Earthworm Uptake Routes and Rates of Ionic Zn and ZnO Nanoparticles at Realistic Concentrations, Traced Using Stable Isotope Labeling

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ABSTRACT: The environmental behavior of ZnO nanoparticles (NPs), their availability to, uptake pathways by, and biokinetics in the earthworm Lumbricus rubellus were investigated using stable isotope labeling. Zinc isotopically enriched to 99.5% in 68Zn (68Zn-E) was used to prepare 68ZnO NPs and a dissolved phase of 68Zn for comparison. These materials enabled tracing of environmentally relevant (below background) NP additions to soil of only 5 mg 68Zn-E kg\(^{-1}\). Uptake routes were isolated by introducing earthworms with sealed and unsealed mouthparts into test soils for up to 72 h. The Zn isotope compositions of the soils, pore waters and earthworms were then determined using multiple collector inductively coupled plasma mass spectrometry. Detection and quantification of 68Zn-E in earthworm tissue was possible after only 4 h of dermal exposure, when the uptake of 68Zn-E had increased the total Zn tissue concentration by 0.03‰. The results demonstrate that at these realistic exposure concentrations there is no distinguishable difference between the uptake of the two forms of Zn by the earthworm L. rubellus, with the dietary pathway accounting for ~95% of total uptake. This stands in contrast to comparable studies where high dosing levels were used and dermal uptake is dominant.

1. INTRODUCTION

The novel properties exhibited by engineered nanoparticles (NPs) have resulted in these materials becoming increasingly prevalent in consumer products. Zinc oxide (ZnO) NPs, for example, have found use in sunscreens, cosmetics and paints due to their UV absorbance properties.\(^1,2\) The increased manufacture and use of engineered NPs will inevitably lead to a corresponding rise in their release into natural environments.\(^3\) Engineered NPs are likely to end up in water treatment plants and be removed as part of the water treatment sludge. In places where such sludge is applied to agricultural land, it is therefore expected that soils will be a major sink. Gottschalk et al.\(^4\) determined predicted environmental concentrations (PECs) for ZnO NPs in European sewage treatment plant sludge of 17.1 mg kg\(^{-1}\), resulting in annual fluxes of 3.25 mg kg\(^{-1}\) to sludge treated soils. For sewage treatment sludge and sludge treated soils Boxall et al.\(^5\) modeled PECs of 2.2–22 mg kg\(^{-1}\) and 3.2–32 mg kg\(^{-1}\), respectively, assuming between 10% and 100% market penetration for ZnO NPs in sunscreens (Figure S1). These PECs are most likely to be overestimates, as recent work has demonstrated that once ZnO NPs enter natural systems they are subject to rapid dissolution or changes in speciation such that they are unlikely to persist.\(^6,7\)

The natural Zn concentrations of soils are typically much higher than the PECs. A report by the US Geological Survey found Zn concentrations of between 10 and 113 mg kg\(^{-1}\) for 38 rural organic rich soils, away from sources of pollution and fertilizer use.\(^8\) A comparable study of British soils from a variety of land classes identified a similar range of Zn levels, with values of 2.5 to 2120 mg kg\(^{-1}\).\(^9\) Taken together, these data imply that the Zn concentrations originating from ZnO NPs in sewage treated soils are likely to be significantly lower than the natural Zn background levels, by a factor of 10 or more (Figure S1).

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Despite this, the unknown behavior and interactions of these materials in the environment has prompted investigations to determine if NPs pose a risk.

A number of such studies have demonstrated that dissolved Zn has a greater toxicity than ZnO NPs even though higher Zn body burdens are commonly observed in exposures to NPs. It has thus been proposed that the release of Zn ions from the dissolution of ZnO NPs is responsible for any toxicity and that “nanospecific” toxic effects are absent.10–12 Furthermore, Hooper et al.10 suggested that conservative risk assessments for ZnO NPs may be conducted by assuming that all Zn from NPs is available in dissolved form. In contrast, other investigations presented evidence for “nanospecific” effects and demonstrated that ZnO NPs exerted greater toxicity for some species than equivalent doses of dissolved Zn.13,14 A proposed mechanism for the observed toxicity is the production of reactive oxygen species as a consequence of the high surface area of the NPs and exposure to UV radiation from sunlight.13,15 Such effects may not be significant for organisms that live in soil, however, due to the shallow penetration depth of UV radiation.16

Notably, previous soil exposures with ZnO NPs were performed at Zn concentrations of 100 to 6400 mg kg⁻¹ and these exposure levels by far exceed the PECs and, in most cases, the high penetration depth of UV radiation.16

In this study, we investigate, over 72 h, the uptake of Zn by the earthworm Lumbricus rubellus from soils dosed with ZnO NPs and soluble ZnCl₂ at an environmentally relevant Zn level of 5 mg kg⁻¹. Zinc is expected to enter the earthworms via two main pathways and both were monitored: (i) dermal uptake by direct contact with soil pore water, whereby it has been shown that considerable exchange of water occurs across the body wall19 and (ii) dietary uptake of dissolved Zn and NPs adsorbed to soil particles, from the earthworm’s burrowing behavior and digestion of organic soil constituents. Given the low Zn dosing level (below background) and short exposure period (≤72 h) employed here, DNA damage and genotoxic effects are not expected and were not investigated. This conclusion is supported by Hu et al.13 who found that ZnO NP soil concentrations of more than 1 g kg⁻¹ were needed to induce DNA damage during a 7-day exposure with the earthworm Eisenia fetida.

Given the low and environmentally realistic exposure levels employed in this study, it is expected that ZnO NPs will undergo rapid dissolution without inhibition by the saturated pore waters that were likely encountered in a recent and comparable investigation, which employed much higher dosing levels.12 A comparative exposure with dissolved Zn was included in our experimental program to further test this hypothesis. If rapid dissolution of ZnO NPs occurs, this should produce similar exposure scenarios for the soils dosed with nanoparticulate and dissolved Zn and, therefore, comparable uptake is expected. The tracing and quantification of the dissolved Zn at the low employed exposure levels was made possible by the use of stable isotope labeling in conjunction with high precision isotope ratio analysis by multiple collector inductively coupled plasma mass spectrometry (MC-ICP-MS).20 While such techniques were previously applied in aquatic exposures,21,22 this investigation represents the first application of stable isotope tracing methods to study the fate of engineered nanomaterials in soils.

2. METHODS

2.1. Materials and Reagents. The well characterized, loamy sand standard test soil LUFASpeyer 2.2 (Sp 2121, Germany, 2009), with a pH of 5.5, a total organic carbon content of 2.09%, a cation exchange capacity of 10.0 mequiv/100 g and a water-holding capacity (WHC) of 46.5%, was used as the exposure medium. The soil was oven-dried at 60 °C overnight prior to use, to eliminate undesired soil fauna.

Zinc that is artificially enriched in the 68Zn isotope (68Zn-E) from the natural abundance of 18.8% to 99.5%, was purchased as a metal powder from Isoflex USA and used for the preparation of 68ZnO NPs and a 68ZnCl₂ solution, hereafter referred to as dissolved 68Zn. AnalAR grade concentrated 14.5 M HNO₃ and 6 M HCl were purified by sub-boiling distillation in a quartz still. Trace element grade Optima 28 M HF was purchased from Fisher Scientific. Milli-Q water of >18 MΩ cm quality (Millipore, UK) and 30% Suprapur H₂O₂ (VWR UK) were used throughout.

The 68ZnO NPs applied in this study were prepared by forced hydrolysis in diethylene glycol (DEG) and are from the same synthesis batch as those previously used by Khan et al.21 Thorough characterization of the 68ZnO NPs was performed to determine the particle size and crystal structure of the material. Based on TEM image analysis the NPs have a primary particle size of 7.8 ± 1.2 nm (n = 100). Full details of the preparation and characterization of the two 68Zn-E forms are presented in the Supporting Information.

2.2. Preparation of the Exposure Medium. Eight soil samples with 600 g dry weight each were prepared. This encompasses two controls and three samples each for treatment with 68ZnO NPs and dissolved 68Zn. The latter were dosed to a 68Zn-E concentration of 5 mg kg⁻¹ by adding 4.04 mL of the corresponding 68Zn-E stock suspension or solution. This is equivalent to the PEC, modeled by Boxall et al.5 for sewage treated soils, assuming a 16% market penetration for ZnO NPs in sunscreens.

The resulting concentration of DEG in the 68ZnO NP spiked soils was 7.5 g kg⁻¹. This concentration was matched in the soils spiked with dissolved 68Zn and one of the control samples by adding 4.04 mL of DEG. At this level, DEG is not expected to significantly alter the solubility or behavior of the 68ZnO NPs once they are mixed into the test soils. This assumption is supported by recent work,23 which demonstrated that the addition of a similar ZnO NP suspension in DEG to an aqueous exposure medium led to rapid agglomeration and dissolution of the NPs, as the stabilizing effect of DEG was soon lost. To investigate whether the presence of DEG had an observable effect on the behavior and mortality of L. rubellus, a separate one-week trial experiment was carried out prior to the main set of exposures. A report on the effects of ethylene glycol, which has a very similar molecular structure to DEG, was shown to result in 25% mortality of the earthworm Eisenia fetida when exposed at a concentration of 20 g kg⁻¹.24

Thorough mixing of the 68Zn-E and DEG with a spatula, followed by the addition of water to achieve a moisture content of 27% (w/w), corresponding to 50% of the maximum WHC, were performed to achieve a homogeneous distribution of the reagents. The soils were then left to equilibrate for 2 days prior to the introduction of earthworms.
2.3. Test Organisms. The test organism in this study was the earthworm <i>Lumbricus rubellus</i>. The specimens were collected from the field (Lasebo BV, Nijkerkerveen, The Netherlands) and maintained in a laboratory culture. Worms were kept in moist soil and fed with horse manure free of any pharmaceuticals. Two days prior to exposure, adult worms with developed clitellum and wet weights ranging from 1.08 to 2.02 g were transferred to clean LuFa 2.2 soil at 50% of WHC for acclimatization and kept in the dark at a temperature of 12 ± 2 °C.

After 2 days acclimatization, 90 earthworms were rinsed with distilled water, blotted dry on filter paper, placed in Petri dishes lined with moist filter paper and allowed to void their guts for a further 2 days, with the filter paper being changed after 24 h to avoid coprophagy. For <i>L. rubellus</i>, egestion for 48 h has been shown to be sufficient for depuration.24 Six control earthworms were selected at this stage, which were used to confirm the natural Zn isotope composition of unexposed acclimatized earthworms. The remaining 84 individuals were split into 2 groups of 42. One of the groups was prevented from oral ingestion so that only dermal uptake through the body wall was possible. This was achieved by removing mucus from the mouth region before sealing by dipping into medical histacyrl glue (Braun aesculap, Germany) using the method described by Vijver et al.25 The glue was allowed to dry before introducing the earthworms into the test soil.

2.4. Exposure Protocol and Sampling. After acclimatization and prior to introduction of the earthworms, representative soil samples of ~2 g were taken and oven-dried at 60 °C to obtain the dry weights of the samples that are needed for the determination of Zn concentrations. Soil pore water was collected prior to and following completion of the exposure period. To this end, ~30 g soil aliquots from each of the eight 600 g treatments were placed in 50 mL Falcon tubes and saturated to their maximum WHC. The tubes were centrifuged (J2-HC, Beckman Coulter, California) for 90 min at 4000 g to separate the soil detritus from the pore water before pipetting the latter from the top of the tube. The pore water samples were then frozen for storage prior to analysis.

To make optimal use of the available 66ZnO NPs, the biokinetic experiments were conducted in two stages over a 6-day period, whereby earthworms were sampled after 4, 8, and 24 h in stage 1 and after 36, 48, and 72 h in stage 2. With this approach, it was possible to collect samples at twice as many sampling time points in comparison to a single stage procedure. A full outline of the exposure protocol and sample timing is given in Table S1. In the first exposure stage, three sealed and unsealed earthworms each, with a mean wet weight of 1.62 ± 0.21 g (1 sd; range = 1.15−2.02 g), were added to each of the eight soil replicates, one control dosed with DEG and one nondosed control. One sealed and one unsealed earthworm were removed from the test soils dosed with 68Zn-E after 4, 8, and 24 h of exposure. The sealed and unsealed earthworms were easily distinguishable as the glue over the mouthparts of the sealed specimens was readily visible (Figure S2). Once the first exposure stage was completed, a second separate set of three sealed and three unsealed earthworms, with a mean wet weight of 1.40 ± 0.25 g (1 sd; range = 1.11−2.21 g) were added back into each test soil. For this second exposure set, one sealed, and one unsealed earthworm each were removed from the containers at 36, 48, and 72 h. The earthworms exposed to the control soils were removed after 72 h.

Throughout the 6-day exposure, the burrowing behavior and appearance of the earthworms was recorded. After sampling, the earthworms were prepared and allowed to void their guts as outlined above. None of the sealed earthworms excreted soil particles, confirming that ingestion was successfully prevented. The earthworms were snap-frozen and freeze-dried for 2 days, and then, the dried tissue was weighed.

2.5. Sample Preparation and Isotopic Analysis. The samples of soil, pore water and <i>L. rubellus</i> were prepared for analysis in the clean room facilities of the Imperial College MAGIC Laboratories. First a microwave digestion procedure was used to obtain sample solutions, which were processed through a one-step anion exchange column chemistry procedure for removal of the sample matrix to obtain pure Zn sample solutions for isotopic analysis.26 Full details of the digestion and chemical separation protocols can be found in the Supporting Information. The subsequent isotopic analyses were also conducted in the MAGIC Laboratories using a Nu Plasma HR MC-ICP-MS instrument, following methods outlined by Larner and Rehákemper.20

2.6. Data Reduction. The measured diagnostic 68Zn/66Zn ratio (R<sub>meas</sub>) of a sample is a function of the proportion of 68Zn-E that is present relative to the natural Zn background, and the degree of isotopic enrichment (for the 68Zn-E). As the isotope compositions of the enriched and the natural endmember are known (Table S3), the proportion of 68Zn-E to total Zn (fr<sub>en</sub>) in a sample can be calculated as

\[
fr_{en} = \left( \frac{68\text{Ab}_{\text{meas}} - (R_{\text{meas}} \times 68\text{Ab}_{\text{nat}})}{(R_{\text{meas}} \times 68\text{Ab}_{\text{nat}}) - 68\text{Ab}_{\text{nat}}} + 1 \right)^{-1}
\]

Here, Ab with a prefix of 68 or 66 refers to the relative (molar) abundance of the corresponding isotope in 68Zn-E or natural Zn, as identified by the suffix en or nat, respectively.

From eq 1 the abundance of each isotope of Zn in a sample can be calculated. For example, the isotopic abundance of 66Zn in a sample (66Ab<sub>meas</sub>) can be determined by

\[
66\text{Ab}_{\text{meas}} = (fr_{en} \times 66\text{Ab}_{\text{nat}}) + (1 - fr_{en}) \times 66\text{Ab}_{\text{nat}}
\]

In this study, the molar amount of 68Zn for each sample (N<sub>68Zn_m</sub>) was determined by comparing the measured 68Zn ion beam intensity to a suitable calibration curve, whereby admixed Cu was used as internal standard. The total molar amount of Zn in the sample (N<sub>Zn_m</sub>) was then calculated by dividing the molar amount of 68Zn in the sample by the isotopic abundance of 66Zn:

\[
N_{Zn_m} = \frac{N_{68Zn_m}}{66\text{Ab}_{\text{meas}}}
\]

The molar amounts of 68Zn-E (N<sub>68Zn</sub>) and natural Zn (N<sub>Zn_en</sub>) were derived using the proportion of 68Zn-E in the sample (eq 1):

\[
N_{Zn_en} = N_{Zn_m} \times fr_{en}
\]

The mass amounts of 68Zn-E (M<sub>68Zn</sub>) and natural Zn (M<sub>Zn_en</sub>) in the sample were determined by multiplying the molar amount by the corresponding atomic weight of 68Zn-E (AtW<sub>68Zn</sub>) or natural Zn (AtW<sub>Zn_en</sub>) (Table S2).

\[
M_{Zn_m} = N_{Zn_m} \times \text{AtW}_{\text{68Zn}}
\]

\[
M_{Zn_en} = N_{Zn_en} \times \text{AtW}_{\text{Zn_en}}
\]

To detect and quantify the presence of 68Zn-E in a sample, the isotopic analysis must identify an analytically resolvable deviation from the natural 66Zn/68Zn ratio. The smallest clearly resolvable isotopic deviation in 66Zn/68Zn is thereby determined by the precision of the analytical technique and the level of natural isotopic variability. The former is best defined by the between-run (external) precision of standard solution analyses, which typically featured a reproducibility of better than ±0.10‰ (2 sd). The latter is constrained by the data obtained for the control earthworm specimens, which provided a mean value with an uncertainty of ±0.13‰ (2 sd, n = 8). Based on error propagation, a deviation in Zn/<sup>68</sup>Zn (between sample and natural value) of only 0.16% is therefore analytically resolvable at the 2sd level. This is equivalent to an increase in the total Zn concentration of a sample, from the addition of 68Zn-E, of only 0.03%.

2.7. Kinetic Modeling. The rate constants for Zn uptake from soil (k<sub>i</sub>, in g<sub>soil</sub> g<sub>soil</sub>−1 h<sup>−1</sup>) and elimination (k<sub>e</sub>, h<sup>−1</sup>) were estimated by applying a one-compartment first-order kinetic model to the data for the 72-h uptake phase.27
Here \( [68\text{Zn-E}]_{\text{worm}} \) is the concentration of \( 68\text{Zn-E} \) in the earthworm tissue at a given time \( t \) (in hours), and \( [68\text{Zn-E}]_{\text{soil}} \) is the \( 68\text{Zn-E} \) concentration of the dry soil. While we employ the initial \( 68\text{Zn-E} \) concentration of the soil for these calculations, it is reasonable to assume that this value remained essentially constant, due to the large \( 68\text{Zn-E} \) budget of the reservoir. For comparison, the uptake rate constant \( k_{\text{soil}} \) (in mL tissue \( \text{g}^{-1} \text{h}^{-1} \)) was determined for the sealed earthworms using the \( 68\text{Zn-E} \) concentration of the pore water sampled after 72 h (eq 8) in place of \( [68\text{Zn-E}]_{\text{soil}} \) (eq 8). Unlike \( [68\text{Zn-E}]_{\text{soil}} \), the \( 68\text{Zn-E} \) concentrations of the pore waters changed during the exposure (Table S1). However, given that rapid dissolution of the \( 68\text{ZnO} \) NPs was confirmed, and comparable pore water \( 68\text{Zn-E} \) concentrations were observed in the exposures with \( 68\text{ZnO} \) NPs and dissolved \( 68\text{Zn} \), the application of \( [68\text{Zn-E}]_{\text{water}} \) in the calculations provides a reasonable, first-order characterization of the system. The use of a longer exposure period to achieve steady state tissue burdens, or a separate elimination period to determine the elimination rate constant, would involve sealed worms suffering from starvation. Our approach was therefore deemed more suitable for the kinetic modeling.

The uptake and elimination rate constants obtained with eq 8 can be used to calculate the kinetic bioaccumulation factor, BAF\(_{\text{soil-water}}\), from the ratio of the uptake and elimination rate constants:

\[
\text{BAF}_{\text{soil-water}} = k_1/k_2
\]

A bioaccumulation factor can also be determined for any time point from the ratio of the \( 68\text{Zn-E} \) concentrations in the earthworms relative to the surrounding medium. Here, BAFs were calculated for the 72-h exposure period, using the earthworm tissue burdens determined at 72 h (eq 11) and the (essentially constant) \( 68\text{Zn-E} \) concentrations of the soils or the \( 68\text{Zn-E} \) contents of the pore waters as measured at the 72-h time point:

\[
\text{BAF}_{\text{soil-water}} = \frac{[68\text{Zn-E}]_{\text{worm}}}{[68\text{Zn-E}]_{\text{soil}}}
\]

However, BAFs are most reasonably determined from the steady state tissue burdens that are obtained in exposures, which are long enough to establish a full dynamic equilibrium between \( 68\text{Zn} \) uptake and elimination. The calculated uptake and elimination rate constants allowed the modeling of a 30-day exposure period to estimate steady state \( 68\text{Zn-E} \) tissue burdens (eq 12). The steady state BAFs can then determined from the ratio of \( 68\text{Zn-E} \) concentrations of the soils or as determined for pore water after 72 h of exposure:

\[
\text{BAF}_{\text{soil-water}} = \frac{[68\text{Zn-E}]_{\text{worm}}}{[68\text{Zn-E}]_{\text{soil}}}
\]

Investigation of the biokinetics revealed that the uptake rate constants \( k_1 \) for \( 68\text{Zn-E} \) accumulation by \( L. \text{rubellus} \) from soil are an order of magnitude higher for the unsealed compared to sealed worms (Table 1) and this difference is significant based on a generalized likelihood ratio test. In contrast, the \( k_1 \) and \( k_{\text{soil}} \) values determined for uptake of the two forms of \( 68\text{Zn-E} \) in exposures of unsealed or sealed earthworms were similar, whereby \( k_{\text{soil}} \) is about twice as large as \( k_1 \). Furthermore, the calculated elimination rate constants were also identical, within uncertainty, for the two forms of \( 68\text{Zn-E} \), regardless of whether the exposed earthworms were sealed or unsealed (Table 1).

The results from the modeled 30-day exposure (Figure S5) show that for dermal-only uptake, steady state \( 68\text{Zn-E} \) tissue concentrations of \( \sim 0.3 \text{ mg kg}^{-1} \) are reached after approximately 1 week for exposures to both \( 68\text{ZnO} \) NP and dissolved \( 68\text{Zn} \). A longer period of 2 to 3 weeks is required until a dynamic equilibrium is achieved for the exposures with dermal and dietary accumulation. In this case, the modeling suggests steady state \( 68\text{Zn-E} \) tissue concentrations of \( \sim 11.5 \) and \( \sim 16.5 \text{ mg kg}^{-1} \) for the \( 68\text{ZnO} \) NP and dissolved \( 68\text{Zn} \) exposures, respectively.

Figure 1. Enriched \( 68\text{Zn-E} \) concentrations in earthworms exposed to \( 68\text{ZnO} \) NPs (red squares and dashed line) and dissolved \( 68\text{Zn} \) (blue circles and solid line): (a) unsealed earthworms (dermal and oral uptake) and (b) sealed earthworms (dermal uptake only). The best-fit lines were calculated using the modeled results and the uptake and elimination rate constants given in Table 1.

\[
(\text{eq 8})
\]

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3. RESULTS

3.1. Soil and Pore Water. The control samples all showed \( 68\text{Zn} \) and \( 66\text{Zn} \) isotope ratios that are within the expected natural variability and natural \( Zn \) concentrations that are comparable to the exposed samples (Tables S4 and S5). The soil samples dosed with \( 68\text{ZnO} \) NP and dissolved \( 68\text{Zn} \) had initial measured dry weight \( 68\text{Zn-E} \) concentrations of 3.58 ± 0.75 (1 sd, \( n = 3 \)) and 3.51 ± 1.13 mg kg\(^{-1}\) (1 sd, \( n = 3 \)), respectively (Table S5). These concentrations are identical, within uncertainty, and show a recovery of >70% for the \( 68\text{Zn-E} \) that was added to the soils at 5 mg kg\(^{-1}\). The pore water samples collected after 2 days of acclimatization from soils spiked with \( 68\text{ZnO} \) NPs and dissolved \( 68\text{Zn} \), had \( 68\text{Zn-E} \) concentrations of 61.6 ± 14.9 (1 sd, \( n = 3 \)) and 53.6 ± 6.9 μg L\(^{-1}\) (1 sd, \( n = 3 \)). At the end of the 6-day experiment, the pore water concentrations had increased to 96.2 ± 6.3 (1 sd, \( n = 3 \)) and 119.7 ± 0.2 μg L\(^{-1}\) (1 sd, \( n = 3 \)) for the \( 68\text{ZnO} \) NP and the dissolved \( 68\text{Zn} \) exposure, respectively (Table S1 and Figure S3).
Although owing to the low levels of $^{68}$Zn-E used and the nature of the experiments, these diﬀerences are smaller than the errors associated with the modeling and the measured data (Table 2). This demonstrates that $^{68}$Zn-E from ZnO NPs is as bioavailable as the ionic form of Zn. In contrast, the low Zn dosing levels employed in previous studies are lower than the respective values determined from the modeled steady state $^{68}$Zn-E tissue concentrations. For example, dermal-and oral uptake is associated with BAF$_{soil}$ and BAF$_{water}$ values representing the rate of loss of accumulated $^{68}$Zn-E and the fit of the modeled trend to the data plotted in Figure 1, respectively. Both are therefore independent of the [Zn-E]$_{soil}$ and [Zn-E]$_{water}$ concentrations used to determine $k_1$ and $k_{1-water}$.

“The quoted uncertainties for the uptake and elimination rate constants are ±1 sd. Sealed earthworms. Unsealed earthworms. The $k_1$ and $R^2$ values represent the rate of loss of accumulated $^{68}$Zn-E and the fit of the modeled trend to the data plotted in Figure 1, respectively. Both are therefore independent of the [Zn-E]$_{soil}$ and [Zn-E]$_{water}$ concentrations used to determine $k_1$ and $k_{1-water}$.

### Table 2. Summary of Bioaccumulation Factors (BAF) Determined for Uptake of $^{68}$Zn-E by Earthworms, As Defined in Eqs 10–14

| BAF$_{soil-water}$ | BAF$_{soil}$ | BAF$_{water}$ |
|-------------------|-------------|---------------|
| $^{68}$ZnO NPs     | 0.06        | 2.33          |
| BAF$_{water}$-ss   | 0.10$^{a}$  | 12.6$^{a}$    |
| BAF$_{water}$-ss   | 2.84$^{d}$  | 142.2$^{d}$   |

Sealed earthworms. Unsealed earthworms. BAF$_{soil}$ = BAF$_{soil-water}$ BAF$_{water}$ = BAF$_{water}$ Labeled as average BAF for exposures of unsealed earthworms in soil at pH 6.4 by Heggelund et al. Labeled as average BCF for exposures of unsealed earthworms in soil at pH 6.4 by Heggelund et al.

(Figure S5). These results were used to calculate the steady state BAFs of Table 2.

As expected, the BAF values calculated from the $^{68}$Zn-E tissue concentrations measured at 72 h (BAF$_{soil-water}$) are lower than the respective values determined from the modeled steady state tissue burdens (BAF$_{soil-water}$), and these diﬀerences are larger for unsealed earthworms with combined dermal and oral uptake of $^{68}$Zn-E (Table 2). Furthermore, all BAF data obtained for exposures with $^{68}$ZnO NP are either identical or remarkably similar to the results acquired for experiments with added dissolved $^{68}$Zn (Table 2). Notably, the BAF$_{water}$ results for the dermal-only uptake of sealed earthworms are also very similar to the BAF$_{soil}$ data for dermal and oral uptake, using both the 72-h and the modeled steady state $^{68}$Zn-E tissue concentrations. For example, dermal-only uptake yields BAF$_{water}$ ≈ 2.8 and 3.7, while dermal and oral uptake is associated with BAF$_{soil}$ ≈ 3.4 and 4.9 (Table 2).

### 4. DISCUSSION

#### 4.1. Zinc Behavior in Soil

Care was taken to achieve a homogeneous distribution of added $^{68}$Zn-E in the test soils but this was not specifically tested. However, the data obtained for three soils indicate that a good distribution was achieved, although owing to the low levels of $^{68}$Zn-E used and the nature of soil, some minor heterogeneity (of ±20 to 30%) is difficult to avoid (Table S5).

After the 2-day acclimatization period, the $^{68}$Zn-E concentrations in the two exposure systems were identical, within error, for both soils and pore waters (Table S5). The after the 6-day exposure period, the $^{68}$Zn-E concentration of the pore water had increased for both treatments, whereby the end point concentration was ~24% higher when dissolved $^{68}$Zn was added (Table S5). This supports the suggestion that at these low soil concentrations the NPs underwent rapid dissolution during the acclimatization period and that the two forms of Zn display similar environmental behavior and partitioning, with comparable amounts binding to the available soil surfaces and entering the pore water phase.

These results can be compared to the study of Heggelund et al., where soils were dosed with ZnO NPs and ZnCl$_2$ at much higher Zn concentrations of 238–2500 mg kg$^{-1}$ (Figure S1). After acclimatization for 10 days, these authors found that soils treated with ZnCl$_2$ had pore water Zn concentrations that were 20–50-fold higher compared to equivalent soils dosed with ZnO NPs. The largest diﬀerences were hereby seen at high levels of Zn addition. The NPs used by these workers had a nominal particle size of 30 nm, larger than the particles used here, and therefore dissolution rates are expected to be slower. However, the larger particle size is unlikely to be primarily responsible for the distinct Zn pore water concentrations. Rather, the distinct Zn partitioning observed by Heggelund et al. is most likely a consequence of the high Zn dosing levels that were employed and not characteristic of the Zn behavior observed at the lower, environmentally relevant, concentrations used here.

Furthermore, it was observed in previous studies that the treatment of soils with high concentrations of ZnO NPs or dissolved Zn can influence the soil pH. Such changes in pH will have an impact on soil toxicity, hence making it difficult to compare any eﬀects and toxicity between the two metal forms, as the pH modulating eﬀects diﬀer between the NP and ionic forms of Zn. In contrast, the low Zn dosing levels employed in this study will be buffered by the soil and are not expected to have any impact on soil chemistry.

#### 4.2. Zinc Uptake and Accumulation

In this study, uptake of the two $^{68}$Zn-E forms by the earthworm L. rubellus yielded essentially indistinguishable results. However, accumulation of $^{68}$Zn-E in the unsealed worms leads to $^{68}$Zn-E tissue concentrations that are about a factor of 20 higher than in sealed specimens (Figure 1 and Table S5). This demonstrates that $^{68}$Zn-E from ZnO NPs is as bioavailable as the ionic form and that oral ingestion constitutes the major uptake pathway, while uptake through the body wall plays a minor but nonetheless significant role contributing about 5% of total uptake. The exposure of the earthworms in two sequential stages, whereby the second set of earthworms was added to the
In this study, the major pathway for uptake of Zn into earthworm tissue was oral ingestion of pore water and soil. Dermal uptake accounted for only ~5% of the total Zn uptake, primarily via absorption of dissolved Zn associated with pore water. Importantly, the results of this study demonstrate that the behavior and bioavailability of ZnO NPs, when introduced at an environmentally relevant concentration, are indistinguishable from dissolved Zn. This implies that nanocotoxicology hazard studies which employ high exposure concentrations, may lead to conclusions, for example on element uptake forms and kinetics, that are not directly relevant for environmentally relevant scenarios. Thus, any nanospecific effects found at high exposure levels may indeed not be relevant for risk assessments. However, the determination of changes in NP speciation is technically challenging, as the detection limits of the X-ray based techniques that are employed for such studies are much higher in comparison to the isotope tracing methods used here.

**ASSOCIATED CONTENT**

*Supporting Information*

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.est.5b03413.
Natural Zn concentrations and predicted environmental concentrations of ZnO NPs (Figure S1), details of the synthesis and characterization of the $^{65}$Zn-E forms, outline of the experimental protocol and sample timing (Table S1), images of earthworms with sealed mouthparts (Figure S2), sample preparation and chemical separation protocols (Table S2), isotope abundances and atomic weights of natural Zn and $^{65}$Zn-E (Table S3), experimental results for control samples (Table S4), summary of experimental results for exposed samples (Table S5), concentration of $^{65}$Zn-E in pore waters (Figure S3), concentration of natural Zn and $^{65}$Zn-E in exposed samples (Figure S4), and modeled uptake of $^{68}$Zn-E over a period of 30 days (Figure S5) (PDF).

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**Notes**
The authors declare no competing financial interest.

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