Improving In-Stream Nutrient Routines in Water Quality Models Using Stable Isotope Tracers: A Review and Synthesis

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ABSTRACT. Water quality models serve as an economically feasible alternative to quantify fluxes of nutrient pollution and to simulate effective mitigation strategies; however, their applicability is often questioned due to broad uncertainties in model structure and parameterization, leading to uncertain outputs. We argue that reduction of uncertainty is partially achieved by integrating stable isotope data streams within the water quality model architecture. This article outlines the use of stable isotopes as a response variable within water quality models to improve the model boundary conditions associated with nutrient source provenance, constrain model parameterization, and elucidate shortcomings in the model structure. To assist researchers in future modeling efforts, we provide an overview of stable isotope theory; review isotopic signatures and applications for relevant carbon, nitrogen, and phosphorus pools; identify biotic and abiotic processes that impact isotope transfer between pools; review existing models that have incorporated stable isotope signatures; and highlight recommendations based on synthesis of existing knowledge. Broadly, we find existing applications that use isotopes have high efficacy for reducing water quality model uncertainty. We make recommendations toward the future use of sediment stable isotope signatures, given their integrative capacity and practical analytical process. We also detail a method to incorporate stable isotopes into multi-objective modeling frameworks. Finally, we encourage watershed modelers to work closely with isotope geochemists to ensure proper integration of stable isotopes into in-stream nutrient fate and transport routines in water quality models.

Keywords. Isotopes, Nutrients, Uncertainty analysis, Water quality modeling, Watershed.

Deterministic water quality models provide an economically feasible approach to quantify fluxes and transformations of nutrients and for scenario analysis of dynamic management, land use, and climate conditions. Nevertheless, the reliability of such models to assist with management decisions is questioned due to compounding uncertainties regarding in-stream transformation rates of contaminants (Beven, 2006; Rode et al., 2010; Robson, 2014; Yen et al., 2014; Wellen et al., 2015; Han and Zheng, 2016). It is the general sentiment in the hydrology and water quality community that researchers need to reduce uncertainty within water quality models. In this article, we work toward this goal by providing a review and synthesis of how stable isotope tracers can reduce uncertainty in these applications.

High uncertainty within water quality modeling is likely an artifact of the historical development of water quality models and continued advancements in perceptual understanding of fluvial biogeochemistry. Following a historical period that saw the development of hydrologic and biogeochemical functions from data collected at the hillslope-plot scale and stream-reach scales in the early 1970s and 1980s, watershed water quality modeling saw rapid advancement via computational capabilities in the 1990s and 2000s to address growing environmental issues related to nutrients (e.g., estuary seasonal hypoxia). Computational advancement allowed several modeling characteristics to take shape, including the ability to inexpensively incorporate spatially explicit data, perform computations at a different resolution or environment than originally envisioned, and couple water, particulate, and dissolved phases within single numerical model formulations. However, the computational advancement of water quality models has not necessarily negated the conceptual representation of in-stream physics and biogeochemistry. For example, conceptual models have been shown to be quite powerful for understanding fluxes from watersheds (Ford et al., 2017). However, new monitoring and measurement capabilities have shown researchers that coupled physio-biochemical processes may vary from the original hydrologic and biogeochemical functions in models. Furthermore, computational advancements have shifted parameterization of models away from inputs and parameters consistent with their original scale of observation and have produced numerous likely inputs and parameter sets within modeling frameworks (e.g., equifinality, as described below). As a result, computational abilities have outweighed the modeler’s ability to constrain input and parameter values.
and have promoted large posterior solution spaces, resulting in high uncertainty. Such uncertainty should be accounted for when reporting and analyzing the results of water quality models.

Given the need to constrain input and parameter values and prevent erroneous model parameterization, innovative data streams should be incorporated into water quality models. Integration of stable isotopes for carbon (C), nitrogen (N), and phosphorus (P) compounds within the model architecture provides one such measurement tool to assist with model uncertainty reduction. This assertion follows recent successes in using water isotope measurements to help parameterize model boundary conditions, reduce model uncertainty due to equifinality, and improve numerical representation of processes within hydrologic model structure (Seibert and McDonnell, 2002; McGuire and McDonnell, 2007; McDonnell and Beven, 2014; Windhorst et al., 2014; Soulsby et al., 2015; Yamanaka and Ma, 2017). In this light, this review article synthesizes the utility of stable isotopes within water quality models to reduce the uncertainty contributed by overparameterization in numerical model estimates, given the ability of stable isotopes to be measured with relatively high precision and accuracy. Our focus is on in-stream biogeochemical modeling of macronutrients, namely C, N, and P, but at the same time it is well-realized that accurate representation of water and solids (i.e., sediment) within streams is a precursor to predicting C, N, and P fluxes and transformations.

We show recent literature evidence that coupling stable isotopes within watershed water quality modeling helps with improving the data inputs associated with: (1) providing boundary conditions of the models, (2) constraining model parameterization, and (3) elucidating improvements needed within the conceptual and numerical representation of processes, i.e., the model structure. The efficacy of stable isotopes for this uncertainty reduction goal is noteworthy, given that recent attention on watershed water quality modeling uncertainty has highlighted these same inaccuracies (i.e., problems with precision and accuracy of input and calibration measurements, uncertainty in parameter specification, and the problem of inaccurate model structure) as three major sources of uncertainty within models (Guzman et al., 2015).

Providing Boundary Conditions

Boundary condition refers to the source contributions of C, N, and P phases that need to be considered within watershed water quality modeling. For example, within a nutrient focused model, the boundary condition inputs refer to the spectrum of potential nutrient inputs, such as N and P from agricultural and urban sources (Xue et al., 2009; Young et al., 2009; Kendall et al., 2010). As another example, within a sediment C focused model, the boundary condition inputs refer to the spectrum of potential sediment C inputs, such as inorganic C, terrestrial particulate C, and autochthonous particulate C (Fox and Ford, 2016; Husic et al., 2017a). The use of stable C, N, and P-bound isotopes to elucidate the boundary condition inputs within the fabric of watershed water quality modeling is perhaps the most obvious coupling of isotopes with the models, given the widely used data-driven unmixing analysis for apportioning source contributions of both dissolved and particulate phases. Source apportionment has existed as a standalone method; therefore, coupling this method to assist with boundary conditions within water quality models seems natural. For these reasons, several studies have used stable isotopes to assist with boundary conditions within numerical models (Hong et al., 2014; Sebestyen et al., 2014; Xue et al., 2014; Fox and Martin, 2014; Ford and Fox, 2015; Husic et al., 2017b).

Constraining Model Parameterization

Constraining parameter uncertainty is another prominent problem with in-stream models, especially as the level of model complexity via coupling of processes and phases (i.e., dissolved, particulate, water) increases. For such models, the broad range of parameters leads to large posterior solution spaces for fluxes and transformations. Parameter specification uncertainty is robustly reflected by the concept of equifinality, which refers to the potential for a posterior solution space of acceptable calibrations to be met by multiple parameterizations, or realizations (Beven, 2006; Adiyanti et al., 2016). The Generalized Likelihood Uncertainty Estimation (GLUE) framework provides a means to quantify equifinality and is applied using Monte Carlo-based realizations of a global parameter space and evaluation of the subsequent solutions against measured data to create a posterior solution space (Beven and Binley, 1992; Dean et al., 2009; Jin et al., 2010; Gong et al., 2011; Shen et al., 2012; Ford and Fox, 2017). The acceptance into such a solution space depends on evaluation of measured and modeled data using statistical metrics such as Nash-Sutcliffe efficiency, percent bias, and ratio of the root mean square error to the standard deviation of measured data, e.g., Moriasi et al. (2007). While we commend the excellent work of researchers in quantifying this uncertainty, it has been shown that stable isotopes may also be coupled with water quality models to further reduce such uncertainty (Adiyanti et al., 2016; Ford et al., 2017). In many ways, elucidation of parameterization via stable isotopes within watershed water quality modeling is another highly conceivable method, given the long history of stable isotopes to elucidate reactions (Sharp, 2007). Essentially, stable isotopes may balance mass that couple biogeochemical reactions within their structure may be added to the elemental mass balances of water quality models, as described in the “Overview of Stable Isotopes” section. These added equations are often accompanied with few new unknowns or insensitive unknowns; therefore, a stable isotope data stream may assist with model parameterization. For these reasons, several studies have used stable isotopes to help with parameterizing water quality models (Tobias and Böhlike, 2011; Van Engelund et al., 2012; Hong et al., 2014; Fox and Martin, 2014; Ford and Fox, 2015; Adiyanti et al., 2016; Ford et al., 2017).

Elucidating Model Improvements

Elucidating improvements in model structure reflects a third opportunity where stable isotopes may assist with advancing research. As the complexity of nutrient cycling continues to unravel through contemporary measurement techniques, it is recognized that numerical model error can be associated with epistemic uncertainties. Regarding epistemic uncertainty, model structure errors may stem from
simplified conceptual models, the equations and algorithms used to reflect that conceptualization, and instabilities of the numerical scheme (Borah and Bera, 2003; Guzman et al., 2015). Recent critiques of water quality models have pointed to a need for improving in-stream biogeochemical simulations (Rode et al., 2010; Robson, 2014; Wellen et al., 2015). As an example, advanced deterministic models that reflect in-stream C and nutrient fate and transport (e.g., AQUATOX, QUAL2K, and WASP) conceptualize the benthos as a two-layer system (1 mm aerobic and 10 cm anaerobic) in which all particulate organic matter is contained in the anaerobic layer and is not subjected to erosion-deposition dynamics (Di Toro, 2001; Wool et al., 2006; Chapra et al., 2008; Park et al., 2008). This conceptualization was well-validated for large, slow-moving waterbodies; however, for turbulent low-order and low-gradient streams, recent research has highlighted the importance of a dynamic 5 to 10 mm aerobic sediment layer (i.e., the surficial fine-grained laminae) that controls the seasonality of benthic C and N dynamics (Droppo et al., 2001; Walling et al., 2006; Russo and Fox, 2012; Ford and Fox, 2014, 2015, 2017; Fox et al., 2014). As models become more robust, unique tools and approaches are needed that rigorously test our conceptualization of in-stream fate and transport. Stable isotopes coupled within water quality modeling may be used through iterations to enhance or test the validity of the model structure (Tobias and Böhlke, 2011; Hong et al., 2014; Sebestyen et al., 2014; Ford et al., 2017).

This review explains the utility of stable isotopes in improving existing water quality model predictions and reducing uncertainty by improving in-stream nutrient fate and transport routines, specifically by (1) providing boundary conditions of the models, (2) constraining model parameterization, and (3) elucidating improvements needed within the model structure. To support the use of stable isotopes for these goals within water quality models, we provide a sequential and comprehensive review of stable isotopes within the fabric of water quality models. First, we define and explain stable isotope theory for modeling-focused researchers who have had minimal exposure to isotope signatures. Second, we describe the pools, measurements, and applications of stable isotope signatures related to C, N, and P cycles in streams. Third, we describe the ability of the isotopes to elucidate sources and transformations so that modelers can understand the breadth of possibilities of where the isotopes are applicable in the stream environment. Fourth, we review watershed water quality modeling studies that have coupled stable isotopes and show how these studies have used the isotopes to reduce uncertainty associated with (1) providing boundary conditions of the models, (2) constraining model parameterization, and (3) elucidating improvements needed within the model structure. Fifth, we provide recommendations to watershed water quality modelers for coupling isotopes into the fabric of the modeling architecture.

**Overview of Stable Isotopes**

Stable isotopes of a given element have identical chemical properties except for a difference in atomic mass, which is caused by the variable number of neutrons in the nucleus. Carbon, nitrogen, oxygen, and hydrogen all have heavy and light stable isotopes, and the relative abundance of the heavy isotope is measured with high precision using isotope ratio mass spectrometry. The relative abundance of heavy to light isotopes for different oxidation states of an element (e.g., ammonium, nitrate, nitrite) is indicated by the widely used delta (δ) notation. In the determination of isotopic ratios, the relative differences between a sample and a reference standard may be ascertained with high precision. The delta notation (δ) was developed by McKinney et al. (1950) to report stable isotope data and is generically defined in equations 1 and 2:

$$\delta = \left( \frac{R_{\text{sample}} - R_{\text{standard}}}{R_{\text{standard}}} \right) \times 1000 \quad (1)$$

where $R$ is the ratio of the abundance of the heavy to light isotopes, $\text{sample}$ is the sample, and $\text{standard}$ is the reference standard that has a known isotope ratio. $R$ is defined explicitly as:

$$R = \frac{[^m X]}{[^o X]} \quad (2)$$

where $^m X$ is the heavy isotope, and $^o X$ is the light isotope.

The unit of measurement for δ values is reported in per mil or parts per thousand, represented as ‰, which reflects the relatively low abundance of heavy isotopes in the natural environment. A positive δ value indicates that the ratio of heavy to light isotopes is greater in the sample than in the standard, and vice versa for a negative δ value.

Stable isotopes are particularly effective for fingerprinting sources and quantifying rates of biogeochemical transformations due to the preferential use of lighter isotopes in a process termed isotope fractionation. Fractionation is characterized by either equilibrium or kinetic isotope effects (Sharp, 2007). In equilibrium isotope-exchange reactions, the forward and backward reaction rates of any single isotope are equal. Kinetic isotope effects cause isotope fractionation to happen when the system is not in isotopic equilibrium and the forward and backward reaction rates are not equal. In kinetic isotope fractionation, the reaction rates are factors of the isotope masses and their vibrational energy; bonds between the light isotopes break more easily than the heavy isotopes, which have stiffer bonds. This results in the preferential use of lighter isotopes during processes because less energy is required to break the lighter bonds. Fractionation processes that are not metabolically driven or kinetically controlled are associated with either an isotope fractionation factor (α) or an enrichment factor (ε), which is determined either analytically or experimentally. These values are directly related to one another through equations 3, 4, and 5:

$$\alpha_{A-B} = \frac{R_A}{R_B} \quad (3)$$

where $\alpha_{A-B}$ is the partitioning of stable isotopes between two substances $A$ and $B$ (fig. 1), and $R$ is the ratio of heavy to light isotopes, as described by equation 1, calculated for each substance. This equation is expressed as:
\[
\delta_{\text{MX}} = \frac{1000 + \delta_A}{1000 + \delta_B} \tag{4}
\]

where \(\delta\) is the relative abundance, as described by equation 2, calculated for each substance. The fractionation factor \((\alpha)\) is then related to the enrichment factor \((\varepsilon)\) as:

\[
\varepsilon = (\alpha - 1) \times 1000 \tag{5}
\]

Using the \(\delta\) values and fractional contributions of known sources coupled with the \(\varepsilon\) values and rates of reactions, the resulting value of a product is estimated. Namely, the famous Rayleigh formulation (Kendall and Caldwell, 1998) shown in equation 6 is used and coupled to isotope mass balance considerations in separate processes in which a product is removed from a reactant. The Rayleigh equation is used to describe isotopic fractionation processes under the following assumptions: (1) in a mixed system, material is continuously removed that contains molecules of at least two isotopic species (e.g., water with \(^{18}\text{O}\) and \(^{16}\text{O}\)), (2) the fractionation associated with the removal process at any instant may be described by the fractionation factor and the enrichment factor, and (3) the fractionation factor and enrichment factor remain constant during the process (Kendall and Caldwell, 1998). The Rayleigh equation may be described as:

\[
\delta^M_{X_b} = \delta^M_{X_A} - \varepsilon_{\text{rxn}} \ln(f_{B-A}) \tag{6}
\]

where \(M\) is the atomic mass of the isotope, \(X\) is the isotope, \(A\) and \(B\) are the two substances, \(\text{rxn}\) is the reaction process or pathway of removal, and \(f\) is the fraction remaining after the process occurs. Application of this equation becomes invalid under transient kinetic fractionation, which occurs when the reactions leading to fractionation do not follow first-order kinetics (Maggi and Riley, 2009). In general, this limitation may be assumed to have minor impact for nutrient-rich systems and would not be rate-limiting in terms of the lack of availability of the lighter isotope during removal.

Equation 6 is a suitable general definition of the enrichment process, but it may be expanded to accurately represent the dynamics of the system. Multiple inputs across a specified control volume will result in a mixing of sources, as shown in figure 1 (left box). To more accurately represent the upstream conditions, \(\delta^M_{X_i}\) may be broken into a summation incorporating the weighted average of each of the unique source inputs (e.g., the three-source mixing example in fig. 1) as:

\[
\delta^M_{X_A} = \sum_{l=1}^{k} \delta^M_{X_l}(W_l) \tag{7}
\]

where \(l\) is the source identifier, \(k\) is total number of sources, and \(W_l\) is the fraction of element \(X\) from source \(l\). Furthering this concept of multiple factors influencing the overall \(\delta^M_{X}\) value, figure 1 (right box) provides a generic definition of the processes of isotope fractionation to impact stream isotope signatures in a generic stream reach with a generic isotope tracer. Prior to entering the stream at input \(A\), there is an abundance of the light isotope in contrast to the heavy isotope. As the substance flows through the stream channel, different biogeochemical processes (e.g., \(\varepsilon_1\) and \(\varepsilon_2\)) occur that preferentially use the lighter isotope in contrast to the heavier isotope. These reactions impact the mass and isotope composition of the outputs depending on the magnitude of the process and the preference for the lighter isotope. As shown in output \(B\) of figure 1, the size of the substance pool decreases and the ratio of heavy to light isotopes increases relative to input \(A\) because of the fractionation processes \((\varepsilon_1\) and \(\varepsilon_2\)). The influence of the different biogeochemical processes and fractionation factors may be reflected in the general expression of equation 6 as:

\[
\delta^M_{X_B} = \delta^M_{X_A} - \sum_{o=1}^{p} \varepsilon_o \ln(f_o) \tag{8}
\]

where \(o\) is the enrichment factor identifier, and \(p\) is total number of fractionation processes.

We may represent the isotope source mixing and fraction-
ation processes dynamically by discretizing the system spatially and temporally. Merging equations 7 and 8 and assuming constant enrichment factors through time and space, we can use the following finite difference approximation for the stable isotope mass balance:

$$\delta^M X_{B,i} = \sum_{i=1}^{k} \delta^M X_{i,i} / (W_{i,i}) - \sum_{o=1}^{p} \delta^M n_{i,o}$$

(9)

where $i$ is the timestep identifier, and $j$ is the reach identifier. In this definition, the mass of an element remaining in a stream reach from a previous timestep is considered a source and is accounted for in the first summation term.

**Oversight of C, N, and P Stable Isotopes in Fluvial Systems**

Isotope signatures have been widely used by environmental and water resource engineers as well as aquatic biogeochemists to study C, N, and P dynamics in streams and rivers (table 1). This section describes (1) the pools of C, N, and P species, (2) the isotope signatures used to study C, N, and P dynamics in streams, and (3) some of the applications for which isotopes have been commonly used.

**Carbon**

Primary forms of C in fluvial ecosystems include dissolved organic C (DOC), particulate organic C (POC), and dissolved inorganic C (DIC) in the form of dissolved carbonates (Hope et al., 1994). Briefly, DIC occurs as CO$_3^{2-}$, HCO$_3^-$, H$_2$CO$_3$, and dissolved CO$_2$, collectively forming the carbonate system. POC and DOC are C from organic compounds, including terrestrial leaf litter and detritus, autochthonous biomass, and biota. POC is distinguished from DOC by size classification, i.e., the solid matter that is retained on a 0.45 µm filter. For the purposes of this article, POC is further classified as fine POC (silt and clay sized particles, or $d < 53$ µm) and coarse POC (sand, cobble, or gravel sized particles, or $d > 53$ µm). DOC is primarily composed of fulvic and humic acids leached from upland soils and benthic organic matter.

Carbon exists in three isotopic forms, with $^{12}$C and $^{13}$C as stable isotopes and $^{14}$C as the radioactive isotope; only the stable forms are considered here. Carbon isotopic signatures are readily measured for all forms using well-accepted methods and are reported as the relative abundance of $^{13}$C to $^{12}$C for a sample as:

$$\delta^{13}C_{\text{sample}} = \left( \frac{\text{C}^{13}C_{\text{sample}} - \text{C}^{13}C_{\text{VPDB}}}{\text{C}^{13}C_{\text{VPDB}}} \right) \times 1000$$

(10)

where VPDB is the reference standard Vienna Pee Dee Belemnite.

Well accepted methods exist to measure $\delta^{13}$C of all three pools: $\delta^{13}$CDIC has been used as a tracer of C pathways, biotic uptake and regeneration, and atmospheric exchange rates (e.g., Doctor et al., 2008; Throckmorton et al., 2015); $\delta^{13}$CDOC has been used in a wide variety of applications, including quantitative apportionment of allochthonous versus autochthonous organic matter (Grey et al., 2001; Zah et al., 2001; Kritzberg et al., 2004; Doi, 2009; Lau et al., 2009), provision of information on trophic linkages (Rosenfeld and Roff, 1992; Zah et al., 2001; Doi, 2009; Lau et al., 2009), and characterization of nutrient sources and terrestrial inputs (Thornton and McManus, 1994; Palmer et al., 2001; Hood et al., 2005); $\delta^{13}$CPOC has commonly been used as a fingerprint for sediment source apportionment (Papanicolaou et al., 2003; Fox and Papanicolaou, 2007; Fox, 2009; Jacinthe et al., 2009; Imberger et al., 2014), as a metric to partition terrestrial versus allochthonous organic matter contributions in suspended loads (e.g., Kendall et al., 2001), and as a metric to provide insight into organic matter quality (Ford et al., 2015a; Fox and Ford, 2016; Lu et al., 2016).

**Nitrogen**

Prevailing pools of N in fluvial ecosystems include dissolved organic N (DON), dissolved inorganic N (DIN), and particulate organic N (PON). The distinctions between DOC and POC also apply to DON and PON. Regarding DIN, nitrate (NO$_3^-$) and ammonium (NH$_4^+$) are of the largest pools; however, nitrite (NO$_2^-$) may often also exist in measurable quantities in the water column. Nevertheless, NO$_2^-$ is an intermediate step in the nitrification process and, in general, is rapidly converted to nitrate (Kendall and Caldwell, 1998). Nitrogen has two stable isotopes ($^{15}$N and $^{14}$N); hence, stable isotope signatures reflect the relative abundance of $^{15}$N/$^{14}$N as:

<p>| Table 1. Measurable C, N, and O isotope signatures and their relevance to significant C, N, and P phases in stream and riverine environments. |</p>
<table>
<thead>
<tr>
<th>Pool</th>
<th>C</th>
<th>N</th>
<th>O</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbon</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dissolved inorganic carbon (H$_2$CO$_3$, HCO$_3^-$, and CO$_2$)</td>
<td>X</td>
<td>-</td>
<td>-</td>
<td>Doctor et al., 2008; Gammons et al., 2011; Rounick et al., 1982</td>
</tr>
<tr>
<td>Particulate organic carbon (POC)</td>
<td>X</td>
<td>-</td>
<td>-</td>
<td>Zah et al., 2001; Kendall et al., 2001; Kao and Liu, 2000</td>
</tr>
<tr>
<td>Dissolved organic carbon (DOC)</td>
<td>X</td>
<td>-</td>
<td>-</td>
<td>Palmer et al., 2001; Schiff et al., 1990; Raymond et al., 2007</td>
</tr>
<tr>
<td>Nitrogen</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nitrate (NO$_3^-$) and nitrite (NO$_2^-$)</td>
<td>-</td>
<td>X</td>
<td>X</td>
<td>Fukada et al., 2003; Pardo et al., 2004; Chang et al., 2002; Kaown et al., 2009</td>
</tr>
<tr>
<td>Ammonium (NH$_4^+$)</td>
<td>-</td>
<td>X</td>
<td>-</td>
<td>Webster and Heymsfield, 2003; Peterson et al., 2001; Ashkenas et al., 2004</td>
</tr>
<tr>
<td>Particulate organic nitrogen (PON)</td>
<td>-</td>
<td>X</td>
<td>-</td>
<td>Kendall et al., 2001; Angradi, 1994; Sarà et al., 2004</td>
</tr>
<tr>
<td>Phosphorus</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dissolved reactive phosphate (PO$_4^{3-}$)</td>
<td>-</td>
<td>-</td>
<td>X</td>
<td>Young et al., 2009; Elsburry et al., 2009; Davies et al., 2014</td>
</tr>
<tr>
<td>Soil and sediment extractable phosphate</td>
<td>-</td>
<td>-</td>
<td>X</td>
<td>Tamburini et al., 2012, 2014; Pistocchi et al., 2017</td>
</tr>
</tbody>
</table>
In this section, we highlight the efficacy of isotope measurements to reflect water quality processes for C, N, and P cycles. Figures 2 through 4 highlight the biotic (figs. 2a, 3a, 4a) and abiotic (figs. 2b, 3b, 4b) processes that impact stream C (Ford and Fox, 2015), N (Peterson et al., 2001; Birgand et
al., 2007; Ford et al., 2017), and P (Withers and Jarvie, 2008). We recognize that processes are often a mixture of biological, chemical, and physical mechanisms (e.g., biochemical reduction of nitrate to dinitrogen gas); therefore, for the purposes of this study, we make the distinction between biotic (biological and biochemical) and abiotic (nonbiological chemical and physical) processes. We highlight the impacts on atmospheric, water, biota, and sediment pools by showing the isotope fractionations and flux contributions to and from each pool.

**CARBON**

Biotic uptake of autochthonous biomass and mineralization of organic matter by endogenous and heterotrophic respiration are the primary biotic mechanisms impacting fluvial organic C cycling (fig. 2a; Ford and Fox, 2014, 2017; Hotchkiss and Hall, 2015). Stabilization is a process in which DIC from the streamwater pool is assimilated in autochthonous biomass, and then more complex organic C compounds are decomposed to fine sediments that have slower rates of decomposition, i.e., compounds that are more recalcitrant to biotic mineralization (Lane et al., 2013). Autochthonous biota, including benthic algae, macrophytes, and phytoplankton, fix dissolved inorganic C into particulate organic C during photosynthesis. Regarding uptake, the C isotopic signature of stabilized autochthonous organic matter is typically low in $^{13}$C relative to allochthonous matter due to $^{13}$C of DIC that is depleted relative to atmospheric CO$_2$ and has a high isotope fractionation value ($e$ between 15‰ and 25‰) (Sharp, 2007; Tobias and Böhlke, 2011; Ford and Fox, 2015). Sediment decomposition and mineralization of organic C result in a loss from the sediment or biota pool that is added to the dissolved inorganic pool and may occur in either aerobic or anaerobic conditions. Sediment C regener-

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**Figure 2.** Depiction of (a) biotic and (b) abiotic processes impacting dissolved, biotic, and sediment C isotope pools. Where applicable, processes include a range of typical fractionation factors observed in the literature. Mass balance Rayleigh-like equations (extending eq. 9) are shown for the environmentally relevant pools often considered in water quality models.
ation through oxidation of organic matter to CO₂ imparts a small fractionation compared to the autochthonous fractionation on the DIC pool (ε < 2‰) (Jacinthe et al., 2009; Ford and Fox, 2015). Degradation of organic matter to methane under anaerobic conditions may be important in landscapes such as peat bogs, resulting in fractionations of 5‰ to 10‰ (Galand et al., 2010).

Prominent abiotic processes impacting the fluvial C cycle include CO₂ flux across the air-water interface, mineral precipitation and dissolution, and hydrodynamic alterations to benthic sediment and biota pools (fig. 2b). CO₂ often evades the stream channel and acts as a source to the atmosphere due to the high rates of mineralization in soil water and benthic sediments that lead to excess partial pressures of CO₂ in stream water. Both equilibrium (ε = 1‰) and kinetic evasion (ε = 2‰) fractionations result from DIC exchange with the atmosphere. Precipitation of dissolved inorganic C is a prominent potential sink for DIC and is balanced by mineral processes. Results from Tobias and Böhlke (2011) highlighted carbonate precipitation as an equally important sink to primary production in a low-order stream in an agroecosystem. While algal uptake exerts a strong kinetic isotopic fractionation on the dissolved inorganic pool, precipitation-dissolution imparts a small equilibrium fractionation (ε < 1‰) (Mook, 2006; Tobias and Böhlke, 2011). The erosion-deposition dynamics of sediment are well documented to impact benthic C isotopic signatures, which reflect sediment C quantity and quality (Ford et al., 2015a). Newly deposited sediments are mixed with existing sediments through turbulent advection of the overlying streamwater into the benthos (Russo and Fox, 2012; Ford and Fox, 2014). The level of mixing is scale-dependent, but in low to mid-order streams with high prominence of fine-cohesive sediments, sediment within the surficial fine-grained laminae of the streamed surface is typically well-mixed (Droppo et al., 2000). Fluvial sloughing of algal biomass has the potential to impact sediment isotope compositions, especially in low DIC systems where fractionations due to autochthonous growth in response to biotic population disequilibrium have a larger footprint on the DIC isotope pool (Ford and Fox, 2015). The dynamics for site-specific conditions are discussed further in the “Review of Stable Isotopes in Water Quality Modeling” section.

NITROGEN

Practically all N fractionation takes place through biologically mediated pathways, including the aforementioned autochthonous growth, heterotrophic and endogenous respiration (mineralization), nitrification, and denitrification (fig. 3a; Sharp 2007). Regarding autotrophic assimilation of N species, biotic algal uptake of N imparts a fractionation on its DIN source of 6‰ to 13‰ for NO₃ (Neeboda et al., 2003; Kendall et al., 2007) and 0‰ to 27‰ for NH₄ (Fogel and Cifuentes, 1993; Kendall et al., 2007). However, fractionations for ammonium are likely small (ε < 4‰) in most aquatic systems with low ammonium concentrations (Fogel and Cifuentes, 1993; Kendall et al., 2007). Regarding N remineralization of organic N to ammonium fractionations are typically negligible, with ε ±1‰ (Kendall et al., 2007).

The benefit of N isotopes to reflect in-stream biotic cycling is recognized from the high fractionations reported for N and O isotopes during dissolved inorganic transformation in nitrification and denitrification processes (Kendall et al., 2007). Nitrification is the two-step aerobic oxidation of ammonium (NH₄⁺) to nitrite (NO₂⁻) and then to nitrate (NO₃⁻). As previously mentioned, δ¹⁵N and δ¹⁸O of nitrate are measured using the dual-isotope approach. With regard to δ¹⁵N, researchers have found that the first step (NH₄⁺ to NO₂⁻) is often the rate-determining step in ammonium-rich systems and occurs very slowly, resulting in large fractionations on the ammonium N pool with ε values ranging from 14‰ to 38‰ (Mariotti et al., 1981; Casciotti et al., 2003; Kendall et al., 2007). In ammonium-limited systems, the fractionation of the N isotope is relatively small. Further, the second step (NO₂⁻ to NO₃⁻) is rapid and typically does not result in a net fractionation. With regard to δ¹⁸O of nitrate, the oxygen isotope composition will generally reflect a mixture of the oxygen isotope signatures of water and dissolved oxygen; however, the level of fractionation is not well understood (Kendall et al., 2007). For denitrification, or the anaerobic reduction of nitrate to N-based gaseous byproducts, enrichment factors of δ¹⁵N range from 1‰ to 18‰ depending on where denitrification occurs (i.e., water column, benthos, riparian zone) (Brandes and Devol, 2002; Sebilo et al., 2003; Lehmann et al., 2004; Sigman et al., 2005; Kendall et al., 2007).

Abiotic processes controlling N cycling and isotope signatures in-stream not only include the aforementioned hydrodynamic and hydraulic factors (analogous to C) but also chemi-physical sorption of DIN to benthic sediments. Abiotic adsorption of ammonium is widely recognized as a transient N sink, with reported apparent equilibrium fractionations ranging from 1‰ to 11‰ (Delwiche and Steyn, 1970; Karamanos and Rennie, 1978; Bernot and Dodds, 2005; Böhlke et al., 2006). Abiotic adsorption of nitrate in streams is not currently part of the perceptual model (Peterson et al., 2001; Birgand et al., 2007; Ford and Fox, 2017); however, evidence exists for nitrate adsorption to variably charged sesquioxides in benthic sediments, analogous to processes reported in soils (Ford et al., 2015b). Given the limited understanding of the magnitude and significance of this flux, the isotopic fractionation is not well understood; therefore, future work is needed to test the significance of the sorption mechanism and identify potential ranges of isotope fractionation under differing sediment and streamwater chemistry.

PHOSPHORUS

Regarding biotic processes, the primary mechanism leading to changes in δ¹⁸OPO₄ is associated with microbial mediated recycling of orthophosphate. Enzymatic breaking of the P-O bond during microbial cycling of orthophosphate drives the phosphate signature toward a temperature-dependent equilibrium fractionation value, with δ¹⁸O following regeneration to the water column (Young et al., 2009; Davies et al., 2014). Therefore, in areas where microbial P cycling is rapid (e.g., benthic biofilms), the δ¹⁸OPO₄ of DRP reflects a mixture of its source signature and rates of microbial P regeneration. Regarding sediment and biota P, we did not find...
information on fractionation associated with uptake or mineralization on the sediment or biota pools; however, techniques for measuring sediment PO₄ signatures are relatively new and do not explicitly distinguish between organic and inorganic P sources (Pistocchi et al., 2017). Abiotic processes, including erosion-deposition, precipitation-dissolution, and sorption-desorption, are more significant for fluvial P cycling than for C and N, which stems from the high sorption capacity of cohesive soils. Soil P may be highly stratified in adsorbed inorganic P; hence, erosion-deposition dynamics are important in fluvial ecosystems (Jarvie et al., 2014). Authigenic production of orthophosphate occurs through co-precipitation with calcite, precipitation with iron and hydroxide in oxic pore waters, and precipitation as vivianite under anaerobic, eutrophic conditions (Withers and Jarvie, 2008, and references within). The mineral growth process is rapid, and fractionation effects between mineral and dissolved phosphate are low; hence, the δ¹⁸O signature of authigenic P commonly reflects its phosphate source, and vice versa for dissolution (Joshi et al., 2015). P uptake through sorption is widely acknowledged within streams and may be a significantly higher sink of P as compared with algal assimilation (Withers and Jarvie, 2008). Further, P desorption may become a prominent source of legacy P under specific redox conditions in agroecosystems, which tend to retain rich stores of P in benthic sediments (Jarvie et al., 2014; Joshi et al., 2015; Baker et al., 2017). Oxygen isotope signatures of phosphate are not subjected to equilibrium fractionations under abiotic processes; hence, phosphate adsorbed to sediment surfaces should reflect its inorganic P source and its regenerated product (Davies et al., 2014).
REVIEW OF STABLE ISOTOPES IN WATER QUALITY MODELING

Coupling of stable isotopes within water quality models is in its infancy within the water resources community, and there are likely many permutations of coupling that may be performed in future research and model development. Nevertheless, based on our review of previous research as well as research advancements in recent years, we highlight three common themes defined earlier in this article, i.e., that stable isotopes are coupled with water quality models to (1) improve data inputs associated with boundary conditions of the models, (2) constrain model parameterization associated with equifinality, and (3) elucidate improvements needed within the model structure. Table 2 highlights the relevant watershed water quality modeling studies reported in the literature. Specifically, we provide summaries of how each study addresses one or more of the themes. As will be shown, at least one of the uncertainty-associated components was overcome when the researchers coupled stable isotopes within their watershed water quality modeling. To our knowledge, these studies represent an exhaustive list of water quality modeling applications that incorporate stable isotopes of C and N. In this manner, current use of stable isotopes in water quality modeling has highlighted their utility for improving reliability and reducing equifinality in hydrologic and water quality model simulations. We have separated this section into C and N isotope applications because no applications for P have been performed to date.

CARBON

Dissolved Carbon

Tobias and Böhlke (2011) quantified the relative amounts of biological and geochemical controls on DIC cycling and
flux within a 1 km first-order agricultural stream reach using daily $\delta^{13}C$ of DIC and $\delta^{18}O$ of $O_2$ applied to a finite-differencing mass balance model. Their use of $\delta^{18}O_{O_2}$ was to aid in constraining interpretations of the $\delta^{13}C_{DIC}$ and DIC measurements. Their logic was that when chemical and isotope modeling is applied in combination with daily observations, there would be an improvement in the overall mechanistic understanding of the diel fluctuations and environmental factors that influence DIC fate and transport. The model output contrasted with the collected data in that the model $\delta^{13}C_{DIC}$ estimates were too high and did not reproduce cation cycles. The values of input parameters needed to reproduce accurate output values were unrealistically high, and the insensitivity of the $\delta^{13}C_{DIC}$ variation to carbonate reaction suggested that the indicator acted as a poor indicator of diel processes except for photosynthesis rates in highly productive systems.

Stable isotope signatures of DIC ($\delta^{13}C_{DIC}$) have recently been implemented in marine and estuarine environments to reduce equifinality. Van Engeland et al. (2012) investigated model uncertainty reduction through inclusion of $\delta^{13}C_{DIC}$ results for injected mesocosm experiments into a marine nitro-
elucidate the impact of reclamation practices on sediment
bank erosion parameters. The source uniqueness and time-
iment delivery ratio for reclaimed mining soils, and stream-
ecosystems with pronounced sediment storage), Ford et al.

approach to estimate time-varying sediment source contribu-
tions within the parametric modeling framework. The
additions of the $\delta^{13}$C tracers constrained uncertainty of
biogeochemical transformations of the model-predicted
rates and fluxes associated with C mass balance. Evaluation
of the NPZD model with and without isotope calibration data
was performed. The authors found that calibrations using
solely concentration data exhibited higher standard deviations
of uncertain parameters, strong correlations between
fitted parameters (suggesting parameter value dependence),
and inaccurate estimates of zooplankton grazing and detritus
sinking rates as compared with multi-objective calibration
with concentration and stable isotope response variables.
Quantitatively, the authors provide evidence of this through a
higher multicollinearity index for the reduced (no isotope)
dataset relative to the full model evaluation dataset (values
of 3.43 and 1.64, respectively).

Adiyanti et al. (2016) collected high spatial resolution
data in a subtropical estuary over five sampling campaigns
and analyzed samples for dissolved inorganic, dissolved or-
nic, and POC isotope signatures. The authors used a mixed
1-D, 3-D modeling approach that coupled hydrodynamics
with C biogeochemistry for the estuary and used DIC and
DOC isotope and concentration measures as model response
variables. The authors highlight that the addition of the iso-
tope response variables allowed better constraint for bioge-
chemical process parameters as compared to using Markov
chain Monte Carlo optimization without the isotopes. Pa-
rameter space constraint was observed because of sensitive
fractionation effects on the isotope response variables that
led to rejection of implausible model outputs. The authors
highlight the utility of the approach for advancing C budget-
ing by using the model to describe spatial variability of the
trophic state within the estuary.

**Particulate Carbon**

Sediment particulate C isotope signatures ($\delta^{13}$C$_{PPOC}$) have
been used to improve model calibration and parameteriza-
tion for conservative and non-conservative tracer behavior.
Fox and Martin (2014) used stable sediment C and N iso-
topes of sediment to separate forest, reclaimed mine, and
streambank sources and to highlight the efficacy of coupling
stable isotope fingerprinting with deterministic sediment
yield modeling in mixed-use landscapes. Carbon isotopes
were used in conjunction with N isotopes as a dual-tracer
approach to estimate time-varying sediment source contribu-
tions within the watersheds, subsequently acting as an addi-
tional response variable in sediment yield model evaluation.
The authors used the added isotope-based response variable
to calibrate the sediment transport capacity coefficient, sed-
iment delivery ratio for reclaimed mining soils, and stream-
bank erosion parameters. The source uniqueness and time-
varying nature of the forest source allowed the authors to
elucidate the impact of reclamation practices on sediment
yield with their model. The authors’ study was found to be
applicable for steep-gradient watersheds with relatively con-
servative tracers (in-stream) due to low residence time.

For non-conservative systems (e.g., low-gradient agro-
cosystems with pronounced sediment storage), Ford et al.
(2015a) used stable C isotopes of transported sediments
($\delta^{13}$C$_{POC}$) to constrain a reach-scale C fate and transport
model that considered benthic autochthonous and terrestrial
C sources. A deterministic C mass balance model for benthic
sediment, algae, and DIC pools (ISOFLOC) was coupled to
a sediment storage and transport model to assess the impact
of algae on the fluvial C budget. Stable isotope mass bal-
cances were simulated for each C pool, and eight years of am-
bient concentrations of fine POC and C isotope data were
used to evaluate the model. The isotope response variable
was found to be highly sensitive to the critical shear stress
of algae and the algal POC source (DIC) and its time-varying
isotope signature. As a result, calibration using the isotope
submodel reduced uncertainty of sloughed algal fluxes by
80%. These highly dependent relationships between bioge-
chemical processes, physical processes, and the ability of
stable isotopes to reflect these processes highlight the im-
portance of ambient isotope response variables to account
for non-conservative contaminant behavior in complex flu-
vial systems.

Sediment stable isotopes of C have also been effective-
ly used to establish boundary conditions of sediment C sources.
Husic et al. (2017a, 2017b) applied sediment C fingerprint-
ing at the upstream monitoring station of a phreatic karst
duinct in central Kentucky. The authors separated sediment
C fractions from surface stream autochthonous detritus, la-
bile terrestrial soil C, and relatively recalcitrant soil C
sources. Given the variability of biological turnover rates of
these C pools and the subsequent implications for water
quality in perennial springs that serve as drinking water sup-
plies, the authors highlight the potential utility of the ap-
proach. Further, the authors discuss the enhanced adoption
of sediment fingerprinting within the hydrologic and water
quality community, highlighting the natural linkage to the
water quality modeling community.

**NITROGEN**

**Dissolved Nitrogen**

Xue et al. (2014) used unmixed isotope inputs for a model
that was unlike the other studies reviewed here because the
model was not a physically based mechanistic model; how-
ever, it included isotopes as inputs for a decision tree model.
The study used two years of monthly $\delta^{15}$N$_{NO3}$ and $\delta^{18}$O$_{NO3}$
data from a multitude of sampling locations as inputs for a
mixing model (SIAR) to determine nitrate source apportion-
ment. The study also assessed the effectiveness of isotopic
data as input in a decision tree model that used physicochem-
ical data. In decision tree models, a critical component of
their construction is the split selection, which involves
choosing the best option to proceed with in the model. The
decision tree model was simulated with and without iso-
otope data, and the isotope data did not improve the performance
of the decision tree model. The authors speculated that this
could have been due to the complex land use of the study site
that resulted in scattered nitrate isotope values. However, the
authors posited that an opportunity exists to use $\delta^{15}$N$_{NO3}$
and $\delta^{18}$O$_{NO3}$ data to cultivate a dependable nitrate polluting ac-
tivity classification.

Sebestyen et al. (2014) used a dual-isotope method of ni-
trate ($\delta^{15}$N$_{NO3}$ and $\delta^{18}$O$_{NO3}$) to study N cycling and source contributions during autumn in a forested stream ecosystem. Their study combined the interactions among biogeochemical processes, N source allocation, and flow paths to investigate how these components affect N variation. Modeling consisted of streamwater and solute mass balances and stable isotope mass balances with Rayleigh fractions. Inclusion of the stable isotopes improved the constraint of stream biochemical reactions and source contributions. Model estimates suggested that in-stream transformations retained 72% of the nitrate entering the stream channel. Further, through the isotope mass balance approach, the study found higher inputs of unprocessed atmospheric nitrate than what is commonly acknowledged for non-snowmelt periods in forested landscapes.

**Particulate Nitrogen**

Fox et al. (2010) focused on modeling sediment transport and sediment source apportionment using N stable isotopes ($\delta^{15}$N$_{PN}$). Their study used N stable isotopes in sediment to aid in differentiating sediment sources and modeling sediment transport because of the effectiveness of $\delta^{15}$N in separating sediment sources in watersheds that contain vegetation with similar photosynthetic pathways. Nevertheless, the authors found that $\delta^{15}$N$_{PN}$ (and sediment N) varied substantially due to physical and biogeochemical processes impacting the transient storage zones in sediments. While their study did not examine robust uncertainty analysis for biogeochemical parameters, it highlighted the efficacy of the stable isotopes to help establish inputs from upland and bank sediment sources and highlighted the importance of the fate of N isotope tracers in productive agroecosystems.

Building on Fox et al. (2010) and Ford and Fox (2015), Ford et al. (2017) developed a reach-scale N model to simulate in-stream N fate and transport in low-gradient agroecosystems. The N model that includes stable N isotope subroutines is known as TRANSFER (Technology for Removable Annual Nitrogen in Streams For Ecosystem Restoration). The authors coupled N mass balances for dissolved and particulate phases to the previously developed ISOFLOC model (see the preceding “Particulate Carbon” section) and included an N stable isotope mass balance equation for each of the elemental mass balances. During model evaluation of a case study, the authors found that fine PN isotope signatures ($\delta^{15}$N$_{PN}$) were sensitive to sediment sources and non-conservative in-stream sediment N generation from autochthonous material and organic N degradation (and hence isotopic signatures of DIN). As a result, the authors reduced equifinality of the estimates of transient DIN removal via algal sloughing and permanent removal via denitrification. Their results showed that reduction of uncertainty by combining sediment elemental and isotope calibration parameters to DIN concentrations resulted in a 67% reduction from the original parameter solution space for downstream DIN flux estimates. This is compared to a 44% reduction from the original parameter solution space when calibrating with DIN concentrations alone. The reduced equifinality elucidated the significance of the transient DIN store and the potential for overestimation of denitrification during sensitive timeframes (e.g., late summer/early fall), when sloughed algal biomass may potentially fuel harmful and nuisance algal blooms (HNABs) downstream. In addition, the case study revealed disagreement between measured and modeled results for the isotope response variable during winter/spring, potentially highlighting limitations in existing perceptual models for in-stream N fate and transport, such as the lack of inclusion of abiotic mobilization/demobilization.

Hong et al. (2014) used $\delta^{15}$N signatures in dolphins to determine methyl mercury (CH$_3$Hg) dietary exposure in Sarasota Bay. The N stable isotope aided in identifying where mercury loading was present and how it was being discharged into the bay system. When one bioconcentration factor in lower trophic level organisms and one biomagnification rate were coupled with a predetermined $\delta^{15}$N, the mercury distributions in the ecosystem were successfully reproduced. This relationship enabled modeling of the fate, transport, and bioaccumulation of monomethyl mercury within the waterbody.

**RECOMMENDATIONS FOR ISOTOPES IN WATER QUALITY MODELS**

Synthesis of the studies in table 2 points to the ability of stable isotopes to constrain uncertainty of hydrologic and water quality models, improve perceptual understanding of in-stream contaminant fate, and establish boundary conditions for in-stream models. Consistent with the themes recognized in the literature review, we provide some recommendations and precautions for water quality modelers to integrate stable isotopes into new and existing models.

**RECOMMENDATION 1:**

*Use Isotopes of Sediments Because They Integrate Processes, Reflect Source Contributions, and Are Inexpensive to Measure*

We perceive high utility in the integration of sediment stable isotopes into in-stream routines in water quality modeling frameworks, given the following factors: the integrative capacity of benthic sediments, the abundance of sediment stable isotope data from watershed sediment source apportionment, the utility of stable isotopes to improve water quality modeling structure and uncertainty reduction, and the now inexpensive costs associated with stable isotope analyses of solids. Sediment fingerprinting has been a popular method for sediment source apportionment over the past 20 years (Collins et al., 1998; Fox and Papanicolaou, 2007; Davis and Fox, 2009). Sediment source apportionment using stable C and N isotopes is limited by the fate of the organic matter in the system (Davis and Fox, 2009; Koiter et al., 2013). In part, this non-conservative behavior reflects processes such as the stabilization of algal biomass through algal decomposition to fine sediment and integration into the benthos, and the sorption-desorption of N phases onto fine sediment aggregates (see the “Processes Impacting Stream Isotope Compositions” section). Therefore, sediment fingerprints reflect not only the upland organic matter and sediment sources but also the rates of processes and the dissolved inorganic nutrient species. For this reason, we foresee high utility in integrating the widespread measurements of C and
N isotopes that have been collected across a broad range of landscapes to test and improve water quality models. Further, we recommend integration of sediment stable isotope measurements into routine water quality monitoring efforts because the sampling equipment is easy to build using standard household items, the data reflect temporal and spatially integrated measures of in-stream transported sediment C and N signatures over the course of an event (Phillips et al., 2000), and the data are relatively inexpensive to process and analyze in the laboratory. We caution that sediment stable isotope signatures should be used as a supplement, not as a replacement, for concentration response variables in water quality modeling. Nevertheless, we foresee that the low cost and relatively low processing time for analysis makes this added response variable a plausible supplementary data collection effort in watershed-based monitoring and modeling programs.

RECOMMENDATION 2: Modelers Should Use Multi-Objective Calibration when Using Isotopes

We highlight the importance of using isotope response variables in multi-objective calibration frameworks to reduce issues with model equifinality. Multi-objective calibration refers to the process of using a set number of weighted numerical metrics that target specific aspects of goodness-of-fit between model results and measured data (Van Griensven and Bauwens, 2003; Rode et al., 2007; Ford and Fox, 2015; Haas et al., 2016). A thrifty approach that has greatly extended the utility of existing concentration data is to use time-varying, multi-objective calibration, whereby calibration statistics are calculated for specific periods to target calibrating parameters when they have heightened sensitivity, i.e., baseflow versus event flow, seasonal performance, and rising versus falling limb of the chemograph (e.g., Haas et al., 2016). Such sensitivities may be identified using time-varying global sensitivity analysis approaches (Reusser et al., 2011; Muleta, 2012; Wang et al., 2013; Hermann et al., 2013; Ford and Fox, 2015). Nevertheless, issues persist with using concentration-based measures because they may be insensitive to nutrient residence times, i.e., transient storage and discriminating rates of in-stream processes (e.g., Jarvis et al., 2014; Ford and Fox, 2017). In this light, integration of isotope response variables within a multi-objective framework may be highly valuable, especially given the widely recognized utility to reduce equifinality, as discussed in the “Review of Stable Isotopes in Water Quality Modeling” section.

RECOMMENDATION 3: Water Quality Modelers Should Work Collaboratively with Isotope Geochemists

With continued advancements in isotope measurement techniques and technology, watershed modelers need to work closely with isotope geochemists to integrate stable isotope measurements into water quality modeling frameworks. From a management perspective, engineers need high-resolution data, especially during storm fluxes, to accurately characterize loadings and source contributions of nutrient fluxes at the watershed scale. Current measurement techniques for grab sample analysis for isotopic measurements of dissolved nutrients are rather expensive, labor intensive, and limit the economic feasibility of high-resolution measurements. Nevertheless, we have seen a rise in in situ technologies, and researchers now have the capability to obtain high-resolution measurements of $\delta^{18}$O$_{H_{2}O}$ and $\delta^{13}$C$_{DIC}$. As these technologies continue to extend to other nutrients (e.g., nitrate) and become more affordable, it will be important for watershed modelers to understand the limitations and applicability of the high-resolution data streams, which will require close collaboration with isotope geochemists. We foresee high utility for water quality model frameworks that use high-resolution isotope sensing to inform practical watershed management decisions.

Further, the cutting-edge work that has been conducted on $\delta^{18}$O$_{PO_{4}}$ DRP and PIP over the past decade and the lack of ambient tracers of P source fate and transport (Jarvie et al., 2014; Williams et al., 2016) suggest a need to assess the efficacy of $\delta^{18}$O$_{PO_{4}}$ in water quality modeling frameworks. Several challenges exist that will require interdisciplinary collaboration to recognize the full potential of the oxygen isotope signature of phosphate as a tool for informing water quality models. Regarding dissolved inorganic phosphate, a current barrier is the large sample volume needed to precipitate an adequate mass of Ag$_3$PO$_4$ for isotope analysis, given the low ambient DRP concentrations (McLaughlin et al., 2004; Young et al., 2009; Pistocchi et al., 2017). In addition, based on existing datasets, it is not clear that the approach may robustly distinguish between nonpoint pollution sources, which has led to suggestions of database expansion of P source characterization in freshwater ecosystems (Young et al., 2009; Davies et al., 2014). A third limitation is that most existing methods for soil and sediment extraction are not pool specific (see Haney et al., 2013, for the exception) and typically reflect adsorbed phosphate, dissolution of phosphate-bearing precipitates, and mineralized organic matter (Tamburini et al., 2014; Davies et al., 2014; Pistocchi et al., 2017). Despite these limitations, isotope measurement provides valuable information on biological processing of P that is not otherwise measurable with existing methods (Pistocchi et al., 2017). We foresee $\delta^{18}$O$_{PO_{4}}$ to hold great promise for numerical model advancement, and we foresee that concurrent advancement of water quality modeling technology with analytical techniques may lead to more robust management of P in fluvial landscapes.

CONCLUSIONS

While model uncertainty continues to be a major challenge facing scientists and engineers, stable isotopes are promising tools for improving in-stream nutrient fate and transport routines in water quality models. This is an exciting time for water quality modelers, as new data streams, such as stable isotopes, offer the promise of constraining model uncertainty. This review highlighted the ability of stable isotopes to (1) improve estimates of boundary conditions, (2) reduce model equifinality, and (3) elucidate model improvement needs by identifying deficiencies in perceptual or numerical model frameworks. As a final note, regarding
the reduction of model equifinality, we highlight the importance for modelers to provide quantitative evidence of uncertainty reduction in future applications. This quantitative evidence is often missing in recent studies, given the emphasis of the studies on establishing new methodologies and showing their efficacy. This effort should be commended; nevertheless, future studies might report quantitative evidence to help researchers understand when the extra data stream and modeling effort are most useful and when they are not. We foresee that such quantitative evidence will also provide modelers with a metric to inform cost-benefit analyses associated with model data collection decisions.

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REFERENCES


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