phase.

The uptake and depuration studies at different pH values demonstrate the importance of exposure medium pH for predicting ionisable chemical uptake into non-target organisms. At

the end of the 48 h uptake phase 47 and 37 fold differences were seen between the internal concentrations in *L. variegatus* in the highest and lowest pH treatments for diclofenac and fluoxetine, respectively whereas the fraction of ionisation only changed by 3 and 2%, respectively. These experimental results demonstrate that the uptake of ionisable chemicals can, in certain circumstances, be extremely sensitive to changes in exposure medium pH where the pKa value of a chemical falls within the environmentally relevant pH range of 2.2 – 9.8.¹⁰ The hydrophobicity and value of the chemical pKa in relation to environmental pH ranges is important – for diclofenac which is an acid with a pKa towards the lower end of typical environmental pH, large differences in uptake and toxicity might be expected across environmental pH values whereas for triclosan, which is a hydrophobic acid with a pKa towards the upper end of the environmental pH range, lower variability in uptake might be expected as the chemical will not be as extensively ionised and the neutral form would be the dominant species.

Evaluation of the modelling approach

The first order one compartment model was successfully fitted (success being indicated by Nash-Sutcliffe Efficiency values well above 0 (See Supporting Information Section 4 Table S2) to the uptake and depuration measurements for the diclofenac, fluoxetine and triclosan treatments for pH 5.5 and 8.5 (Figure 2). Resulting uptake parameters for the neutral and ionised species and the combined depuration rates of the chemicals, obtained using both the full and minimised methods, are provided in Table 1. For diclofenac and fluoxetine the uptake rate constants for the ionised form of the molecules were more than three orders of magnitude lower than the corresponding neutral form. Despite the fact that it is typically assumed that the uptake of the ionic form of a molecule is lower than the neutral form (as observed for diclofenac and fluoxetine), for triclosan, this was found not to be the case with the uptake rates for the neutral and ionised forms of the molecule being similar. The triclosan findings are similar to previous observations into the uptake of chlorinated phenols at fish

gills where uptake at different pH values was similar even though the degree of ionisation of the chlorinated phenols at the pH values studied was very different.²³

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The full and the minimised methods provided similar predictions demonstrating that the minimised approach can be used to derive uptake and depuration rate constants for the neutral and ionic species of a molecule and thus reduce the amount of experimental effort (by a factor of four) needed in studies of this type. Predictions of concentrations in L. variegatus based on the derived K_{in-neut} and K_{in-ion} values to estimate uptake from water for the other pH conditions (i.e. pH 7 and 7.4) are shown in Figure 2. Overall the model performed best for triclosan followed by fluoxetine and diclofenac. Whilst predictions, using rate constants derived using both the full and minimised approaches, accurately matched observations for triclosan (Nash-Sutcliffe Efficiency > 0.56) the model significantly underestimated internal concentrations of fluoxetine and diclofenac at pH 7.4 (Nash-Sutcliffe Efficiency < 0) and fluoxetine at pH 7.0 when rate constants using the full approach were used and significantly underestimated internal concentrations of diclofenac and fluoxetine at pH 7.4 when rate constants obtained using the minimised approach were used. Even so, predicted internal concentrations of fluoxetine concentrations, obtained using the model, were within a factor of two of experimental values while internal concentration predictions for diclofenac were within a factor of four of experimental values. Given the large observed range seen in the uptake experiments for fluoxetine (a 37 fold difference between pH 5.5 and pH 8.50) and diclofenac (a 47 fold difference between pH 5.5 and 8.5), the model predictions seem reasonable and useful for use in spatial environmental risk assessments. Comparison of the model predictions of internal concentrations of the chemicals for the sediment studies with measured concentrations in L. variegatus (Figure 3) showed that the approach worked reasonably well for diclofenac and fluoxetine with predictions being less than a factor of five lower than empirical observations. For triclosan, however, predicted concentrations in the worms were 10-15 times lower than the experimental observations.

The mismatch for all three chemicals might be explained by differences between the

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physico-chemical characteristics of the sediment pore water and the media used in the water-only studies. Previous studies with neutral organic chemicals have shown that dissolved organic carbon, which will be present in the sediment water, can enhance the diffusive mass transfer of compounds.²⁴ It is also possible that the sorption coefficients used. which were obtained using OECD-type batch sorption studies using one concentration, do not reflect the actual sorption behaviour of the chemical in the sediment-worm system. For example, sorption isotherms may deviate from linearity at high concentrations. Concentrations used in the sorption experiments were however low (15 - 22.6 nM) and within an order of magnitude of concentrations observed in the aqueous phase of the sediment uptake studies (diclofenac 21.5-23.5 nM; fluoxetine 5.30-5.82 nM; and triclosan 1.64-3.5 nM). The differences for triclosan may also be partly explained by the fact that sediment ingestion has been shown to play a role in the uptake of this chemical by L. variegatus from this sediment 20 while this has been shown to not be an important uptake route for diclofenac and fluoxetine. The impacts of pore water chemistry on both sorption and uptake and, for selected chemicals, ingestion may therefore need to be considered in the future in order to develop approaches to better assess uptake from sediments across a landscape. Finally, it is important to recognise that the studies presented here measured levels of radioactivity in the different components of the system over time. It is possible, that once accumulated, the study chemicals were metabolised to some degree. Differences in properties of the transformation products compared to the parent chemical may also contribute to the mismatch between predictions and experimental observations. Previous work we have performed into the uptake and metabolism of unlabelled compounds indicates that diclofenac is non-metabolised by the worms which fluoxetine and triclosan may be metabolised to some degree.²⁵

Implications for environmental risk assessment

The application of the combined experimental and modelling approach to characterise internal concentrations worked reasonably well for all chemicals in the water-only studies

and for diclofenac and fluoxetine in the sediment studies. The results indicate that, if data
are available on exposure medium pH and concentrations in sediment for a landscape, then
by deriving uptake and depuration rate constants for two pH conditions, it will be possible to
establish the internal concentration of ionisable chemicals to within a factor of 4.0 in water
and sediment dwelling invertebrates across the landscape over time. Further work on a
wider variety of ionisable chemicals and sediments is however needed in order to test the
general applicability of the approach. The generation of a more extensive dataset on uptake
and depuration constants of neutral and ionic species of molecules could, in the longer term,
result in the development of models that allow prediction of uptake across a landscape
based on chemical structure alone.
Incorporation of our approach into current risk assessment practices offers a move towards
making risk assessment more representative of the natural environment. For example in the
UK, typical stream water pH ranges from $5.2-8.4.^{10}$ This increase of over 3 pH units will
result in changes in the ionised fraction of chemicals. For example at pH 5.2, diclofenac will
be 93.9 % ionised in comparison to complete ionisation (100%) at pH 8.4. Based on the
relationship between uptake rates for ionised and neutral diclofenac presented in this study,
if lumbricids occur in sediments across these pH ranges, the uptake of diclofenac in $\it L.$
variegatus across UK streams could vary by up to a factor of 168. For fluoxetine the
differences in uptake would differ by up to a factor of 68, whereas for triclosan only small
differences in uptake might be expected across UK streams (factor 1.15). Taking into
account the increased pH variation across European streams (pH range of $2.2-9.8$) these
factors dramatically increase for diclofenac (3035) and fluoxetine (749), but stay almost
constant for triclosan (1.20). Establishing which sites are of greatest concern based on pH
data will allow for targeting monitoring and a more comprehensive evaluation of the risks.
The combined experimental and modelling approach can be used to predict the internal
concentration of ionisable chemicals across a wide spatial scale in water-sediment systems
covering a broad range of pH values and sediment sorption coefficients

Even with extensive monitoring data demonstrating the presence of ionisable chemicals in the aquatic environment, very little data currently exists with regards to measurements of these chemicals in biota, and even fewer studies have demonstrated the uptake of ionisable chemicals in water-sediment systems. There is a real need to understand the uptake of ionisable chemicals in water and sediment systems to fully understand the risks these chemicals may pose to non-target organisms.²⁶ The model and experimental model parameterisation approach presented in this paper offers a way to fill this knowledge gap by generating data on the internal concentration of selected ionisable chemicals in invertebrates such as *L. variegatus* and other aquatic species. The results clearly demonstrate that ionisable chemical uptake is sensitive to changes in exposure medium pH and this needs to be considered when evaluating the risk of such chemicals in aquatic systems. The modelling approach presented could be a very useful tool for assessing the risks of ionisable compounds to benthic organisms at the landscape scale in the future.

Acknowledgements

We would like to thank Unilever who funded MVK's work on the project. LJCs contribution to the study received support from the EU/EFPiA Innovative Medicines Initiative Joint Undertaking (iPiE Grant Number: 115735). AAs contribution to the study received support from the Innovate UK-funded VFETL project. We would also like to thank two anonymous reviewers for very useful insights on an earlier version of this manuscript.

Supporting Information

Detailed information on sorption of study compounds to sediment, mass balance calculations are provided in the Supporting Information. This material is available free of charge at http://pubs.acs.org.

369	Table legends
370	Table 1: Physico-chemical properties of the compounds tested, range of measured pH
371	values during each study, and derived uptake and depuration rate constants. Standard
372	deviations are shown in the parentheses.
373	

Figure	legends
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Figure 1: Schematic diagramme of the modelling approach for estimating uptake of an					
ionisable chemical, AB, into an aquatic invertebrate. $K_{\text{in-neut}}$ = uptake rate constant for the					
neutral species; $K_{\text{in-ion}}$ = uptake rate constant for the ionised species; K_{out} = elimination rate					
constant for the combination of the neutral and ionised species; $pH_{\text{M}} = pH$ of the external					
media; pH_O = internal pH of the organism.					
Figure 2: Measured (points) and simulated (lines) internal concentration of three ionisable					
chemicals in Lumbriculus variegatus in water at different pH values. Dashed lines indicate					
the 95 $\%$ confidence interval of the simulations following the parameter estimation with the					
full data set (black) or according to the minimised design method (blue).					
Figure 3: Comparison of measured (points) and predicted (lines) internal chemical					
concentrations over time in sediment-exposed <i>Lumbriculus variegatus</i> . Dashed lines indicate					
the 95 $\%$ confidence interval of the simulations following the parameter estimation with the					
full data set (black) or according to the minimised design method (blue)					

Table 1

		Diclofenac	Fluoxetine	Triclosan
Log Kow		4.06	4.09	5.17
рКа		4.01	9.62	7.90
Acid/Base		Acid	Base	Acid
Measured pH	Water pH 5.5	5.5-5.7	5.4-5.6	5.4-5.6
	Water pH 7.0	7.0-7.2	6.8-7.9	7.0-7.2
	Water pH 7.4	7.6-8.3	7.5-7.6	7.9-8.3
	Water pH 8.5	7.4-8.9	7.7-8.7	7.7-8.6
	Sediment	7.67	7.67	7.67
f _{ion} used for	Water pH 5.5	0.9731 (0.0043)	1 (0.0004)	0 (0.0004)
modelling	Water pH 7.0	0.9993 (0.0001)	0.9959 (0.0053)	0.1445 (0.0146)
_	Water pH 7.4	0.9998 (0.0001)	0.9915 (0.0014)	0.6076 (0.1522)
	Water pH 8.5 Sediment pH	1 (0.0001)	0.9458 (0.0278)	0.7213 (0.1055)
	7.64	0.9999 (0.001)	0.9889 (0.001)	0.3706 (0.001)
K _{in-neut} (L kg ⁻¹ d ⁻¹)	Full data set	17811 (585)	14203 (747)	1119 (76.4)
K_{in-ion} (L kg ⁻¹ d ⁻¹)		5.75 (0.97)	11.43 (1.42)	1181 (50.9)
$K_{out}(L kg^{-1} d^{-1})$		0.86 (0.18)	0.25 (0.09)	0.01 (0.035)
K _{in-neut} (L kg ⁻¹ d ⁻¹)	Minimised design	14342 (1007)	9735 (1130)	1084 (162)
K_{in-ion} (L kg ⁻¹ d ⁻¹)	acoign	6.82 (0.27)	8.54 (0.99)	1302 (156)
$K_{out}(L kg^{-1} d^{-1})$		0.634 (0.04)	0.039 (0.06)	0.00001 (0.059)

Figure 1

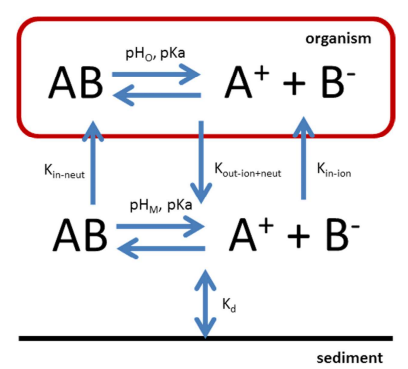


Figure 2

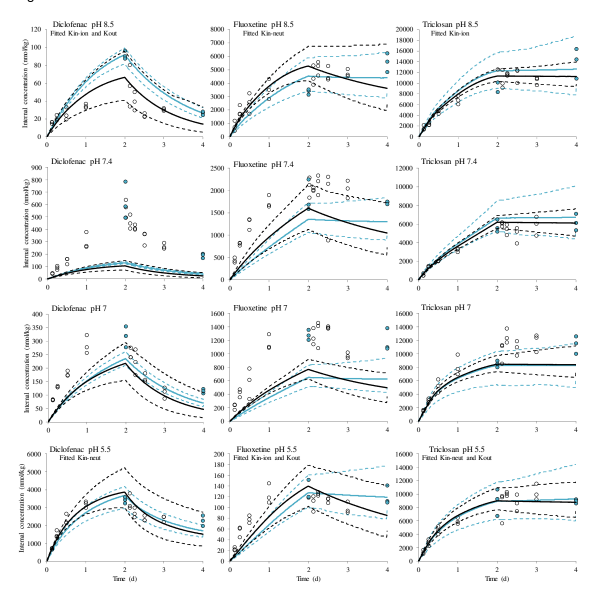
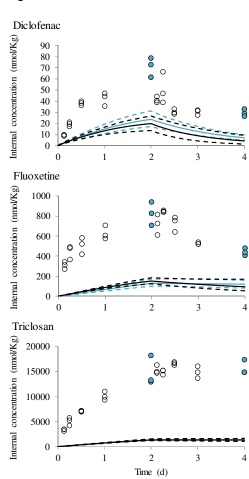


Figure 3



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