Rivers transport the elements, minerals, chemicals, and pollutants produced in their upstream basins. A sample from a river is a mixture of all of its upstream sources, making it challenging to pinpoint the contribution from each individual source. Here, we show how a nested sample design and convex optimization can be used to efficiently unmix downstream samples of a well-mixed, conservative tracer into the contributions of their upstream sources. Our approach is significantly faster than previous methods. We represent the river’s sub-catchments, defined by sampling sites, using a directed acyclic graph. This graph is used to build a convex optimization problem which, thanks to its convexity, can be quickly solved to global optimality—in under a second on desktop hardware for datasets of ~100 samples or fewer. Uncertainties in the upstream predictions can be generated using Monte Carlo resampling. We provide an open-source implementation of this approach in Python. The inputs required are straightforward: a table containing sample locations and observed tracer concentrations, along with a D8 flow-direction raster map. As a case study, we use this method to map the elemental geochemistry of sediment sources for rivers draining the Cairngorms mountains, UK. This method could be extended to non-conservative and non-steady state tracers. We also show, theoretically, how multiple tracers could be simultaneously inverted to recover upstream run-off or erosion rates as well as source concentrations. Overall, this approach can provide valuable insights to researchers in various fields, including water quality, geochemical exploration, geochemistry, hydrology, and wastewater epidemiology.

**Keywords** Pollution Apportionment · Convex Optimisation · Hydrochemistry · Geochemistry · Tracers · Wastewater Epidemiology

1 Introduction

Rivers transport the elements, minerals, chemicals, and pollutants that are produced in their upstream basins; we refer to these, collectively, as tracers. When a water or sediment sample is collected from a specific point in a river, we can analyze the concentrations of these tracers to estimate their upstream production rates. This information is valuable in different ways depending on the type of tracer being measured. For pollutants, it can help us assess and target mitigation strategies, while for different ways, learning about the geology and hydrology of an area. However, a single sample of a river can’t provide much useful information because all of the different sources have been mixed together by the river. If multiple samples are taken upstream and downstream, we show that a computational technique called convex optimization can be used to “unmix” the river’s waters revealing where the elements have come from. This approach is accurate and fast enough that we can run it hundreds of times with small changes to the sampled values to get a sense of the uncertainty. Though we speak about our method in terms of rivers here, it could be useful in other contexts, such as determining where pollutants or tracers of disease outbreaks are entering sewage networks.
other on both the main channel and in its tributaries) as a directed acyclic graph (p. 574; Sedgewick and Wayne 2011) constrained by the river network. We use this graph to pose unmixing as a convex optimization problem. For a dataset composed of ∼100 samples our method solves the unmixing problem in less than a second. This is ∼10^6 times faster than previous methods. The computation time of our method also scales nearly linearly with the number of samples over a range of problem sizes. The speed of our method makes it easy to quantify uncertainty through Monte Carlo resampling and to explore the impact of changing parameters such as regularization strength. Such calculations can be performed easily on standard desktop hardware and do not require the use of high-performance computing or clusters.

We present our new method as an open-source Python package (github.com/r-barnes/faster-unmixer). The implementation requires only a geospatial raster of "DB8" flow-directions (which are widely available or can be generated from digital elevation data using GIS tools) and a data table of samples giving observed tracer concentrations and the location of sample sites. Our software can be used to analyze any dataset of nested, conservative tracers, in both sediment and water, across a drainage network.

1.1 Background

Lipp et al. (2021) previously worked to unmix observations of sediment geochemistry to determine the geochemical composition of source regions. First, the concentration of an element in downstream sediments was described as an integral of the concentration of the element in the upstream area. This forward model was then inverted. To overcome non-uniqueness in the solutions, the model was regularized by seeking the smoothest upstream concentration map that best fit observations downstream. Whilst this scheme was successfully used for geochemical mapping in northern Australia and for quantifying the sources of heavy metal pollution in the River Clyde, UK (Eschenfelder et al. 2023; Lipp et al. 2023), it has a number of limitations.

The primary limitation is that this scheme was solved as a general non-linear optimization problem. Solvers for such problems are unable to guarantee that they have found a globally optimal solution. Additionally, these solvers (e.g., Nelder–Mead, Powell’s Method) can be computationally expensive relative to those used to solve other types of problems such as convex optimization, linear-program and least-squares (Press et al. 1992; Boyd and Vandenberghe 2004). This computational slowness (hours for tens of samples on standard desktop hardware), makes it impractical to explore hyper-parameters or quantify uncertainty via Monte Carlo methods.

2 Methods

2.1 Intuition

If the concentration of elements, minerals, chemicals, isotopes, or pollutants (herein referred to generically as tracers) is sampled at a single point on a river, what can we say about the upstream processes that generated this sample? If we have observed a tracer concentration \( d \) in the water or sediment then, assuming we have no other information, a parsimonious interpretation is that the source concentration, \( c \), is the same as the observed concentration, \( d \), at the sample site (i.e., \( c = d \)) and that the tracer is generated uniformly across the whole upstream area (Figure 1a).

We consider a situation where we’ve taken two samples from the river, one downstream of the other. We call the upstream Sample U and the downstream Sample D (Figure 1b). As before, we can’t infer much about the production rates upstream of Sample U: whatever the value of the concentration \( d_U \) is, it is assumed to be generated uniformly by all the area upstream of Sample U, and so \( c_U = d_U \). However, we can now say more about the source concentration in the area between the two samples, \( c_D \).

The observed concentration at Sample D is:

\[
d_D = \frac{c_U q_U + c_D q_D}{q_U + q_D},
\]

(1)
a weighted mixture of the sources upstream of U and in the area between U and D. In this expression, \( q \) refers to the flux of water or sediment added in the sub-basin of a given sample. For example, \( q_D \) is the flux generated between Sample U and Sample D. It’s also important to note that this relation is only true if the tracer is conservative and well-mixed; that is, this is only true if the tracer does not get removed from the water as it flows downstream (via, e.g., photodegradation, consumption, flocculation and settling, and/or absorption) and the production and transport of the tracer is in steady-state.

Increasing the number of samples we collect means we can identify where tracers are coming from in the catchment in greater spatial detail. For example, imagine we take a sample on the tributary, Sample T, that joins the river between Sample U and Sample D (Figure 1c). We can see again that \( c_T = d_T \), and again \( c_U = d_U \). However, now, Sample D combines information from the areas defined by Samples U and D as well as the new Sample T. Therefore,

\[
d_D = \frac{c_U q_U + c_D q_D + c_T q_T}{q_U + q_D + q_T},
\]

(2)

The question we are interested in is, given downstream observations, how can we solve these expressions for the upstream tracer sources? If \( d_U, d_D, \) and \( d_T \) are exact measurements, we...
We now formally describe how to set up and solve this problem. At first, it may seem that this problem can be solved with a least-squares approach by minimizing the squared distance between predicted and observed concentrations (see A). However, least-squares is actually an inappropriate way to measure differences for this situation as real concentration measurements typically follow log-normal distributions spanning orders of magnitude (e.g., Gaillardet et al. 1999). This effect is a particular concern in situations where pollutants from a point source are diluted as they flow downstream. For example, the addition of the tributary’s flow \( q_T \) to the main river in Figure 1 may increase \( q_D \) significantly. More generally, as we move a distance \( L \) downstream, Hack’s Law says that we should expect the basin area (and its flow) to increase as \( \sim L^{1.6} \) (Hack 1957). Therefore any tracer discharged from a point source upstream, will naturally dilute as a function of \( L^{1.6} \) as it is transported downstream. If we use least-squares to solve such a case we will end up weighting the undiluted, upstream samples much more heavily than diluted, downstream samples. Therefore, it is more sensible to solve these equations by instead considering the relative difference (e.g., the log-ratio) between the observed and actual concentrations. This is similar to the well-known need to normalize data in machine learning (LeCun et al. 2012).

We now formally describe how to set up and solve this problem for any set of observations on a drainage network. We first consider how to build a “forward” model that translates a given set of upstream concentrations into a suite of concentrations at sample sites. We then demonstrate how to “invert” this forward model to translate a set of downstream observations into a set of predicted upstream concentrations.

2.2 Forward model

We assume that we have made nested (upstream, downstream, and tributary) observations of concentrations of a well-mixed, conservative tracer at sample-sites within a basin (e.g., Figure 2a). Each sample site becomes a node in a directed acyclic graph (DAG) which we term a “sample network”. The topology of this graph (Figure 2b) reflects the topology of the drainage basin (Figure 2a).

Each node of the DAG has several attributes:

- \( a, \text{[L}^2\text{]} \) e.g., \( \text{km}^2 \): the sub-basin area, a constant. The area of each sub-basin uniquely defined by the sample. In Figure 2 this is indicated with dashed lines.
- \( \phi, \text{[M L}^{-2} \text{T}^{-1}] \) e.g., \( \text{kg/km}^2\text{yr} \): the material export-rate, a parameter. This quantity is the total amount of water or sediment produced in the sub-basin that reaches the most downstream node in the network. If there is no storage of material in the network this can be considered equivalent to the average run-off or sediment generation rate of the sub-basin.
- \( q, \text{[M T}^{-1}] \) e.g., \( \text{kg/yr} \): the material flux of the sub-basin, a dummy variable, given by \( q = a \phi \).
- \( d, \text{unitless e.g., mg/kg} \): the observed concentration, a parameter. This is the concentration of the tracer observed in the water or sediment at the sample site. This is a function of all the nodes in the network upstream of the sample point.
- \( c, \text{unitless e.g., mg/kg} \): the generated concentration, a variable, which we would like to estimate by solving the unmixing problem. This quantity is the average concentration of the tracer in material being exported by the sub-basin. For example, in Figure 2, \( c \) is zero for all nodes except for node 2. The tracer exported from node 2 is transported, and diluted downstream meaning that the observed concentration \( d \) is highest at node 2 and decreases downstream at sites 4 and 6. Identifying \( c \) for each node in a network, given point observations of \( d \) is the goal of this study.

As the tracer is assumed to be conservative and in steady state we can build expressions for the components of \( d \) as flux-weighted mixtures of all the nodes upstream in the network:

\[
d_i = \frac{\sum_{j \in U_i} q_j c_j}{\sum_{j \in U_i} q_j},
\]

where \( U_i \) is the set of nodes upstream of node \( i \), including itself. Note that for “leaf” nodes (a node with no upstream nodes) the only contributor to the flux is the unique upstream area of that node and so the concentration at the sample site is simply the same as the source concentration. As an example, the concentration at node 4 is:

\[
d_4 = \frac{c_1 q_2 + c_2 q_2 + c_4 q_4}{q_1 + q_2 + q_4}.
\]

Thus, assuming that \( a \) and \( \phi \) are known, it is possible to translate a vector of source concentration values \( c \) into a vector of predicted concentrations at sample sites \( d_{\text{pred}} \).

2.3 Inverse model

We now consider the inverse problem: what is the vector of source concentrations \( c \) that best fits a set of downstream tracer observations \( d_{\text{obs}} \), given a sample network, \( N \). We pose this question as an optimization problem, seeking to minimize the total misfit between \( d_{\text{obs}} \) and \( d_{\text{pred}} \) (calculated using Equation 3) with respect to \( c \) across all nodes in the network. We therefore solve

\[
\text{minimize}_{w.r.t. c} \sum_{\in N} \text{misfit}(d_{\text{obs},i}, d_{\text{pred},i})
\]

\[
= \text{minimize}_{w.r.t. c} \sum_{\in N} \text{misfit}(d_{\text{obs},i}, \frac{\sum_{j \in U_i} q_j c_j}{\sum_{j \in U_i} q_j})
\]

given a misfit function between two concentration values, and where \( i \) is the index of nodes in the network \( N \). A naive
Figure 2: Sample sites on a drainage network and their representation as a directed acyclic graph. a) Theoretical drainage network with six sample sites, 1 through 6. Each sample defines a sub-basin indicated by dashed lines. Colour indicates the concentration \( d \) of some tracer, which is sourced in the sub-basin of sample 2 then transported and diluted downstream. b) The sample network in panel a, represented as a directed acyclic graph.

Figure 3: Visualising two relative misfit terms. Red curve shows log-ratio misfit between predicted concentrations \( d_{\text{pred}} \) and an arbitrarily chosen observed concentration \( d_{\text{obs}} \) value of 30. Blue curve is the misfit between the same values but using the convex misfit term chosen in this study. The log-ratio curve is non-convex. Note that the convex misfit function has a minimum value of 1, not 0 like the log-ratio misfit.

choice for the misfit function would be the Euclidean distance \( \sqrt{(d_{\text{pred}} - d_{\text{obs}})^2} \). In fact, if such a Euclidean misfit function is chosen, the minimization problem can be solved using a linear least-squares approach as discussed above and in A. However, concentration data typically span many orders of magnitude, and as such the Euclidean distance over-weights observations of high concentration relative to observations of low concentration. Additionally, because concentration data are technically compositional, the information they contain is relative not absolute (Aitchison 1986).

To better extract these relative differences, a more appropriate misfit function would be the absolute value of the log-ratio of the two concentrations: \( |\log(d_{\text{pred}}/d_{\text{obs}})| \) (Aitchison 1986). This function is minimized at \( d_{\text{pred}} = d_{\text{obs}} \), and is symmetrical with respect to relative changes in value. However, this also has a drawback: it is a non-convex function. Functions which are non-convex are challenging to minimize because there is no general algorithm for finding their global minimum, though local minima can be identified (Boyd and Vandenberghe 2004). In contrast, for convex functions the global minimum is the same as the local minimum and an extensive set of robust tools are available to find it. We therefore define the following convex misfit function that penalises relative differences between \( d_{\text{pred}} \) and \( d_{\text{obs}} \):

\[
\text{misfit}(d_{\text{pred}}, d_{\text{obs}}) = \max\left(\frac{d_{\text{obs}}}{d_{\text{pred}}}, \frac{d_{\text{pred}}}{d_{\text{obs}}}\right). \tag{5}
\]

This misfit function is visualised in Figure 3. As \( \frac{d_{\text{obs}}}{d_{\text{pred}}} \) and \( \frac{d_{\text{pred}}}{d_{\text{obs}}} \) are convex and affine respectively for \( d_{\text{pred}} > 0 \), their maximum is also convex over this range. We note that Equation 5 is non-differentiable at \( d_{\text{pred}} = d_{\text{obs}} \), but this is not relevant for our approach. Note that Equation 5 takes its minimum value at the same point as \( |\log(d_{\text{pred}}/d_{\text{obs}})| \); thus, any solution which...
minimizes the convex function Equation 5 would also minimize \( \log(d_{\text{pred}}/d_{\text{obs}}) \). Although this function is undefined if either \( d_{\text{pred}} \) or \( d_{\text{obs}} \) are equal to zero, concentration observations in nature are rarely truly equal to zero; instead, they mostly represent values below detection limits. However, this limitation does mean that any zero values in a dataset ought be removed or replaced with small values prior to analysis (e.g., Martín-Fernández et al. 2003).

We can therefore pose a convex minimization problem to return the source concentrations, \( \mathbf{c} \) that best fit a set of observations downstream:

\[
\text{minimize}_{\mathbf{c}} \quad \sum_{i \in N} \max \left( \frac{d_{\text{obs},i}}{d_{\text{pred},i}}, \frac{d_{\text{pred},i}}{d_{\text{obs},i}} \right) \quad (6)
\]

\[
= \text{minimize}_{\mathbf{c}} \quad \sum_{i \in N} \max \left( \frac{d_{\text{obs},i} \sum_{j \in U_i} q_i}{\sum_{j \in U_i} q_{i,j}}, \frac{d_{\text{obs},i}}{\sum_{j \in U_i} q_{i,j}} \right) \quad (7)
\]

### 2.4 Regularization

Measurements of concentrations are always accompanied by a degree of uncertainty that reflects both analytical noise, and also cases where the modelling assumptions are not fully satisfied, for example, imperfect mixing of the water column, hydrodynamic sorting of sediment, or non-conservative behaviour (e.g., Bouchez et al. 2010; Stutenbecker et al. 2023). To prevent the optimization from over-fitting this noise we regularize our solution by penalising relative deviations in \( \mathbf{c} \) from the mean observed concentration. We opt to penalize differences from the geometric as opposed to the arithmetic mean of the observed concentrations,

\[
\tilde{d} = \left( \prod_{i=1}^{n} d_{\text{obs},i} \right)^{\frac{1}{n}}.
\]

We use the geometric mean because, as discussed above, concentration observations typically span many orders of magnitude and the arithmetic mean, if chosen, would overweight high concentrations. This regularization prevents solutions from containing implausibly extreme concentrations. We utilize the same misfit function as above to quantify relative deviations from the mean. Thus, we can add regularization terms to Equation 7:

\[
\text{minimize}_{\mathbf{c}} \quad \sum_{i \in N} \max \left( \frac{d_{\text{obs},i}}{d_{\text{pred},i}}, \frac{d_{\text{pred},i}}{d_{\text{obs},i}} \right) + \lambda \sum_{i \in N} \max \left( \frac{\tilde{d}}{c_i}, \frac{c_i}{\tilde{d}} \right) \quad (8)
\]

where \( \lambda \) is a hyper-parameter controlling the strength of regularization. The value of \( \lambda \) can be calibrated using various methods. One approach is to analyze the trade-off between data misfit and the total deviations of the model from the mean observed value. Then, the model with the smallest deviations that still provides an adequate fit to the data is selected (the Occam’s or “elbow” approach; Constable et al. 1987). Another method is to utilize an unseen “validation” subset of the data to calibrate the value of \( \lambda \), or similar cross-validation approaches.

Previous unmixing approaches regularized by penalizing the “roughness” of the source concentration (Lipp et al. 2021). Roughness was calculated as the first derivative of \( \mathbf{c} \) with respect to spatial coordinates. We opt not to use this approach here for two reasons. First, we haven’t yet found a convex relative difference term between two variables that is compatible with the other constraints of our problem; Equation 5 is only convex if one argument is a constant. Thus, calculating the relative difference between two spatially adjacent \( \mathbf{c} \) variables is not, at present, possible without violating the convexity of our optimization problem. Secondly, prior authors solved for \( \mathbf{c} \) on a regular grid over the modelled domain for which roughness has a more intuitive interpretation. By contrast, we now solve for \( \mathbf{c} \) defined on discrete sub-basins which are irregular in shape (Figure 2). Whilst it is possible to conceive of definitions of roughness on such a domain (e.g., normalising the difference between adjacent nodes by the length of shared boundaries) it is not clear that these always represent sensible terms to penalize.

### 2.5 Disjoint Networks

Note that the neither of the optimization problems described by Equations 7 and 8 are sensitive to whether or not the directed acyclic network is fully-connected. If two disjoint river networks with fully separate basins are combined in the same optimization problem the solution to that problem is the same as if the networks were solved separately.

### 3 Implementation

We provide an open-source python implementation of the above method at github.com/r-barnes/faster-unmixer and archived at zenodo.org/record/8299812. The code is well-documented (63% comments) with unit tests, examples, and installation instructions. A package “funmixer” is generated which can be used to easily create and solve instances of the above problem given a D8 raster of flow-directions and a data table of sample locations and concentrations. Here we describe our implementation in more detail.

#### 3.1 Building sample networks

The first stage for solving the inverse problem is building the sample network data structure. To build this, we require a D8 raster (O’Callaghan and Mark 1984) of the modelled flow-directions for the study-region and a data table containing the spatial coordinates of the sample sites. The D8 hydrological model assumes only convergent flow across a network, i.e., each cell in the raster is able to donate flow to only one neighbouring node. As a result, our method is unable to consider networks which have strongly divergent flow, though it could be modified to do so since even divergent flow still forms a directed acyclic network.

To build a sample network, we iterate upstream from sink cells, which are assumed to occur at the boundary of the D8 and at natural sinks within the domain (e.g., lakes). At each cell, we check to see if our location corresponds to one of the provided sample-sites. If it does, we generate a new node in the sample-network. We then iterate through the neighbours of this cell, and if they donate flow to it, we assign them a label to indicate that these nodes donate their flow uniquely to that sample-site. We then recursively continue to search upstream of these donor cells...
labelling cells as we go. If we reach a cell which corresponds to a sample site we change the label we apply to the cells upstream of that site and the process continues.

To allow this approach to work even for very large D8 rasters, we implement it in C++ using tools from the RichDEM terrain analysis software (Barnes 2016b); however, the resulting sample network is passed back to Python where it is represented using the networkx package (Hagberg et al. 2008). Conceptually, our method is general to any acyclic network; the use of rasters and D8 is only a convenience meant to make the method accessible for the significant use-case of analysing data gathered on river networks.

The geographic locations of river sample sites and hydrologic stations are often misaligned to channels extracted from digital elevation models (Lindsay et al. 2008). As a result, observations are often automatically or manually snapped to the nearest channel cell in a digital elevation model (DEM), for example using GIS software. We strongly recommend performing such snapping prior to creating the sample network and using our method. If samples are not aligned to channel cells beforehand, samples may not produce a connected network or instead have unrealistically small drainage areas. We also recommend that the D8 raster be hydrologically conditioned prior to building the network such that modelled channel locations align closely with real observed channels. This hydrological conditioning often removes artificial depressions that prevent continuous flow-lines across a landscape. Algorithms to perform this sink-filling are widely available (e.g., Barnes et al. 2014; Barnes 2016a). Hydrologically conditioned D8 and DEMs are also increasingly publicly available at a variety of resolutions (e.g., Hydrosheds; www.hydrosheds.org).

3.2 Solving the unmixing problem

To solve the optimization problem posed by Equation 8 we take the networkx representation of the sample network from §3.1 and associate each node with a variable \( c \) for its concentration, which we wish to solve for. We also associate each node with a parameter \( \phi \) for its export rate and a constant \( a \) for its area. We therefore distinguish between variables, values we wish to solve for, and parameters, values which we set but would like to modify, and constants, whose values are fixed. We also add a dummy variable \( d \), the predicted concentration at the sample site, which is initially set to zero, but which we describe in terms of the concentrations of the upstream nodes, as in Equation 3. We then implement the forward model from §2.2 using a topological sort (p. 575; Sedgewick and Wayne 2011) on the networkx sample network. As the sort moves from upstream to downstream we propagate concentrations and fluxes downstream building Equation 3 for each node as it is reached. Finally, we extract the dummy variables \( d \) into a vector \( \mathbf{d}_{\text{pred}} \) and combine these with the observed values held in the \( \mathbf{d}_{\text{obs}} \) parameters to build the objective function from Equation 7. Optionally, regularization terms are built by creating a list of the differences between the \( c \) parameters and the mean observation parameters. If regularization is chosen, this requires the regularization strength \( \lambda \) to be defined giving us the optimization problem defined by Equation 8.

To improve numerical accuracy, we internally normalize the values of the areas, export-rates, and observed concentrations such that they are all approximately equal in absolute value. We do this by dividing each of these values by the geometric mean of all the values before calculations. The final calculated concentrations are then multiplied by these values to recover the original units. This normalization does however increase numerical accuracy when the problem is passed to solvers, below. Another consequence of this is that concentrations, areas and export rates can be provided in any units and our method will still produce the same results.

We implement the variables, parameters, objective function, and optimization problem from above using Python’s CVXPY package, a domain-specific language for disciplined convex programming (Grant et al. 2006; Diamond and Boyd 2016). The introduction of parameters means that we need to use CVXPY’s Disciplined Parametrized Programming extensions (Agrawal et al. 2019).

Once the problem has been expressed to CVXPY, CVXPY must perform an operation called canonicalization. Canonicalization ensures that the problem is convex, and therefore solvable, while converting the many variables and parameters in the problem into an intermediate representation suitable for passing to a solver. At present canonicalization in CVXPY is a very time-consuming operation relative to solving. Fortunately, if only the values of the parameters (in this case, the observation values, export-rates and regularization strength) are changed the intermediate representation can be updated without having to do canonicalization again. In practice this means that, once a problem has been composed, the export rates, observed concentration values, and regularization strength can be adjusted and subsequent solves occur rapidly. This allows for very efficient uncertainty propagation using Monte Carlo inversion as well as efficient calibration of the regularization strength parameter. An example of this is detailed below.

Solvers are the programs that find the solutions to a given optimization problem. A variety of solvers can be used for a given problem and CVXPY facilitates this by providing a unified way of writing problems. For our problem, we experiment with the SCS, ECOS, and Gurobi solvers (Domahidi et al. 2013; O’Donoghue et al. 2016; Gurobi 2023). A comparison of the performance of these solvers is detailed in §3.4. The solvers typically complete their work very quickly, even for large problems, and CVXPY translates their solutions back into a convenient form. Solvers often have a number of settings that trade accuracy for speed. In the tests described in §3.3 the accuracy requirements for ECOS had to be reduced from its default settings for it to converge. SCS was run with its default settings and Gurobi was run in a high-accuracy mode.

Note that solvers often perform preprocessing which allows them to separate out the components of disjoint problems. In practice this means that several disconnected river networks can be solved at once without having a significant impact on solution times (as shown in the case study below).
3.3 Method verification

To verify the numerical accuracy of our method we ran it on randomly-generated sample networks with known concentration values. Using the forward model (§2.2), we generated synthetic observations. Using our method, we inverted these observations to try to recover the original concentrations. We found that we can recover the original concentrations to within a specified tolerance thereby verifying that our method is accurate. We performed this test for each of three types of networks: random trees, full R-ary trees, and balanced trees. These three tree types cover a variety of structures likely to be represented in real datasets. The latter two networks required a branching factor to be specified, which determines how many upstream nodes flow into each downstream node. This branching factor was also chosen randomly. We assigned each node in each network a random concentration and random sub-basin area, both of which were drawn from log-uniform distributions. Subsequently, we generated a suite of synthetic concentration observations by mixing along the network using the forward model from §2.2. Finally, we solved the optimization problem from Equation 7 (without regularization) and compared what we recovered to the initial synthetic input. The software accompanying this paper implements the above test using Python’s Hypothesis testing library (MacIver et al. 2019). Unlike a standard unit test, which checks that a given input produces an expected output, Hypothesis implements property-based testing. In this form of testing a large number of random inputs are passed to a function and we test whether some known property holds for each of the inputs. This form of testing can often identify errors that a single, manually-created unit test would miss. In this case, the property we test is that, for any randomly-generated network and set of concentrations and areas, we can recover these values after simulating how they would propagate downstream through the network.

We draw concentrations and areas uniformly from log-uniform distributions over the range [1, 100] (i.e., over two orders of magnitude), the network size uniformly from the range [2, 100], and the branching factor uniformly from the range [1, 5]. For each trial scenario we used both the ECOS and GUROBI solvers to solve the optimization problem posed by Equation 7. We find that under these conditions we are able to successfully recover the upstream concentration for each upstream node to an accuracy of better than 1%. This test is included in the code repository.

Because we normalize absolute values by their geometric mean (§3.2) this test is valid for any datasets where sub-basin area and source concentration range across two orders of magnitude, regardless of the absolute values. Additionally, because the flux from each node is the product of sub-basin area and source concentration (for a fixed export rate), the above test corresponds to networks where the tracer flux from the node varies over four orders of magnitude. For sample campaigns where sub-basin areas are approximately equal in size, we find that we can draw source concentrations from an even greater range and still achieve the same level of accuracy. The concentration values we expect to encounter in a given problem can vary widely, and may vary beyond the specific range tested here. We therefore recommend conducting more specific simulated tests that match a specific problem’s characteristics in order to better quantify the accuracy of the recovered results—this is a best practice for any statistical analysis.

3.4 Benchmarking runtime

Our method performs quickly across a range of network sizes, including sizes larger than any sampling campaign we are aware of. To demonstrate this, we generate and solve random problems, as described in §3.3. For simplicity we consider only full R-ary trees with a branching factor of three (i.e., ternary trees). We run our benchmark for the SCS, ECOS and GUROBI solvers on networks up to a maximum size of 500 nodes. For each network size, ten trials were performed and the average and standard deviation of the runtimes were calculated. The results are shown in Figure 4. Calculations were performed on a 12th Gen Intel Core i9-12900HK processor with 64 GB RAM. A script to reproduce a version of this figure is provided in our code repository.

Our results indicate that on a standard desktop it is possible to compile and solve the unmixing problem on networks of 500 nodes—significantly larger than most geochemical datasets—in under a minute. For networks of less than 100 nodes, solutions can be found in under a second. For the first solve, ECOS and GUROBI show approximately linear scaling for networks up to about 100 nodes, with log-log slopes of 1.04 and 1.02 respectively. For networks larger than this, the compilation time begins to increase at a greater rate. SCS shows a more complex scaling.

Although the time taken for the first solve is thousands of times faster than existing methods, it is still slower than the time taken for subsequent solves of the same pre-compiled problem—those solves for which only parameter values are changed. Our benchmark shows that even for the largest networks we consider, subsequent solves take on the order of a second or less. For networks larger than ten nodes, the subsequent solves of both ECOS and GUROBI scale approximately linearly with respect to network size, with log-log slopes of 0.98 and 0.97 respectively. ECOS and SCS are faster than GUROBI. SCS shows a more complex scaling and is less consistent than the other solvers.

The total runtimes in Figure 4 include both the time spent in canonicalization as well as the time spent in the solvers. As discussed in §3.2 and demonstrated in the figure by the solver times (dashed lines), canonicalization dominates the runtime and canonicalizing the problem for the initial solve takes significantly longer than subsequent solves of the same problem using modified parameters. The amount of time spent in the solvers themselves is orders of magnitude less than the time spent building the problem. This shows that the compilation of the problem itself is limiting.

This limitation can be overcome, allowing very large problems to be solved, by using faster optimization languages such as Julia’s Convex.jl library (Udell et al. 2014). We test this on randomly-generated ternary trees and observe that Convex.jl, using Gurobi as a solver, takes 32 s to solve a 59,049 node tree and 334 s to solve a 177,147 node tree. Although this performance is attractive, the Python code we’ve used for the rest of this study is much more flexible and easier to use than its Julia counterpart.
Figure 4: Scaling of algorithm solve time against network size. Runtime tests performed on synthetic R-ary networks of branching factor three up to 500 nodes in size. Areas and concentrations of each node randomly varied between one and 100. For each network size, ten repeats were performed and averages and standard deviations were calculated. Different coloured lines indicate different solvers. Solid colours with solid lines indicate total runtimes for first solve of a problem including problem compilation. Faded colours with solid lines are total runtimes for all subsequent solves of a precompiled problem. Dashed lines indicate how much time is spent in the solvers themselves.
Runtime is not the only factor that should be considered in finding solutions to an optimization problem. We notice that Gurobi almost always finds optimal solutions to the problems we give it. In contrast, ECOS often finds solutions which it cannot guarantee are accurate, but which, in practice, are close to the values Gurobi finds in the cases we’ve checked. SCS is often unable to find optimal accurate solutions and quits early if it determines this. In Figure 4 this effect accounts for the wide variance of SCS run-times. For its accuracy, consistent convergence, and speed, we prefer to use the Gurobi solver for our method. However, ECOS is the default solver we use with our implementation because it is freely available whereas Gurobi requires a license (academic licenses are free at the time of writing).

4 CASE STUDY: CAIRNGORMS, SCOTLAND, UK

4.1 Background

To demonstrate the method we apply it to a river sediment chemistry dataset collected from rivers draining the Cairngorms mountains, UK (Figure 5a). This dataset was first presented in Lipp et al. (2020, 2021), where it was used to test the unmixing method discussed in §1.1. The dataset consists of fine-grained (< 150 μm fraction) samples gathered from 63 sites on five rivers (Spey, Dee, Deveron, Don, & Tay) draining NE Scotland (Figure 5b). The sample networks corresponding to this dataset are shown in Figure 5c. The region contains a diverse range of sedimentary, metamorphic and igneous rocks resulting in a diversity of geochemistry in source regions (5d). An advantage of applying the method in this region is that the source regions of the sampled rivers are already geochemically very well constrained by the G-BASE geochemical survey (Johnson et al. 2005) which gathered first order stream sediment samples at a density of ~1 sample per 2 km² (Figure 5e). The G-BASE data can therefore be used as an independent dataset to test our predictions of upstream concentrations. The primary control on source region chemistry is the composition of the underlying lithology (see Figures 5d–e).

4.2 Regularization

As this real dataset incorporates some sampling noise, we opt to regularize the problem to prevent overfitting. This regularization will result in solutions that are stable and meaningful despite noise in the input data. However, regularization does require us to calibrate the regularization parameter $\lambda$. Identifying a regularization parameter is in general a challenging task across all fields, and a full discussion is beyond the scope of this manuscript. We take an “Occam’s razor” style approach seeking the set of upstream concentrations with the smallest number of deviations from the mean observed value that is still able to adequately fit all of the observations downstream. We determine this by performing 100 inversions each with a different regularization parameter sampled equally between $10^{-3}$ and $10^{3}$. For each $\lambda$ we calculate the global root mean squared (RMS) error between the observed and predicted concentrations (i.e., between $d_{\text{obs}}$ and $d_{\text{pred}}$). Plotting this RMS against $\lambda$ results in the trade-off plot shown in Figure 6. We chose a value ($\lambda = 10^{-1}$) that lies at the “elbow” of the trade-off plot, a standard approach to choosing the $\lambda$ parameter.

4.3 Monte Carlo inversion

The uncertainty in the downstream observations was constrained by Lipp et al. (2020) by gathering duplicate samples at four randomly chosen sample sites. Concentration differences between the duplicate samples are controlled by the heterogeneity within a sample-site and also uncertainty introduced by the sampling process. Analytical uncertainty introduced in the laboratory process was noted to be negligible in comparison. For Magnesium, the duplicates on average showed a relative uncertainty (i.e., $\sigma/\mu$, where $\sigma$ is the standard deviation and $\mu$ is the mean) of 3.53%. To propagate this downstream uncertainty upstream we performed uncertainty quantification with Monte Carlo sampling by resampling the downstream data 1000 times from a distribution defined by the observed value and the relative uncertainty. We invert each of these resampled datasets and calculate the geometric mean and standard deviation of the upstream predictions.

4.4 Results

The central estimate and uncertainty for Mg are shown in Figure 7a–b. We can assess the accuracy of these predictions by comparison to the independent G-BASE geochemical survey. The predicted upstream concentration in each subcatchment represents the mixture of all sources of the element within the subcatchment. To provide an independent estimate of this mixture, we calculate the arithmetic mean concentration of all G-BASE samples (Figure 5e) within each sub-basin, resulting in the map shown in Figure 7c. We define the uncertainty on the G-BASE estimate for sub-basin concentration as the standard error of the calculated mean (i.e., $\sigma/\sqrt{N}$). Only one small sub-basin in the lower reaches of the Tay basin contained no G-BASE samples and therefore the predictions for this sub-basin can not be independently validated. Visual comparison of the upstream predictions (Figure 7a) and the independent G-BASE dataset (Figure 7c) indicate a strong similarity. Visualising the misfit spatially (Figure 3d) shows close accordance for most of the study area barring a few sub-catchments to the south. The cross-plot is shown in Figure 7e. A linear-regression through these data, weighted by upstream-area, returns a slope sub-parallel to the 1:1 line with an $R^2$ value of 0.63 and a root mean squared misfit of 0.13. This model is able to closely fit the concentration downstream shown in Figure 7f. Overall we conclude that the inversion scheme is able to generate accurate upstream concentrations with associated uncertainties.

A major advance of this study is the ability to efficiently invert the problem repeatedly for different parameter and data values. For example, the entire procedure described above including the calibration of the regularization parameter and the Monte Carlo calculation of uncertainties was performed in ~ 30 seconds on a standard desktop computer, despite performing over 1000 separate inversions. In studies incorporating larger datasets, each of these wholly independent inversions could be performed in parallel for further speed-ups. Previous methods took hours to perform a single inversion, which precluded Monte Carlo resampling.
Using convex optimization to efficiently apportion tracer and pollutant sources.

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**Map a**: Elevation / m

**Map b**: Map of Scotland

**Map c**: Flowchart showing the distribution of rivers

**Map d**: Map of Mg concentration

**Map e**: Map of Mg concentration with color scale

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Submitted manuscript; not peer-reviewed.
As a consequence of this generality, our method could be applied to any constituent in sediment or water that is well-mixed and conservative. The possibility of extending to non-conservative and time-varying tracers is discussed below. Additionally, the concentration of cosmogenic nuclides (e.g., $^{10}$Be, $^{26}$Al) in river sands are used to quantify the average erosion rate (Blanckenburg 2005). By unmixing nested cosmogenic nuclide samples, erosion rates could therefore be quantified across catchments at a greater spatial resolution.

Finally, whilst we consider only river networks in this study, our approach can be applied to unmixing tracers along any other structure that can be represented as a directed acyclic graph, such as municipal sewerage networks. For example, waste-water epidemiology uses observations of disease tracers such as the genetic remnants of SARS-Cov-2 in sewerage systems to identify the severity and timing of outbreaks (Sims and Kasprzyk-Hordern 2020). If a pollutant is measured at multiple nested sites across a basin, the presented scheme can be used to apportion the pollution into the contribution from each sub-basin. This approach could be particularly useful for characterising diffuse pollutant sources, which are typically challenging to identify and quantify (Bowes et al. 2008). This inverse approach to apportionment contrasts with the more common forward modelling approaches to estimating pollutant contributions e.g., by assuming a relationship between land-use and pollutant production (e.g., “export coefficients”; Johnes 1996).

The elemental and isotopic chemistry of river water and sediments is also frequently measured to understand controls on chemical erosion rates in upstream regions (e.g., Bufe et al. 2021); however, it is known that mixing of chemically heterogeneous source regions obscures the primary source region signature (Baronas et al. 2017; Torres et al. 2017). The scheme presented here can be used to account for these mixing effects allowing for more accurate estimates of upstream chemical weathering rates. This in turn can be used to better constrain their underlying driving mechanisms. Additionally, the concentration of cosmogenic nuclides (e.g., $^{10}$Be, $^{26}$Al) in river sands are used to quantify the average erosion rate (Blanckenburg 2005). By unmixing nested cosmogenic nuclide samples, erosion rates could therefore be quantified across catchments at a greater spatial resolution.

5 Discussion

5.1 Applications

The method we present is designed to be deliberately general and can be applied to any constituent in sediment or water that is well-mixed and conservative. The possibility of extending to non-conservative and time-varying tracers is discussed below. As a consequence of this generality, our method could be applied to a number of problems in different disciplines. For example, in the field of geochemical exploration, nested sampling campaigns are frequently conducted for river and stream sediments. These samples are then used to infer the geochemistry of the upstream lithology for mineral exploration or for setting environmental baselines (Garrett et al. 2008). This unmixing procedure can be applied to such data to accurately reconstruct the geochemistry of the sediment source regions (e.g., Lipp et al. 2023).

The method could also be used in pollutant apportionment studies. If a pollutant is measured at multiple nested sites across a basin, the presented scheme can be used to apportion the pollutant into the contribution from each sub-basin. This approach could be particularly useful for characterising diffuse pollutant sources, which are typically challenging to identify and quantify (Bowes et al. 2008). This inverse approach to apportionment contrasts with the more common forward modelling approaches to estimating pollutant contributions e.g., by assuming a relationship between land-use and pollutant production (e.g., “export coefficients”; Johnes 1996).

Finally, whilst we consider only river networks in this study, our approach can be applied to unmixing tracers along any other structure that can be represented as a directed acyclic graph, such as municipal sewerage networks. For example, waste-water epidemiology uses observations of disease tracers such as the genetic remnants of SARS-Cov-2 in sewerage systems to identify the severity and timing of outbreaks (Sims and Kasprzyk-Hordern 2020). If a suite of samples were taken on a sewerage network (e.g., Nourinejad et al. 2021) our approach could be used to isolate where in the network SARS-Cov-2 is entering the system and, therefore, to estimate where, geographically, the outbreak is occurring. A similar strategy could be applied to other diseases such as polio, malaria, HIV, tuberculosis, measles, and zika (Kilaru et al. 2023). We will explore this further in future work.

5.2 Tributaries, drainage areas and designing sampling campaigns

Our method works by solving for the source contribution from the sub-basin defined by individual samples. As a result, our method is agnostic to whether and where any specific tributaries may be joining other channels within that sub-basin. For example, in Figure 1, Samples 3 & 5 lie on a tributary upstream of...
Sample 6. If these samples are excluded, the sub-basin defined by Sample 6 includes the entire catchment of this tributary. As a result, the calculated source concentration for sub-basin 6 is the average production rate of a large area. Such a large sub-basin may not provide useful information.

This result has implications for how to sensibly design sampling campaigns to recover source concentrations. If the goal is equal coverage across a basin, samples should be gathered such that the resulting sub-basins are approximately equal in area. Such a campaign would likely involve nested sampling into tributary basins. Simply sampling along the main channel could result in irregular, large, sub-basins which are not diagnostic. Consequently, it would be a challenge to disambiguate between what is generated on the main channel versus what is added to it by tributaries. Algorithms to identify suitable sample sites for finite sampling resources given these constraints would be an interesting further avenue of research.

5.3 Solving for optimal export-rates

To solve the optimization problem discussed in this manuscript (i.e., Equation 7) the relative export rate of water or sediment from each sub-basin needs to be known or assumed. By default, our implementation assumes that the export rates of all the sub-basins are the same. If estimates of the run-off rate or sediment export-rate are known for each sub-basin (for example, from gauging data) these values can instead be used for more accurate results. However, we note that previous studies have found that synthetic variations in export rate have a limited impact on the results of the inversion (Lipp et al. 2021, 2023). This insensitivity was attributed to the fact that within a sampled river basin, geochemical concentrations typically show much greater variability than sediment/water export rates. Nonetheless, an interesting question is whether optimal estimates of the export rate parameters, \( \phi \), can also be recovered from downstream observations. In other words, can a set of concentration observations be jointly inverted for both sub-basin source concentration and the export-rate. We now explore how this could be posed as an inverse, convex optimization problem.

The convex minimization expression in Equation 7 can be reformulated as a misfit function of the export rates, and a vector of tracer observations:

\[
F(\phi, \mathbf{d}_{obs}) = \min_{\phi \in \mathbb{R}^n} \sum_{i \in \mathbb{N}} \text{misfit}\left(d_{obs,i}, \frac{\sum_{j \in U_i} \phi_j a_j}{\sum_{j \in U_i} \phi_j a_j}\right),
\]

where “min.” indicates the minimum value of the expression. This function, \( F \), therefore maps a vector of export-rates and a vector of tracer concentrations onto a single value representing the misfit to the data. Note that because Equation 7 is convex, we can conclude that \( F \) must map onto a single value for the vectors \( \phi, \mathbf{d}_{obs} \) (i.e., for \( n \) observations, \( F : \mathbb{R}^n \times \mathbb{R}^n \rightarrow \mathbb{R} \)).

For a particular dataset each sample-site may have observations of many different tracers. For example, methods for determining elemental concentrations in water and sediment typically return the concentrations of many elements at once. We can therefore construct a matrix of concentration observations, \( \mathbf{D}_{\text{exm}} \), of \( m \) different tracers at \( n \) different sample-sites. Each column of \( \mathbf{D} \) is a vector equivalent to \( \mathbf{d}_{obs} \). However, in each of these cases, whilst \( \mathbf{d}_{obs} \) is different for each tracer, the export rate of the sub-basin is always the same. This is because whilst the particular value of \( d \) will be controlled by the tracer sources (e.g., lithology, pollutants, etc.) the total material export is controlled by independent processes such as erosion or rainfall rate. Collectively, all of these independent tracers provide some constraint on the upstream export rates. We are looking for the vector of export rates, \( \phi \), that minimizes this expression, which gives, finally, the minimization problem:

\[
\min_{w.r.t. \phi} \sum_{j=1}^{m} F(\phi, \mathbf{D}_j).
\]

Solving this expression could, potentially, be a novel approach to yield important geomorphic and hydrologic information. For example, if tracers in sediment are analyzed, the solution could provide, jointly, the relative upstream erosion rate in each sub-catchment as well as the source region geochemistry just from a set of point observations in a network. In the case of water, it would return upstream run-off as well as hydrochemical information. Equation 10 could be considered as a special instance of the general “explicit” mixing problem of Weltje (1997), which solves for mixture proportions and source region concentrations, applied to mixtures upon a network.

Unfortunately we have not yet found a convex approach to solving Equation 10, suggesting it may be challenging to optimize. Additionally, the mapping \( F \), and the resulting misfit space, depends on the structure of the sample network as well as the dataset of geochemical observations. Consequently, a full exploration of this optimization problem, including instances when it is easily solved, is beyond the scope of this manuscript. Nonetheless, we state it here as a potential interesting avenue for future work.

5.4 Comparison to prior mixing models

Methods to do some form of unmixing of hydrochemical and sedimentary geochemical observations do already exist in a
variety of forms (e.g., Christophersen et al. 1990; Weltje 1997; Stock et al. 2018; Kemeny and Torres 2021). The most common form is to assume that the observations are mixtures of known source compositions (generally referred to as “endmembers”). For example, Kemeny and Torres (2021) describe the chemistry of river water as a mixture of water derived from weathering of a discrete set of lithologies, each with a distinct chemical composition (e.g., carbonates, silicates, and evaporites). This formulation of the unmixing operation can be solved with least-squares. This type of approach is widely used in hydrology (e.g., Christophersen et al. 1990) and for determining the sources of sediment pollution, where it is termed “sediment fingerprinting” (Walling 2005; Collins et al. 2020). However, the requirement for a priori knowledge of likely endmembers can be a significant burden. This either requires making significant assumptions about sources or the gathering of further data to constrain the endmembers. Approaches have therefore been developed to determine both the likely endmembers and mixing proportions concurrently (Weltje 1997; Carrera et al. 2004; Valder et al. 2012; Shaughnessy et al. 2021).

However, none of these approaches, unlike the one presented here, consider the spatial distribution of the samples. Our approach is therefore unique in considering the topology of a drainage network when applied to observations nested on a network. The method presented here is most similar to the methods developed by Carraro et al. (2018, 2020) for mapping the sources of downstream eDNA observations in river basins. However, Carraro’s approach requires continuous datasets of auxiliary variables across the studied region such as lithology or temperature, on which it trains an optimal transfer function. In cases where auxiliary data is absent, or not relevant to the target tracer (such as a point-source pollutant), such an approach cannot be used. By contrast, our approach requires only very limited upstream information.

5.5 Further work

The approach as defined here is only able to consider tracers which can be considered conservative on the timescales of mixing. However, it could be extended to take into account non-conservative tracers under certain assumptions. For example, if the concentration of a non-conservative tracer can be modelled to decay exponentially with time spent in transport, this can be added as a prefactor to the mixing calculation, making assumptions regarding transit times. Carraro et al. (2018) take such an approach for determining the sources of non-conservative eDNA in drainage basins. This approach would require the calibration of this decay hyper-parameter potentially using field measurements (e.g., Guiliet et al. 2019). Elsewise, it could be calibrated using a held-out subset similar to the regularization parameter. Such an approach could be used for determining the sources of a number of significant non-conservative pollutants such as nitrate and phosphate.

At present we consider only well-mixed tracers, whose concentration does not vary significantly in time. This assumption is valid only when the sources of a tracer vary slowly with respect to the tracer travel-time. However, hydrochemical data is observed to sometimes vary transiently over storm-events (Knapp et al. 2020). Additionally, a pollutant released from a point-source for a small duration may also result in transient increases in concentration downstream (e.g., waste water released from sewage overflows; Munro et al. 2019). Consequently, a scheme able to partition the sources of tracers in space and time given nested time-series observations would be desirable (see Benettin et al. 2022). This development would require both time-series hydrochemical observations (e.g., using in-situ sensors; Rode et al. 2016) and a model of the drainage network hydrodynamics. Alternatively, composite samples could be gathered over an extended period of time, averaging out temporal variability (e.g., Cassidy et al. 2018).

6 Conclusions

We describe a very efficient method for unmixing nested concentration data on drainage networks to determine the location and magnitude of the tracer sources. Our approach is applicable to well-mixed conservative tracers in both river sediment or water. We show that the unmixing problem can be posed as a convex optimization problem which can thus be solved very quickly while being guaranteed to find a single global solution. This method also scales well to large problems. The rapid solve time allows for uncertainties in the upstream concentration to be calculated using Monte Carlo sampling. Our method can be used for apportioning pollutant sources within river basins, for more accurately constraining geochemical processes, or for generating geochemical maps. Beyond rivers, this approach could be used to analyze nested concentration data on other kinds of networks such as sewerage networks in the field of wastewater epidemiology. We provide an open-source Python implementation of our method that is benchmarked, unit-tested, and accompanied by examples. This implementation requires only a raster of hydrological flow-directions and a data-table giving sample-site locations and associated concentration data. We believe our network-based approach can be extended to consider tracers that are non-conservative, and not in steady-state. Our method therefore provides a strong framework for a wider inverse approach for analysing geochemical data gathered on drainage networks, as well as in other settings.

Open Research

Our Python package and relevant data is available from github.com/r-barnes/faster-unmixer and archived at zenodo.org/record/8299812.

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References


A LINEAR LEAST-SQUARES SOLUTION TO UNMIXING ON A NETWORK

In the main manuscript we use our unmixing method to recover the magnitude of a set of source concentrations from their observed counterparts using convex optimization, penalising relative differences between predicted and observed concentrations. In some scenarios, it may be valid to instead penalize absolute differences between tracer observations. For example, isotopic geochemical data typically has very low log-variances and so absolute differences between observations may be an appropriate approximation (Blondes et al. 2016). Here we show that if absolute differences are penalised, the unmixing problem can be posed as a linear least-squares matrix inversion.

We consider the directed acyclic graph shown in Figure 5 with sample sites 1–6. This network of sample sites, G, has observed concentrations \( \mathbf{d} = [d_1, d_2, \ldots, d_6] \), source concentrations \( \mathbf{c} = [c_1, c_2, \ldots, c_6] \), and total fluxes \( \mathbf{q} = [q_1, q_2, \ldots, q_6] \). As described in Equation 3, we can construct expressions for \( \mathbf{d} \) as mixtures of the source concentrations \( \mathbf{c} \), weighted by the fluxes \( \mathbf{q} \):

\[

d_1 = c_1 \\
d_2 = c_2 \\
d_3 = c_3 \\
d_4 = \frac{q_1 c_1 + q_2 c_2 + q_4 c_4}{q_1 + q_2 + q_4} \\
d_5 = \frac{q_3 c_3 + q_5 c_5}{q_3 + q_5} \\
d_6 = \frac{q_1 c_1 + q_2 c_2 + q_3 c_3 + q_4 c_4 + q_5 c_5 + q_6 c_6}{q_1 + q_2 + q_3 + q_4 + q_5 + q_6}
\]

This set of linear equations can be represented in matrix form:

\[

d = \begin{pmatrix}
1 & 0 & 0 & 0 & 0 & 0 \\
0 & 1 & 0 & 0 & 0 & 0 \\
0 & 0 & 1 & 0 & 0 & 0 \\
\frac{q_1}{\sum q_1} & \frac{q_2}{\sum q_1} & 0 & \frac{q_4}{\sum q_1} & 0 & 0 \\
0 & 0 & \frac{q_3}{\sum q_3} & \frac{q_5}{\sum q_3} & 0 & 0 \\
\frac{q_1}{\sum q_1} & \frac{q_2}{\sum q_1} & \frac{q_3}{\sum q_3} & \frac{q_4}{\sum q_4} & \frac{q_5}{\sum q_5} & \frac{q_6}{\sum q_6}
\end{pmatrix}
\begin{pmatrix}
c_1 \\
c_2 \\
c_3 \\
c_4 \\
c_5 \\
c_6
\end{pmatrix}
\]

which is the linear matrix equation \( \mathbf{d} = \mathbf{Mc} \), that can be inverted for \( \mathbf{c} \), if \( \mathbf{M} \) is known. We therefore seek an expression for each entry in the matrix \( \mathbf{M} \).

Note that \( M_{ij} \) is non-zero only when either \( i = j \) or node \( j \) is upstream of node \( i \). We therefore define a 0-1 matrix \( \Theta \) which indicates whether an entry in \( \mathbf{M} \) is zero or not. \( \Theta \) is simply the path matrix of the graph \( G \) where every node also has a self-edge; that is \( A_{ij} = 1 \) if a path exists from node \( j \) to node \( i \) and every node has a path to itself. With \( \Theta \) defined we can thus build the following expression for \( \mathbf{M} \):

\[
M_{ij} = \frac{\Theta_{ij} q_j}{\sum_{k=1}^{n} \Theta_{ik} q_k}.
\]