



OPEN

Received: 17 September 2018 Accepted: 18 December 2018 Published online: 15 February 2019

Impact of post-hatching maturation on the pharmacokinetics of paracetamol in zebrafish larvae

Rob C. van Wijk 1, Elke H. J. Krekels 1, Vasudev Kantae 1, Amy C. Harms 1, Thomas Hankemeier 1, Piet H. van der Graaf 2, & Herman P. Spaink 1,

Zebrafish larvae are increasingly used in pharmacological and toxicological studies, but it is often overlooked that internal exposure to exogenous compounds, rather than the incubation medium concentration, is driving observed effects. Moreover, as the zebrafish larva is a developing organism, continuous physiological changes impact pharmacokinetic or toxicokinetic processes like the absorption and elimination of exogenous compounds, influencing the interpretation of observations and conclusions drawn from experiments at different larval ages. Here, using paracetamol as paradigm compound, mathematical modelling is used to quantify absorption and elimination rates from internal exposure over time profiles after waterborne treatment, as well as changes in these parameters in post-hatching larvae of 3, 4, and 5 days post fertilisation (dpf). An increase of 106% in absorption rate was observed between 3 and 4 dpf, but no further increase at 5 dpf, and an increase of 17.5% in elimination rate for each dpf. Paracetamol clearance, determined from elimination rate constants and reported total larval volumes of 253, 263, and 300 nL at 3, 4, and 5 dpf respectively, correlates best with higher vertebrates at 5 dpf. This suggests that when studying direct effects of exogenous compounds, experiments with zebrafish larvae are best performed at 5 dpf.

The zebrafish (*Danio rerio*), especially the zebrafish larva, is increasingly used in drug discovery and early drug development, and toxicological screens^{1,2}. It is a data and resource efficient vertebrate model organism³, that shows 70% genetic homology with humans⁴. Its many advantages include high fecundity and small larval size which is ideal for high-throughput experiments⁵. Additionally, transparency in early life stages enables optical imaging to study *in vivo* effects of exogenous compounds observable by brightfield or fluorescence microscopy. Moreover, it is ethically preferable to perform *in vivo* experiments in the lowest vertebrate, like for example the zebrafish. Additionally, no ethical approval is necessary for studies on larvae before they start independent feeding^{6,7}. Experiments in zebrafish larvae bridge the gap between *in vitro* research and *in vivo* preclinical mammal studies as they combine experimental efficiency of cell cultures and organoids with the opportunity to study whole vertebrate organism, including all on- and off-target effects, which will improve extrapolation of observations to higher vertebrates.

In pharmacological and toxicological research with aquatic species, the studied compounds are usually dissolved in the incubation medium (i.e. waterborne treatment). The relationship between the medium concentration of the exogenous compound and its internal exposure is essential for reliable interpretation of the observed results⁸⁻¹², since it is the internal concentration that drives pharmacological and toxicological effects. Because target engagement, which is responsible for the response to exogenous compounds, depends on the pharmacokinetics or toxicokinetics of internal exposure over time, longitudinal data of exposure over time is needed for reliable interpretation of observed effects¹³⁻¹⁵. It is well documented that ignoring this critical issue leads to poor outcomes in drug discovery research¹⁶.

To derive internal exposure based on the external concentration of the compounds, for example based on physicochemical properties, has been shown to be very challenging ^{17–19}. Measuring this essential internal exposure is also a challenge due to the small size of zebrafish larvae and the subsequently required very sensitive quantification methods. Recently however, we demonstrated the technical feasibility of measuring pharmacokinetics

¹Systems Biomedicine and Pharmacology, Leiden Academic Centre for Drug Research (LACDR), Leiden University, Leiden, The Netherlands. ²Certara QSP, Canterbury Innovation House, Canterbury, UK. ³Animal Sciences and Health, Institute of Biology Leiden (IBL), Leiden University, Leiden, The Netherlands. Correspondence and requests for materials should be addressed to H.P.S. (email: h.p.spaink@biology.leidenuniv.nl)

in zebrafish by developing a profile of internal amount over time for zebrafish of 3 days post fertilisation (dpf), using paracetamol (acetaminophen) as paradigm compound²⁰. In this analysis, mathematical modelling was used to describe the pharmacokinetics by quantifying the absorption rate constant, and elimination rate constant which reflects both metabolism and excretion, processes that in addition to the distribution drive the internal exposure.

Although experiments with larva have many advantages, studying an organism during its development will require understanding of the effect of maturation on the studied feature. Zebrafish development is rapid, showing embryogenesis within 3 dpf²¹, liver budding from 1 dpf and growth from 2 dpf²², development of a functional renal system after 2 dpf²³, and presence of a gastro-intestinal (GI) tract from 1–4 dpf²⁴, reaching adulthood in 3 months²⁵. These developmental and maturation processes in the first days post fertilisation are expected to have an impact on the absorption and elimination of compounds. As most experiments in the field of pharmacology and toxicology are performed during these first days¹, it is essential to understand and quantify the impact of development, and to know what difference a single experimental day makes on the internal exposure of exogenous compounds. This is especially the case when studying direct, short-term effects of exogenous compounds.

Using paracetamol as paradigm compound, our aim is therefore to use mathematical modelling to quantify absorption and elimination rate constants from internal exposure over time profiles after waterborne treatment in post-hatching zebrafish larvae of 3 to 5 dpf, and to characterise the impact of development on these parameters using post fertilisation age as marker.

Methods

Chemicals. Paracetamol and paracetamol-D4 internal standard were purchased from Sigma (Sigma-Aldrich Chemie B.V., Zwijndrecht, The Netherlands). UPLC/MS grade MeOH was purchased from Biosolve (Biosolve B.V., Valkenswaard, The Netherlands). Purified water (H₂O) was retrieved from PURELAB (Veolia Water Technologies B.V., Ede, The Netherlands).

Zebrafish husbandry. All experiments were planned and executed in compliance with European regulation⁶. Handling and maintenance of zebrafish and zebrafish larvae was in accordance with international standard protocols²⁶. Adult wild type AB/TL zebrafish were set-up for overnight family cross breeding, separated by sex. Next morning, males and females were combined in breeding tanks with inserts and after 20 minutes eggs were collected. This way, time of fertilisation was controlled. Fertilised eggs were kept at 28 °C in petri dishes in embryo medium (demineralised containing $60\,\mu\text{g/mL}$ Instant Ocean sea salts; Sera, Heinsberg, Germany) which was refreshed daily.

Experimental design. The experimental design of Kantae *et al.* performed in larvae of 3 dpf²⁰, was repeated with larvae of 4 and 5 dpf in samples of n = 5 zebrafish larvae. In short, two experiments were performed, one in which larvae were continuously treated with 1 mM waterborne paracetamol in embryo medium (treatment medium) for 0–180 minutes and one in which the larvae were treated with treatment medium for 60 minutes and then washed with embryo medium using Netwell inserts filters (Corning Life Sciences B.V., Amsterdam, The Netherlands) and transferred to drug-free medium for a washout period of 60–240 minutes. After the designated constant waterborne treatment or washout period, the larvae were washed with 20/80 MeOH/H₂O (v/v) using Netwell inserts, transferred to Safe-Lock tubes (Eppendorf Nedeland B.V., Nijmegen, The Netherlands), excess volume was removed and $100 \,\mu$ L 90/10 MeOH/H₂O with 45 pg/uL paracetamol-D4 internal standard was added. The sample was snap frozen in liquid nitrogen and stored at $-80\,^{\circ}$ C until quantification. Measurements at all time points were performed at least *in triplo*.

Additionally, to ensure paracetamol concentrations in the treatment medium remained constant throughout the duration of the experiment, treatment medium from a set-up with and without 3, 4, and 5 dpf larvae was sampled at 180 minutes and compared to 1 mM paracetamol solution in H_2O and to fresh treatment medium, all in triplo. Samples were frozen at $-80\,^{\circ}C$ until quantification.

Measurements of internal exposure. The method to quantify internal paracetamol exposure were described earlier by Kantae *et al.*²⁰. In short, samples were lysed by iterations of snap freezing the solution with the larvae in liquid nitrogen and submerging the sample in a sonication bath until a homogeneous suspension was obtained. These suspensions were centrifuged at $16,000\,g$ for $10\,minutes$ and $90\,\mu$ L supernatant was added to $72\,\mu$ L H₂O to reach $50/50\,MeOH/H_2O\,(v/v)$ to be injected into the ultra-performance liquid chromatography (UPLC) system (Acquity, Waters Chromatography B.V., Etten-Leur, The Netherlands) linked to a quadrupole-ion trap MS/MS (QTRAP, AB Sciex B.V., Nieuwerkerk aan den IJssel, The Netherlands) with an electrospray ionisation source in positive mode. Development criteria were 90-100% accuracy and relative standard deviations less than 10% as measure of precision. Paracetamol concentrations were determined through a calibration curve in matrix ranging from 0.09 to $180\,pg/uL$, and calculated to total paracetamol amount in pmole per zebrafish larva. Treatment medium samples were diluted with H₂O to fall within the academic calibration range from 0.05 to $100\,pg/uL$ with a final internal standard concentration of $25\,pg/uL$ paracetamol-D4.

Pharmacokinetic modelling. To quantify the physiological processes driving the internal exposure of paracetamol, a mathematical model was developed using non-linear mixed effects (NLME) modelling, which combines the quantification of non-random trends in the data called fixed effects as well as random variability known as random effects. NLME modelling was performed using the First Order Conditional Estimation (FOCE) algorithm in NONMEM $(v.7.3)^{27}$, which was operated through the interfaces Pirana $(v.2.9.6)^{28}$ and PsN $(v.4.7.0)^{29}$. Graphical output was generated using R $(v.3.4.2)^{30}$ running through the RStudio (v.1.1.383), RStudio Inc, Boston, Massachusetts, USA) interface.

Age	k _e (min ⁻¹)	Total volume (nL)	CL (nL/h)
3 dpf	0.0193	253	292.3
4 dpf	0.0226	263	356.8
5 dpf	0.0266	300	478.1

Table 1. Paracetamol elimination rate constant (k_e), reported total larval volume³³, and derived absolute paracetamol clearance (CL) for 3, 4, and 5 dpf larvae.

A one and two compartment model was tested. In case of the two compartment model, the sum of the amounts in both compartments were fitted to the observed total amounts, while elimination was only limited to one compartment. Absorption of paracetamol was estimated as a zero-order process, assuming the paracetamol concentration in the incubation medium to remain constant over time. For the elimination estimation both first-order linear and saturable non-linear Michaelis Menten processes were tested.

Quantification of the residual error was tested as additive, proportional, and a combination of additive and proportional error. As the larvae were lysed to quantify internal exposure, only single observations were obtained from a larval sample. As a result inter-individual variability in internal exposure caused by individual variability in model parameters could not be distinguished from residual variability caused by experimental or analytical error.

Quantification of the correlation between model parameters and larval age, was tested with both continuous (Equations 1 and 2) and discrete (Equation 3) functions:

$$P = P_{base} \cdot (1 + slope \cdot (age - ref)) \tag{1}$$

$$P = P_{hase} \cdot (1 + slope)^{age-ref} \tag{2}$$

$$P = \begin{cases} P_1 & age = 3 dpf \\ P_2 & age = 4 dpf \\ P_3 & age = 5 dpf \end{cases}$$

$$(3)$$

where P is the parameter of interest, P_{base} is the base value at the reference age ref, and P_1 , P_2 and P_3 are different functions or estimates of the parameter of interest for their respective conditions.

For the continuous relationship, a linear function (Equation 1) or power function (Equation 2) was tested to describe the relationship between age and parameter values. In the discrete function (Equation 3) different parameter values were estimated for larvae older and/or younger than a specified reference age.

For model selection, the likelihood ratio test was used between nested models³¹, assuming a χ^2 distribution and using a significance level of p < 0.01. Additional selection criteria were successful minimisation, estimates of parameter values with 3 or more significant digits and relative standard errors below 50%, and biologically plausibility of the parameter estimates. Graphically, model accuracy was assessed using goodness-of-fit plots, consisting of observed versus predicted plots and conditional weighted residuals (CWRES) versus time or predicted paracetamol amounts, which should show no bias over time or predicted paracetamol amounts³². Stability of paracetamol concentrations in treatment medium in the control experiment were normalized to H₂O control and tested by non-parametric Kruskall-Wallis test, as the data were not normally distributed, with level of significance of 0.05.

Comparison of paracetamol clearance to higher vertebrates. The degree of correlation between para cetamol clearance in zebrafish larvae with higher vertebrates was assessed by calculating paracetamol clearance values in the larvae by multiplying the obtained elimination rate constants with previously reported total larval volumes at corresponding ages³³ that are provided in Table 1. This assumes the distribution volume to be equal to the total volume of the larva and a homogenous distribution of the compound throughout the whole larva. The paracetamol clearance values in the larvae at different ages, were graphically compared to reported paracetamol clearance values in higher vertebrates²⁰, in a plot of clearance values versus bodyweight of the species. The bodyweight of the larvae was derived from their volume, assuming a density of 0.997 g/mL²⁰. A linear least squares regression with 95% confidence interval of the log transformed clearance and log transformed bodyweight was calculated in R, based on clearance values obtained in mature individuals of the different species included in the graph.

Results

Measurements of internal exposure. The observed internal exposure of paracetamol expressed as total amount per larva over time is shown in Fig. 1 for larvae of 3, 4, and 5 dpf for both the constant waterborne treatment and the washout experiment. It can be seen that steady state of internal paracetamol exposure is reached between 100 and 120 minutes of constant waterborne treatment, meaning an equilibrium between paracetamol amounts absorbed and eliminated per time unit has been reached. Steady state exposure in the constant waterborne treatment experiment increased in larvae between 3 and 4 dpf, while it remained relatively constant in larvae between 4 and 5 dpf. The washout experiment showed a mono-exponential decline of the paracetamol amount per larva after the larvae were transferred to paracetamol-free medium. The steepness of this curve, reflecting the elimination rate, increases in larvae with increasing age. The dataset is available through the DDMoRe Repository, Model ID DDMODEL00000294 (http://repository.ddmore.foundation/model/DDMODEL00000294). The stability of the paracetamol concentration in the treatment medium was not impacted by the experimental set-up (Supplementary Fig. 1).

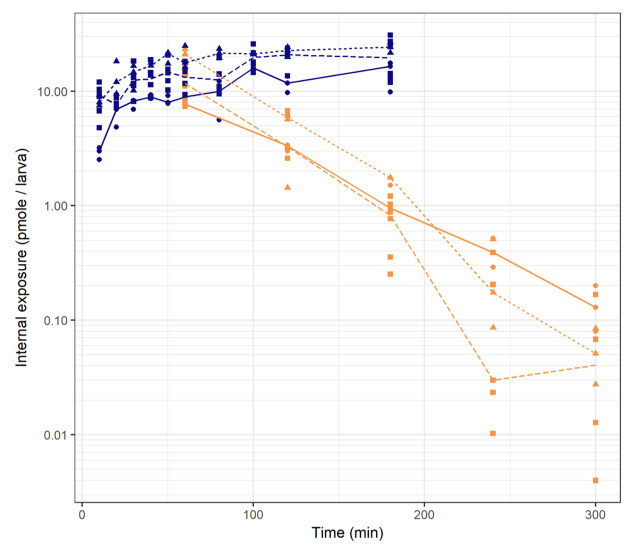


Figure 1. Raw data of internal exposure of paracetamol amounts over time for zebrafish larvae of 3 dpf (solid line, closed circles), 4 dpf (dotted line, closed triangles), and 5 dpf (dashed line, closed squares) for both constant waterborne drug treatment experiment (blue) or washout experiment (orange). Datapoints are total amount per larva from a pooled sample with n=5. The lines are connecting the median values at each time point.

Pharmacokinetic modelling. Based on the selection criteria, a one-compartment model with zero-order absorption and first-order elimination best fitted the observed profiles of paracetamol amounts in zebrafish larvae over time for both experiments. A combination of additive and proportional error model was found to describe residual variability best, with the variance of the proportional error being 0.109 corresponding to 33% and the variance of the additive residual unexplained error being 0.00844 pmole/larva. A schematic and mathematical representation of this model is provided in Fig. 2 and Equation 4, respectively. The final model included a discrete relationship between age and the absorption rate constant (Equation 5) and a power relationship between age and the elimination rate constant (Equation 6).

$$\frac{dA}{dt} = k_a - k_e \cdot A \tag{4}$$

$$k_{a} = \begin{cases} k_{a,base} & age = 3 \ dpf \\ k_{a,base} \cdot (1 + factor_{a}) \ 3 \ dpf < age \le 5 \ dpf \end{cases}$$
 (5)

$$k_e = k_{e,base} \cdot (1 + slope_e)^{age - 3dpf} \tag{6}$$

where A is the paracetamol amount in a single larva, k_a is the zero-order absorption rate constant, $k_{a,base}$ is the base value of the absorption rate constant at the reference age of 3 dpf, and factor_a describes the fractional increase

	Parameter value	Relative standard error (%)
Structural parameters		
k _{a,base} (pmole/min)	0.289	4
factor _a (-)	1.06	14
$k_{e,base} (min^{-1})$	0.0193	5
slope _e (–)	0.175	18
Stochastic parameters		
Variance of proportional residual error (-)	0.109	14
Variance of additive residual error (pmole/larva)	0.00844	48

Table 2. Obtained model parameter values and their relative standard error.



Figure 2. Schematic representation of the final model to describe the total amount of paracetamol in zebrafish larvae over time. ka = zero-order absorption rate constant, A = amount of paracetamol in one larva, ke = first-order elimination rate constant.

in the absorption rate constant in zebrafish larvae that are older than 3 days, k_e is the first-order elimination rate constant, $k_{e,base}$ is the base value of the elimination rate constant at the reference age of 3 dpf, and slope_e is the estimated slope in the relationships between the elimination rate constant and age.

The obtained parameter values are presented in Table 2 and final model code is available through the DDMoRe Repository, Model ID DDMODEL00000294 (http://repository.ddmore.foundation/model/DDMODEL00000294).

According to the obtained results, at 3 dpf the value of the zero-order absorption rate constant of paracetamol is 0.289 pmole/min and the first-order elimination rate constant is 0.0193 min $^{-1}$. The absorption rate constant was found to be statistically significantly (p < 1e-10) increased between 3 and 4 dpf by 106% in the final model, but the difference in this parameter between larvae of 4 dpf and 5 dpf was found not to be statistically significant (p = 0.46). The elimination rate constant was found to statistically significantly (p < 1e-6) increase between all three ages. In the final model, the elimination rate constant increased by 17.5% per day, resulting in an elimination rate constant of 0.0226 min $^{-1}$ and 0.0266 min $^{-1}$ for larvae of 4 and 5 dpf respectively.

The model predicted concentration-time profile per age and experiment together with the observed concentrations are shown in Fig. 3, showing good agreement between observed and predicted concentrations. The diagnostic goodness-of-fit plots further confirmed good accuracy of the model predictions (Supplementary Figs 2–4). The relative standard error values of the obtained structural model parameters are well below 20%, indicating good precision of these estimates.

Comparison of paracetamol clearance to higher vertebrates. Paracetamol clearance and previously reported larval volume for 3, 4, and 5 dpf larvae are shown in Table 1. Figure 4 shows the correlation between paracetamol clearance and bodyweight for 13 species including the zebrafish. This plot has previously been reported including the results of zebrafish larvae of 3 dpf only²⁰ and now includes also the clearance values for 4 and 5 dpf larvae. It can be seen that the older and heavier larvae show a closer correlation with the higher vertebrates, as they are positioned closer to the 95% confidence interval of the allometric relationship between bodyweight and paracetamol clearance as established based on data from mature individuals only. They do remain below the confidence interval, as do the data points obtained in paediatric human studies (red triangles).

Discussion

Here the impact of development on the pharmacokinetic or toxicokinetic processes of absorption and elimination through post fertilisation age as marker was quantified by mathematical modelling based on the profiles of internal exposure over time after waterborne treatment in post-hatching zebrafish larvae of 3, 4, and 5 dpf. The absorption of paracetamol was shown to increase 106% between 3 and 4 dpf, but not to significantly further increase at 5 dpf, while paracetamol elimination increased 17.5% per day in this 3 to 5 days post fertilisation period.

Within the mathematical model, the relationships between age and the absorption and elimination rate constants were parameterised with values for larvae at 3 dpf as reference values. These values are comparable to the values reported previously for zebrafish larvae of 3 dpf alone²⁰. The doubling of the absorption rate between 3 and 4 dpf, can be explained by the opening of the GI tract, which is a discrete event completing with the opening of the anus at 4 dpf^{7,24}. From that moment, instead of only transdermal or trans-gill absorption, the larvae will also ingest the exogenous compound orally. Recently examined absorption of the antihistamine diphenhydramine between zebrafish embryos and larvae showed an chorion-independent increase in absorption between 2 and 4 dpf and are in concordance with our results here³⁴. The absorption rate constant did not increase further between

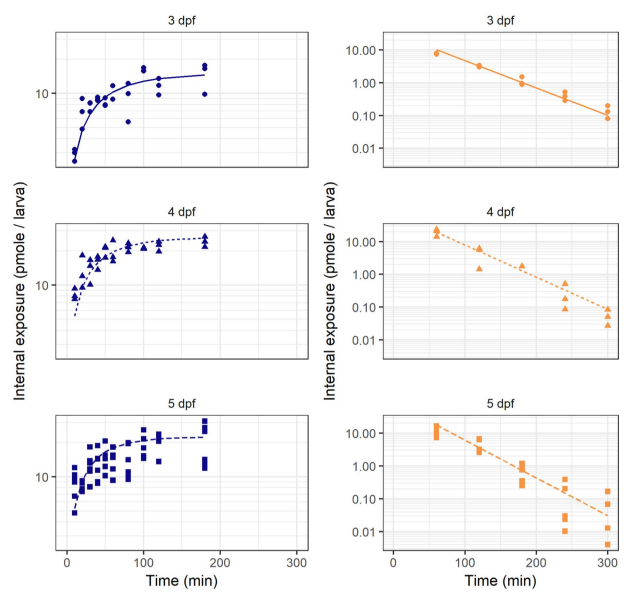


Figure 3. Model fit (lines) through observed paracetamol amounts over time (symbols) in larvae of 3 dpf (solid line, closed circles), 4 dpf (dotted line, closed triangles), and 5 dpf (dashed line, closed squares) for both the constant waterborne drug treatment experiment (blue, left panels) and the washout experiment (orange, right panels).

4 and 5 dpf, suggesting that potential other processes that add to the absorption of paracetamol, do not show maturational changes in the age range studied here.

The 17.5% increase in the elimination rate constant between each of the three post fertilisation days is expected to result from the continuous growth of eliminating organs like the liver and kidneys, as well as continuous maturation of enzymatic processes 22 . Indeed, the clearance values of immature organisms of both the zebrafish and human are lower than expected based on bodyweight alone, but these values do move towards the regression line with increasing age (Fig. 4), with the larval clearance being 35, 41, and 49% of the lower bound of the 95% confidence interval of the extrapolated clearance calculated based on the values of higher vertebrates for larvae of 3, 4, and 5 dpf respectively. It has to be kept in mind that for comparison to higher vertebrates, the absolute clearance in the zebrafish larvae was calculated based on total larval volume, assuming a homogenous distribution over the total body of the larvae, because information on the distribution volume of paracetamol in fish is not available in literature. Given that the distribution volume of paracetamol has been reported to range from 0.8–0.9 L/kg 35,36 in humans, this assumption seems to be reasonable, although further research into the distribution of this compound in zebrafish larvae is required. If the true distribution volume is larger, the calculated clearance values would also be proportionally larger and fall within the 95% confidence interval, or vice versa. Another factor that may contribute to deviations of the clearance values in zebrafish larvae from the regression line could be the fact that fish are poikilotherms, for which lower metabolic rates have been reported 37 .

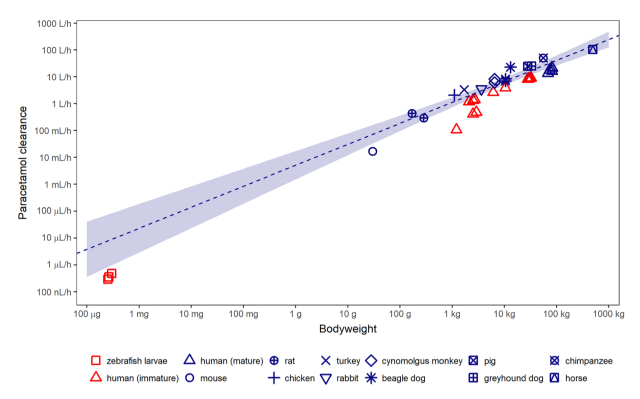


Figure 4. Allometric relationship between paracetamol clearance and bodyweight of 13 vertebrate species including the zebrafish larvae at three different ages. Blue and red symbols show mature or immature individuals of the species, respectively. Allometric relationship (dashed blue line) and 95% confidence interval (shaded area) are determined based on data in mature organisms only. Adapted with permission from Kantae *et al.*²⁰.

From our results it is clear that the age of the larvae during experiments with waterborne treatment influences internal exposure of exogenous compounds, at least for our paradigm compound. Because the internal exposure of the exogenous compound drives its pharmacological or toxicological effect, it can be expected that the age will also impact the observed effects resulting from the treatment. The age of the larvae used to investigate the direct effects of exogenous compounds is therefore an important experimental design consideration. To determine which age to include, three criteria are of importance. Firstly, the internal exposure of the studied compound should be high enough to yield an effect and prevent false negatives. Secondly, the larval metabolic capacity should be large enough to biotransform exogenous compounds to their active metabolites as drug metabolites can also be biologically active. Thirdly, extrapolating observations to higher vertebrates, for instance to improve study design of mammal studies based on translation of clearance, benefits from a direct comparison of the pharmacokinetic processes between species. Based on the results of the paracetamol study presented here, we propose experiments for the testing of direct effects of exogenous compounds to be performed in zebrafish larvae at 5 dpf, because absorption is highest at 4 and 5 dpf, while the metabolic capacity at 5 dpf is largest and clearance at that time resembles clearance of higher vertebrates most within the age range that still falls within the ethical constraints for experiments in zebrafish larvae.

Conclusion

In conclusion, it is of importance to quantify internal exposure over time when testing exogenous compounds by waterborne treatment in zebrafish larvae. The opening of the GI-tract will likely result in increased absorption, which is seen here between 3 and 4 dpf when absorption of paracetamol more than doubles. Continuous growth of eliminating organs as well as maturation of enzymatic processes lead to increased elimination, which is 17.5% daily for paracetamol between 3 and 5 day post fertilisation. To increase internal exposure to parent compounds and metabolites in short term exposure studies, we therefore recommend careful consideration of zebrafish age in experimental design when these pharmacokinetic or toxicokinetic processes are of relevance to the research question. Based on our results with paracetamol, using 5 dpf zebrafish larvae may be preferable for studying direct short-term effects of exogenous compounds.

Data Availability

The full dataset and model file are available through the DDMoRe Repository, Model ID DDMODEL00000294 (http://repository.ddmore.foundation/model/DDMODEL00000294).

References

- 1. Rennekamp, A. J. & Peterson, R. T. 15 years of zebrafish chemical screening. Curr. Opin. Chem. Biol. 24, 58–70 (2015).
- 2. Peterson, R. T. & Macrae, C. A. Systematic approaches to toxicology in the zebrafish. *Annu. Rev. Pharmacol. Toxicol.* **52**, 433–53 (2012).
- 3. Zon, L. I. & Peterson, R. T. In vivo drug discovery in the zebrafish. Nat Rev Drug Discov 4, 35-44 (2005).

- 4. Howe, K. et al. The zebrafish reference genome sequence and its relationship to the human genome. Nature 496, 498–503 (2013).
- Schulthess, P. & Van Wijk, R.C. et al. Outside-in systems pharmacology combines innovative computational methods with highthroughput whole vertebrate studies. CPT Pharmacometrics Syst. Pharmacol. 7, 285–287 (2018).
- 6. Council Directive 2010/63/EU on the protection of animals used for scientific purposes. Off. J. Eur. Union L276/33 (2010).
- 7. Strähle, U. *et al.* Zebrafish embryos as an alternative to animal experiments Å commentary on the definition of the onset of protected life stages in animal welfare regulations. *Reprod. Toxicol.* 33, 128–132 (2012).
- 8. Damalas, D. E., Bletsou, A. A., Agalou, A., Beis, D. & Thomaidis, N. S. Assessment of the acute toxicity, uptake and biotransformation potential of benzotriazoles in zebrafish (Danio rerio) larvae combining HILIC-with RPLC-HRMS for high-throughput identification. *Environ. Sci. Technol.* 52, 6023–6031 (2018).
- 9. Achenbach, J. C. et al. Analysis of the uptake, metabolism, and behavioral effects of cannabinoids on zebrafish larvae. Zebrafish 15, 349–360 (2018).
- Kühnert, A., Vogs, C., Altenburger, R. & Küster, E. The internal concentration of organic substances in fish embryos a toxicokinetic approach. Environ. Toxicol. Chem. 32, 1819–1827 (2013).
- 11. Kühnert, A. et al. Biotransformation in the zebrafish embryo –temporal gene transcription changes of cytochrome P450 enzymes and internal exposure dynamics of the AhR binding xenobiotic benz[a]anthracene. Environ. Pollut. 230, 1–11 (2017).
- 12. Brox, S., Seiwert, B., Küster, E. & Reemtsma, T. Toxicokinetics of polar chemicals in zebrafish embryo (Danio rerio): influence of physicochemical properties and of biological processes. *Environ. Sci. Technol.* **50**, 10264–10272 (2016).
- 13. Li, Y. et al. Dissolved organic matter affects both bioconcentration kinetics and steady-state concentrations of polycyclic aromatic hydrocarbons in zebrafish (Danio rerio). Sci. Total Environ. 639, 648–656 (2018).
- 14. Liu, H. et al. Pharmacokinetics and effects of tetrabromobisphenol a (TBBPA) to early life stages of zebrafish (Danio rerio). Chemosphere 190, 243–252 (2018).
- 15. Van Wijk, R. C., Krekels, E. H. J., Hankemeier, T., Spaink, H. P. & Van der Graaf, P. H. Systems pharmacology of hepatic metabolism in zebrafish larvae. *Drug Discov. Today Dis. Model.* 22, 27–34 (2017).
- 16. Morgan, P. et al. Can the flow of medicines be improved? Fundamental pharmacokinetic and pharmacological principles toward improving Phase II survival. *Drug Discov. Today* 17, 419–424 (2012).
- 17. Geier, M. C. *et al.* Systematic developmental neurotoxicity assessment of a representative PAH Superfund mixture using zebrafish. *Toxicol. Appl. Pharmacol.* **354**, 115–125 (2018).
- 18. Diekmann, H. & Hill, A. Zebrafish as a platform for *in vivo* drug discovery ADMETox in zebrafish. *Drug Discov. Today Dis. Model.* 10, e31–e35 (2013).
- Ordas, A. et al. Testing tuberculosis drug efficacy in a zebrafish high-throughput translational medicine screen. Antimicrob. Agents Chemother. 59, 753–762 (2015).
- Kantae, V. et al. Pharmacokinetic modeling of paracetamol uptake and clearance in zebrafish larvae: Expanding the allometric scale in vertebrates with five orders of magnitude. Zebrafish 13, 504–510 (2016).
- 21. Kimmel, C. B., Ballard, W. W., Kimmel, S. R., Ullmann, B. & Schilling, T. F. Stages of embryonic development of the zebrafish. *Dev. Dyn.* 203, 253–310 (1995).
- 22. Tao, T. & Peng, J. Liver development in zebrafish (Danio rerio). J. Genet. Genomics 36, 325-334 (2009).
- 23. Gehrig, J., Pandey, G. & Westhoff, J. H. Zebrafish as a model for drug screening in genetic kidney diseases. Front. Pediatr. 6, 183 (2018).
- 24. Ng, A. N. Y. *et al.* Formation of the digestive system in zebrafish: III. Intestinal epithelium morphogenesis. *Dev. Biol.* **286**, 114–135 (2005).
- 25. Parichy, D. M., Elizondo, M. R., Mills, M. G., Gordon, T. N. & Engeszer, R. E. Normal table of post-embryonic zebrafish development: staging by externally visible anatomy of the living fish. *Dev. Dyn.* 238, 2975–3015 (2009).
- 26. Westerfield, M. The Zebrafish Book. A Guide For The Laboratory Use Of Zebrafish (Danio Rerio). (University of Oregon Press, 2000).
- Beal, S., Sheiner, L., Boeckmann, A. & Bauer, R. J. (eds) NONMEM 7.3.0 Users Guides. ICON Development Solutions, Hanover, MD, USA. (1989–2013).
- 28. Keizer, R., Van Benten, M., Beijnen, J., Schellens, J. & Huitema, A. Pirana and PCluster: a modeling environment and cluster infrastructure for NONMEM. Comput Methods Programs Biomed 101, 72–79 (2011).
- Lindbom, L., Pihlgren, P. & Jonsson, E. PsNtoolkit—a collection of computer intensive statistical methods for non-linear mixed effect modeling using NONMEM. Comput Methods Programs Biomed 79, 241–257 (2005).
- 30. R Core Team. R: A language and environment for statistical computing. R Found. Stat. Comput. Vienna, Austria (2014).
- 31. Mould, D. R. & Upton, R. N. Basic concepts in population modeling, simulation, and model-based drug development-part 2: introduction to pharmacokinetic modeling methods. *CPT pharmacometrics Syst. Pharmacol.* 2, e38 (2013).
- 32. Nguyen, T. H. T. et al. Model evaluation of continuous data pharmacometric models: metrics and graphics. CPT Pharmacometrics Syst. Pharmacol. 6, 87–109 (2017).
- 33. Guo, Y., Veneman, W. J., Spaink, H. P. & Verbeek, F. J. Three-dimensional reconstruction and measurements of zebrafish larvae from high-throughput axial-view *in vivo* imaging. *Biomed. Opt. Express* 8, 2611 (2017).
- 34. Kristofco, L. A., Haddad, S. P., Chambliss, C. K. & Brooks, B. W. Differential uptake of and sensitivity to diphenhydramine in embryonic and larval zebrafish. *Environ. Toxicol. Chem.* 37, 1175–1181 (2017).
- 35. Reith, D. et al. Simultaneous modelling of the Michaelis-Menten kinetics of paracetamol sulphation and glucuronidation. Clin. Exp. Pharmacol. Physiol. 36, 35–42 (2009).
- 36. Prescott, L. F. Kinetics and metabolism of paracetamol and phenacetin. Br. J. Clin. Pharmacol. 10, 291-298 (1980).
- 37. White, C. R., Phillips, N. F. & Seymour, R. S. The scaling and temperature dependence of vertebrate metabolism. *Biol. Lett.* 2, 125–7 (2006).

Acknowledgements

The authors thank Sebastiaan Goulooze for peer-reviewing scripts for data handling and mathematical modelling.

Author Contributions

R.C.v.W., E.H.J.K., T.H., P.H.v.d.G. and H.P.S. planned experiments and interpreted results, R.C.v.W. executed experiments and data analysis, V.K. and A.H. assisted in drug quantification. R.C.v.W. wrote the manuscript. All co-authors reviewed the manuscript.

Additional Information

Supplementary information accompanies this paper at https://doi.org/10.1038/s41598-019-38530-w.

Competing Interests: The authors declare no competing interests.

Publisher's note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit https://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2019