

# QUANTIFYING EXCESS HEAVY METAL CONCENTRATIONS IN DRAINAGE BASINS USING CONSERVATIVE MIXING MODELLING

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## ABSTRACT

High concentrations of heavy metals and other pollutants in river sediments can have detrimental effects on the health of ecosystems and humans. As such, determining the composition of river sediments throughout drainage basins is an essential aspect of environmental monitoring. Well characterised natural baseline concentrations are important information with which observations can be compared to identify potentially harmful pollutants. In this study, forward and inverse mixing models are used to map natural baselines and elemental concentrations in river sediments in the upper reaches of the Clyde drainage basin, UK. Continuous baselines are generated using forward mixing models parameterised with 1185 measurements of elemental concentrations from first-order streams. Calculated baselines are compared to concentrations measured at 60 localities along the main channel of the Clyde river. For a range of major and trace elements (e.g. Mg, Sr, K), the downstream observations are in close agreement with the baseline concentrations predicted from conservative mixing. However, some heavy metal concentrations (Pb, Cu, Zn) tend to exceed the predicted baseline concentrations. Assuming conservative mixing, the excess elemental masses required to match these observations are calculated by inverting ('unmixing'), the concentrations of downstream samples. We tentatively suggest that anthropogenic activity inserts 9.7 Mg/yr of Pb, 1.5 Mg/yr of Cu and 5.7 Mg/yr of Zn into the sediments of the river Clyde. Finally, a strategy for mapping toxic elemental concentrations along rivers and limitations on model resolution imposed by the spatial distribution of the data are discussed.

**Keywords** River pollution · Clyde, Scotland · Geochemical modelling · Heavy metal · Mixing

## 1 INTRODUCTION

It is generally agreed that the composition of river sediments depends principally on the mixture of elements in upstream source regions (e.g. Ercolani et al. 2019; Caracciolo 2020; Lipp et al. 2020). A corollary is that the composition of river sediments provide information about source region compositions (e.g. Lipp et al. 2021). Weathering, cation exchange and sorting play moderating roles (e.g. Bouchez et al. 2011; Lupker et al. 2012; Tipper et al. 2021). We build on these basic assertions to, first, demonstrate how simple conservative mixing models can be used to generate 'natural' baseline concentrations of elements using high-resolution geochemical maps of source regions. We demonstrate this in a case-study focused on the upper reaches of the Clyde river, Scotland. Second, we show how, using these environmental baselines, excess, likely anthropogenic, contributions to sediment heavy metal (Pb, Cu, Zn) concentrations, which are often associated with poor ecological health, can be quantified.

This study has three parts. First, we generate predictions from conservative mixing models that assume the composition of river sediments is set by natural 'geologic' input. Formally, the measured concentrations of first-order stream sediments are integrated with respect to upstream area to predict compositions of river sediments downstream. These predicted downstream concentrations are used as a natural baseline concentration to which observations can be compared. For some elements (e.g. Mg, Sr, K), observed concentrations are tightly distributed around the baseline predictions. For some heavy metals however (Pb, Cu,

Zn) we observe discrepancies between observed and baseline concentrations along the Clyde river. Consequently, we, secondly, quantify the excess, likely anthropogenic, input required to explain observed discrepancies. Finally, we use these results to tentatively identify loci where heavy metal concentrations exceed toxic limits along the Clyde river.

### 1.1 Clyde drainage basin

We focus on the upper reaches of the Clyde river, Scotland, where the composition of sediments at 60 localities along the main channel upstream of Glasgow were measured by the British Geological Survey (BGS) as part of the Clyde Urban Super Project (CUSP; Figure 1; Fordyce et al. 2017). The upper Clyde Basin covers an area of 1873 km<sup>2</sup> and is mainly composed of sedimentary rocks: sandstone and wackes (Figure 1b). Felsic and mafic intrusions can be found throughout the region including ~ 10 km<sup>2</sup> of extrusive igneous rock southwest of Glasgow. This region has been home to intensive mining and related industry for the last several centuries (Skillen 1987). Economic decline after World War I saw much of the industry abandoning the city, leaving behind high rates of chronic disease and one of the lowest life expectancies in Europe (Kintrea and Madgin 2019). Environmental remediation was a key aspect of the regions successful revitalisation (Maver 2019). The CUSP, a large scale geochemical surveying campaign to understand the current state of the environment in the Clyde Basin and estuary<sup>2</sup>, was created to support these efforts (Campbell et al. 2010;

<sup>2</sup>see: <https://www.bgs.ac.uk/geology-projects/cusp/>

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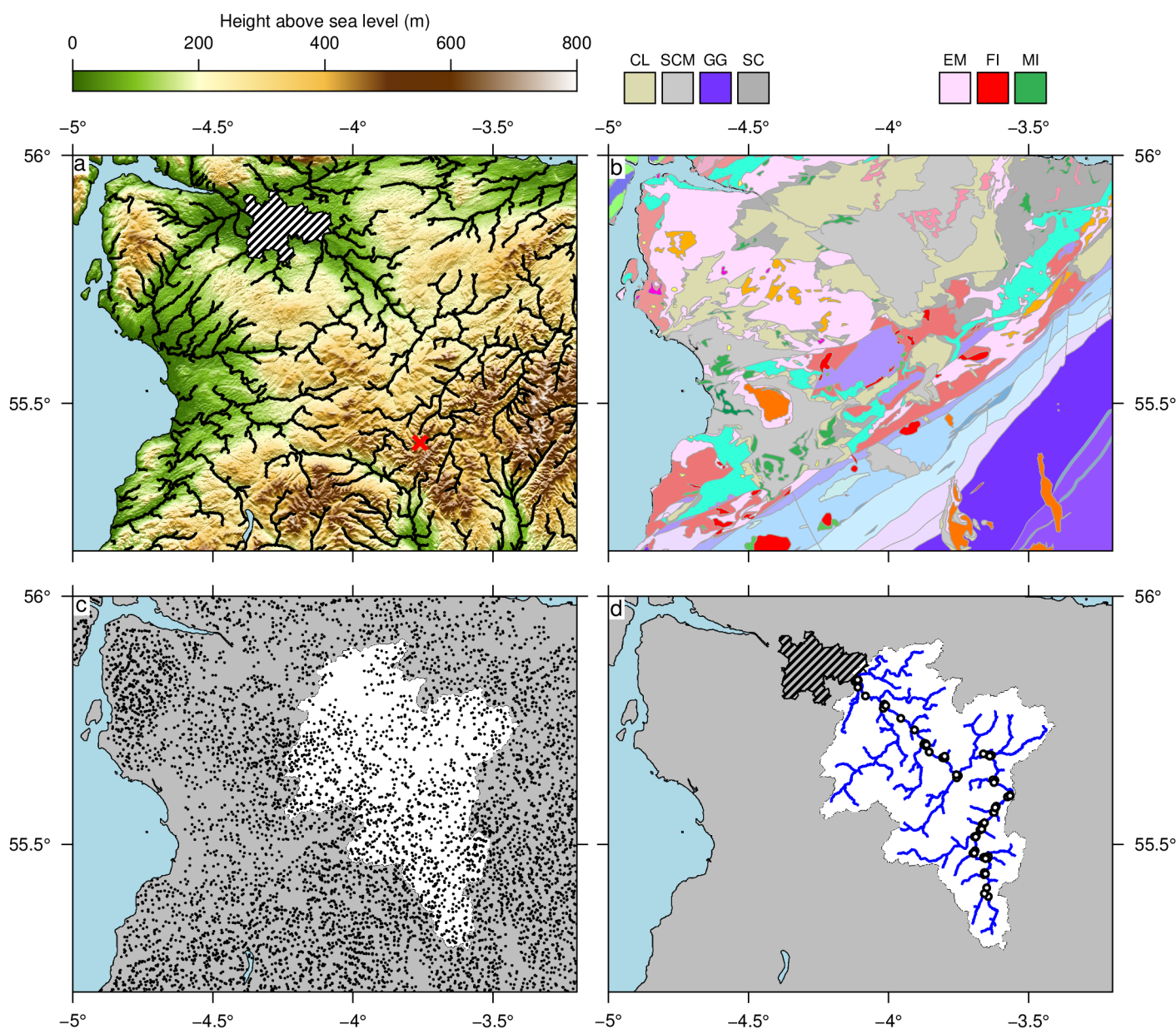


Figure 1: **Study area: Clyde basin, Scotland.** (a) Topography (SRTM 1 arc second) and extracted drainage network (Farr and Kobrick 2000; De Jager and Vogt 2010). Hashed polygon = Glasgow city area. Red cross = Lowther Hills Pb mining area. (b) Simplified geologic map. CL: Clackamannan Group (fluvial sandstones), SCM: Scottish Coal Measures (Mud/Silt/Sandstone), GG: Gala Group (Wackestone), SC: Strathclyde Group (estuarine sedimentary rock), EM: Extrusive Mafic lava/tuff, FI: Felsic Intrusions, MI: Mafic Intrusions. (c) Black circles = G-BASE samples of first-order stream sediments. White polygon = Upper Clyde drainage basin, the focus of this study. (d) White circles = geochemical samples along Clyde river collected as part of the Clyde Urban Super Project (CUSP).

Appleton et al. 2013; Morrison et al. 2014). In ten years, this campaign gathered 5943 stream, soil and water samples and analysed them for a range of parameters including elemental chemistry (Fordyce et al. 2017).

In this study, river sediment samples collected along the main channel of the Clyde during the CUSP are combined with first-order stream samples from the BGS’s Geochemical Baseline Survey of the Environment (G-BASE) to quantify heavy metal pollution in the region through conservative mixing models.

## 2 DATA

### 2.1 G-BASE

Two pre-existing geochemical data sets are used in this study. First, elemental concentrations in stream sediments, recorded in G-BASE, are used to quantify source region composition. G-BASE was a multi-decade (1960-2014) sampling campaign with the goal of generating a geochemical baseline for the UK (Johnson et al. 2005). Sediment samples from first-order streams were obtained with an average sampling density of 2 km<sup>2</sup>. Full sampling and analytical protocols for G-BASE are given in Johnson et al. (2018b) and Johnson et al. (2018a). Figure 1c shows the location of the 1185 samples within the Clyde basin; note that the Glasgow urban area has the lowest sampling density. Through sieving, the grain-size fraction < 150 μm was extracted and analysed using X-ray Fluorescence (XRF) and direct reading optical emission spectroscopy. Figure 2 shows that the elemental concentrations of first-order streams are highly correlated with underlying lithologic units (e.g. high Mg in the south-east coincides with layers of wacke). We consider stream sediment compositions as source region concentrations in this study because chemical weathering has already occurred prior to the sediment reaching the streams.

### 2.2 Upper Clyde Sediments

The second dataset is a suite of sediment elemental concentrations measured at 60 localities along the main Clyde channel, and select tributaries. These samples were collected as part of the Upper River Clyde Sediment Survey, a section of the CUSP. These river bed samples were collected along the main trunk of the River Clyde and at the mouths of major tributaries (Smedley et al. 2017). They have been analysed in accordance with G-BASE protocols (grain-size fraction < 150 μm). Their distribution is shown in Figure 1d. Note that additional CUSP samples were also collected further downstream along the Clyde (Jones et al. 2017). These samples are not considered further here because a different grainsize fraction (< 2 mm) relative to the rest of the dataset was used.

## 3 METHODS

Two computational methods are used in this study. First, continuous concentrations of elements along rivers (‘natural’ baselines) are predicted using a simple conservative mixing (forward) model (Lipp et al. 2020). Secondly, the CUSP (higher-order river) samples are inverted (‘unmixed’) to predict concentrations of elements in upstream source regions. For both the forward and inverse model a stream network must be defined. First, the

SRTM1s DEM is downsampled onto a 100 × 100 m grid and loaded into the Python package LandLab 2.3.0 (Hobley et al. 2017; Barnhart et al. 2020). Secondly, the topography is filled using the priority flood method to avoid artificial depressions (Barnes et al. 2014). Finally, the D8 flow-routing algorithm is used to generate the stream network from the SRTM 1s topographic data (O’Callaghan and Mark 1984; Farr and Kobrick 2000). Rivers in this study are defined by drainage areas > 8 km<sup>2</sup>.

### 3.1 Forward Model

Assuming conservative mixing, there are two controls on the downstream concentration  $D(x)$  of an element, upstream source concentration,  $C(x)$ , and the vertical erosion rate  $\partial z/\partial t$ , which yields the following relation

$$D(x) = \frac{1}{\int_A \partial z/\partial t dA} \int_A \frac{\partial z}{\partial t} C(x) dA \quad (1)$$

where  $x$  is a location along a streamline,  $A$  is the upstream area,  $z$  is elevation and  $t$  is time. In compact notation  $D(x) = F(C)$ . In other words, river sediment compositions are a function of upstream source region compositions propagated and mixed downstream. Forward modelling in the Cairngorms, Scotland, suggests that different erosional models have a minor impact on the conservative mixing of stream sediments (Lipp et al. 2020). Hence we assume a constant erosion rate  $\partial z/\partial t = k$  throughout space. This simplifies Equation 1 to

$$D(x) = \frac{1}{|A(x)|} \int_A C(x) dA, \quad (2)$$

where  $|A(x)|$  is the total area upstream. Note that in the following sections we have dropped the  $x$  notation for simplicity.

To generate forward predictions of downstream geochemistry, the composition of sediments entering the drainage network needs to be known. To generate a continuous input field, the G-BASE samples are interpolated to a 100 × 100 m grid using continuous curvature splines (Smith and Wessel 1990). This grid is then used to parameterise the forward model. Equation 2 is implemented using LandLab and solved with the interpolated G-BASE grid in an equal-area projection.

### 3.2 Inverse Model

The inverse model seeks smoothly varying source region concentrations that yield low residual misfits to observed concentrations downstream. Discrete samples of downstream composition along the river network are inverted for upstream source geochemistry  $C$ . To find optimal source region compositions (i.e. the smoothest best-fitting model), the misfit between known observations  $D$  and theoretical downstream compositions  $F(C)$ , calculated from mixing of the proposed source, are minimised. In vector-space this problem can be written as

$$\|\log(\mathbf{F}(C)) - \log(\mathbf{D})\|^2 = \sum_i [\log(F(C)_i) - \log(D_{obs,i})]^2, \quad (3)$$

where bold letters denote vectors. This problem is likely to always be underdetermined, i.e. fewer observations than predictions (Lipp et al. 2021). Therefore, we seek the smoothest

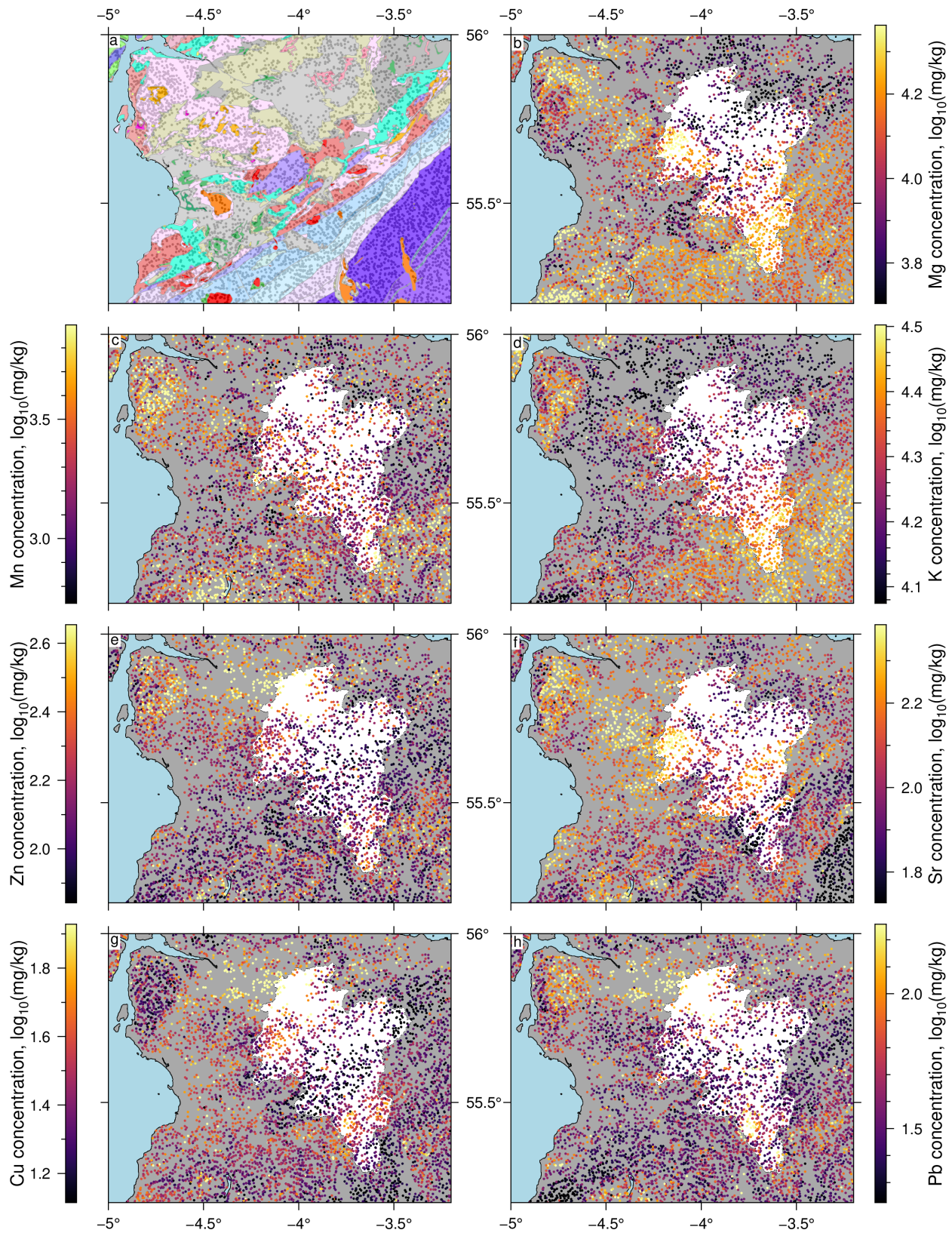


Figure 2: **Source region elemental concentrations.** (a) Geologic map and G-BASE sample localities in first-order streams (grey circles; see Figure 1). (b) Concentrations of Mg in G-BASE samples (coloured circles). White polygon = upper Clyde catchment. (c) Mn. (d) K. (e) Zn. (f) Sr. (g) Cu. (h) Pb.

model of upstream concentrations that generate theoretical concentrations downstream that have lowest residual misfit to the CUSP observations (e.g. Parker 1994). The following objective function is minimised,

$$X(\mathbf{C}) = \|\log(F(\mathbf{C})) - \log(\mathbf{D})\|^2 + \lambda \left( \left\| \frac{\delta \log \mathbf{C}}{\delta x} \right\|^2 + \left\| \frac{\delta \log \mathbf{C}}{\delta y} \right\|^2 \right), \quad (4)$$

where the first summand,  $h$ , is misfit. Model roughness,  $r$ , is calculated within the parentheses of the second summand.  $\lambda$  is a hyper-parameter controlling the effect roughness will have on the solution; increasing  $\lambda$  will create smoother solutions that fit the data less well. To calibrate  $\lambda$  we utilise Occam's approach to identify values that generate the smoothest models that adequately fit the data (i.e. low  $h$  and  $r$  values). Formally, we identify optimal  $\lambda$  values as those that yield the highest curvature of  $h(r)$  i.e. the 'elbow' in Parker's L-curves (see Supporting Information; Parker 1994).

As this inverse problem is non-linear with no obvious analytical solution, Equation 4 is minimised numerically using Powell's conjugate gradient method as implemented in SciPy (Powell 1964; Press et al. 1986). The inversion scheme was run on the Imperial College High-Performance Computing cluster with an inversion node resolution of  $1.5 \times 1.5$  km, taking between 2 and 72 hours to complete depending on the smoothing parameter used.

### 3.3 Quantifying excess concentrations

Calculating differences between predicted concentrations, from the forward and the inverse models, provides a means to quantify excess elemental masses that cannot be accounted for by the 'geologic' or natural contributions measured by G-BASE. The parsimonious procedure used to quantify excess elemental masses is as follows. First, the total measured suspended sedimentary flux of the Clyde river ( $0.11 \times 10^9$  kg yr<sup>-1</sup>; Milliman and Farnsworth 2011), is divided by the total drainage area to yield an average erosion rate of 0.059 kg m<sup>-2</sup> yr<sup>-1</sup>. The total eroded mass,  $M(x)$ , at each point in the drainage network is calculated by integrating the average erosion rate with respect to upstream area. Finally, the following equation is used to convert this mass flux into the mass flux of each element along the drainage network,

$$m(x) = C(x)M(x), \quad (5)$$

where  $C$  is the predicted fractional (i.e.  $0 < C(x) \leq 1$ ) concentration of a given element. This calculation is performed for both the inverse and forward model predictions. The difference between the two calculated mass fluxes gives the expected excess mass of a given element, relative to the natural baseline. Note that as the measured sedimentary flux inserted into this equation only considers the suspended load, calculated masses are an underestimate of the actual values. Alternative mass fluxes and erosion rates could be inserted into Equation 5 should they become available. Note that we are implicitly assuming that all grainsize fractions behave in the same way as the studied grainsize fraction.

## 4 RESULTS

### 4.1 Forward model predictions of river compositions: Towards continuous natural baselines

Figure 3 shows Mg, K, Sr and Mn concentrations predicted from the forward model,  $F(C)$ , alongside observed concentrations,  $D$ . These elements cover a large range of concentrations, as well as different chemical affinities. K and Sr are both generally associated with felsic rocks but occur as, respectively, major and trace elements. Mg and Mn contrastingly are mafic major and trace elements. Mg, K and Sr all tightly cluster around the 1:1 line in Figure 3b, indicating minimal deviation relative to the predicted natural baseline. The root-mean-squared misfit (rms) values are low,  $< 0.25$ , for these four elements; rms, is defined as

$$rms = \left[ \frac{1}{N} \sum_{i=1}^N [\log(F(C)_i) - \log(D_i)]^2 \right]^{1/2}, \quad (6)$$

where  $N$  is the number of downstream observations. A closer inspection of the distribution of Mn misfits shows that they are roughly symmetrically distributed around a misfit of 0 (see orange histogram in Figure 3c). Figure 3c shows the predictions along the main River Clyde (lines) and the observations (symbols). These results indicate that the concentrations of these elements are largely in accordance with their predicted natural baselines. Similarly accurate predictions of As, Ca, Co, Cr, Mn, Ti and Zr concentrations are shown in Supplementary Information and in a previous study of rivers draining the Cairngorms, Scotland (Lipp et al. 2020). Collectively they show that conservative mixing models can be used to approximate the natural baseline concentration of a wide-range of elements, should appropriate source region chemistry data be available. Figures showing the interpolated G-BASE measurements used to define source region compositions are given in Supplementary Information.

Observed heavy metal (Pb, Cu and Zn) concentrations, however, exceed calculated natural baselines (see Figure 4). Figure 4b shows that Cu and Zn concentrations are consistently under-predicted by 'natural' baselines calculated from the measured upstream G-BASE concentrations. Pb deviates most strongly with residuals that are broadly uniformly distributed and an rms misfit of 0.58 (see Figure 4c).

### 4.2 Inverse modelling to estimate excess (non-geologic) contributors to river compositions

To estimate heavy metal concentrations in the upper Clyde basin that best-fit observations downstream, we use the inverse model to 'unmix' the CUSP samples. We stress that this approach assumes conservative mixing and seeks smooth solutions (Figure 5b, e, h). Ideally, this procedure would make use of widely distributed samples to constrain the distribution of concentrations across the basin (cf. Lipp et al. 2021). As most of the samples available here lie along the main trunk of the Clyde river there is little resolving power in tributary watersheds (see Figures 16-17 in Supporting Information). We therefore focus on the best-fitting downstream river profiles for the upper Clyde. Figures 6-8 show the results for Pb, Cu, Zn. In these examples, optimal damping parameter ( $\lambda$ ) values for Pb, Zn and Cu are  $10^{0.2}$ ,  $10^{-0.2}$  and  $10^{-0.2}$ , respectively (see Figure 18 in Supporting Information). Unsurprisingly, the inverse models yield lower

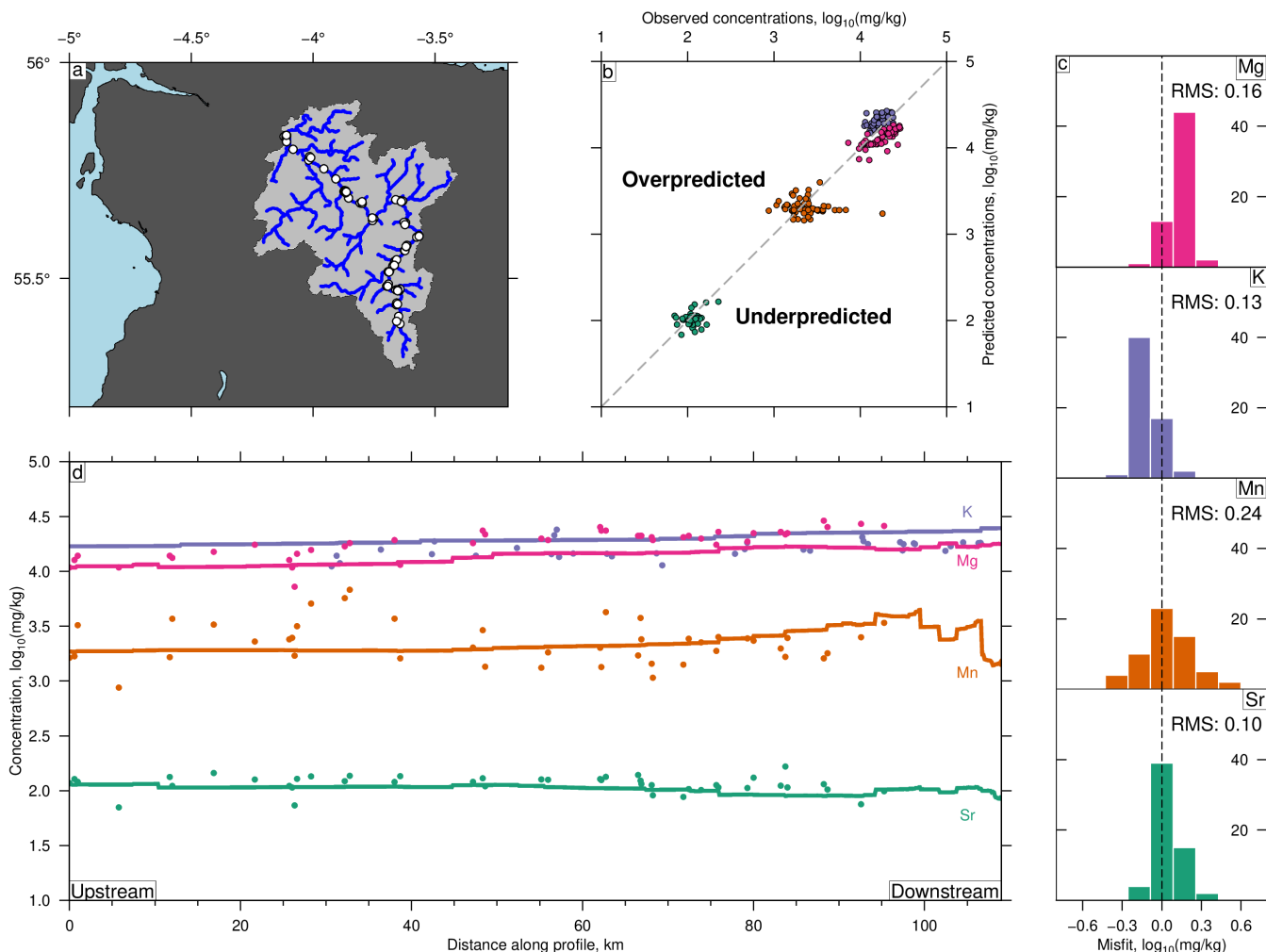


Figure 3: **Calculated baselines for select elements.** (a) Blue = upper Clyde drainage network; white circles = loci of 60 CUSP samples used to test model predictions. (b) Observed (CUSP) vs. predicted concentrations at sample sites for Mg (pink), K (purple), Mn (orange), Sr (dark green); predictions were generated from forward model (solid lines in panel d; Equation 2; see body text; Supplementary Figure 7); dashed line = 1:1 relationship. (c) Difference,  $\log(D_i) - \log(F(C)_i)$ , between observations and predictions. Bin size =  $0.15 \log_{10}(\text{mg/kg})$ ; ordinates give absolute count ( $N = 60$ ); dashed line is centred on 0. Root Mean Square (RMS) misfit for each element is annotated. (d) Circles = observed concentrations of labelled elements (CUSP; white circles in panel a); solid lines = predicted elemental concentrations along Clyde river.

residual misfits than the natural (G-BASE-derived) baselines. Predicted concentrations from the best-fitting, smooth, inverse models of Cu and Zn concentrations cluster around the 1:1 line (solid circles), more closely than the natural baseline (cf. empty circles; Figures 7b & 8b). The Pb residuals have a broader symmetrical distribution centred on 0 (Figure 6a).

## 5 DISCUSSION

### 5.1 Quantifying pollution as excess elemental masses

By comparing the two profiles (from the forward and inverse models) it is possible to infer the location of excess elemental mass sources (Equation 5). Figures 6d, 7d and 8d show estimated excess mass of Pb, Cu and Zn as a function of distance along the river. The inverse models predict higher concentrations of heavy metals in almost all of the source areas compared

to predicted natural (geologic) baselines. An exception is for Zn at distances  $> 80$  km, where the inverse model predicts lower concentrations than the forward model (Figure 8c). We note that there are few observations available for inversion at distances  $> 95$  km, as such this result could be an artefact of poor sampling. The largest increase in excess Pb occurs between  $\sim 90 - 45$  km. We interpret this result as an indication of a source of Pb that is not derived from natural erosion and weathering of the local bedrock (recorded by the G-BASE samples). An obvious possible source is the Lowther Hills, which have been a Pb mining area since the early Middle Ages, and have previously been identified as a candidate for Pb pollution of the Clyde (Rowan et al. 1995, red cross in Figure 1a). The Glengonar Waters, which drains the Lowther Hills, meets the Clyde at  $\sim 90$  km (i.e. close to the start of the relatively rapid increase in excess Pb) and was found to have increased Pb-levels

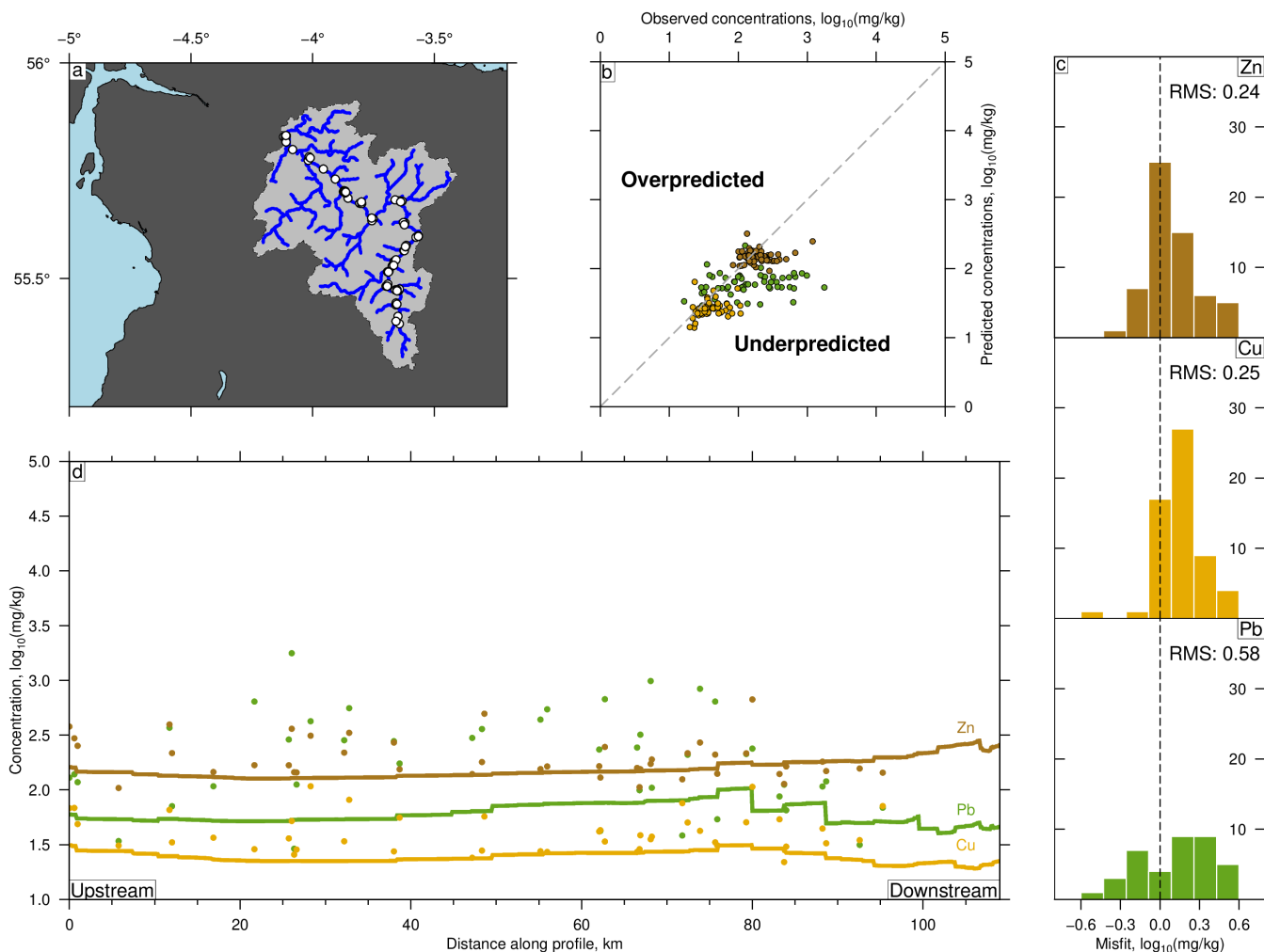


Figure 4: **Calculated baselines for heavy metals.** (a) Blue = upper Clyde drainage network; white circles = CUSP samples. (b) Observed vs. predicted concentrations for Zn (brown), Pb (green), Cu (yellow). Dashed line = 1:1 relationship. (c) Misfit (difference) between observations and predictions. Bin size =  $0.15 \log_{10}(\text{mg/kg})$ . Note annotated root-mean-square misfit (RMS); and tendency for under-prediction. (d) Circles = observed concentrations; solid lines = predicted downstream composition along Clyde river. See Figure 3 for extended caption.

in its sediment in previous studies (Oliver and Naysmith 2011). Similar, but less pronounced steps can be seen for Cu at 45 km and 80 km for Zn.

## 5.2 Identifying toxic concentrations

Heavy metal pollution in river sediments can be dangerous to human health and ecosystems. The “threshold effect level” (TEL) and “predicted effect level” (PEL) have been proposed as guides to ecotoxicity (see e.g. Hudson-Edwards et al. 2008; Oliver and Naysmith 2011). Concentrations of elements in river sediments below TEL are considered unlikely to pose significant hazard to aquatic organisms, while concentrations above PEL are often associated with significant adverse impacts on the ecology of rivers TEL/PEL, mg /kg: Pb: 35/91.3, Cu: 36.7/197, Zn: 123/315; Hudson-Edwards et al. 2008. All three heavy metals examined in this study show elevated levels compared to the predicted natural baselines. For Cu and Zn the continuous predicted concentrations derived by inverting measured concentrations

consistently exceed the TEL. For Zn five samples exceed the PEL, however the inverse result never exceeds it, which we attribute to model damping. Pb consistently exceeds the PEL after the incorporation of tributaries draining the Lowther Hills. We suggest that these methodologies show promise for identifying watersheds where remediation efforts could be focused.

## 5.3 Uncertainties

First, by using the forward model predictions as a natural baseline, we assume that the first-order stream samples are unaffected by any human influences. This assumption is probably reasonable given that G-BASE sampling guidelines explicitly state that samples ought to be taken upstream of contamination sources (Johnson et al. 2005). However, it is possible that some pollution is still present due to diffuse source (i.e. car exhausts, atmospheric fallout). We suggest that combining isotopic ‘fingerprinting’ with the computational methods we present could

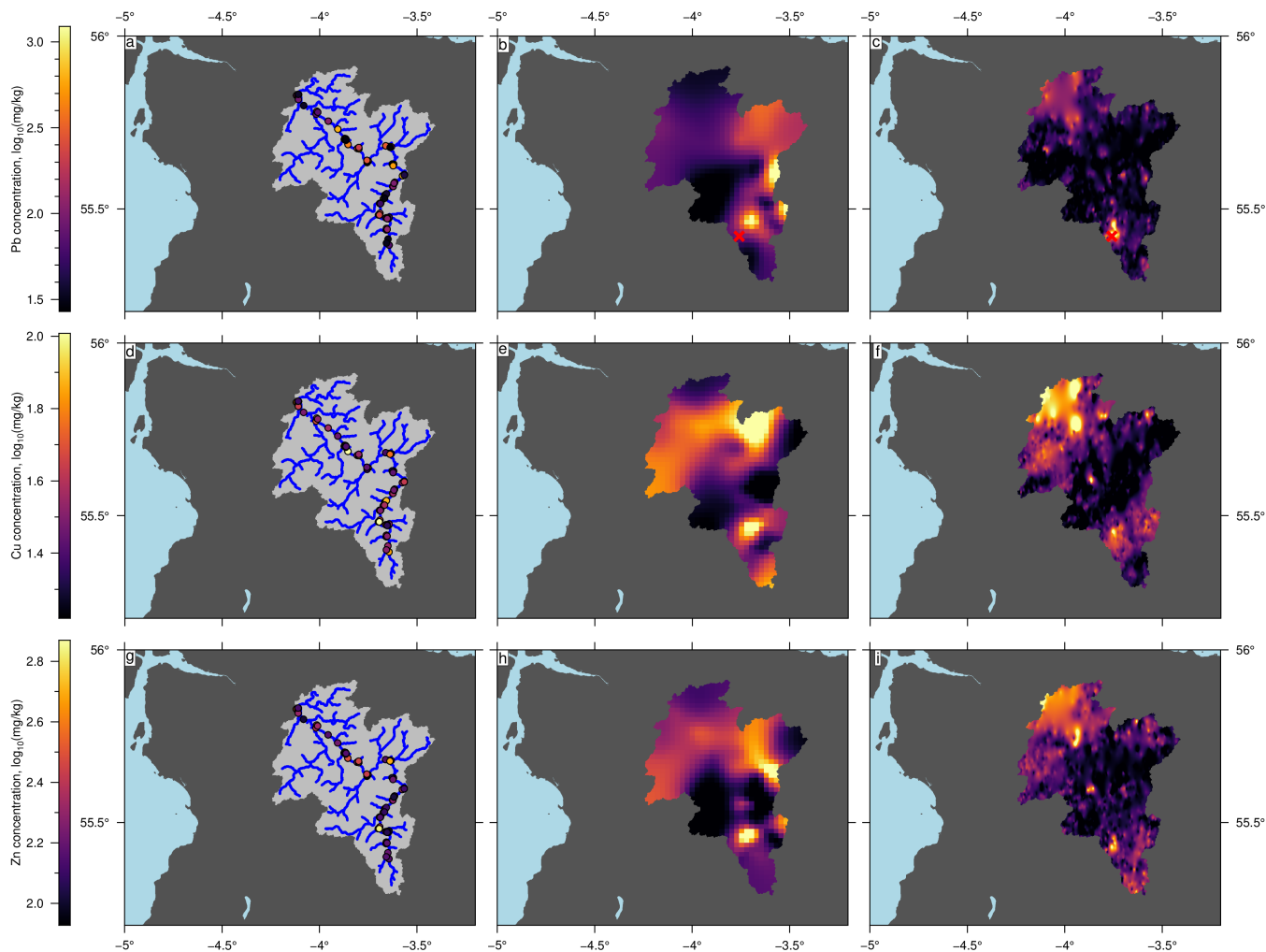


Figure 5: **Source region heavy metal concentrations.** (a, d, g): Coloured circles = Observed concentrations in CUSP samples inverted for upstream (source region) concentrations; light grey polygon/blue lines = drainage basin/network. (b, e, h): Predicted source region concentrations from best-fitting inverse model. (c, f, i): Observed ‘geologic’ source region concentrations interpolated from G-BASE measurements of first-order stream compositions (see Figure 2). (a-c) Pb. (d-f) Cu. (g-i) Zn.

be a fruitful way to identify the origins of measured elemental concentrations.

The largest source of uncertainties in predictions from the inverse model are a consequence of the distribution of downstream samples. Ideally, samples should be taken along the main river and throughout its tributaries, such that the upstream area between each sample is approximately equal. In this particular dataset however, samples were collected almost entirely along the main trunk. Consequently, the inverse model is unable to resolve variations in source concentrations along tributaries (see Supplementary Information: Figure 17). This limitation is straightforward to remedy with additional samples (see e.g. Lipp et al. 2021).

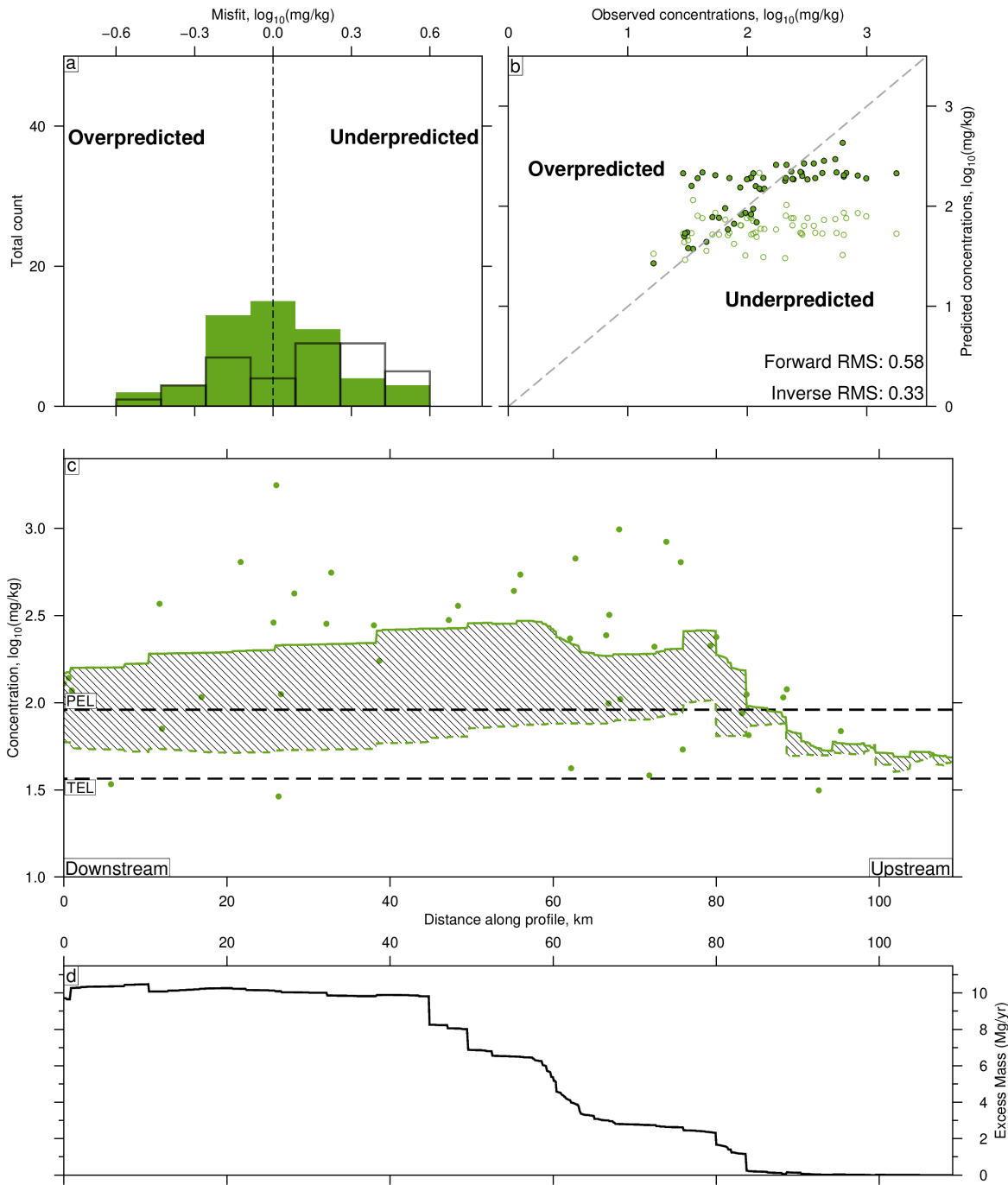
Finally, the methodologies we tested assume that river sediments are well-mixed and that they are ‘instantaneously’ transported from upstream sources. Very recent additional elemental sources, which are yet to be well-mixed into the sediment, may be poorly characterised by the scheme we present. Whilst, more complex

schemes which explicitly take into account time-varying sediment transport (e.g. Palu and Julien 2019) could be considered we think that it is prudent to first test simple, conservative, mixing models. This is particularly true where detailed information about sedimentary fluxes is not available. It is encouraging that simple conservative mixing models reliably predict concentrations downstream for a suite of ‘natural’ elements that have diverse affinities and concentrations.

## 6 CONCLUSIONS

Conservative mixing models are used to calculate natural baselines for the downstream sedimentary chemistry of the Clyde river, UK. These baselines are subsequently compared to predictions from inverse modeling of measured downstream compositions to quantify heavy metal pollution in the upper reaches of the river. The forward models presented use observations from first-order streams to generate continuous natural baselines. Many elements (e.g. Mg, K, Mn, Sr) with a broad range of





**Figure 6: Comparing baseline (forward model) predictions of Pb concentrations to predictions from best-fitting inverse model.** (a) Misfit (difference) between observed and theoretical concentrations at the loci of the 60 CUSP samples (see Figure 5a). Bin size = 0.15 log<sub>10</sub>(mg/kg). Ordinate gives absolute count ( $N = 60$ ). Solid green = results from best-fitting inverse model (Figure 5b). Black = results from forward model with source region parameterised using observed G-BASE first-order stream concentrations (Figure 5c). (b) Observed (CUSP) vs. predicted concentrations at the loci of the CUSP samples; solid/open circles = results from best-fitting inverse model/forward model. RMS = root mean squared misfit between observations and predictions from forward and best-fitting inverse models. (c) Pb concentrations along the main river profile. Circles = observed concentrations (CUSP). Green dashed line = baseline predicted by forward modeling. Solid green line = predictions from best-fitting inverse model. Black dashed lines indicate threshold (TEL) and probable (PEL) effect levels (toxic elemental concentrations) for adverse effects on biological activity (Hudson-Edwards et al. 2008). (d) Calculated excess mass of Pb added to river per year; difference between predictions from best-fitting inverse and forward models. Total calculated excess mass: 9700 kg/yr.

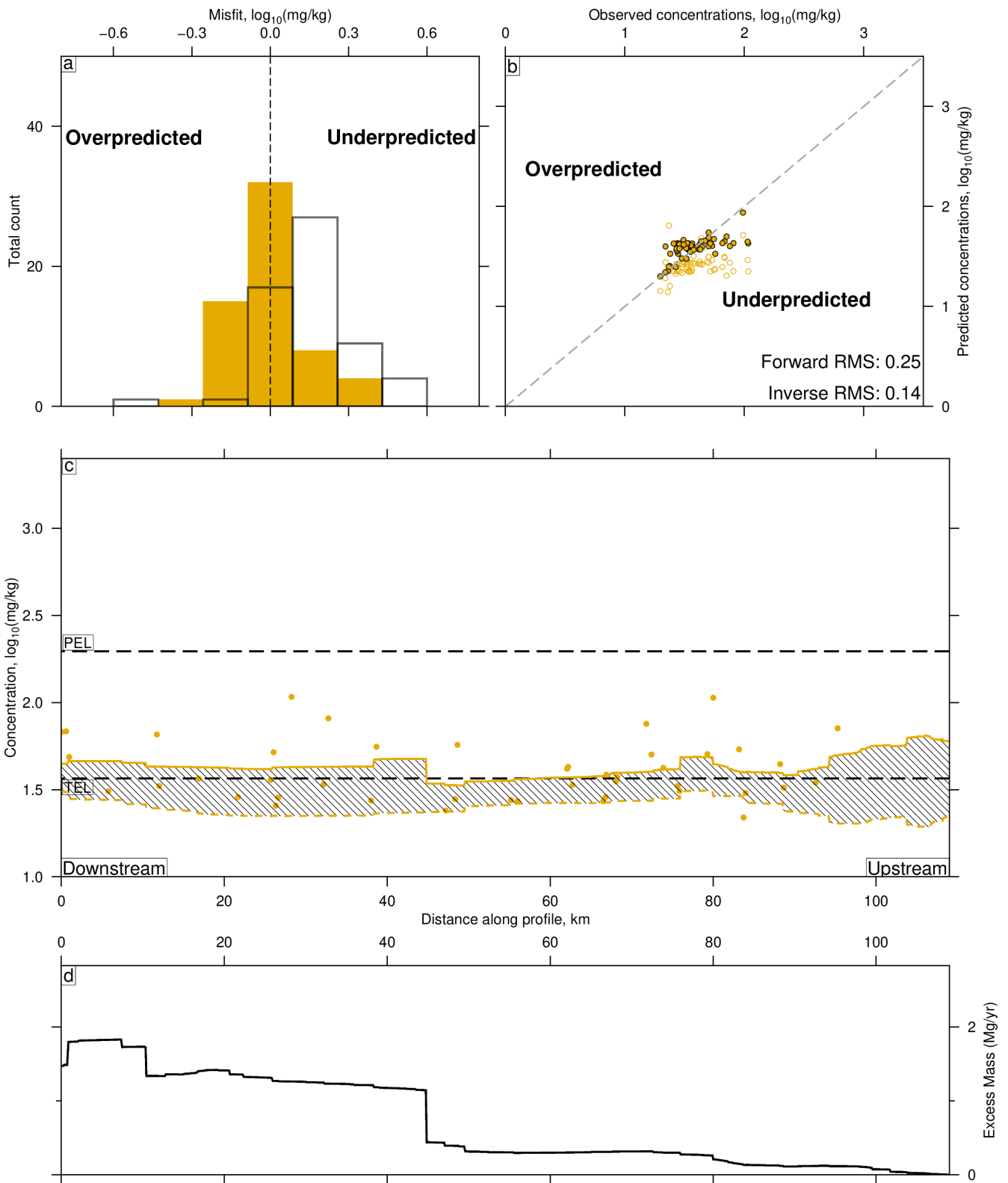


Figure 7: Comparing baseline (forward model) predictions of Cu concentrations to predictions from best-fitting inverse model. (a-d) See Figure 6 for explanation. Total calculated excess mass: 1500 kg/yr.

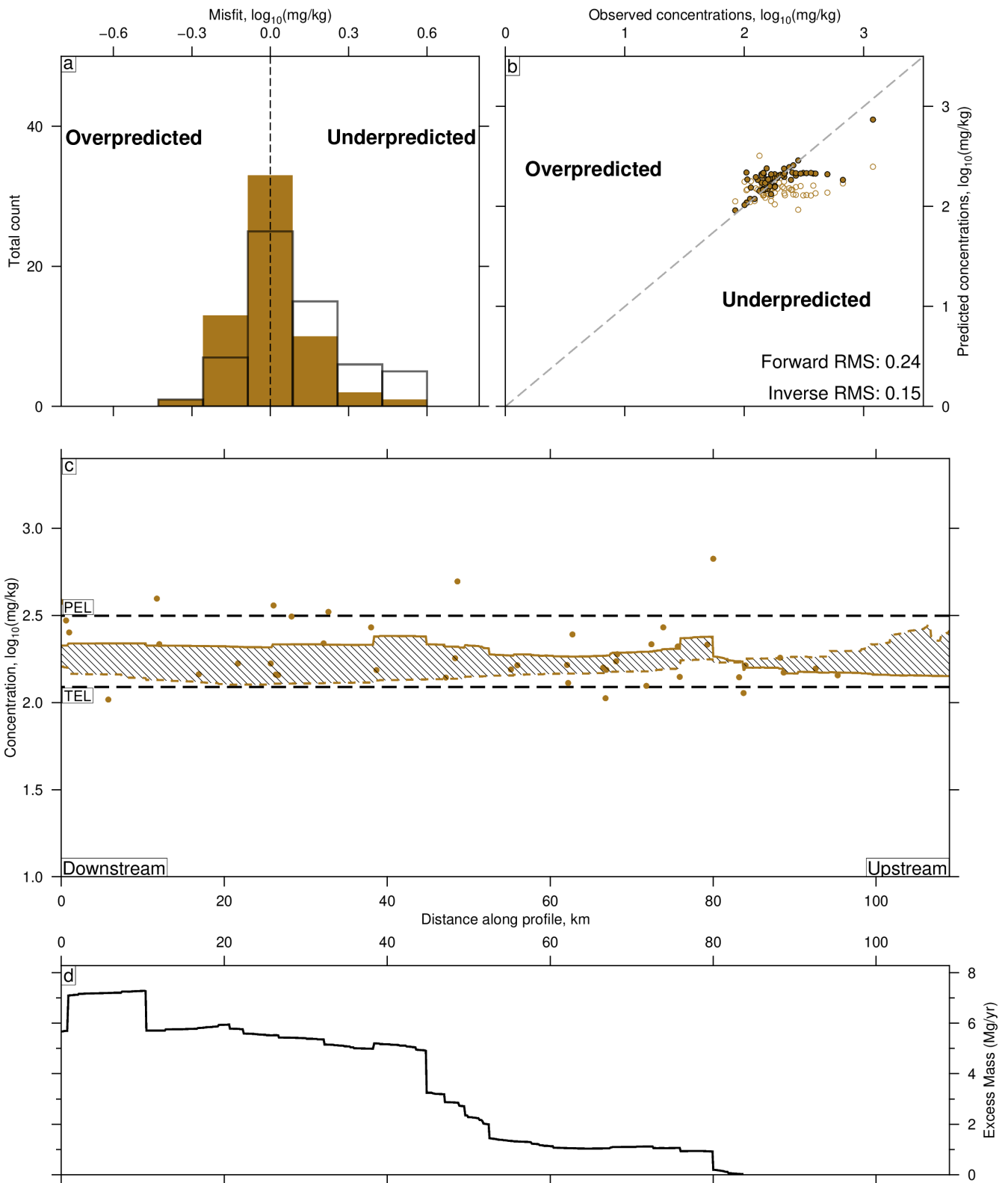


Figure 8: Comparing baseline (forward model) predictions of Zn concentrations to predictions from best-fitting inverse model. (a-d) See Figure 6 for explanation. Total calculated excess mass: 5700 kg/yr.

concentrations and chemical affinities are observed to be closely aligned with their predicted baselines. Key exceptions are the heavy metals Pb, Zn and Cu, which have relatively high residual misfits and are consistently underpredicted by conservative mixing models parameterised with upstream source compositions measured by the BGS in the Geochemical Baseline Survey of the Environment (G-BASE). Elemental concentrations and mixtures in source regions required to generate good fits to measured concentrations downstream along the upper Clyde river are calculated by inverting measurements made at 60 localities during the Clyde Urban Super Project (CUSP). Misfit between observed and predicted downstream concentrations is significantly reduced, and model residuals clusters around a 1:1 relation, if source region concentrations are considerably higher than the ‘natural’ concentrations measured by G-BASE. Comparing predicted heavy metal concentrations from the forward and inverse models allows us to quantify excess (non-geological) contributions to heavy metal concentrations and to tentatively infer sources of pollution. Most notably for Pb, an increase in concentration can be traced back to the Lowther Hills, a historic mining region. Total pollution, i.e. cumulative masses in excess of those predicted by conservative mixing of source concentrations, can be quantified. Using this data and previously defined threshold levels, it is possible to identify foci of toxic elemental concentrations, and sources of pollutants, which we suggest could help focus remediation efforts.

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## SUPPORTING INFORMATION

### Introduction

This section contains additional results from forward and inverse modelling. Figures 1-15 show the results from forward modelling of 15 elements. Figures 16-17 demonstrate the resolution of the inverse model. Supplementary Figure 18 shows results from tests to identify optimal damping parameter values.

### Forward model predictions

Supplementary Figures 1–15 show forward model predictions of the following element concentrations in Scottish rivers, including the Clyde basin: As, Ca, Co, Cr, Cu, K, Mg, Mn, Pb, Sb, Sn, Sr, Ti, Zn and Zr. Equation 3 and associated body text in the main manuscript describe the forward model. Predictions are compared to measured concentrations generated during the Clyde Urban Super Project (CUSP; 60 samples). The CUSP samples were collected using the same methodology as the first-order stream sediment samples collected for G-BASE (source region concentrations; < 150  $\mu\text{m}$  grain size from sieving; see main manuscript). We note for completeness that 16 samples (not shown here) of grain-size fraction < 2 mm were collected from the Clyde estuary bed using Day grab, gravity corer, Mackereth piston corer or Van Veen grab. Six samples were likely also gathered on foot from the riverbank. Their compositions were measured by XRF spectrometry. The location of these samples and their compositions are given in Jones et al. (2017). Given the different sampling strategies used to collect them they are excluded from further analysis in this contribution.

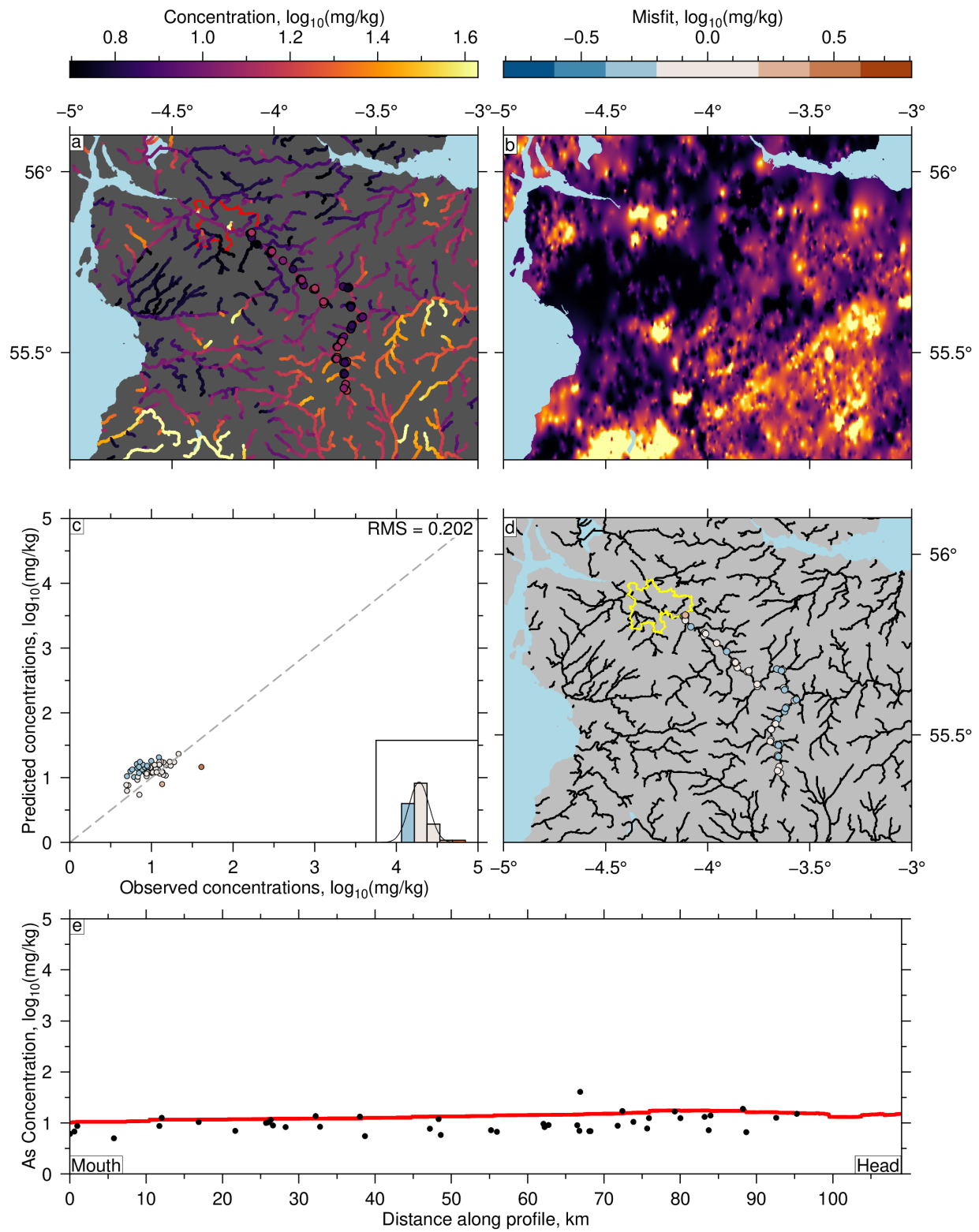
In Supplementary Figures 1–15 (a) circles = measured concentrations, < 150  $\mu\text{m}$  grain size, from CUSP survey. Coloured lines = predicted concentrations along rivers from forward model. Red outlined polygon = Glasgow city limits. (b) Source region concentrations used by forward model to predict concentrations along rivers. Grid is interpolated from G-BASE first order stream samples (< 150  $\mu\text{m}$ ; see Figure 1 in main text). (c) Comparison of observed and predicted concentrations. Dashed line = 1:1 relationship. Inset histogram shows misfit centred on zero; black curve indicates ideal normal distribution. RMS = root mean squared misfit (see Equation 6 in main manuscript). (d) Difference between observed and predicted concentrations. Yellow = Glasgow. (e) Longitudinal river profile showing observed (circles/squares) and predicted (red line) concentrations.

### Inverse model coverage

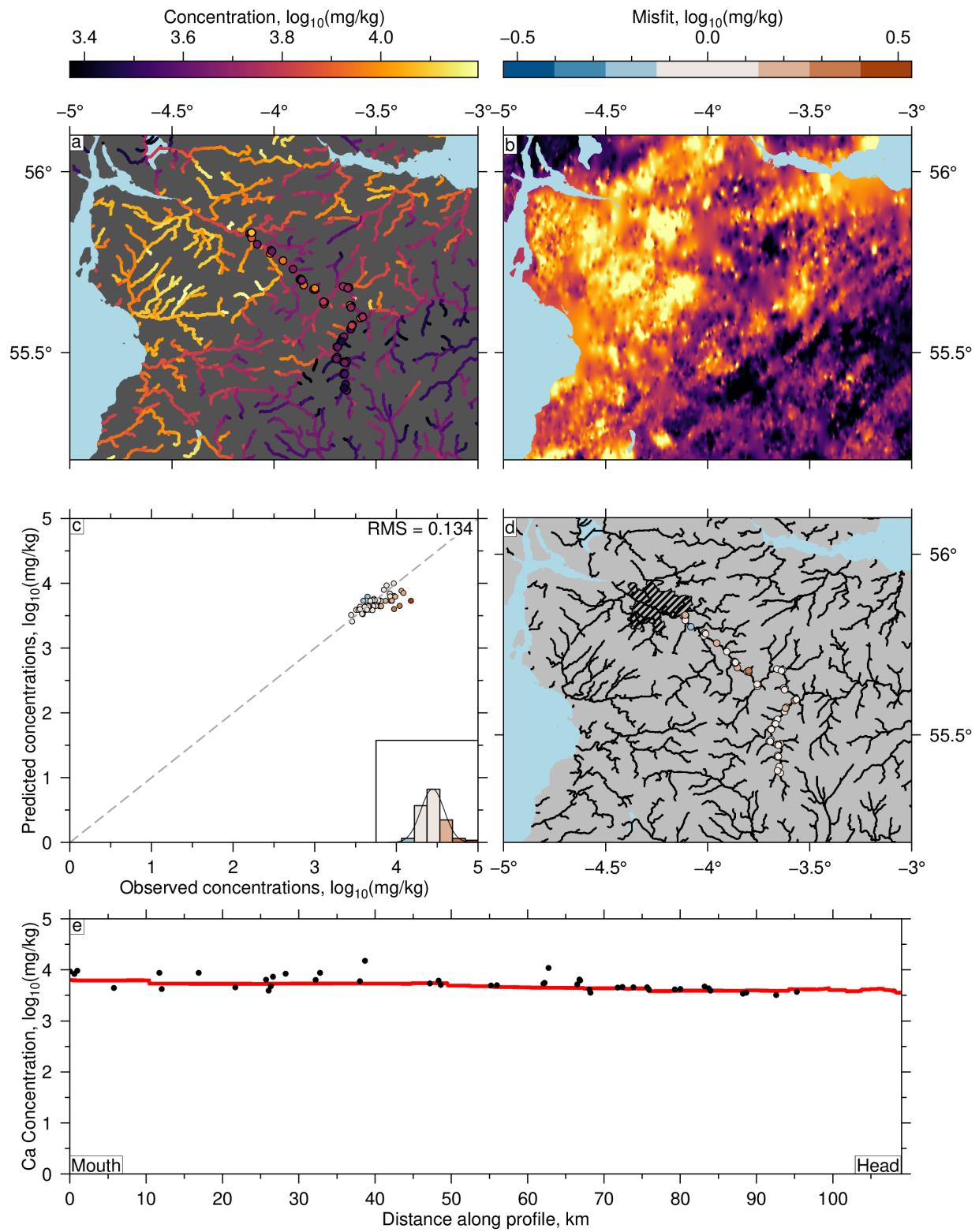
The area upstream of each sample site is a guide to the resolving power of the inverse model. Supplementary Figure 16 shows the area between each sample and the next upstream sample (where present) or the drainage divide. The 60 CUSP samples inverted lie along the main channel, which tends to result in low model coverage (resolution) in tributary basins. Model resolution is highest in the headwaters and lowest along sparsely sampled tributaries downstream. These results are supported by ‘chequerboard’ tests. In these tests a known (synthetic) source is used to calculate compositions at the actual CUSP sample sites using the actual drainage network and the forward model described in the main manuscript. The source concentrations at

the 60 sample sites are then inverted following the methodology described in the main manuscript. The inverse model is not preconditioned using information about the known source; in essence we have discarded the known source compositions and are attempting to recover them from only the concentrations at the 60 sample sites and the actual drainage network. The ability of the inverse model to recover source region concentrations can then be straightforwardly assessed by comparison with the known synthetic source. The results indicate that model resolution is highest in headwaters (Supplementary Figure 17). An optimised sampling strategy would include samples along the tributaries, which would drastically improve the resolution of the inverse model (cf. Lipp et al. 2021, their Figure 5).

Supplementary Figure 18 show an example of the relationship between model roughness and data misfit for Zn (see Equation 4 in main manuscript for details). In this series of tests, the inverse model was reran many times (with same starting conditions, input data, etc.) for a systematic sweep of damping parameter,  $\lambda$ , values. In the example shown in Supplementary Figure 18, the optimal value, i.e. the one that generates the smoothest model that best fits the data is  $10^{-0.2}$ . The ‘l-curves’ for Pb and Cu have a similar functional form and yield optimal  $\lambda$  values of  $10^{0.2}$  and  $10^{-0.2}$ , respectively.

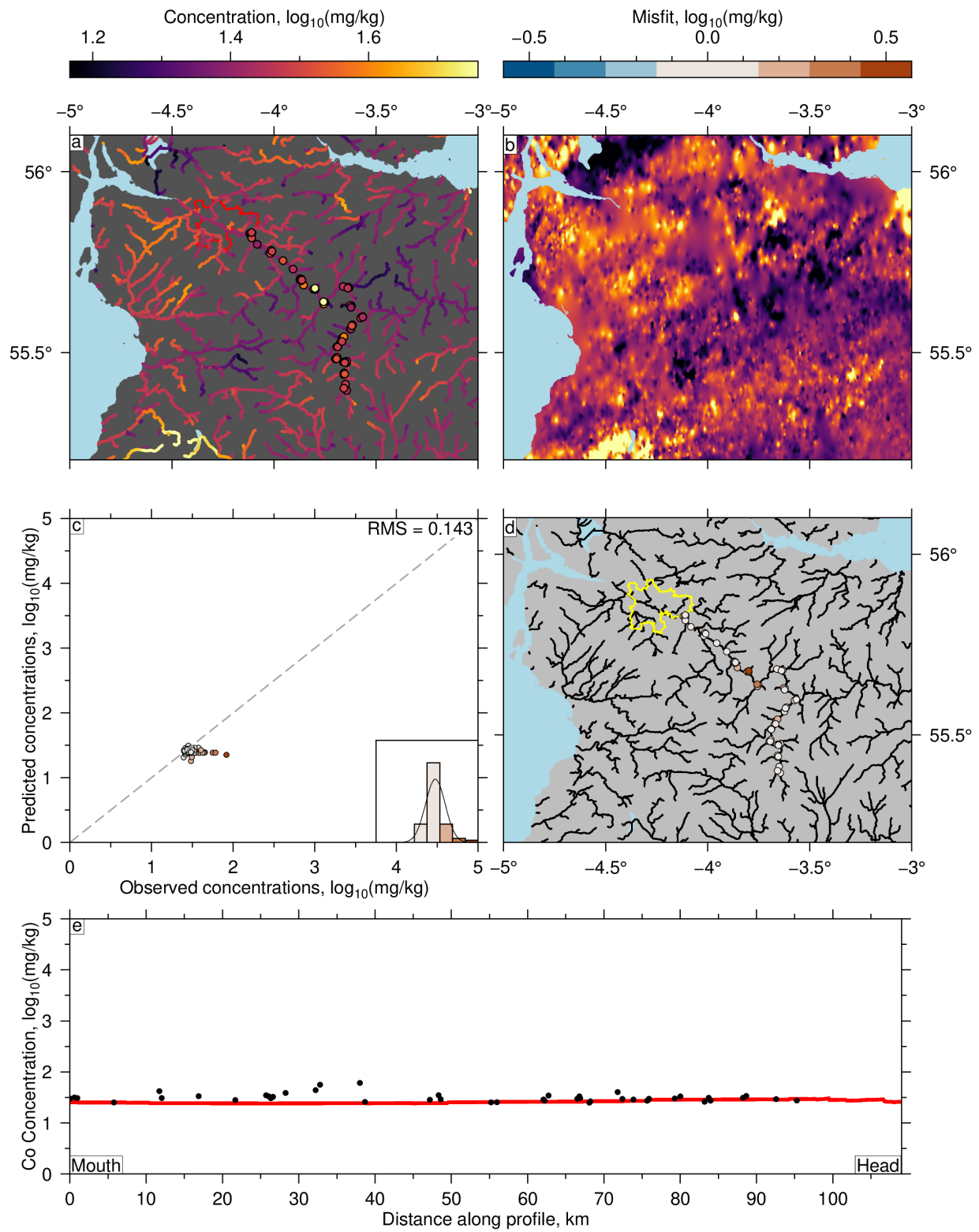


Supplementary Figure 1: **Forward model predictions of arsenic concentration in Scottish rivers.** See body text for details.

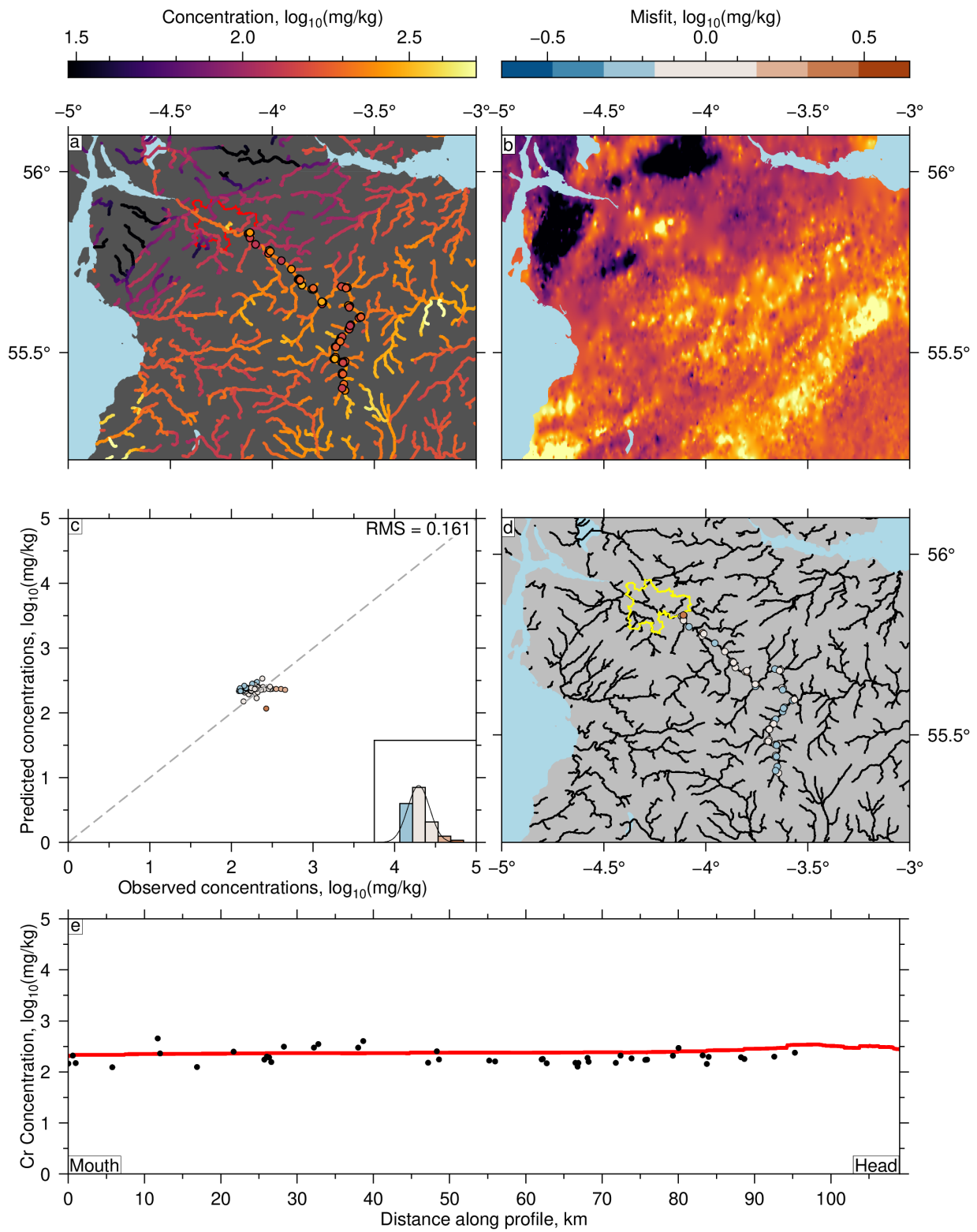


Supplementary Figure 2: **Forward model predictions of calcium concentration in Scottish rivers.**

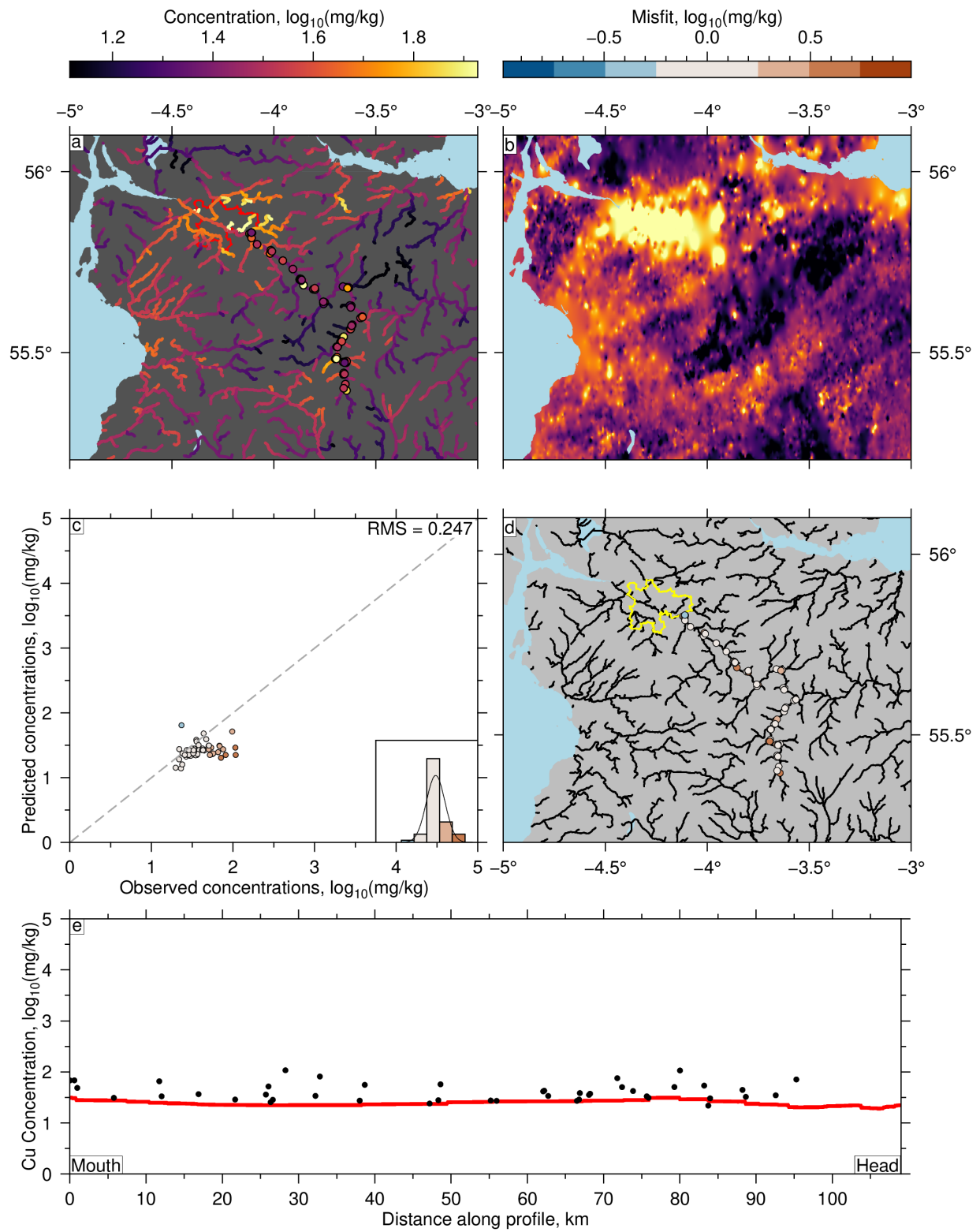




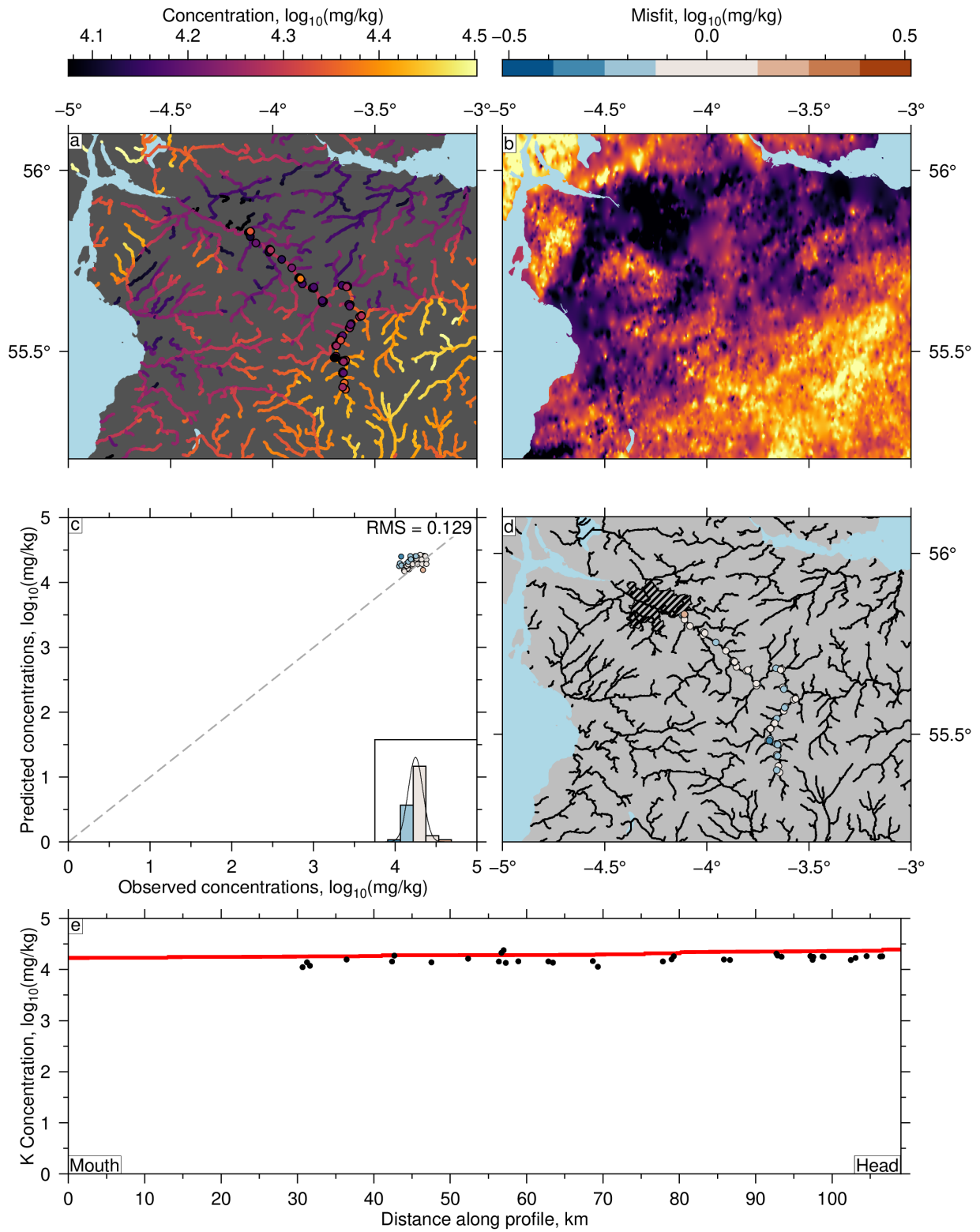
Supplementary Figure 3: Forward model predictions of cobalt concentration in Scottish rivers.



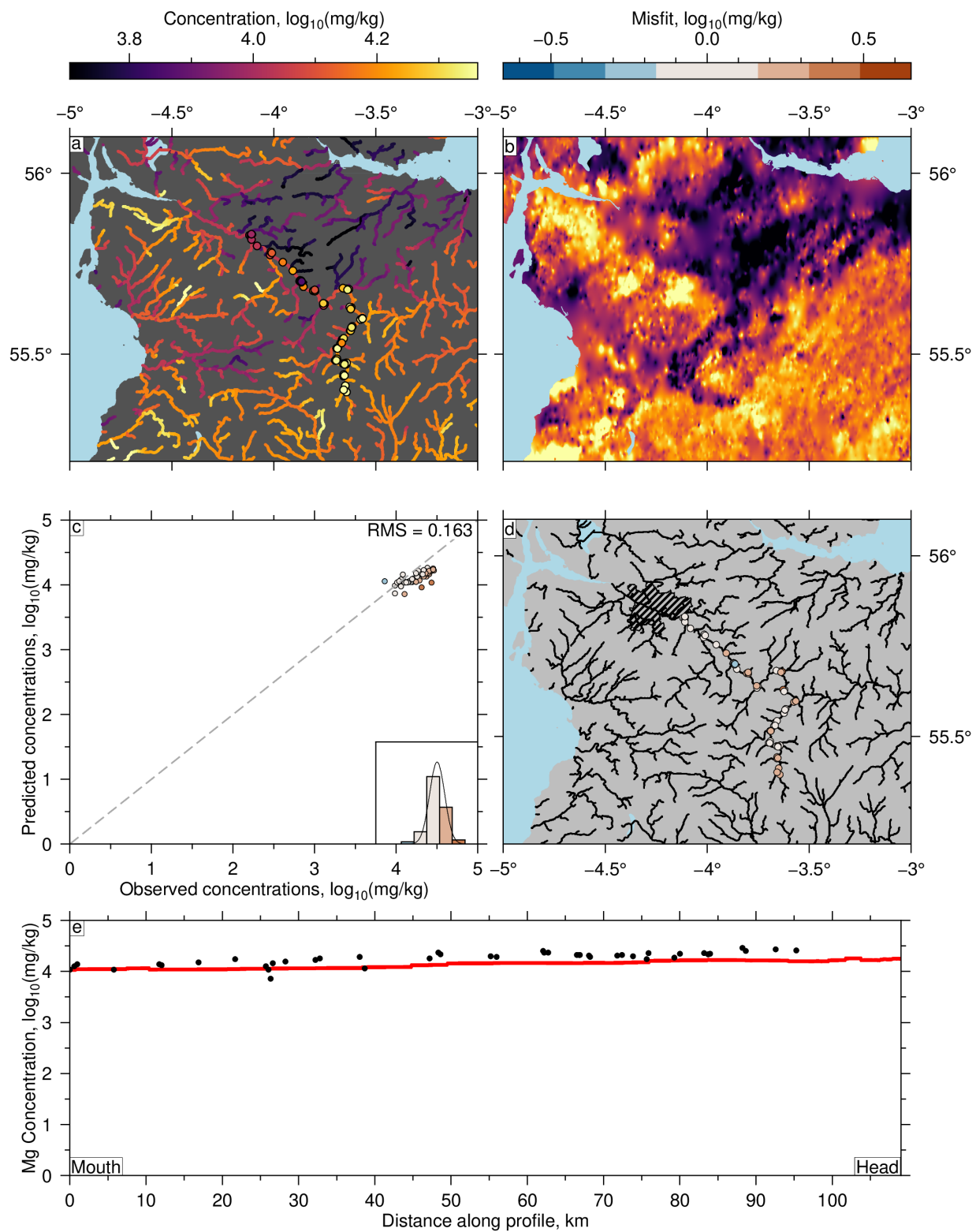
Supplementary Figure 4: **Forward model predictions of chromium concentration in Scottish rivers.**



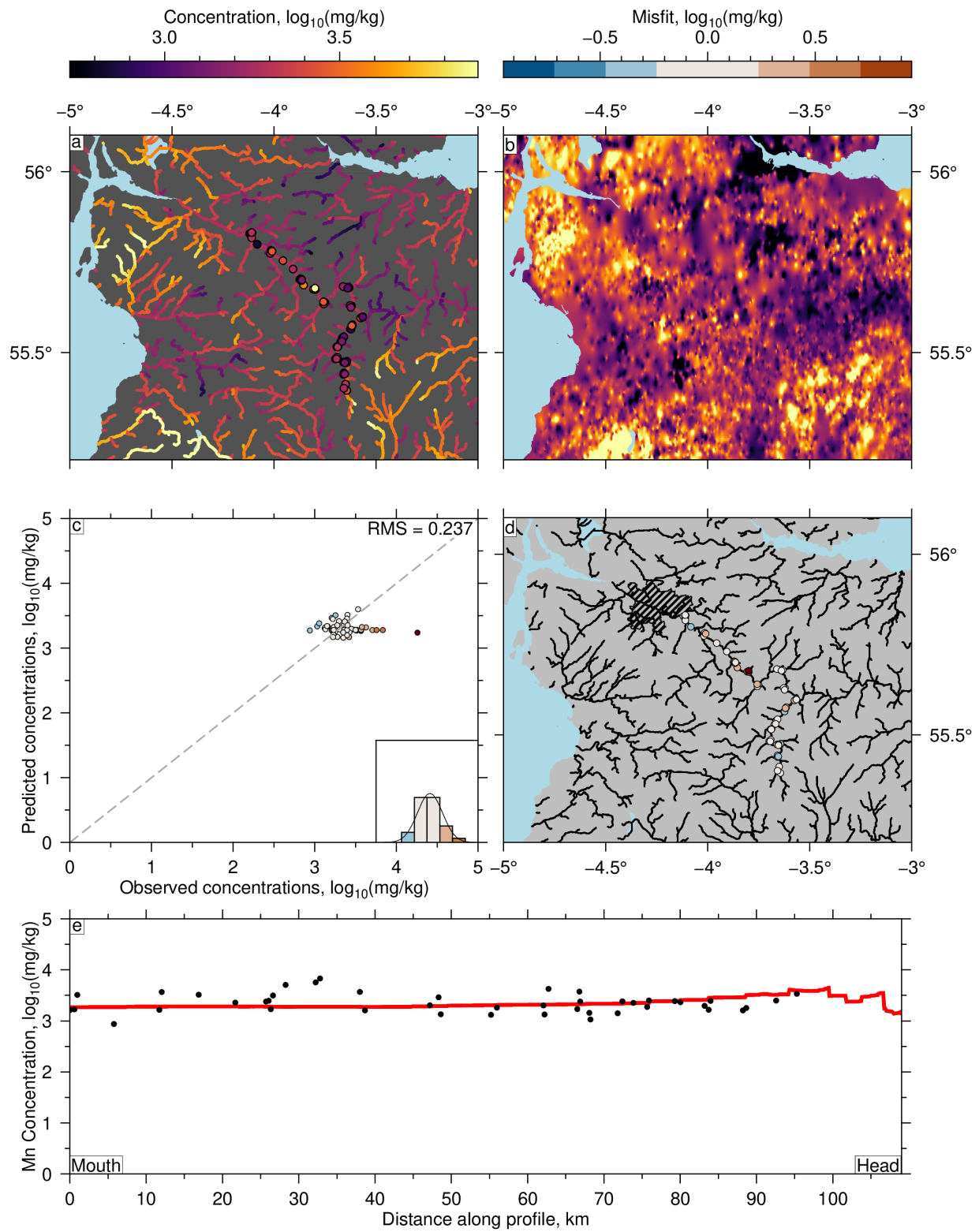
Supplementary Figure 5: Forward model predictions of copper concentration in Scottish rivers.



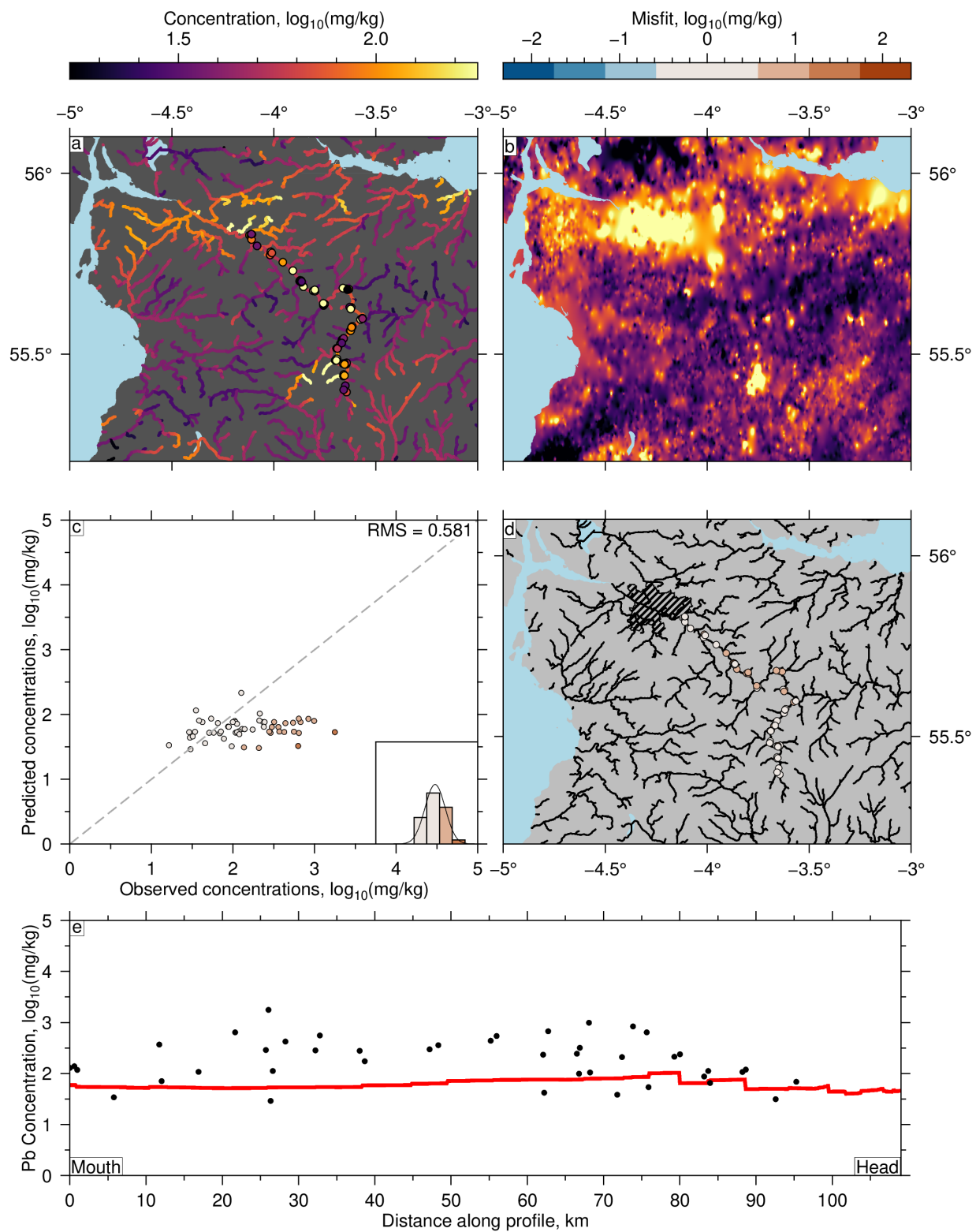
Supplementary Figure 6: **Forward model predictions of potassium concentration in Scottish rivers.**



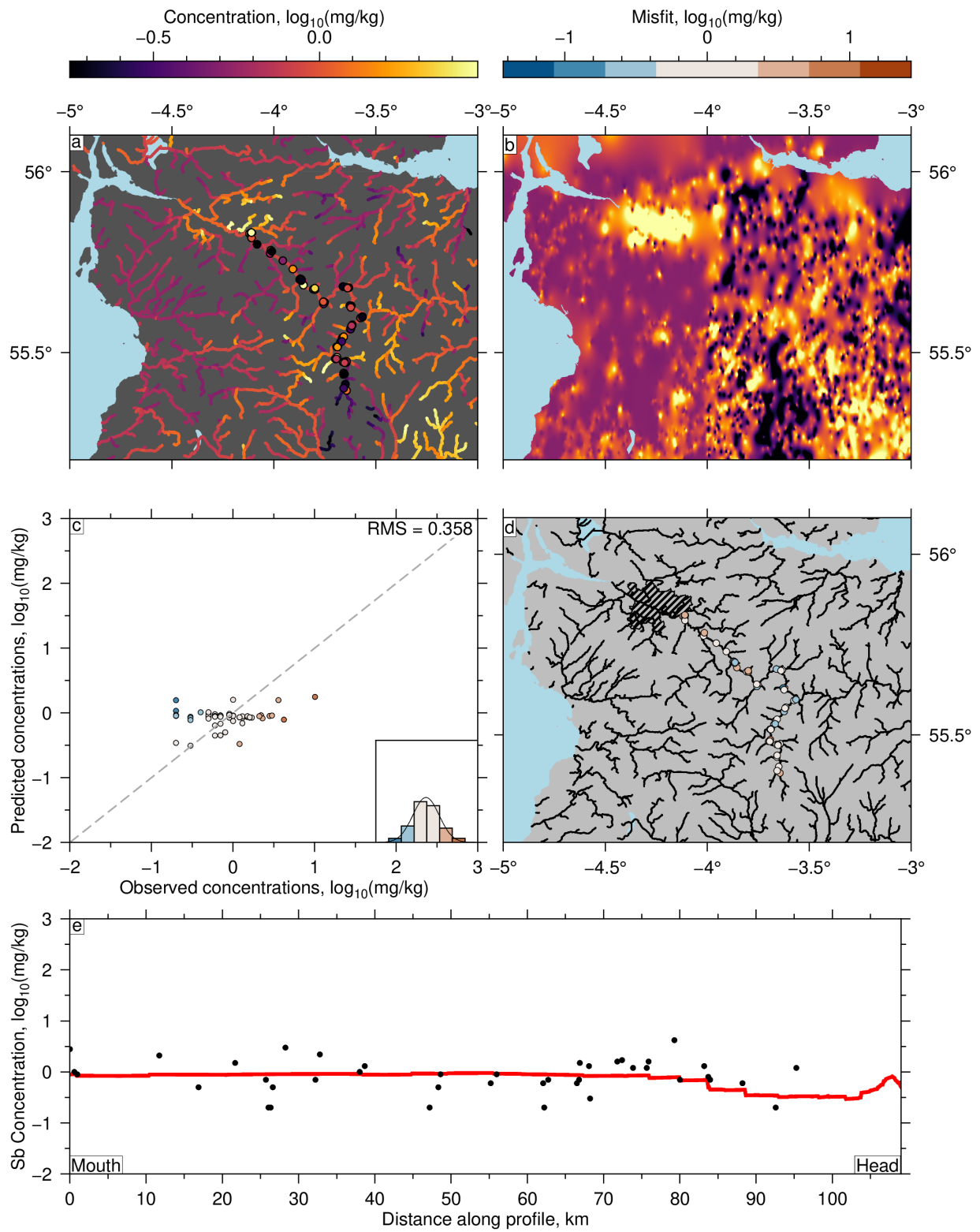
Supplementary Figure 7: Forward model predictions of magnesium concentration in Scottish rivers.



Supplementary Figure 8: Forward model predictions of manganese concentration in Scottish rivers.

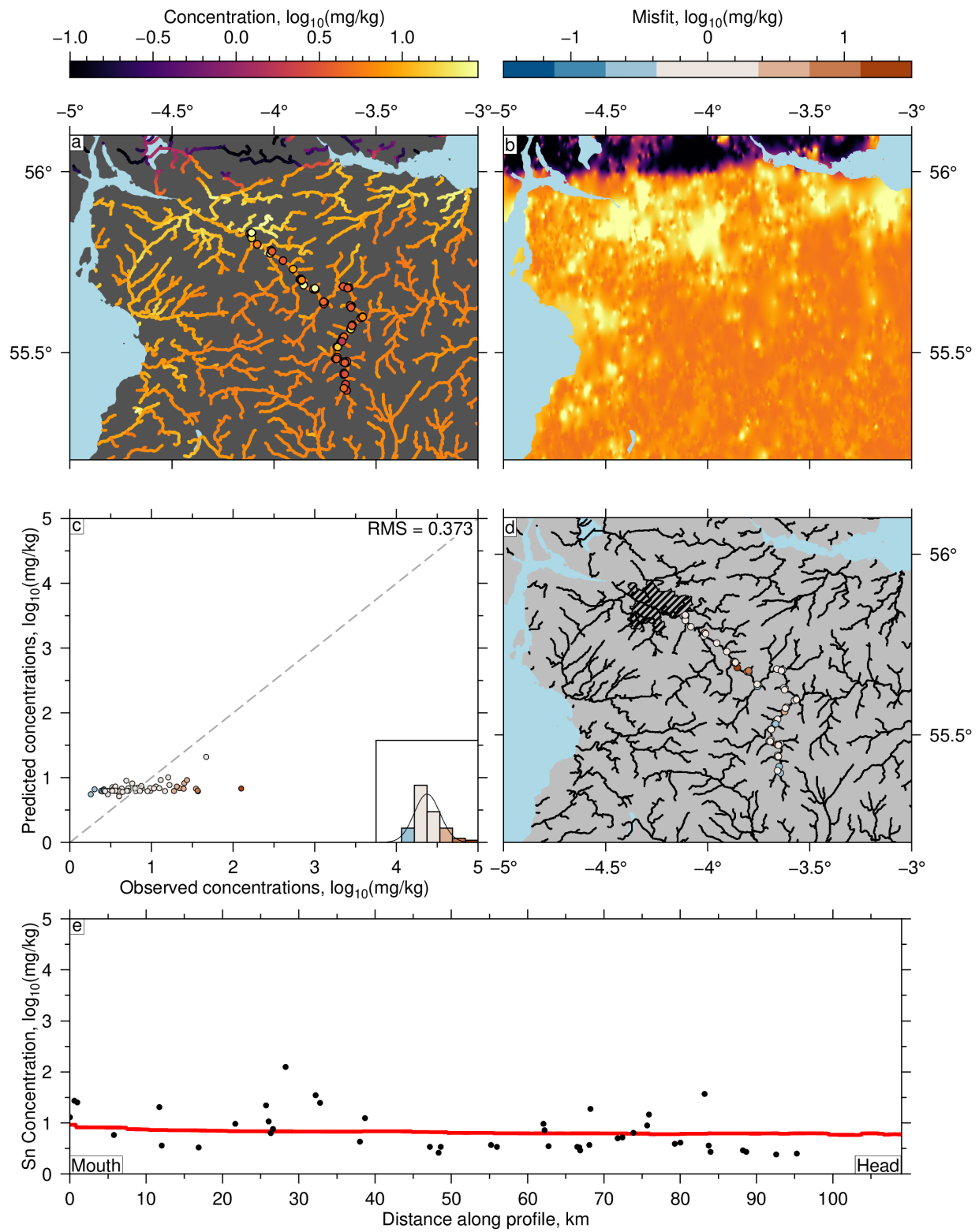


Supplementary Figure 9: **Forward model predictions of lead concentration in Scottish rivers.**

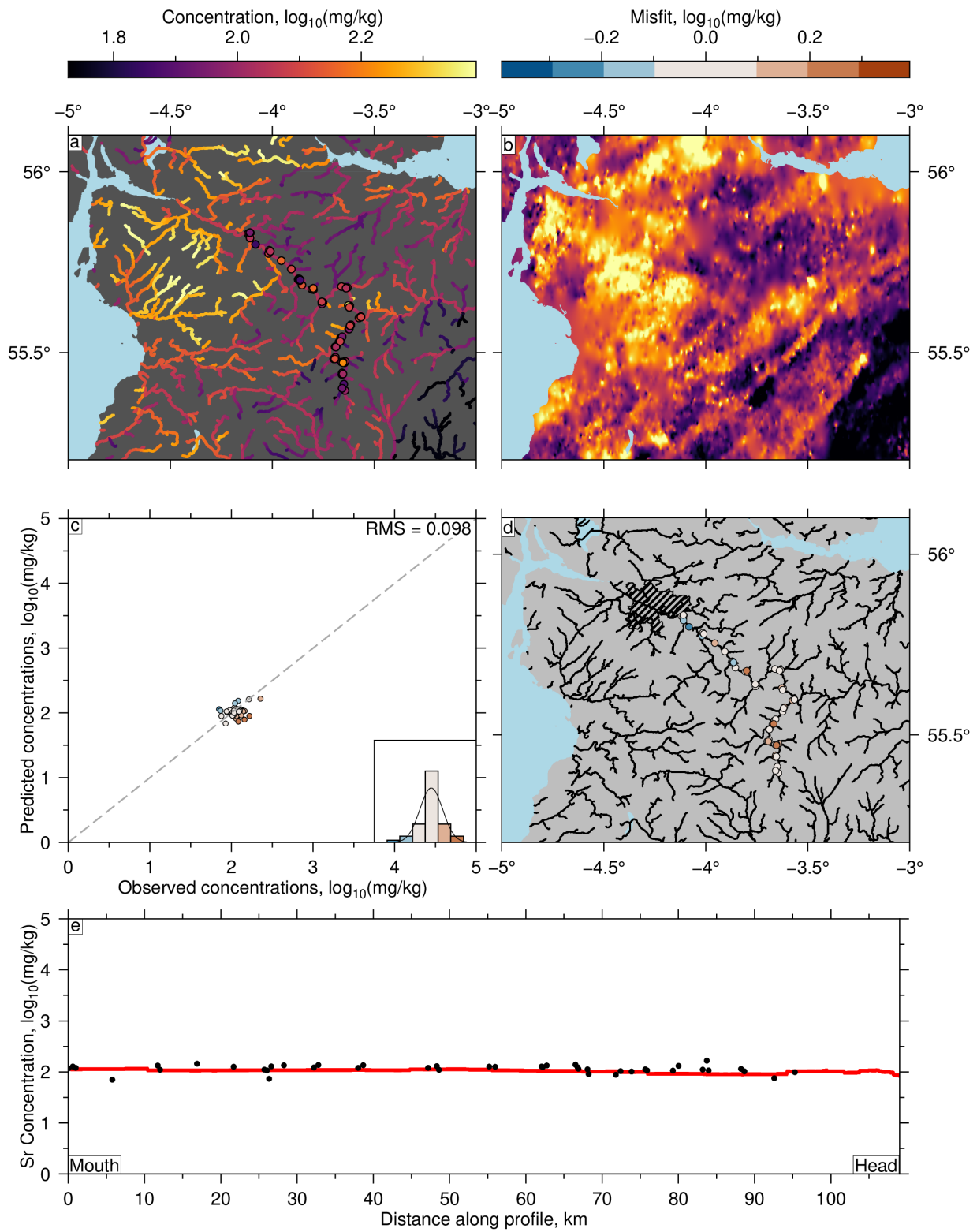


Supplementary Figure 10: **Forward model predictions of antimony concentration in Scottish rivers.**

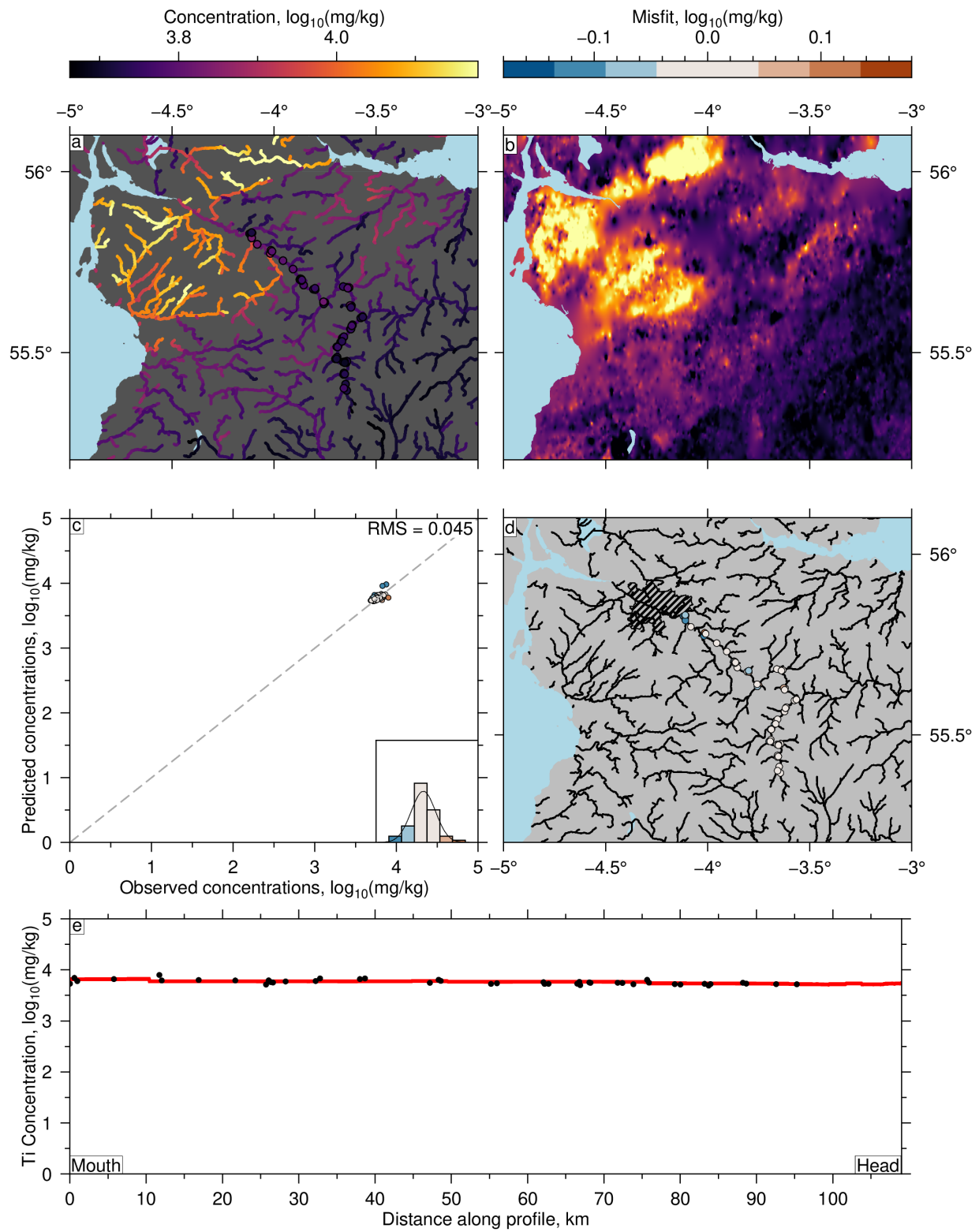




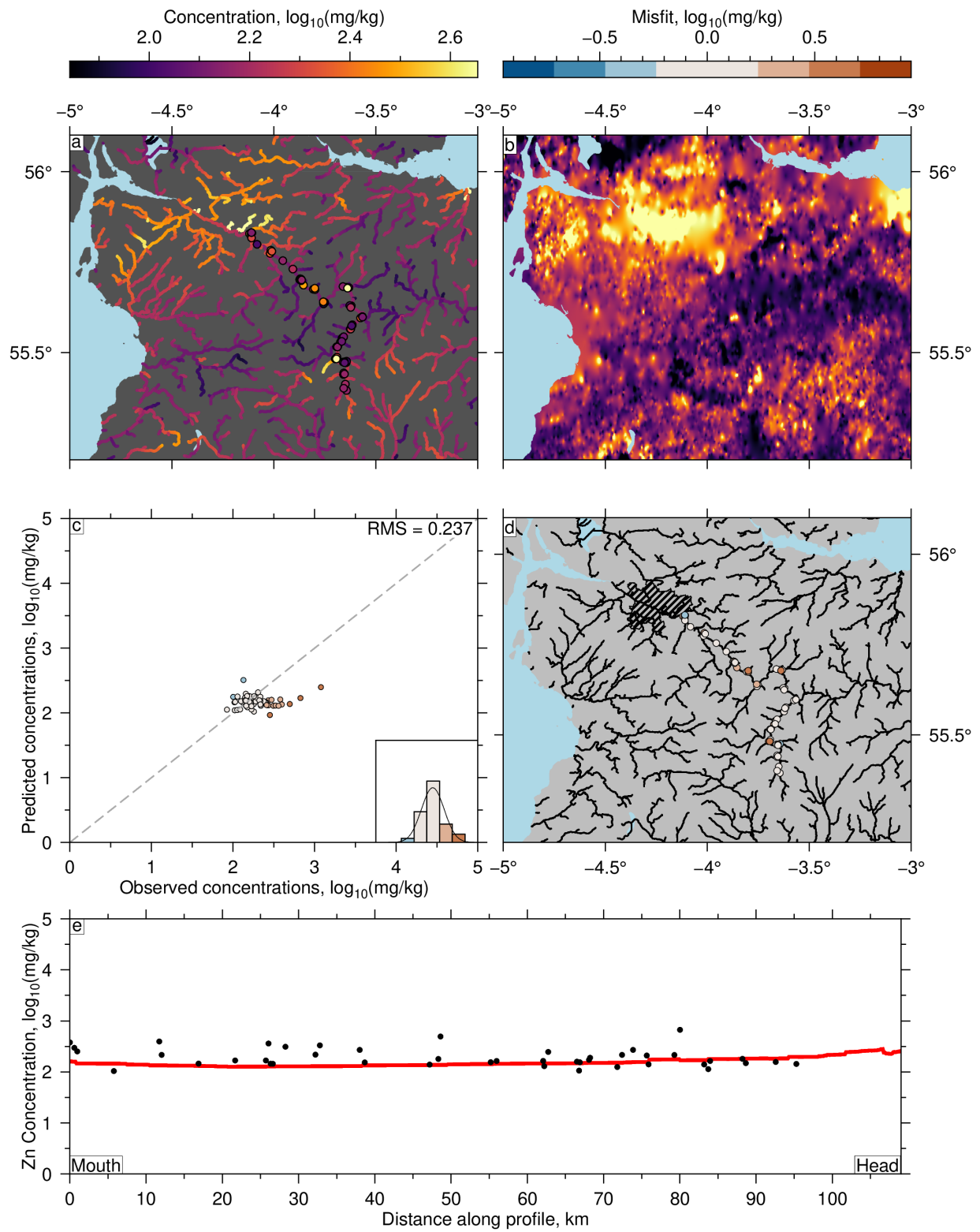
Supplementary Figure 11: **Forward model predictions of tin concentration in Scottish rivers.**



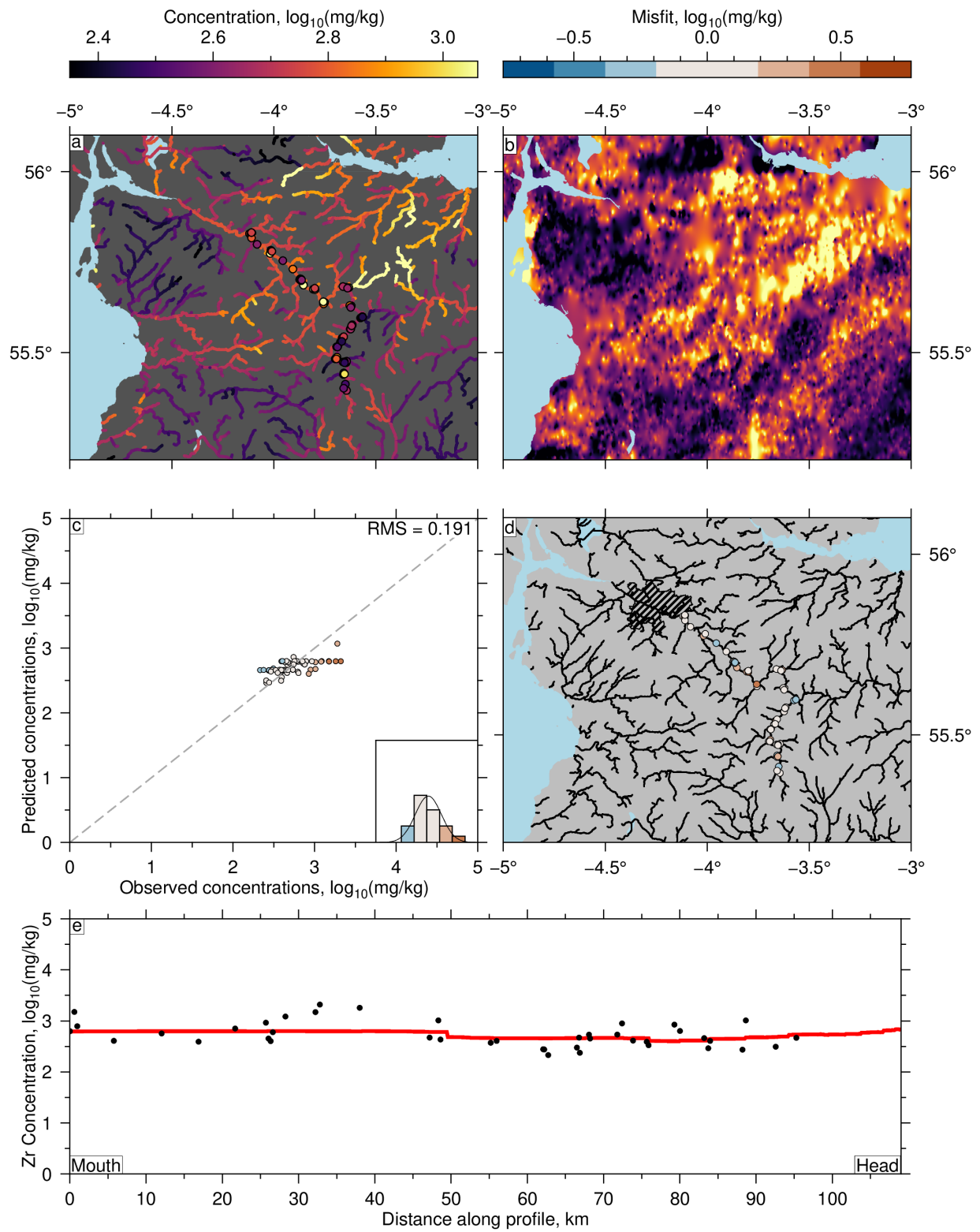
Supplementary Figure 12: **Forward model predictions of strontium concentration in Scottish rivers.**



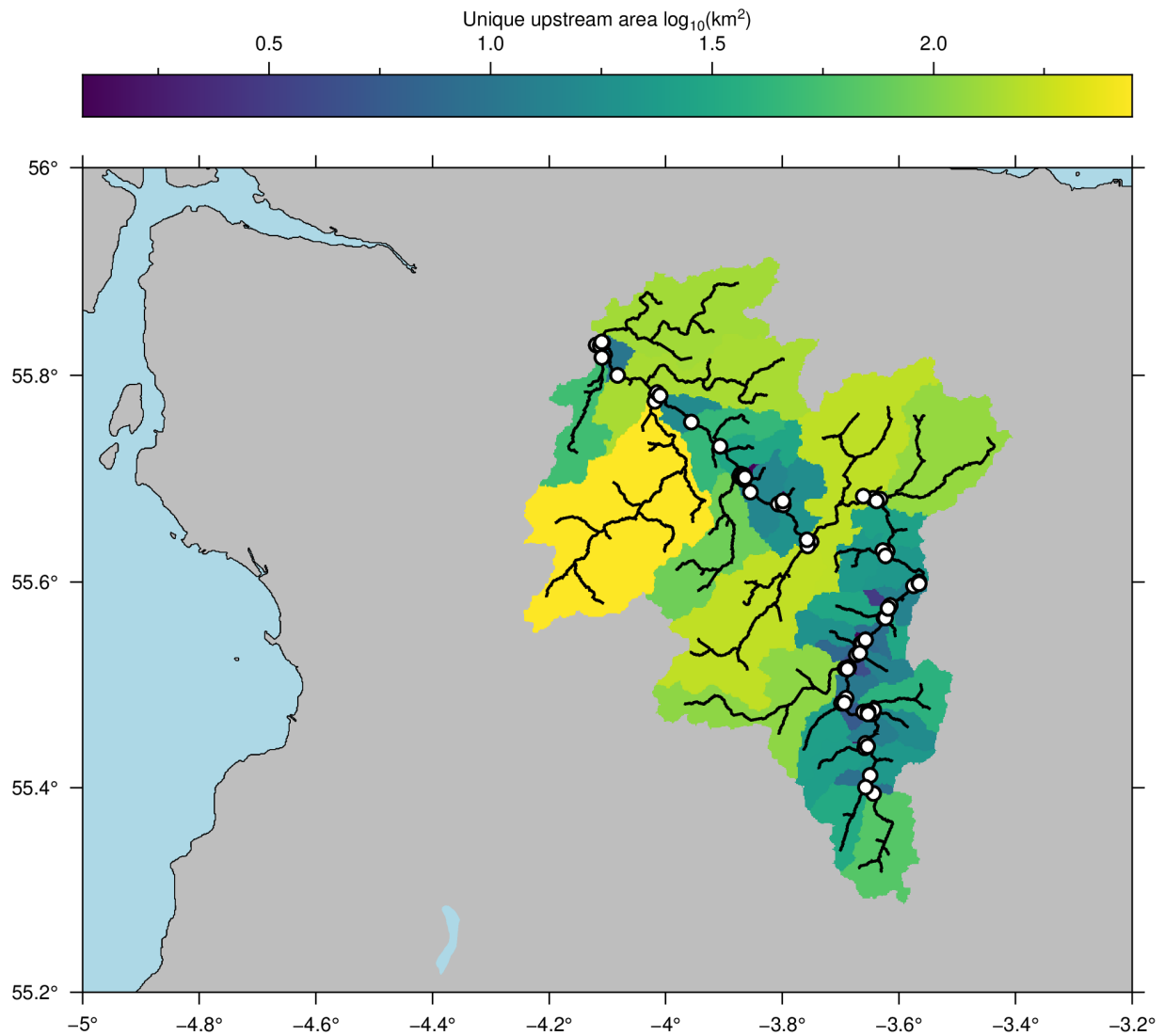
Supplementary Figure 13: **Forward model predictions of titanium concentration in Scottish rivers.**



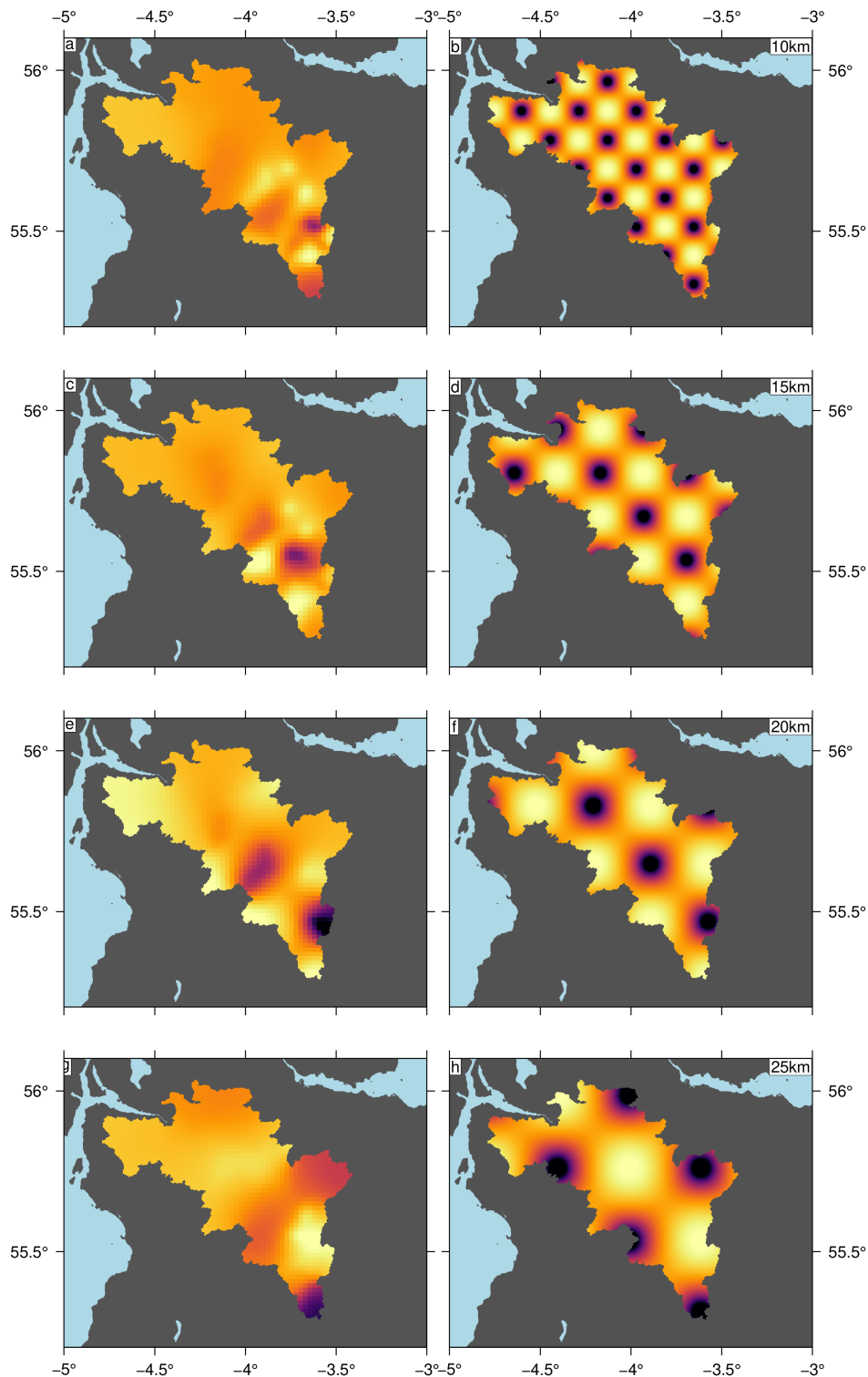
Supplementary Figure 14: **Forward model predictions of zinc concentration in Scottish rivers.**



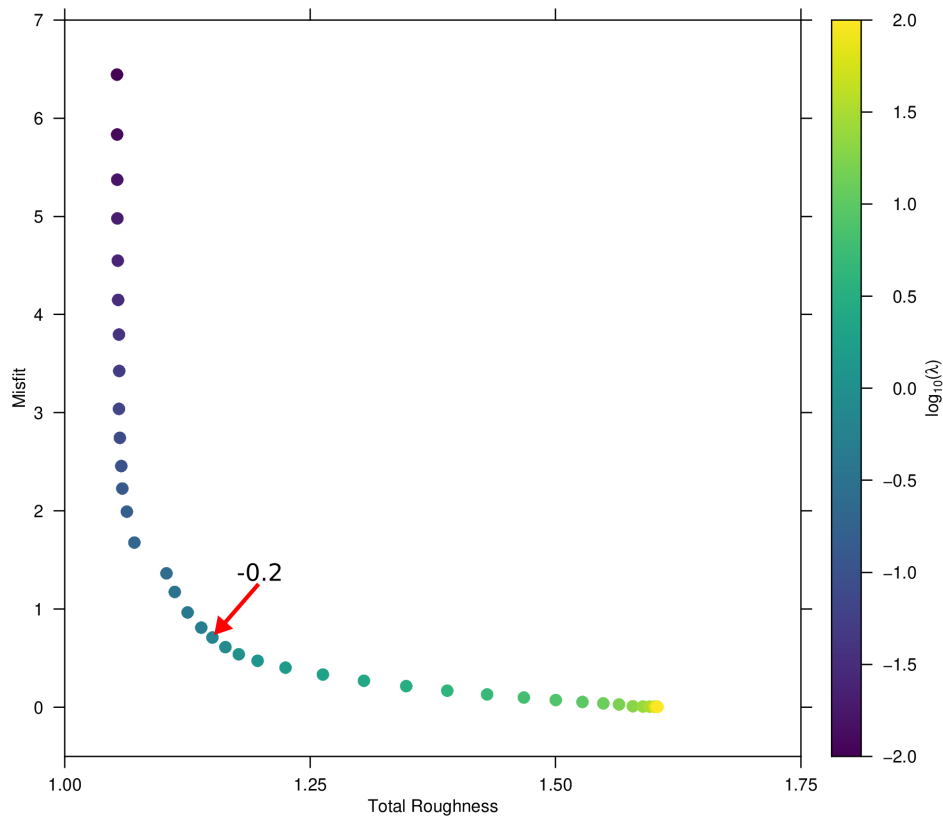
Supplementary Figure 15: **Forward model predictions of zirconium concentration in Scottish rivers.**



Supplementary Figure 16: **Unique upstream area of each sample used for the inversion.** White circles = CUSP samples. Area is given in log scale. Colours are a guide to the resolution of the inverse models. Note highest resolution (dark colours) in headwaters of the main channel and relatively low resolution in large tributaries downstream.



Supplementary Figure 17: **Inverse model resolution tests.** Synthetic ‘chequerboard’ source region concentrations (right column) were used to calculate concentrations at the loci of the 76 CUSP and estuary samples using the forward model described in the main manuscript (Figure 1; Equation 3 in main text). Annotation in top right corner indicates wavelength of chequerboard pattern. The calculated concentrations at the 76 sample localities were then inverted to solve for source region concentrations (left column). The actual drainage network was used in these calculations to test the resolving power of sample distributions and flow paths (cf. Supplementary Figure 16). Note low resolution in large tributaries downstream (cf. Supplementary Figure 16), and decreasing resolving power of short wavelength (< 20 km) changes in concentration.



Supplementary Figure 18: **Trade-off between model roughness and data misfit for Zn.** Example of relationship between misfit and model roughness used to calibrate  $\lambda$ . The optimal  $\lambda$  value ( $10^{-0.2}$ ; smoothest model that best fits the data) is identified at the point of maximum curvature, i.e. the ‘elbow’ of the trade-off curve indicated by the annotated red arrow.