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Interactive comment

Interactive comment on "Reproducible determination of dissolved organic matter photosensitivity" by Alec W. Armstrong et al.

Patrick Neale (Referee)

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General Comments

Measurements of spectral fluoresence have become a standard part of any study of the dynamics and distribution of dissolved organic matter in both freshwater and marine environments. Methods for deriving standardized metrics from fluorescence data are well established, and most studies of DOM dynamics acquire detailed sets of excitation-emission matrices and analyze them using parallel factor analysis (PARAFAC). In the present contribution, Armstrong, et al., argue that the kinetics of fluorescence photosensitivity are also useful properties to characterize DOM, in particular, the magnitude and rate constants of biexponential kinetics. However, several important considerations

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need to be observed if these photosensitivity metrics are to be compared across studies. These are shown in the context of a specialized continuous loop exposure/assay system that the authors have devised, but several are applicable to any photosensitivity study. One is the use of a reference material, SRNOM, to both test the derivation of PARAFAC components and standardize the kinetics for a given exposure set up. Another informative test is the reproducibility of the results in repeated runs including those by different observers, which helps to establish the uncertainty in the estimated kinetic parameters.

I quite agree with these recommendations which are straightforward to implement. For others, it is not clear that they will always contribute to reproducible, or more importantly, interpretable, results. One is the preparation of solid phase extracts of DOM vs using fresh filtrates for the measurements. While extracts are inherently more stable than raw filtrates, the authors demonstrate differences in the photosensitivity kinetics of extracts vs the original filtrates, the reasons for which are not well understood. The authors advise preparing extracts in some situations but not others, this leaves open the question of comparability. The authors also advise using a standard optical density in the sample, yet also show there is a concentration dependence to the kinetics, even for an optically thin exposure conditions. Again, a decision will have to made of whats more important, knowing how the DOM behaves at its natural concentration or being able to compare it to other sources.

The authors also advocate expressing kinetic results as a function of cumulative photon exposure as opposed to time, this will adjust for differences in exposure regime between different studies. In principle, this is a good idea. Any photosensitivity study should include a detailed documentation of the optical setup and exposure measurements such that the results can be expressed either as a function of time or cumulative exposure. Studies do range in how thoroughly exposure conditions are documented. For these kinetic studies, exposure has been measured with a nitrite actinometer. Actinometry is very useful to quantitate the effects within a specific optical set up and is

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used effectively by the authors to monitor exposure between experiments, including those with variable working volume. However, the nitrite actinometer only measures quantum fluxes between 330 and 380 nm. For incident solar UV, this is only about 60 per cent of the spectrum (by quanta). For comparison across all photosensitivity studies, I would suggest a more general exposure metric, e.g. total UV radiation. Given the broadband 330-380 nm quantum flux, total UV can be estimated with reference to the manufacturer's stated spectral distribution. But lamp and component aging can change that (even adjusting for constant power), so the best approach is to measure the spectrum in conjunction with actinometry (admittedly, neither of these are trivial to perform).

In the end, investigators will need more information to decide whether the additional insight gained from defining photosensitivity kinetics will be worth the efforts needed to perform all these standardization steps. This report documents experimental tests that show what are sources of variability in the procedure but does not show what new understanding is actually being gained from the results (vs what can already be gained from other optical/chemical measurements typically made on DOM). As far as reproducibility, there is only one instance of repeat determination on a "real world" sample, the small wetland. Further repeat determinations are needed to demonstrate the general reproducibility of the approach. Also, some modification of how the photosensitive fractions are estimated may be needed if the results are to be used in a comparative context (see specific comments below). Despite that this is still a work in progress, it is quite appropriate to have the authors' study and recommendations in a Biogeosciences discussion article. Comments by a wide range of readers should help refine the ideas proposed and ultimately advance the field.

Specific comments:

Line 140 "Solar exposure" – This was not solar exposure, but a laboratory set up. Better (?) "Total sample exposure"

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Figure 7 and Table 1 – The captions for these describe kL and kSL with units of mol photons m⁻². Those are the units of P, the units of k should be the inverse, m² [mol photons]⁻¹.

Figure 7 and Table 1 apparently show the same data – and both state the statistics of the comparison. Questionable whether both are necessary. Figure 7 caption, however, says see Fig. 5 for data. It appears that Fig. 6 has the data.

Line 350-360 – Factors affecting storage of filtered water. Previous text only mentioned preparation of filtrates using 0.7 μ M glass fiber filters, which allows passage of a substantial bacterial fraction. However, subsequent guidelines caution against used stored material even filtered to 0.2 μ M (some bacteria can even get through those). Please clarify what filtration method was used for the stored sample, but in any case (slow) microbial degradation may be another factor resulting in changes in stored material.

Figure 10 – Caption says see Figure 6 for data, data appears to be in Fig. 9. This correction also applies to the caption for Figure A1. Figure 10 could be misleading to the reader, as a quick inspection may suggest that the spread of the multiple points for a particular sample source (and standard deviation in the table) are indications of reproducibility. Actually, only the SRNOM and three of the small wetland points are repeat determinations on the same sample. Separating the points for different sample dates would avoid this confusion, especially given that later on (section 3.3.2) the authors attribute some of the difference in the wetland results with time to possible differences in sample composition and/or previous solar exposure.

Line 456ff, section 3.3.1 Here the relative abundances of kinetic components are considered as metrics of DOM composition or other influences on photosensitivity. A problem with the comparative use of fL and fSL is that the two parameters are not independent. Assuming a good fit to the actual time course the two fractions will always sum to approximately 1, which means that any variation in one fraction will be reflected in a complementary variation in the other. Thus, changes in, e.g. fL, with sample date or

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location (as later discussed in 3.3.2) could be caused by change in the concentration of labile or semi-labile fraction, or both. If the focus is on the rate parameters, this is not so important, but if the magnitudes of the fractions are to be comparative metrics, the need to be independent of each other. For example, ft could be the actual component score, score normalized to to total fluorescence, or score normalized to DOC.

The authors offer the interpretation that the fractions correspond to "pools" of more or less reactive DOM. However, Murphy et al. (2018) interpret biexponential kinetics as possibly reflecting multiple photoreaction pathways, a fast one involving ROS and slower one related to direct photochemistry. How do the authors reconcile these differing interpretations?

Interactive comment on Biogeosciences Discuss., https://doi.org/10.5194/bg-2020-207, 2020.

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Interactive comment on "Reproducible determination of dissolved organic matter photosensitivity" by Alec W. Armstrong et al.

Anonymous Referee #2

Received and published: 10 September 2020

Overall comment: The study described in the manuscript aims to obtain a generalized method for studying light sensitivity of DOM in different water bodies. Authors suggest that this method may provide additional information on natural DOM quality. In general, I find this approach novel, however, some aspects are missing that would make me believe the method worse trying. The method seems to be very complicated and affected by numerous factors that the reader should keep in mind, but the discussion on the advantages of this method is very abstractive.

Specific comments: I have several more specific comments to the manuscript: Authors mentioned by themselves that pH changes might lead to reactions of flocculation, furthermore, pH may greatly affect EEM and absorbance signals which measured in this study, herewith I have a question, why did the Authors acidify the sample prior to ex-

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Discussion paper



tractions. And why did they use GFFs rather than 0.2 μ m membrane filters? In my understanding, the colloidal fraction of DOM that is most likely to be present in the sample after $0.7\mu m$ filtration is the most susceptible to flocculation. The authors mentioned that natural samples were affected by cold storage. How exactly the sensitivity study was performed? The authors did a comparison study for a humic standard comparing extracted and not extracted solutions and saw a difference in the quality of DOM on EEM. Since guite altered by isolation and freeze-drying standard differ between extracting and non-extracting approach, the question is how would the natural samples differ? To which extent those extracts would be representative of what is actually going on in-situ? Also, the extracts were evaporated and diluted, what method was used for that? Microwave? The Authors describe precisely in details different approaches and drawbacks for the standard humic substance but the methods are lucking the description of the experiments performed on natural water samples. For instance, the reader learns that the natural organic matter was collected in different seasons only in the section "Results and discussion" and so on. Have the authors measured the initial conditions sample in each water bodies? How did EEM differ in terms of Fmax scores? Were they different? Also, for different seasons? I would expect that the Fmax scores would indicate that the composition of DOM is different between those water bodies, therefore I wonder whether the reported method would provide greater advantages than the outcome that DOM between water bodies is different? If the question is rather on the photosensitivity, in my opinion, some kind of discussion should be present on how comparable the results will be to what may happen in situ.

Technical corrections: Fig.9 Two plots for Comp3 For EEM comp. decay there are 4 curves for shallow and deep wetlands, were those replicates, or those different seasons measurements the authors have mentioned in the last chapter?

Interactive comment on Biogeosciences Discuss., https://doi.org/10.5194/bg-2020-207, 2020.

RC1 response

We appreciate the thoughtful and thorough comments, which we believe raise three substantial points – justifying our use of a nitrite actinometer, our suggestions for improving reproducibility and comparability in photochemical degradation studies of DOM, and the novelty of our contribution in terms of its utility vs. existing approaches and our suggested interpretation of the data we generated.

We used a nitrite actinometer to illustrate the importance of controlling photon exposure and the potential limitations of modeling degradation processes as a function of time, shown distinctly in Figure 3. As the referee states, "Any photosensitivity study should include a detailed documentation of the optical setup and exposure measurements such that the results can be expressed either as a function of time or cumulative exposure. Studies do range in how thoroughly exposure conditions are documented [emphasis ours]." We contend it is often difficult to calculate cumulative exposure from reported time alone in the published literature, making comparability difficult, and show that using a nitrite actinometer to calculate and report exposure directly is one way to improve this situation. We agree that other means of calculating and reporting cumulative exposure, such as measuring radiation across the UV spectrum directly, can only address this issue when irradiation design allows for accurate photon dose quantification with a spectroradiometer (e.g. irradiating quartz vessels/spectrophotometer cells with a known pathlength, cells are isolated from each other, ensuring that light is collimated/off axis photons do not hit samples, etc). Many labs have successfully designed systems that allow for very accurate photon dose quantification using solar simulators. For example, Powers and Miller (2015, MarChem, 171) irradiated 10 cm cells vertically in a black water-cooled aluminum block that maintains cells at a set temperature and allows no transfer of light between samples. Additionally, a gray 0.5" PVC plate with holes matching the spacing/interior diameter of the spectrophotometer cells was placed over the block and positioned under various 2" diameter Schott long-bandpass cutoff filters (280 nm to 480 nm) that were evenly spaced in a tray located at the bottom of the solar simulator exposure chamber. Due to the reflective exclusion of high angles at the surface of the Schott glass filters, reflectance at the face of the quartz cell is minimized. Furthermore, the gray plate reduces the entrance diameter and absorbs any off axis photons within the "gershun-like" tube formed by the gray plate, which also removed (or greatly minimized) reflectance at the face of the quartz cells by collimating the light. However, as mentioned in our manuscript, this type of experimental design allows for a limited number of samples, and therefore a limited number of measurements. Because the system described in Powers and Miller (2015) irradiates samples in 10 cm cells, even samples with relatively low CDOM values are not optically thin. While Powers and Miller (2015) used the methods of Hu et al. (2002) to correct "photons absorbed by CDOM, $Q_a(\lambda)$ " for shelf-shading (i.e. $Q_a(\lambda)$ = $E_0(\lambda) \times S(1-\exp(-a_g(\lambda) \times L) \times t)$, where $E_0(\lambda)$ is the measured spectral irradiance entering each cell, S is the irradiated surface area, $a_g(\lambda)$ is the CDOM absorption coefficient, L is the pathlength, and t is irradiation time), it is difficult to verify that these corrections are sufficient. It is possible that there are concentration effects but this is difficult to evaluate due to changes in sample matrix when diluting samples. Furthermore, samples in such a system must also be irradiated with no headspace because bubbles are terrible for optics, leading to oxygen loss during irradiations. While this may not be a problem for short exposures, many photoproducts of interest (i.e. CO₂) require long irradiations to generate a measurable product and it is possible that rates decrease at lower oxygen concentrations (e.g. Gao and Zepp, 1998, ES&T, 32). Lastly, this system allows

for no pH control during irradiations, even though it is well known that pH can change during irradiations (Gao and Zepp, 1998) and photodegradation rates change at different pH values (Timko et al. 2015, Water Res.; Song et al. 2017, ES:P&I, 19). While some researchers have chosen to buffer samples to overcome this issue (e.g. Powers and Miller, 2015), to our knowledge it is still unknown how buffers may or may not impact photodegradation rates.

Therefore, our primary methodological objectives were to establish a reproducible design that would 1. Irradiate samples under "optically-thin" conditions, even at relatively high CDOM values. We therefore chose the quartz spiral flowcell with a 1 mm pathlength, which allows for 100x high CDOM values when compared to a 10 cm spec cell. One very surprising result was that even though SRNOM with DOC concentrations ranging from 10 to 50 mg L⁻¹ degraded at similar rates, there appears to be a concentration dependence at low SRNOM DOC concentrations. This is a subject that we hope to evaluate more in future work. 2. To prevent oxygen loss, the sample is continually returned to an equilibrator, where it is mixed and reoxygenated. 3. While we did not control samples for pH in this study (as mentioned no pH changes when irradiating our samples at pH 3), pH control is possible with a microtitrator and has been described in a previous study (Timko et al. 2015). We realize with this design we have sacrificed accurate photon dose quantification with a spectroradiometer that is possible in designs with very well defined and controlled optics. However, with the goal of reproducibility in mind and limiting problems that occur when irradiating sealed vessels, we believe this system is a valuable tool to use when evaluating "photosensitivity". For instance, it will be a great tool to evaluate some of the questions/issues mentioned above (e.g. do buffers impact photodegradation rates?). Unfortunately, many studies do not control optics or do not use reproducible designs, e.g. irradiating samples in curved vessels on their sides or in flasks. While we cannot calculate spectral photon doses in our system, we thought that using nitrite actinometry as a broadband measurement of UVA photon dose was better to use than irradiation time as mentioned in the manuscript (lines 211-220, 259-268, Figure 3). We thank the reviewer for bringing up these valuable discussion points. Revisions discuss in detail the tradeoffs we made between radiometer-quantifiable optics and sample control (lines 424-430).

The referee raises a few points related to our other suggestions for improving comparability and reproducibility when measuring photodegradation kinetics, stating "I quite agree with these recommendations which are straightforward to implement. For others, it is not clear that they will always contribute to reproducible, or more importantly, interpretable, results." Here we stress that we do not claim our approach provides a universally applicable prescription to ensure reproducible results – rather we hope to illustrate several obstacles to reproducing and comparing studies of photodegradation kinetics, and demonstrate some progress towards improving or at least identifying these sources of uncertainty. The manuscript was revised to convey this distinction more clearly (lines 71-72; 576-578). Specifically, the referee notes "[w]hile extracts are inherently more stable than raw filtrates, the authors demonstrate differences in the photosensitivity kinetics of extracts vs the original filtrates, the reasons for which are not well understood. The authors advise preparing extracts in some situations but not others, this leaves open the question of comparability."

This is in fact exactly our intention – to illustrate obstacles to reproducibility that need to be explicitly addressed when conducting or at least when comparing studies of photodegradation

kinetics. As the referee notes, the differences between extracts and filtered raw water samples are not well understood and worthy of future study. Until these issues are resolved, researchers will have to weigh the advantages and disadvantages of extracts vs. raw water. Here we chose to use only extracts in subsequent trials in order to compare samples collected at different times that were extracted for stable storage, and to limit our inference to differences in photodegradation kinetics arising from DOM composition and not matrix differences. Other studies will have to choose between using extracts vs. raw water in accordance with study priorities. The differences we demonstrate (Figures 6 and 7, Table 1) illustrate underappreciated limitations to reproducibility and identify opportunities for further research into the effects of extraction on photodegradation kinetics and mechanisms of action (e.g. effect of matrix vs. fraction of DOM extracted). Along similar lines the referee notes "The authors also advise using a standard optical density in the sample, yet also show there is a concentration dependence to the kinetics, even for an optically thin exposure conditions. Again, a decision will have to made of whats more important, knowing how the DOM behaves at its natural concentration or being able to compare it to other sources." Again, we are in agreement with the referee here, and believe one of our manuscript's contributions is highlighting these underappreciated sources of variability in DOM photodegradation kinetics. This issue in particular is vexing, as differences between DOM concentrations under conditions meeting conventional thresholds for optically thin solutions are not explained well by currently published literature. We revised the manuscript text to make it more clear that our goals were not to show only reproducible results but to identify underappreciated sources of variability that need to be controlled, and to use these insights to conduct limited trials with comparable results (lines 71-72; 576-578).

The referee's final point concerns the utility of our approach in a wider sense – "In the end, investigators will need more information to decide whether the additional insight gained from defining photosensitivity kinetics will be worth the efforts needed to perform all these standardization steps. This report documents experimental tests that show what are sources of variability in the procedure but does not show what new understanding is actually being gained from the results (vs what can already be gained from other optical/chemical measurements typically made on DOM)." We agree that our approach is not trivial, and some applications may not really benefit from this level of effort. Our primary research goal was to compare the sensitivity of different DOM sources to photochemical degradation processes at environmentally relevant temporal scales. Measuring kinetics of DOM photodegradation changes seemed the best way to pursue this, but we found existing approaches to describing DOM photodegradation kinetics allowed for possible uncertainties that were barriers to meaningful comparison (as discussed in detail above). We believe the methodological suggestions we present are useful to other researchers in this area pursuing similar goals, and our research findings (seasonal and spatial differences in wetland DOM photo-sensitivity, with implications for understanding DOM composition and the other ecological processes that affect it in this setting) demonstrates the potential utility of this approach and illustrates phenomena worthy of further investigation.

Specific concerns:

Line 140 "Solar exposure" – This was not solar exposure, but a laboratory set up. Better (?) "Total sample exposure"

We agree and revised the manuscript accordingly (now line 154).

Figure 7 and Table 1 – The captions for these describe kL and kSL with units of mol photons m-2. Those are the units of P, the units of k should be the inverse, m² [mol photons]⁻¹. We agree and revised the manuscript accordingly (Figure 7, Figure 10, Figure 12, Table A1, Table A2, Table A3 captions).

Figure 7 and Table 1 apparently show the same data – and both state the statistics of the comparison. Questionable whether both are necessary. Figure 7 caption, however, says see Fig. 5 for data. It appears that Fig. 6 has the data.

The referee is correct that Figure 7 caption should refer to Figure 6 instead of Figure 5, and also correct that both Figure 7 and Table 1 show the same data. Readers may find the graphical comparison easier to quickly parse, but the tabular presentation may be more useful to future studies seeking comparison. We think the graphical presentation is more valuable to readers in the main text, and moved Table 1 to an appendix upon revision (now Table A1).

Line 350-360 – Factors affecting storage of filtered water. Previous text only mentioned preparation of filtrates using 0.7 _M glass fiber filters, which allows passage of a substantial bacterial fraction. However, subsequent guidelines caution against used stored material even filtered to 0.2 _M (some bacteria can even get through those). Please clarify what filtration method was used for the stored sample, but in any case (slow) microbial degradation may be another factor resulting in changes in stored material.

We inadvertently failed to note an important step in our methods – all samples, whether raw water or solid-phase extracts redissolved in water, were filtered through syringe-mounted 0.2 µm cellulose acetate filters that were pre-rinsed with > 30 mL ultrapure C-free water. GFF filters were used as a pre-filter immediately after samples were returned to the lab, before either shortterm storage or solid-phase extraction. Furthermore, since 0.7 µm filters were combusted, their true pore size is probably smaller than 0.7 µm (0.3 µm in Navar and Chou, 2003) and therefore be comparable 0.2 µm filters (Nayar and Chou, 2003). We apologize for the omission and ensuing confusion and revised the manuscript to clarify (lines 95-97). We also revised our manuscript to clarify the methods used in the storage time experiment (lines 194-200). The referee notes that microbial degradation is still possible even when using a filter of 0.2 µm pore sizes – we agree (see Brailsford et al., 2017; Luef et al., 2015), and believe this supports our recommendation to either use extracts or keep storage time minimal and highlights the need for more research in this area (revisions to lines 118-120). We also believe microbial degradation/contamination is minimal during exposures under the solar simulator using optically thin conditions (e.g. Stubbins et al. 2017, 122), but revised our manuscript to mention the full range of possibilities that must be accounted for in any similar experimental set up (lines 139-144). This would include discussion of between-sample cleaning with dilute NaOH and occasional flushes of sample lines with isopropanol (followed by extensive water flushes and testing for DOC residue) to prevent microbial contamination of flow lines.

Figure 10 – Caption says see Figure 6 for data, data appears to be in Fig. 9. This correction also applies to the caption for Figure A1. Figure 10 could be misleading to the reader, as a quick inspection may suggest that the spread of the multiple points for a particular sample source (and standard deviation in the table) are indications of reproducibility. Actually, only the SRNOM and three of the small wetland points are repeat determinations on the same sample. Separating the points for different sample dates would avoid this confusion, especially given that

later on (section 3.3.2) the authors attribute some of the difference in the wetland results with time to possible differences in sample composition and/or previous solar exposure. The referee is correct that the captions for Figure 10 and A1 should read "see Fig. 6 for data". We agree that the representation of data in Figure 10 may be confusing. This criticism may also apply to Table 2 (in original submission), where group means and standard deviations for photodegradations of DOM sources with more than one trial are grouped differently – SRNOM represents three actual replicate trials using the same sample source, the three "large wetland" trials represent different samples (collected at same site but at different times), and the five "small wetland" trials represent three replicate trials using the same source and two trials from different sample sources (collected at the same site but at different times). We originally thought it was useful to make the comparison in Figure 10 in this way because of the widespread interest in differentiating DOM sources by setting. By showing fitted parameters as independent data points we thought readers could parse the different types of grouping by reading the text carefully. However, we agree with the referee's concerns – at the very least the sampling dates should be differentiated by using different symbols in the plot (revised Figure 10 and Figure A1). Upon reflection we also believe Table 2 (in original submission) should be changed along these lines – our original intention was to facilitate comparisons between environmental settings, but calculating group means with these data is not appropriate given their different structures. Means for replicated trials are sensible, and it may even be valuable to see how un-replicated trials with samples from different seasons at a site differ, as solely spatial comparisons often do not account for temporal/seasonal variation but readers may still be interested in summarizing data by site. But calculating one mean from data that includes 3 replicated trials of the same sample source and two further distinct samples does not convey useful information. We think publishing tabular data may be useful for future research, but this data would be better represented separately. For size reasons, such a table may be better suited to an appendix. We re-calculated means where appropriate for replicates and listed parameter fits in Tables A2 and A3.

Line 456ff, section 3.3.1 Here the relative abundances of kinetic components are considered as metrics of DOM composition or other influences on photosensitivity. A problem with the comparative use of fL and fSL is that the two parameters are not independent. Assuming a good fit to the actual time course the two fractions will always sum to approximately 1, which means that any variation in one fraction will be reflected in a complementary variation in the other. Thus, changes in, e.g. fL, with sample date or location (as later discussed in 3.3.2) could be caused by change in the concentration of labile or semilabile fraction, or both. If the focus is on the rate parameters, this is not so important, but if the magnitudes of the fractions are to be comparative metrics, the need to be independent of each other. For example, ft could be the actual component score, score normalized to t0 total fluorescence, or score normalized to DOC.

We see the referee's point that fSL and fL are not truly independent. They were fitted as separate coefficients in the nonlinear fitting algorithm applied to the data, but should indeed sum to 1, so either could be expressed as the difference between 1 and its counterpart (i.e. fSL = 1- fL or vice versa). We expressed them separately here in accordance with their separate representation in the equation conventionally used to define the decay model (Equation 2, which Equation 3 then followed), and because they were fitted separately (which would make the choice of which to use to find its counterpart arbitrary). We agree that this causes ambiguities when trying to understand differences in these values between samples, as in the example offered by the referee. We do

believe using these values may be informative, as a major part of our claim to novelty rests on pointing out possible connections between these values and environmental phenomena (such as previous exposure to photodegradation or differences in source material composition) instead of basing inference solely on k parameters. We are not sure how to best resolve this issue. It already represents PARAFAC component fluorescence normalized to time 0 for each sample. Further processing the data to derive a single value (like the ratio of fSL and fL) adds another layer of abstraction to a situation that already involves multiple modeling steps, and seems harder to interpret in a way that materially connects to the phenomena described. The revised manuscript discusses these points (lines 244-247). However, we still think it is valuable to compare fL between samples – even if differences in this value represent multiple possible causes as described by the referee, differences here represent real observed differences in photodegradation behavior. Normalization to [DOC] or to absorbance at some reference wavelength may be useful in this context. We believe this is an area ripe for future research, and view our contribution in part as pointing out the need for attention to previously unexplored dimensions of current paradigms of DOM photodegradation.

The authors offer the interpretation that the fractions correspond to "pools" of more or less reactive DOM. However, Murphy et al. (2018) interpret biexponential kinetics as possibly reflecting multiple photoreaction pathways, a fast one involving ROS and slower one related to direct photochemistry. How do the authors reconcile these differing interpretations? We appreciate the referee's thoughtful reading of our conclusions. Our conceptualization – differences in photodegradation kinetics arising from different relative quantities of "pools" reacting at relatively fast and slow rates – may be unnecessarily general or vague. However, we think the explanation from Murphy et al. (2018) invoked by the referee is not necessarily opposed to the "pools" we ascribe to the model. We consider these "pools" in a general sense. Given the immense complexity of natural DOM, it is unlikely there are two distinct, nonoverlapping compositional types of photo-sensitive DOM. Yet the excellent fit of biexponential models across diverse DOM sources requires some explanation. What we refer to generally as reactive pools could more specifically be the pools of DOM subject to the respective reaction pathways suggested by Murphy et al. and their associated rate constants. Or perhaps it is more appropriate to consider a single "pool" with differing capacities for two classes of reactions. We are not able to definitively conclude what lies behind the biexponential model – but believe our novel attention to the f parameters in the model equation represents the kind of exploration that is needed to make progress in this direction. Our revised manuscript discusses these possibilities and clarifies our use of "pools" (lines 284-285; 497-502)

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RC2 response

We appreciate the thoughtful comments from the referee. The most pressing issue presented in this review is how to justify the effort involved. Our main goal was to compare the sensitivity of different DOM sources to photochemical alteration to better understand potential influences on DOM composition in natural waters. There are many approaches to characterizing the optical properties different DOM sources and an established tradition of photochemical experimentation on DOM. However, DOM in natural waters is subject to many potential transformation processes that may compete and interact on different temporal scales, so reliable comparison of degradation kinetics seemed like the most promising approach to characterize photosensitivity under environmentally relevant photon exposures. For example, changes to DOM composition that slowly over the equivalent of several days of photodegradation may be less important than changes occurring in a few hours in an environment where water is moving or mixing and subject to other processes like microbial transformations. Much of the current manuscript is devoted to the methodological considerations we encountered in our effort to reliably compare degradation kinetics, many of which are also relevant to other approaches used to study DOM photodegradation and thus broadly interesting to this community. Our specific research findings were that wetland DOM photo-sensitivity varied in space and time with potential implications for understanding DOM composition in this setting generally, and that our method identified particular aspects of photodegradation kinetics that might represent links between DOM composition, environmental phenomena, and photodegradation. This represents an initial foray into understanding the role of photodegradation in shaping DOM composition in our study area, and we hope it illustrates aspects of DOM photodegradation that may be applicable elsewhere and interest the broader research community. Our manuscript was revised to clarify the goals and novelty of our work (lines 71-72; 576-578)

The referee raised several methodological questions. We acidified our samples to pH 3 because we had to control pH - pH changes produce fluorescence changes that confound the fluorescence decay due to photodegradation reactions under investigation. We did not wish to use chemical buffers, as some of these may also interfere with fluorescence and may have unknown effects on degradation kinetics. Active control of pH during experiments using autotitration is another option we did not use because the electrode contaminated samples during prolonged irradiations. Therefore, we needed to be sure that pH would be consistent between samples and during photodegradation of each sample. Starting samples at pH 3 allowed the full protonation of most organic acids (Ritchie and Perdue, 2003), which should prevent solution pH change due to the photoproduction of CO₂. We recognize that this itself presents an obstacle to reproducibility or comparison with other studies, as kinetics are greatly affected by pH, but this seemed the most reliable way to avoid pH change during photodegradation to allow comparisons between our samples. Another methodological question involved filtration choices. We appreciate the careful attention of the reviewer here, which helped us discover that we inadvertently failed to note an important step in our methods – all samples, whether raw water or solid-phase extracts redissolved in water, were filtered through syringe-mounted 0.2 µm cellulose acetate filters that were pre-rinsed with > 30 mL ultrapure C-free water. GFF filters were used as a pre-filter immediately after samples were returned to the lab, before either short-term storage or solidphase extraction. We apologize for the omission and ensuing confusion. Additionally, as mentioned to reviewer 1, because 0.7 µm filters were combusted, their true pore size is probably

smaller than $0.7 \mu m$ and likely be comparable $0.2 \mu m$ filters (Nayar and Chou, 2003). We revised the manuscript to clarify our methods (lines 95-97).

The referee's methodological questions included several about comparability of our results to natural, un-processed DOM. These included questions about effects of extraction, the means used during evaporation of methanol extracts, details on the experiment showing effects of storage, use of freeze-dried IHSS standards, and questions about the collection and initial characterization of natural DOM sources. A few of these questions were addressed in the text. The methanol extracts were evaporated in an open vial under a fume hood under a stream of high-purity N2 gas – the gas flow created turbulence in methanol, facilitating its evaporation. We used IHSS SRNOM standard as the DOM source in many of our experiments because it has been extensively studied and provides a benchmark against which other photodegradation kinetics experiments can compare results. Reconstituted freeze-dried SRNOM and its filtered source water showed very similar molar absorptivity at 280 nm and fluorescence index values when collected (Kuhn et al., 2014). While this may not translate to identical kinetics of photodegradation, it suggests optical properties are not radically transformed during freezedrying and reconstitution. We believe the advantages of using a standardized material in wide circulation outweigh any artifact of freeze-drying. The experiment showing the effects of storage on photodegradation kinetics was conducted according to different protocols than following experiments – samples had higher DOC concentrations, and a different flow cell with shorter path length was used. This experiment was the first conducted, chronologically, of those presented here, and its failures of replication motivated further work and helped us develop our protocols. We apologize if introducing these protocols sowed confusion, and edited the manuscript to ensure it is clear which protocols applied to which experiments (lines 194-200). The referee also asked whether EEMs and other measures were available for comparison between the solid-phase extracts used throughout the paper and their original raw water DOM sources. While these are not available (at least in comparable form) for many of the samples used to show the range of variability across a wide gradient of DOM sources (e.g. the stream, coastal ocean, and Sargassum sp.), we did collect EEMs and absorbance spectra for the "large" and "small" wetlands sampled in different seasons at the time of collection. We revised the manuscript to include a comparisons between source water EEMs and extract EEMs from wetlands (Figure A4).

The referee ended their specific comments by requesting "some kind of discussion...on how comparable the results [from extracts studied in the lab] will be to what may happen in situ." We appreciate the utility of this kind of information – inference on the processes at work in the natural setting is the original motivation for our entire effort. While we were able to determine that extracts behaved differently than raw water (demonstrated with SRNOM), we cannot identify the mechanistic basis of these differences with the current evidence. The difference between *in situ* conditions and our lab results might be useful to expound further in a future manuscript, however, this is not a trivial task. For instance, we used solid phase extracts to minimize matrix effects and controlled pH to look at differences in photochemistry between DOM samples and SRNOM. Cations and in particular magnesium, may actually enhance DOM fluorescence signals (Stichak et al. 2019, ES:P&I,21) and pH also influences sample optical properties, as mentioned previously. Metals like iron are known to enhance absorbance measurements and potentially quench fluorescence (Poulin et al. 2014, E&T, 48). Halides may

also enhance photodegradation rates (Grebel et al. 2009). These potential matrix effects (see Li and Hur, 2017 for a detailed discussion) make observed differences in whole water optical properties difficult to interpret. SPE allows for comparisons with minimal or entirely without potential interferences and matrix effects. While sample matrix is indeed different *in situ*, our method may best be described as measuring potential photodegradation of DOM itself, which is why we describe it as photosensitivity. One of our key results shows the utility of our method in this regard – we are able to measure the relative differences in fast degradation processes (described by fL and kL), which may be more environmentally relevant than the slower processes in the context of constantly changing DOM subject to a panoply of other transformation processes and constantly in motion. We thank the referee for asking for this kind of discussion, and revised the manuscript to make the relationships between our lab approach and in situ inferences more clear (lines 542-553).

The referee's final question was about Figure 9. There is an error in this figure that may have produced additional confusion – the left plot is of PARAFAC component 3 while the right plot is of PARAFAC component 4. We corrected this in the revised manuscript. The referee's question concerned the provenance of the multiple curves shown for the wetland samples. These plots show the full data available for all different DOM sources subjected to the final protocol. This includes the three replicates of extracts from the same "small wetland" DOM source and two trials with extracts from "small wetland" DOM collected at different times, three trials with extracts from "large wetland" DOM collected at different times, and the three replicates of SRNOM extracts. As described in our response to RC1, we recognize the need to clarify the different dimensions of these multiple trials for each site/source, especially in Figure 10 and Table 2 (as numbered in original submission, now Tables A2 and A3). It may be useful to further clarify here as well, though as this figure shows rawer data (PARAFAC model component fluorescence change over time) rather than biexponential model parameters for each curve shown here, this may be less pressing for this figure.

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Reproducible determination of dissolved organic matter photosensitivity

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Abstract. Dissolved organic matter (DOM) connects aquatic and terrestrial ecosystems, plays an important role in C and N cycles, and supports aquatic food webs. Understanding DOM chemical composition and reactivity is key to predict its ecological role, but characterization is difficult as natural DOM is comprised of a large but unknown number of distinct molecules. Photochemistry is one of the environmental processes responsible for changing the molecular composition of DOM and DOM composition also defines its susceptibility to photochemical alteration. Reliably differentiating the photosensitivity of DOM from different sources can improve our knowledge of how DOM composition is shaped by photochemical alteration and aid research into photochemistry's role in various DOM transformation processes. Here we describe an approach to measure and compare DOM photosensitivity consistently based on the kinetics of changes in DOM fluorescence during 20h photodegradation experiments. We assess the influence of experimental conditions that might affect reproducibility, discuss our modelling approach, offer guidelines for adopting our methods, and illustrate possible applications for ecological inferences. Central to our approach is the use of a reference material, precise control of conditions, leveraging actinometry to estimate photon dose, and frequent (every 20 minutes) fluorescence and absorbance measurements during exposure to artificial sunlight. We compared DOM from freshwater wetlands, a stream, an estuary, and Sargassum sp. leachate and observed differences in sensitivity that could help identify or explain differences in their composition. Finally, we offer an example applying our approach to compare DOM photosensitivity in two adjacent wetlands as seasonal hydrologic changes alter their DOM sources. Our approach may improve reproducibility when compared to other methods and captures time-resolved changes in optical properties that may have been missed previously.

1 Introduction

The photochemical reactivity of dissolved organic matter (DOM) is inherently linked to its composition and photochemical behavior reflects compositional differences between samples. Several authors have discussed the fundamental processes involved in light absorption by DOM and the phenomena that may follow (Miller, 1998; Sharpless et al., 2014), including loss of absorbance (Del Vecchio and Blough, 2002), production of new substances (Gonsior et al., 2014; Blough and Zepp, 1995; Bushaw et al., 1996; Moran and Zepp, 1997), and loss of fluorescence (Blough and Del Vecchio, 2002). Absorption

30 spectra and derived values such as spectral slopes and their ratios have long been used to characterize DOM (Blough and Del Vecchio, 2002; Helms et al., 2008; Twardowski et al., 2004). Fluorescence measurements arise from only a fraction of chromophoric DOM (CDOM) but are sensitive to small variations in DOM chemical composition (Blough and Del Vecchio, 2002). To the extent that photochemical reactivity is a property of DOM chemical composition (Boyle et al., 2009; Cory et al., 2014; Del Vecchio and Blough, 2004; Gonsior et al., 2009, 2013; Wünsch Urban J. et al., 2017), comparing photosensitivity of different DOM sources or treatments may be a useful tool in the continuing effort to characterize DOM composition and to describe its susceptibility to sunlight-induced degradation.

Research across ecosystem settings has measured changes in optical properties following sunlight or simulated-sunlight irradiation to infer changes in DOM composition. A general discussion of this approach and its bases has been previously published (Hansen et al., 2016; Kujawinski et al., 2004; Sulzberger and Durisch-Kaiser, 2009). Examples of recent research using photochemical changes to make ecologically significant distinctions between DOM samples collected in specific ecosystems have been described in detail elsewhere (Gonsior et al., 2013; Laurion and Mladenov, 2013; McEnroe et al., 2013; Minor et al., 2007). DOM photo-reactivity itself has ecological consequences, affecting overall carbon (C) cycling (Anesio and Granéli, 2003; Obernosterer and Benner, 2004), microbial heterotrophy of DOM (Amado et al., 2015; Cory et al., 2014; Lapierre and del Giorgio, 2014), and algal and submerged plant primary productivity (Arrigo and Brown, 1996; Thrane et al., 2014).

Experimental approaches connecting DOM chemical composition, its optical properties and their photochemical bases, and relevant ecological phenomena typically expose natural DOM samples to natural or simulated sunlight and measure the change in optical properties over time. *In situ* experiments have been used to explore the role of photodegradation relative to other transformations of DOM in aquatic ecosystems but field studies are difficult if not impossible to reproduce (Cory et al., 2014; Groeneveld et al., 2016; Laurion and Mladenov, 2013). Laboratory-based irradiation experiments may allow greater reproducibility and logistical flexibility. Laboratory photodegradation experiments have tested the potential ecological significance of photodegradation and explored the fundamental photochemical mechanisms involved in photobleaching (Chen and Jaffé, 2016; Del Vecchio and Blough, 2002; Goldstone et al., 2004; Hefner et al., 2006). These experiments usually involve simultaneous irradiation of DOM in several sample vials under polychromatic or monochromatic light. Vials are then destructively sampled for DOM measurements at intervals throughout the experiment, or simply compared before and after light exposure. While powerful, these experiments require a trade off in effort between reproducibility and temporal resolution. Replicate vials are often sampled to ensure precision and improve reproducibility, but lamp space is finite, limiting temporal sampling resolution.

Continuous measurement of a single sample undergoing controlled photoirradiation offers an alternative experimental approach. The kinetics of DOM fluorescence loss during photoirradiation experiments have been recently described (Murphy

et al., 2018; Timko et al., 2015). These studies leveraged novel time series of frequent measurements (e.g. every 20 minutes) of fluorescence and UV-Vis absorption which allowed modeling of distinct reactive components. Fluorescence losses were best described by the sum of two exponential decay terms, allowing straightforward and precise modeling of photosensitive fluorescence signals that degraded quickly.

The goal of this study is to compare the photosensitivity of different DOM sources to better understand the links between DOM composition, environmental setting, and photochemical degradation processes. Our first task is to demonstrate the suitability of our approach. In a series of experiments, we explored potential sources of variability in photodegradation kinetics stemming from experimental conditions and methodology. We further develop a previously described experimental setup (Timko et al., 2015), showing results are reproducible under controlled conditions using a common reference material, and suggest a set of best practices for collecting reproducible and high resolution time series of fluorescence measurements during experimental irradiation of a single sample. Then we apply this approach to several natural DOM sources and identify photosensitivity differences that may be ecologically relevant. Finally, we focus on DOM from two wetlands to show how these key differences in photosensitivity metrics may help us link DOM composition to ecological phenomena.

2 Materials and procedures

2.1 Sample materials

We used Suwannee River natural organic matter (SRNOM) obtained from the International Humic Substances Society as a reference material (catalog no. 2R101N, isolated by reverse osmosis; (Green et al., 2014)). Freeze-dried SRNOM was dissolved in Milli-Q water and was prepared less than one week prior to use (hereafter called RO SRNOM). Dilutions approximately corresponded to a dissolved organic carbon (DOC) concentration of 5 mg C l⁻¹. This is well below the [DOC] range found in SRNOM source material before it was extracted, but within the range of other aquatic DOM sources dominated by terrestrially-derived DOM. Additionally, SRNOM solid phase extracts using the Agilent PPL resin were extracted in May 2012 during the same time the SRNOM standard material was isolated, and were prepared directly before irradiation experiments (see details below).

Additional water samples were collected across a variety of aquatic ecosystems to explore the range of our approach and to validate it. Sample sources include two freshwater wetland sites (Caroline County, Maryland, USA), one perennial stream (Parker's Creek, Calvert County, Maryland, USA, collected September 2017), one estuary (Delaware Bay, USA, collected July 2016), and leachate from live *Sargassum sp.* collected in Bermuda in July 2016 (Powers et al., 2019). These samples were 0.7 µM filtered within 24 hours of collection through combusted (500°C) Whatman GF/F filters and acidified to pH 2 using concentrated HCl (Sigma, 32% pure) before solid-phase extraction. The true pore size used in this pre-filter step was probably smaller than 0.7 µm (e.g. 0.3 µm in Nayar and Chou, 2003). All samples, whether whole water or solid-phase extracts

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redissolved in water, were filtered through syringe-mounted 0.2 µm cellulose acetate filters that were pre-rinsed with > 30 mL ultrapure C-free water.

Samples from the two freshwater wetland sites are used in the more detailed comparison presented in Section 3.3 and hence these sites merit additional description. Small topographic depressions are common throughout the interior of Delmarva Peninsula. These depressions persist in this low-elevation, low-relief landscape, and regular seasonal inundation has led to the development of wetland soils and biota in many of these depressions. Depressions on land not drained for agriculture are inundated for several months most years. Some do not exchange water through surface flow with perennial stream networks, while others sustain downstream connections through temporary surface channels for several months in the wettest months of the year (typically late winter-spring). These two sites, referred to as "smaller wetland" and "larger wetland", are adjacent but lie within distinct topographic depressions. Their inundated areas expand and contract with water level fluctuations, and both may go entirely dry at the surface in the summer. If water levels are sufficiently high, their surface waters merge, and a temporary channel may fill and sustain export flow to the perennial stream network. One sampling site is within the smaller depression, which mostly lacks submerged and emergent vegetation and is hemmed closely by trees. The other site is within a larger depression, where surface water is more exposed to light and features a variety of herbaceous submerged and aquatic plants. Experiments were run with DOM from both sites, sampled on three dates (2017-10-05, 2017-12-20, 2018-04-01).

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Except for RO SRNOM samples used to test the effect of solid phase extraction and wetland samples used for the storage time experiment described below, all samples were solid-phase extracted using a proprietary styrene divinyl benzene polymer resin (Agilent PPL Bond Elut) following a procedure described previously (Dittmar et al., 2008). PPL extracts were used because our goal is to develop a reproducible method to compare photochemical behavior of natural organic matter without the influence of the sample matrix. Extracts allow longer storage, isolate organic matter from potentially photosensitive matrices, and capture representative photosensitive organic matter fractions (Murphy et al., 2018). While filtration to 0.2 µm should remove most viable microbes, microbial degradation may still be possible in filtered water if ultra-small microorganisms are present (Brailsford et al., 2017; Luef et al., 2015). Extraction removes this possibility.

Immediately prior to each experiment, 0.5-5 ml of the extract was evaporated under high-purity N2 gas, dissolved in 30 ml

ultrapure C-free Milli-Q water, and diluted to similar CDOM absorbance values to minimize any potential inner filter effects on fluorescence degradation kinetics. Absorbance (A) at 300 nm was used as a benchmark for dilution instead of adjustments based on measured [DOC] because it could be done quickly on the equipment used for the photochemical experiments and allowed consistent correction of inner filtering effects. We adjusted all samples (except for those used in the storage time experiments described below) to a raw absorbance of 0.12 (+ 0.01), which translates to a Napierian absorption coefficient (a) of 27.6 m⁻¹. Delaware Bay samples were too dilute to generate sufficient volume to fill the photoirradiation system, so several sample extracts from throughout the depth profile of a single sample station were combined prior to evaporation.

4 Author's changes

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2.2 Photoirradiation system

We needed our system to irradiate samples under optically thin conditions even at relatively high CDOM concentrations. The photoirradiation system circulates an aqueous sample between a mixing reservoir (i.e. equilibration flask), a solar simulator, and a spectrofluorometer, similar to a system described previously (Timko et al., 2015). Samples were continuously circulated between a central mixing reservoir and system components were connected by PEEK tubing (LEAP PAL Parts & Consumables, 0.0625" OD/0.030" ID). The central reservoir was a 25 mL borosilicate equilibrator flask with a magnetic stir bar constantly rotating at its bottom. Sample gently dripping from flow lines into the equilibrator ensured sample remained oxygenated during photodegradation. A micro gear pump (HNP Mikrosysteme, mzr-4665) was used to pump the sample with almost pulse-less flow through the system at a rate of 1.5±0.1 ml min⁻¹. The spectrophotometer flow cell and equilibrator flask were surrounded by a circulating water jacket set to 25 °C. To prevent contamination or the establishment of microbes that could degrade DOM during experiments, the system was flushed with 0.1 M NaOH between experiments then thoroughly flushed with ultrapure water. Ultrapure water for blanks was circulated for at least 10 minutes before checking absorbance and fluorescence for signs of contamination. If blank contamination persisted after subsequent rinses, the system was flushed with isopropanol and thoroughly rinsed with ultrapure water before checking for contamination by examining optics and testing [DOC].

Samples were irradiated as they were slowly pumped through a custom-built flow cell (SCHOTT Borofloat borosilicate glass, Hellma Analytics, 70 to 85% transmission between 300 and 350 nm, 85% transmission at wavelengths >350 nm), with a total exposure path area of 101 cm² arranged in an Archimedean spiral and returned to the equilibrator flask. This 20x20 cm borosilicate spiral flow cell had a 1 mm deep x 2 mm wide long flow path covering the irradiation area and was located underneath a solar simulator (Oriel Sol2A) with a 1,000 W Xe arc lamp equipped with an air mass (AM) 1.5 filter. Lamp output was checked periodically using an Oriel PV reference cell set to one sun which corresponds here to exactly 1,000 W m ² and lamp power was held constant during irradiation experiments using a Newport 68951 Digital Exposure Controller. Another tubing carried the sample from the equilibrator flask to a temperature-controlled square quartz fluorescence flow cell (1 cm x 1 cm) located within a Horiba Jobin Yvon Aqualog spectrofluorometer. Total sample exposure varied depending on the total volume in the photodegradation system. We controlled volume by completely filling the tubing and flow cells (12.2 mL volume) and adjusting volume added to the equilibration flask. With 10 mL volume added to the equilibrator (our typical experimental conditions), a 20 h irradiation experiment was equivalent to 1.0 day of exposure between 330 - 380 nm at 45 °N latitude in mid-July where one day is ~15.75 h long. For the lowest total volume used here (0.5 mL in the equilibrator, total volume 12.7 mL), photon dose was 1.7 times higher than this estimate. We calculated a mean photon flux of 3.9x10⁻⁵ mol photons m⁻²s⁻¹ for experiments with 10 mL sample added once flow lines were filled (total sample volume 22.2 mL), based on a mean photon exposure of 0.23 µmol photons cm⁻² min⁻¹ (5 trials, standard deviation 0.0045). This calculated flux is based

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on nitrite actinometry and a response bandwidth between 330 and 380 nm (Jankowski et al., 1999, 2000). Average July solar

irradiance was modeled using the System for Transfer of Atmospheric Radiation model (Ruggaber et al., 1994) calculated just below the water surface as described previously (Fichot and Miller, 2010).

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Past experiments revealed the importance of pH control on DOM fluorescence and photodegradation kinetics (Timko et al., 2015). We adjusted initial sample pH to 3.0 (+ 0.2) with HCl but did not control pH by autotitration. At pH 3 natural organic acids should generally be protonated regardless of compositional differences between DOM sources, which should prevent solution pH change due to the photoproduction of CO2 (Ritchie and Perdue, 2003). Starting at pH 3 and equilibrating the sample in an air-filled reaction vessel ensured minimal pH change during irradiation, never changing by more than 0.2 pH units, in line with expectations from work on mechanisms explaining pH decreases during photooxidation (Xie et al., 2004).

2.3 Optical measurements

We used a Horiba Jobin Yvon Aqualog spectrofluorometer to collect time series of UV-Vis absorbance and excitation-emission matrix (EEM) fluorescence spectra throughout experiments. UV-Vis absorbance was measured at 3 nm intervals between 600 and 230 nm. Fluorescence excitation occurred at the same intervals, and emission spectra were recorded from 600 to 230 nm at 8 pixel CCD resolution, or approximately 3.24 nm intervals. EEMs integration times were 1 second. Milli-Q water (18.2 MΩ-cm) adjusted to pH 3 with concentrated HCl was circulated through the system and used as a measurement blank immediately prior to each experiment.

2.4 Experiments

185 Several sets of experiments explored method reproducibility, sensitivities to experimental conditions, and differences between DOM sources. A series of experiments used SRNOM PPL extracts at varying concentrations and volumes added to the photodegradation system to test their influence on degradation kinetics. Different researchers in our group then repeated experiments with SRNOM PPL extracts to test reproducibility. We explored effects of storage time on filtered water sample photodegradation results. We next compared SRNOM PPL extracts and SRNOM reference material reconstituted in ultrapure water (RO SRNOM) to test the effect of extraction on photodegradation kinetics. After examining the methodological sensitivities with SRNOM, we compared the DOM sampled from several contrasting aquatic ecosystems.

Samples were exposed to 20 hours of simulated sunlight, and EEM spectra were collected (using the "Sample Q" feature in Aqualog software) starting immediately before irradiation began with a 17.5 minute interval between each scan, generating a time series of 60 EEM spectra for each experiment. Where applicable, time of EEM collection was converted to cumulative photon exposure (mol photon m⁻²) by multiplying time by calculated photon flux (mol photon m⁻² s⁻¹) using actinometry results generated with the same sample volume.

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200 The experiments testing effects of storage time on photodegradation kinetics require additional explanation. These preceded the other experiments and the experimental setup was modified based on their results. These used filtered water samples (as above) taken from the small and large wetland sites described above, but were collected in November 2017. They were filtered, stored in the dark at 4°C, and run through the photodegradation system undiluted using a 3x3mm quartz flow cell in the spectrofluorometer instead of the 10x10mm cell used for all other experiments. Experiments were run after 5-8, 9-13, and 14-16 days of storage. These results are reported as a function of time rather than photon exposure as no actinometry was collected with an analogous experimental setup.

2.5 Data analyses

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Fluorescence EEM spectra were inner-filter corrected and had 1st order Rayleigh scatter removed by the built-in Aqualog software (based on Origin). Second order Rayleigh scatter was removed using an in-house Matlab toolbox following methods previously described (Zepp et al., 2004). EEM spectra were normalized by dividing fluorescence measurements by the area of the water Raman scatter peak of the water blanks. Data were processed in Matlab R2018a using an in-house toolbox and the drEEM toolbox (Murphy et al., 2013). Absorbance data were converted to absorption coefficients using Eq. 1:

 $a_{(\lambda)} = 2.303A(\lambda)/l \tag{1}$

where a is the absorption coefficient at wavelength λ , A is raw absorbance at wavelength λ , and l is path length in m, here 0.01 (Hu et al., 2002).

We fitted a 4-component parallel factor analysis (PARAFAC) model to data from 3 SRNOM PPL extract experiments (60 EEMs each, 180 EEMs total). PARAFAC models with 3, 4, and 5 components were fitted to the 3 SRNOM PPL extract experiment EEMs. The 4-component model was chosen as it exhibited better component spectral characteristics than the others. Emission spectra from components matched the 4 components identified in similar experiments (Murphy et al., 2018). Split-half validation is often used to validate PARAFAC models fitted to data sets where each EEM represents a different DOM source but may not be appropriate for data sets where EEMs are not independent. Instead, 4-component models were fitted from each of the three SRNOM PPL extract experiments individually to confirm each experiment's data led to the same PARAFAC model, then the model built from all three experiments was compared to each of these. All comparisons were confirmed using Tucker congruence (rex*rem>0.99 for all components in all cases. Wavelengths below 270 nm were excluded due to high leverage on models that led to noisy loading spectra and for ready comparison to the PARAFAC models presented elsewhere (Murphy et al., 2018). The full data set of EEMs from all degradation experiments was then projected onto the 4-component model derived from SRNOM PPL. This allowed standardization of the fluorescence signal loss we wished to model. Fluorescence intensity at the maximum of each component (Fmax) was normalized to the second data point in each degradation experiment time series, as the first points (collected immediately before lamp exposure) were often outliers with aberrant residuals after modelling fluorescence losses (e.g. Eq. 2 and 3).

Previous studies (Murphy et al. 2018, Del Vecchio and Blough 2002) used a bi-exponential model to describe fluorescence loss during photo-exposure as described in Eq. 2:

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$$f_t = f_L e^{-k_L t} + f_{SL} e^{-k_{SL} t}$$
 (2)

where f₁, total fluorescence normalized to the first EEM collected after the solar simulator lamp shutter opened at time t, is the sum of two fluorescence fractions (fi. and fsi.) undergoing decay at different rates (ki. and ksi.) (Murphy et al., 2018; Timko et al., 2015).

240 We modified Eq. 2 to replace time t with cumulative photon dose, assuming lamp photon output is constant throughout each experiment. If it can be properly measured, using cumulative photon exposure instead of time as the independent variable in models of fluorescence loss may allow better comparison of parameters between experiments, researchers, and experimental setups. The model is given in Eq. 3:

$$f_P = f_L e^{-k_L P} + f_{SL} e^{-k_{SL} P} \tag{3}$$

- where f_P is total normalized fluorescence after cumulative photon exposure P (in moles of photons). Other variables are the same as in Eq. 2. Photon dose estimations from nitrite actinometry can be applied to DOM irradiated under the same conditions if those conditions allow for optically thin solutions during exposure. The 1 mm pathlength spiral exposure cell we used should ensure optical thinness even in highly absorbent DOM solutions.
- 250 Results from fitting Eq. 3 are reported as four separate parameters: f_L, k_L, f_{SL}, and k_{SL}. However, f_L and f_{SL} are not independent as they should always sum to 1. They are expressed separately in our results because we believe these f values may be useful for understanding the compositional bases of degradation differences despite the difficulties for interpretation this dependence presents, and because each f value was fitted separately, so modelled fits not always sum exactly to 1.
- 255 R software (v. 3.6.0) was used to fit bi-exponential models using the nlsLM function from the minpack.lm package, and R was also used for significance testing and plotting most results.

3 Results and discussion

3.1 Method optimization and reproducibility

3.1.1 PARAFAC model

260 Our results confirm many of the findings reported by Murphy et al. (2018) in that the fitted PARAFAC model of SRNOM PPL photodegradations produced similar components despite the independent data collection and analysis by different researchers (Fig. 1). Emission maxima for components 1 to 4 were 439, 412, 525, and 452 nm; however, only components 3 and 4 followed the bi-exponential decay pattern. Figure 2 shows an example of fluorescence change in each PARAFAC

component during photodegradation of SRNOM PPL. Component 3 in this study corresponds with F₅₂₀ in Murphy et al., 2018, while Component 4 corresponds to the F450. Matching component spectra to models in the online OpenFluor database confirmed these matches, with Tucker congruence r values over 0.98 for emission spectra for both components. The weaker match between component 4 in this study and F450 in Murphy et al. is driven by differences in the excitation spectra (r = 0.949), but strong correlation between all 4 components in our PARAFAC model and higher information density in low wavelength ranges of excitation spectra could interfere with excitation spectral signal discrimination. Components 1 and 2 in this study 270 did not exhibit bi-exponential decay during photodegradation. In most experiments Component 1 decayed but did not follow a bi-exponential pattern, while Component 2 showed little net change. Differences in PARAFAC component matches and behavior between this study and Murphy et al. (2018) could arise from operating at a different pH (3 here vs. their minimum pH of 4). For example, despite spectral differences, Component 1 behaves similarly to F₄₂₀ in Murphy et al. (2018), which showed less rapid initial decay and a more linear overall pattern as pH decreased from 8 to 4 (see Fig. S4 in Murphy et al., 2018). Further results will focus on components 3 and 4 as they are most sensitive to photodegradation.

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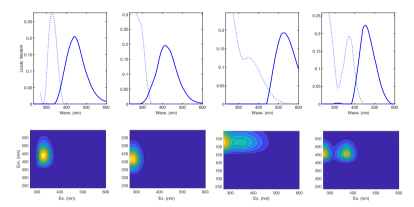


Figure 1. Spectral loadings and contour plots of PARAFAC components modeled from EEMs of SRNOM PPL extract photodegradation time series. The full dataset of all degradation time series EEMs was projected onto this model.

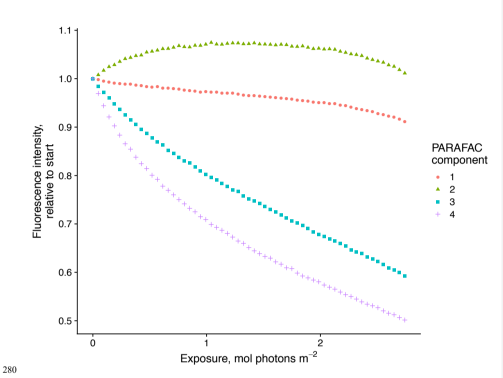


Figure 2. Example of fluorescence change in PARAFAC components during photodegradation. Data show degradation of SRNOM PPL.

3.1.2 Fluorescence loss model fit and utility of model parameter estimates

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Differences in biexponential model parameters between samples may allow reproducible comparisons of natural DOM photosensitivity. This approach has been used before given the excellent fit of this type of model to photodegradation data sets, and biexponential models indeed provided excellent fits to fluorescence losses in PARAFAC components 3 and 4 in our data sets (see Fig. 10 below for an example of fit). The biexponential model represents the sum of two terms, often referred to as labile and semi-labile to reflect the large relative differences in exponential slopes (k_L and k_{SL} in Eq. 2). This model captures loss of 2 pools of fluorescence intensity, possibly arising from 2 pools of DOM fluorophores decreasing in abundance at differing rates, or perhaps a single pool of photoreactive DOM with differing capacities for 2 types of reactions contributing to loss of fluorescence (Murphy et al., 2018). Potential interpretation of these parameter values is discussed in section 3.3.

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3.1.3 SRNOM experiments – experimental conditions and photon dose

Photodegradation kinetics in SRNOM trials were sensitive to many experimental conditions, but most importantly those that affected cumulative photon exposure. Total volume of sample in the system affected degradation kinetics by altering the cumulative photon exposure relative to the abundance of optically active molecules. Figure 3 shows loss of absorbance at 254 nm and loss of fluorescence intensity of components 3 and 4 relative to starting values in experiments where total volume of sample varied. Sample volume predictably affects photon dose relative to the quantity of starting material, because in all trials a fixed volume of the total volume is exposed to light at any time before returning to the mixing vessel. We found that flow rates from 1.5 to 8 mL per minute did not impact photon dose. Expressing loss of absorbance and fluorescence as a function of estimated photon exposure rather than a function of time seems necessary to ensure comparability with other experimental systems, and we will follow this convention where possible.

However, the reader is reminded that actinometers do have limitations (e.g. broadband response measurement) and caveats exist for their successful interpretation. Because CDOM absorption spectra generally increase exponentially with decreasing wavelengths, many experimental designs may violate the requirement that samples are optically thin when irradiated (Hu et al. 2002). The irradiation cell used here has a depth of 1 mm, which should prevent self-shading during photo-exposure at all concentrations tested. Previous work using this system showed that fluorescence loss was independent of SRNOM concentrations between 25 and 100 mg L⁻¹ (Timko et al. 2015). Concentration dependence in photochemistry is often assumed to stem from self-shading alone, and past work has shown the importance of working with "optically thin" solutions or properly correcting for inner filter effects when measuring photochemical behavior. All solutions shown here were considered optically thin at 300 nm and greater wavelengths following the convention that for optically thin solutions,

$$A_T \times L \ll 1 \tag{4}$$

where A_T is total (Napierian) absorption coefficient and L is path length in m (Hu et al., 2002). Although inner-filter corrections can be applied to correct for self-shading in spectrophotometer cells with known geometry (Hu et al. 2002), these corrections cannot be easily applied in other irradiation designs (e.g. vials on their sides and spiral flow cells). The definition for optically thin solutions (Eq. 4) is somewhat vague, so we also tested the dependence of DOM concentration on photodegradation rates.

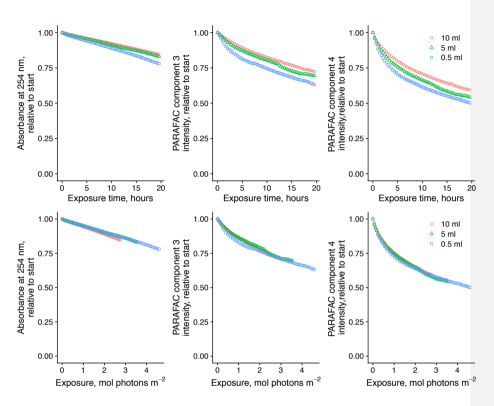


Figure 3. Photodegradation time series of absorbance at 254 nm and fluorescence intensities of PARAFAC component 3 and 4 relative to starting values. Data are shown from experiments with SRNOM PPL that varied volume of sample added to mixing reactor (after filling flow cell lines). Top panels show values as a function of exposure time, while bottom panels show values as a function of cumulative photon exposure calculated from NO_2/NO_3 actinometry.

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Degradation patterns seemed to be sensitive to DOM concentration as well but the effects were less clear (Fig. 4). In general, lower concentrations showed greater overall losses of absorbance and fluorescence. For the two most dilute solutions, PARAFAC C3 loss could not be modeled with a bi-exponential model, in contrast to all other samples throughout our study. Our results suggest either that our solutions experienced self-shading despite meeting the conventional definition of optical thinness, or some other mechanism links CDOM concentration to absorbance or fluorescence degradation kinetics such as

concentration-dependent charge transfer interactions (Sharpless and Blough, 2014). Further work is needed to explain these findings.

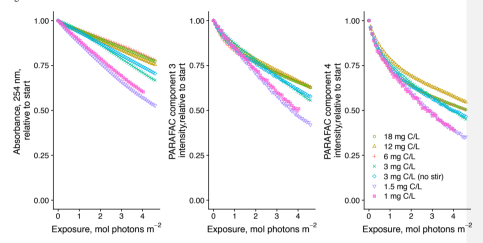


Figure 4. Photodegradation time series of absorbance at 254 nm and fluorescence intensities of PARAFAC component 3 and 4 relative to starting values. Data are shown from experiments with SRNOM PPL that varied approximate DOC concentrations. In all experiments 0.5 ml SRNOM PPL solution was added to mixing reactor after filling flow lines.

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Two researchers followed the same protocols with the same material (SRNOM PPL) as a test of reproducibility due to sample handling. Agreement between researchers was good and results varied to a similar degree as repeated tests by the same researcher (Fig. 5). Two-tailed t-tests were not able to distinguish differences in means between trials run by each researcher for any biexponential model parameters (p-values all greater than 0.10).

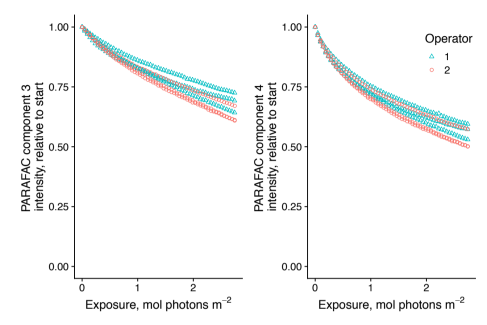


Figure 5. Photodegradation time series of PARAFAC component 3 and 4 fluorescence intensity, relative to starting values. Data are shown from experiments using SRNOM PPL performed by 2 of the authors to test reproducibility of results.

3.1.4 Effects of solid-phase extraction

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Fluorescence degradation from reconstituted RO SRNOM and SRNOM PPL extracts generated the same PARAFAC components. However, the overall loss of modeled components 3 and 4 differed between SRNOM PPL extracts and RO SRNOM, as did kinetics of fluorescence loss (Fig. 6). The differences in fluorescence loss were small but systematic. Two-tailed t-tests of relative fluorescence loss suggested differences between PPL and RO SRNOM in PARAFAC component 4 (p-value < 0.01) with limited support for differences in component 3 (p-value = 0.06) and no support for differences in absorbance loss (p-value = 0.3 for 254 nm). Projecting the data onto a PARAFAC model built from RO SRNOM degradation data instead of SRNOM PPL data did not affect these results. Fitted model parameters from Eq. 3 suggest these differences stem from the kinetics of the semi-labile fluorescence pool, with possible differences in the relative starting abundances of the labile vs. semi-labile pools (Fig. 7 and Table A1). Rate constants of the labile pool did not vary for either PARAFAC component, suggesting extraction did not affect behavior of this pool, so studies focusing on this pool should not be affected by PPL extraction. Capturing changes in this pool is one of the explicit advantages of our experimental system, and future work on environmental

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photo-reactivity may focus on this time scale as photochemical reactions in the environment are often driven by initial rates (Powers and Miller, 2015). However, slower degradation processes or longer irradiations may be affected by extraction.

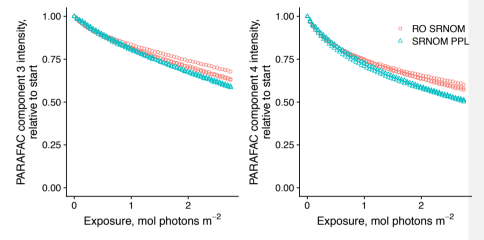


Figure 6. Photodegradation time series of PARAFAC component 3 and 4 fluorescence intensity, relative to starting values. Data are shown from 3 replicates of both RO SRNOM and SRNOM PPL.

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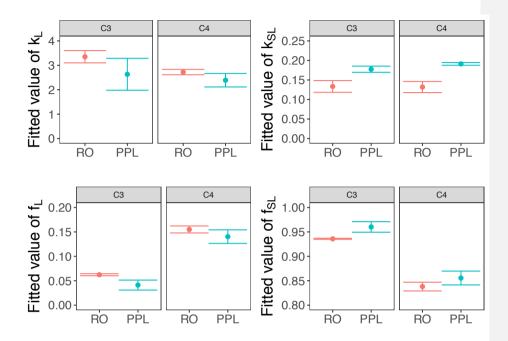


Figure 7. Fitted biexponential model parameters (Eq. 3) from the time series of loss of PARAFAC components 3 and 4 in irradiation experiments comparing RO SRNOM to PPL SRNOM (see Fig. 6 for data). f is unitless and k is m^2 [mol photons]⁻¹. C3 and C4 denote PARAFAC components 3 and 4. Error bars represent mean \pm standard deviation from three experiments. Two-tailed t-tests suggest differences in ks_L for both components (p = 0.020 in component 3, p = 0.015 in component 4), while f_L and f_{SL} may differ (p = 0.065 and 0.058) in component 3.

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Shared PARAFAC components suggest PPL extraction did not strongly alter the compositional bases of fluorescence photosensitivity in the RO SRNOM, but the differences in losses suggest researchers should take care when comparing extracts to original samples in future photodegradation kinetics studies. We are not sure what gave rise to these differences, but the RO SRNOM likely contains much more highly polar compounds such as (poly)saccharides and related compounds (e.g. glycosates). Differences between PPL and RO samples here are probably not due to variation in photon dose, as volume and initial absorbance were equal across samples. If concentration of fluorophores affects degradation kinetics, differing fluorophore concentrations between our PPL extracts and whole SRNOM could explain the discrepancy. Even though we adjusted all samples to similar starting absorbance, selective enrichment or dilution of absorbing or fluorescing compounds in extracts could affect the mechanism responsible for any concentration dependence. Differences in electronic coupling and

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Moved down [1]: 1. Fitted biexponential model parameters (Eq. 3) for comparison between RO and PPL SRNOM, p-values are from two-saided t-test of difference in means; n = 3 for both RO SRNOM and SRNOM PPL. fis unitless and k is

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390 charge-transfer abilities (Del Vecchio and Blough, 2004; Sharpless and Blough, 2014) could arise in extracts and affect fluorescence degradation kinetics. RO SRNOM may present matrix effects relative to extracted SRNOM PPL, as metals and other possible interferences are still present (albeit at much lower concentrations relative to DOC than in source water) despite the cation exchange and desalting treatments that accompanied the original reverse osmosis isolation (Kuhn et al., 2014).

Preliminary experiments showed photochemical behavior of filtered whole water (before extraction) was affected by cold storage duration, precluding reproducible experiments on samples collected at different times. Unstable behavior was observed over time in whole water wetland samples with high DOC concentrations (15-40 mg/L) in experiments run without dilution using a 3x3 mm flow cell in the spectrofluorometer (Fig. 8). While DOM absorbance seems stable in seawater samples after storage at 4° C up to 1 year (Swan et al., 2009), concentrated DOM in inland waters may be unstable in cold storage conditions, affecting its optical properties or responses to photoirradiation. Further work is required to understand the cause of this behavior, but losses of DOC and changes to optical properties during cold storage of samples have been reported elsewhere (Peacock et al., 2015). High DOC concentrations may also promote flocculation (von Wachenfeldt and Tranvik, 2008), which is known to specifically involve CDOM (Wachenfeldt et al., 2009). DOM or matrix composition may also affect storage stability. Differing degrees of instability between sources (e.g. in Fig. 8 large wetland sample becomes noisier than small wetland) suggest differences in DOM chemical composition may affect sensitivity to storage. With high concentrations of

stability. Differing degrees of instability between sources (e.g. in Fig. 8 large wetland sample becomes noisier than small wetland) suggest differences in DOM chemical composition may affect sensitivity to storage. With high concentrations of DOM, reproducible use of whole water samples instead of extracts may be possible if irradiation experiments are conducted within a week of sample collection, or if samples are diluted prior to storage, but this will require further investigation and would assume no major differences in the sample matrix.

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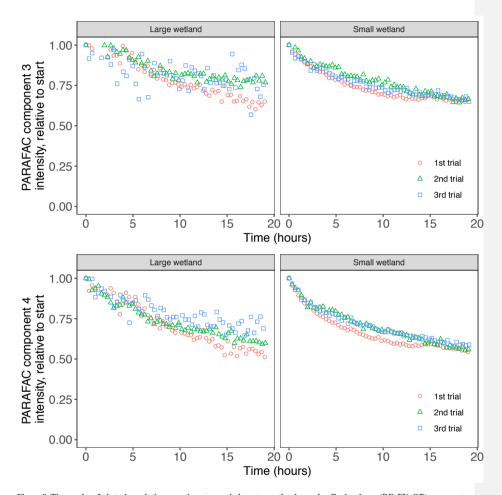


Figure 8. Time series of photodegradation experiments on whole water wetland samples. Each column (DF, FN, QB) represents samples from a different wetland source. Three experiments were run with aliquots drawn from a water sample from each wetland, and results seemed to change with storage time. First experiments with each wetland water source were run 5-8 days after sample collection, second experiments were run 9-13 days after sample collection, and third experiments were run 14-16 days after collection. The trend in most cases toward lower relative photosensitivity in measured variables and in some cases increasing data noise as samples aged informed the decision to use solid phase extracts to improve reproducibility.

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3.1.5 Guidelines for photodegradation fluorescence kinetics experiments

It has been established that initial pH and pH change during photodegradation affects fluorescence photodegradation kinetics (Timko et al., 2015). We chose to conduct experiments at pH 3 because control by autotitration was not possible during these experiments due to contamination from the pH probe, and starting at pH 3 ensured minimal pH change during photodegradation. If research goals do not explicitly include understanding effects of pH during photodegradation, we recommend bringing all samples to the same starting pH and controlling pH during the course of photodegradation experiments, or starting experiments at pH 3 and ensuring change during the experiment is minimal.

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Using a reference material allows consistency within and between research labs. We recommend using SRNOM as it has been widely studied and characterized (Green et al., 2014). Comparing total absorbance and fluorescence loss and degradation kinetics of SRNOM to DOM sources of interest will allow more meaningful comparison between lab groups. Repeated experiments with the same standard can identify sources of error and quantify variability due to experimental procedures. Checking this variability against variability among repeated measurements of a sample may allow common variability to be estimated and thus reduce the need for replication in future runs with similar DOM sources. We also used SRNOM (after solid phase extraction) as the basis for our PARAFAC model of fluorescence change during photodegradation and projected this model onto the rest of our data set, standardizing fluorescence losses between DOM sources to the same signal.

435 For research into compositional changes in DOM during photodegradation, test materials should be brought to similar starting absorbance. We adjusted all samples to a raw absorbance of 0.12 at 300 nm (with a 1 cm path length), but this may be difficult or less ecologically meaningful with naturally dilute (e.g. ocean) or concentrated (e.g. leachates) DOM sources. If possible,

on photochemical rates. In our system,

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Photon dose obviously affects degradation kinetics. Our experimental system offered several procedural choices that could affect photon dose, including volume of sample in the system and lamp intensity. Researchers should carefully control these parameters and ensure their procedures are generating reproducible results by running several replicated experiments with a reference material. We encourage repeating this process with multiple individuals within a lab to understand the impact of individual methodological choices on results (e.g. gravimetric measurement of volume added vs. pipetting, preparation of samples). We strongly encourage at least reporting actinometry results or assumed actinometry for the experimental conditions used in order to better compare photon doses across studies and in the environment. While the additional work of actinometry is not trivial, we believe this represents one way to improve reproducibility of degradation kinetics that avoids the limitations of using time alone. Even this approach could be improved – our actinometer did not directly measure radiation across the UV spectrum, which could allow more accurate quantification of cumulative photon dose. Striking a balance between effort

testing different DOM concentrations for the same sample is recommended in order to establish any concentration dependence

required and reproducibility is difficult, but we believe our work illustrates some of the limitations of conventional approaches where photon exposure cannot be reliably calculated, and we hope our efforts inspire alternative approaches to overcoming these limitations. Ideally, samples should be irradiated under optically thin conditions when actinometry measurements or other approaches can be used to estimate photon doses for kinetic modelling (e.g. using Eq. 3 instead of 2).

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Photodegradation is affected by both DOM composition and matrix conditions. While we found that the same PARAFAC model captured fluorescence decay in both SRNOM and solid phase extracts of SRNOM (as in (Murphy et al., 2018)), extraction did affect total fluorescence loss and its kinetics. Storage of water samples for greater than two weeks led to changes in fluorescence loss patterns, even when filtered to 0.2 µm (Fig. 8). We expect this may be due to the high DOC concentrations used in those experiments, as these may be more susceptible to flocculation or other aggregation processes than dilute samples, but further work would be required to test this. We recommend using extracts with greater storage stability to allow comparison over time, unless all experiments can be conducted shortly after sample collection or previous experience shows that the optical properties of the DOM in question are stable for the duration of storage. Comparisons of kinetics between extracts and whole water samples should be made with care, but experiments using such comparisons may help disentangle the role of DOM chemical composition from other matrix effects in determining photodegradation behavior and sensitivity. Matrix effects may be especially important for extrapolating lab photodegradation findings to inferences at ecosystem scales. For example, if the approach described here is used to investigate longitudinal changes in DOM photosensitivity along a river network, tying these findings to residence times and photon doses in the field would be difficult without considering light attenuation by inorganic chromophores and particles. Matrix constituents may also fundamentally alter the photosensitivity of DOM by participating in charge-transfer processes. We recommend using DOM isolated from its matrix by extraction here not because it is a sufficient approach to understand these phenomena, but as a foundation to explore this complexity. More work is needed to understand the relative influence of DOM and matrix compositions on photodegradation kinetics.

3.2 Photosensitivity differences between DOM sources

We compared several DOM sources in order to see whether high resolution fluorescence time series could reveal differences in photosensitivity between sources. Fig. 9 shows the degradation of PARAFAC components 3 and 4 relative to starting intensities in samples from different DOM sources. Both components showed potentially divergent decay patterns among DOM sources, with Sargassum leachate starkly diverging from bulk DOM sources. Fitted biexponential model parameters of decay in PARAFAC components 3 and 4 are shown in Tables A2 and A3, with parameters from component 4 plotted in Fig.
10 (similar plot for component 3 can be found in Appendix A, Fig. A1). We did not conduct repeated trials with every DOM source shown here due to logistical constraints, but t-tests on three trials each with SRNOM and one of the wetland samples supported potential differences in f_L and f_{SL} in both PARAFAC components, and possible differences in k_{SL} in component 3. Notably, these two DOM sources had biexponential parameter values that were among the most similar compared to other

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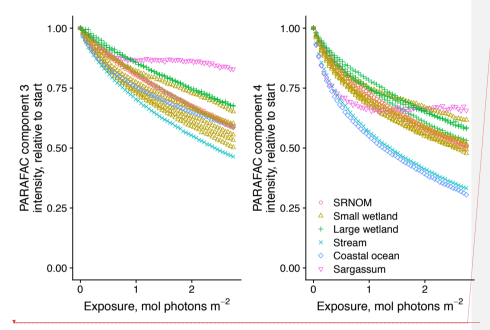
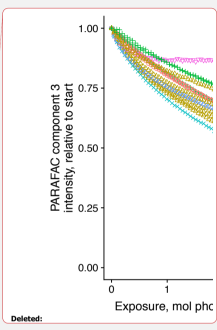


Figure 9. Photodegradation time series of PARAFAC component 3 and 4 fluorescence intensity, relative to starting values. Data are shown from experiments using PPL extracts from different DOM sources (see Methods for source descriptions). "Large wetland" and "Small wetland" samples use the same symbol for samples from each source, including samples collected on different dates.



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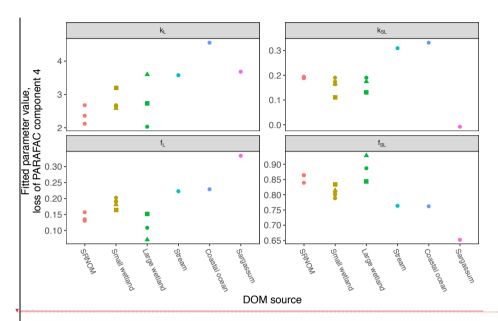
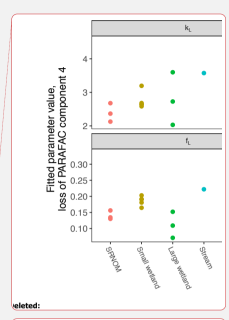


Figure 10. Fitted biexponential model parameters (Eq. 3) from the time series of PARAFAC component 4 (see Fig. 2 for data). f is unitless and k is m² [mol photons]-1. For wetland samples, shapes represent different sampling dates (circles are 2017-10-04, triangles are 2017-12-20, and squares are 2018-04-01).

The outlier in our comparison of DOM sources was *Sargassum* leachate extract, which was expected given the unique composition and the presence of phlorotannins (Powers et al., 2019). The natural DOM used in a previous study (Murphy et al., 2018) that yielded PARAFAC components appearing in all photodegradation experiments did not include leachates, only natural bulk DOM. Interestingly, this sample alone showed little or very slow semi-labile fluorescence loss with total fluorescence loss of projected PARAFAC components 3 and 4 dominated by rapid initial loss. Future studies using leaf or soil/sediment leachates, or lysed algal cells, or other putative sources of natural DOM instead of bulk natural DOM itself need to test this modelling approach more thoroughly to ensure it is appropriate, but using other leachate sources may highlight the compositional basis of the semi-labile fluorescence decay that seems ubiquitous in bulk natural DOM but absent in *Sargassum* leachate here.



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Moved down [5]: Where n > 1, means are shown with standard deviations in parentheses. SRNOM PPL trials are three experiments using the same sample source, small wetland trials include three trials with the same sample source (to test non-SRNOM system stability) and two trials with samples from other dates, and large wetland trials include one trial each from three sampling dates. p-values of biexponential model fits all below 1x10*6 except for ks. for Sargassum, p = 0.016. Model parameters for every individual trial can be found in associated data set (Armstrong, 2020).¶

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530 3.3 Photosensitivity and ecological inference

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3.3.1 Interpreting biexponential model parameters

In other studies (e.g. Murphy et al., 2018; Timko et al., 2015) the rate parameters k_L and k_{SL} have received the most attention, as different average rates of change in fluorescence governed by these rate constants may indicate differences in DOM chemical composition, matrix composition, environmental conditions (if experiments are performed in situ), or experimental conditions, making these values potentially useful metrics of compositional differences between DOM sources. However, differences in loss of fluorescence between samples may also arise from differing relative abundances of two "pools" of at the beginning of the time series. Figure 11 shows degradation time series from two experiments, along with fitted model parameters. These experiments compare DOM sampled in October 2017 from the two freshwater wetlands in Maryland. Figure 11 shows loss of PARAFAC component 3 (see Fig. A2 in Appendix A for a similar plot showing loss of component 4). The model fits are shown against the data in upper panels, while the modelled fits for each of the two terms from Eq. 3 $(f_L e^{-k_L P})$ and $f_{cl}e^{-k_{SL}P}$ are plotted separately against the data in lower panels. This visualization is useful to weigh the contribution of differing rate parameters (kL and ksL) against the relative abundance of their respective fractions (fL and fsL) at the onset of the experiment in determining overall differences in photodegradation behavior between samples. Component 3 loss models show similar k_L values but different relative fractions of the "fast" pool of fluorescence loss at the start of the experiment. Differences in these starting fractions between samples may play a role in overall differences in degradation kinetics in component 4 as well. It is crucial to note that the chemical interpretation of these modelled fits is not clear. "Pools" of fluorescence in different relative abundances that decay at different rates may not map directly onto different groups of fluorophores. This behavior may stem from differences in the capacity for two classes of photochemical reactions - where k describes the reaction rates and f describes the relative capacity of the sample to undergo the corresponding reaction at the outset of the experiment. Further work is needed to understand what gives rise to relative differences between f terms in different samples, though as noted f_L and f_{SL} are not independent in the model presented here. This highlights one of the strengths of our approach – the ability to capture optical properties of DOM that change very quickly during photodegradation. The modelled labile portion of fluorescence contributes negligibly to total fluorescence after receiving between 0.5 and 1.2 moles of photons per square meter, (3-10 hours of irradiation with our experimental setup). Future work relating the photon dose required to reach this point and the environmental conditions affecting this dose in natural DOM could improve knowledge of DOM origins, residence times, and interactions with other degradation processes.

3.3.2 Linking photosensitivity to DOM sources in dynamic ecosystems

High resolution photodegradation experiments of natural DOM can reveal fundamental photophysical behavior of ecological importance. We believe the approach described here can help unravel sources or light exposure histories of DOM in natural settings. One of our overall goals is to determine relative photosensitivity among samples. The biexponential models that fit experimental photodegradation data may help with these comparisons. For example, in the two wetland samples compared in

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Fig. 11, distinct patterns of photodegradation suggest distinct DOM composition. DOM fluorescence in the larger wetland had relatively less "fast" decaying fluorescence in photosensitive PARAFAC components (parameter ft.) than the smaller wetland, These wetlands are depressions located less than 100 m from each other, but with isolated surface water during the October 2017 sampling. They differ in basin size, canopy cover, and vegetation communities. Our data and fitted model parameters suggest that DOM in the larger wetland has either previously been exposed to sunlight that has depleted the potential for "fast" decaying fluorescence, or that differences in source material or other processing of DOM pools in each wetland have given 570 rise to relatively less photosensitive material in the larger wetland. In winter, water levels rose in each depression, and eventually both depressions were connected by surface flow from the larger to smaller wetland. Photosensitivity differences show DOM composition and reactivity are affected by these phenomena. Figure 12 compares biexponential model parameters in samples from each wetland depression taken in October 2017, December 2017, and April 2018. This is an especially dynamic period in the seasonal cycles that affect DOM in this area - the October sampling is just before deciduous leaves senesce and fall, and the December sampling occurred less than a month before rising surface water levels connected the two depressions. Figure 12 shows that we may be able to capture the effects of ecosystem phenomena on DOM sensitivity. kL values for both PARAFAC components do not show any obvious pattern, while k_{SL} values are very similar at each sampling site for all three dates but may be changing between dates due to some shift in DOM composition over time affecting both sites. The most obvious pattern is in fL and fSL. These differ between sites in October and December, suggesting that despite their proximity, conditions at these sites differ enough to affect DOM photosensitivity in their surface water. The larger depression has less of the faster-decaying fluorescence, either due to differences in the source of the material on the landscape or depletion relative to the smaller depression reflecting greater light exposure and natural degradation. These differences are homogenized in April, when surface water mixing (and shorter residence times in surface storage due to export) means sitespecific processes are less influential in shaping DOM composition.

These photosensitivity differences may have consequences for other ecosystem processes. For example, if low f_L at the time of sampling reflects high rates of photodegradation in wetland surface water, photopriming may contribute to microbial heterotrophy. Or wetland DOM with high f_L may influence downstream ecosystems, if DOM exported to stream networks is then susceptible to photodegradation which alters its lability to heterotrophs (Judd et al., 2007) or promotes flocculation (Helms et al., 2013). The sensitivity of our approach may also allow revisiting questions of longitudinal dynamics of light exposure in stream systems (Larson et al., 2007).

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Photodegradation of DOM extracts in the lab does not replicate in situ photodegradation of DOM in surface waters. However, in situ photodegradation of DOM in surface water is extremely convoluted – the complexity of DOM chemical composition, surface water matrix composition, simultaneous ecological processes that also alter DOM composition, and the natural dynamism of surface water systems are intertwined and make it difficult to understand the role of photodegradation of DOM in surface water ecosystems. Our approach represents one step in the direction of disentangling this story, but leaves many

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questions unanswered. We demonstrated several sources of potential variability in degradation kinetics that require more attention, any of which may affect our understanding of different influences on in situ photodegradation and its ecological consequences. Further research is required to understand how differences in DOM composition alone (as isolated in our work with extracts) interact with matrix composition (Grebel et al., 2009; Poulin et al., 2014; Timko et al., 2015; T. Stirchak et al., 2019), and how these reactivity differences affect other DOM transformation processes (Amado et al., 2015; Chen and Jaffé, 2016; Lønborg et al., 2016) and ecosystem- or macrosystem-scale or biogeochemistry (Anderson et al., 2019; Pickard et al., 2017; Rutledge et al., 2010).

This example is not conclusive for these sites but is presented to illustrate the possible uses of our method. Clearly much more research is needed to explain the observed differences in photodegradation kinetics between these two wetlands and test these hypotheses, ideally with more detailed data on DOM composition associated with differing photosensitivity. Regardless, our approach can complement established techniques for describing DOM such as bulk optical properties, ultrahigh resolution mass spectrometry, or nuclear magnetic resonance, and could be combined with other experimental approaches probing natural DOM sources and transformations.

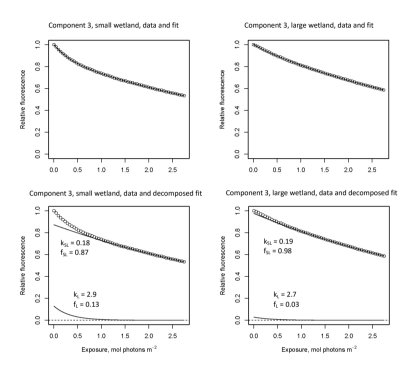
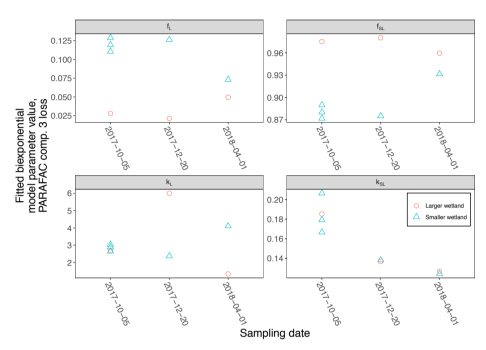


Figure 11. Data and model fit of PARAFAC component 3 loss in experiments with two wetland samples. Top panels show data and model fit (Eq. 3) while bottom panels decompose the fitted model into its two summed terms, $f_L e^{-k_L P}$ and $f_{SL} e^{-k_{SL} P}$, or labile and semi-labile terms.



620 Figure 12. Fitted biexponential model parameters (Eq. 3) from the time series of PARAFAC component 3, comparing DOM from large and small wetland sampling sites collected on different dates. f is unitless and k is m² [mol photons].

4 Conclusion

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Photodegradation experiments have improved our understanding of the role of DOM light sensitivity in ecological processes. As researchers continue to explore related questions and experiments proliferate, it is important to use approaches that constrain the influence of experimental conditions and promise reproducible or at least comparable results. Our method allows reproducible and relatively short experiments that capture photosensitivity differences between varying sources of natural DOM on time scales relevant for investigating degradation processes in the environment. This approach can be used to ensure experiments conducted at different times or by different researchers can be compared. Our work illustrates several obstacles to reproducing and comparing studies of photodegradation kinetics, highlights underappreciated sources of uncertainty, and offers an approach that improves upon past methodological limitations. It also captures distinct fast dynamics that differ

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Appendix A: Additional tables and figures

Table A1. Fitted biexponential model parameters (Eq. 3) for comparison between RO and PPL SRNOM, p-values are from two-sided t-test of difference in means; n = 3 for both RO SRNOM and SRNOM PPL, f is unitless and k is m² [mol photons]⁻¹.

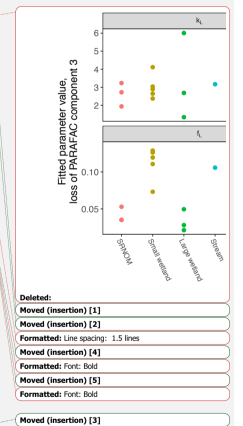
PARAFAC component	Biexponential parameter	RO SRNOM Mean (SD)	SRNOM PPL Mean (SD)	t-test p-value
<u>3</u>	$\underline{\mathbf{k}_{\mathrm{L}}}$	3.35 (0.252)	2.63 (0.654)	<u>0.18</u>
<u>3</u>	<u>fr</u>	0.0623 (0.00207)	0.0411 (0.0102)	0.065
<u>3</u>	$\underline{\mathbf{k}_{\mathrm{SL}}}$	0.133 (0.0150)	0.177 (0.00792)	0.02
<u>3</u>	$\underline{f_{\mathrm{SL}}}$	0.936 (0.00113)	0.960 (0.0107)	0.058
<u>4</u>	<u>k</u> _	2.72 (0.110)	2.39 (0.276)	0.16
<u>4</u>	<u>fr</u>	0.155 (0.00717)	0.140 (0.0139)	<u>0.2</u>
<u>4</u>	$\underline{kS_L}$	0.132 (0.143)	0.191 (0.00346)	0.015
<u>4</u>	<u>fS</u> L	0.838 (0.00897)	0.856 (0.0142)	0.16

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Table A2 Fitted biexponential model parameters (Eq. 3) for different DOM sources. f is unitless and k is m^2 [mol photons] Where $n \ge 1$, means are shown with standard deviations in parentheses. SRNOM PPL trials are three experiments using the same sample source, small wetland trials include three trials with the same sample source (to test non-SRNOM system stability) and two trials with samples from other dates, and large wetland trials include one trial each from three sampling dates, p-values of biexponential model fits all below 1×10^6 except for $k_{\rm SD}$ for Sargassum, p = 0.016. Model parameters for every individual trial can be found in associated data set (Armstrong, 2020).

PARAFAC component	Biexponential parameter	$\frac{SRNOM\ PPL}{(n=3)}$	$\frac{Stream}{(n=1)}$	$\frac{\text{Coastal ocean}}{(n=1)}$	$\frac{Sargassum}{(n=1)}$
<u>3</u>	<u>k</u> _	2.63 (0.654)	3.17	<u>2.84</u>	<u>5.4</u>
<u>3</u>	$\underline{\mathbf{f}_{L}}$	0.0411 (0.0102)	0.106	<u>0.135</u>	<u>0.11</u>
<u>3</u>	<u>k</u> sl	0.177 (0.00792)	0.237	0.137	0.0194
<u>3</u>	$\underline{\mathbf{f}_{\mathrm{SL}}}$	0.960 (0.0107)	0.887	0.857	0.889
<u>4</u>	$\underline{\mathbf{k}_{\mathrm{L}}}$	2.39 (0.276)	3.58	4.55	3.68
<u>4</u>	<u>fL</u>	0.140 (0.0139)	0.222	0.229	0.334
<u>4</u>	$\underline{\mathbf{k}_{\mathtt{SL}}}$	0.191 (0.00346)	0.308	0.332	-0.00803
<u>4</u>	<u>fS</u> L	0.856 (0.0142)	0.764	0.762	0.653



<u>Table A3</u>. Fitted biexponential model parameters (Eq. 3) for wetland DOM samples. f is unitless and k is m^2 [mol photons]⁻¹. Where $n \ge 1$, means are shown with standard deviations in parentheses, p-values of biexponential model fits all below 1×10^{-6} . Model parameters for every individual trial can be found in associated data set (Armstrong, 2020).

655

PARAFAC component	Biexponential parameter	$\frac{\text{Small wetland.}}{2017-10-05}$ $\frac{(n=3)}{}$	$\frac{\text{Small wetland,}}{2017-12-20}$ $\frac{(n=1)}{}$	$\frac{\text{Small wetland,}}{2018-04-01}$ $\frac{(n=1)}{}$	$\frac{\text{Large wetland,}}{2017-10-05}$ $\frac{(n=1)}{}$	$\frac{\text{Large wetland,}}{2017-12-20}$ $\frac{(n=1)}{}$	$\frac{\text{Large wetland.}}{2018-04-01}$ $\frac{(n=1)}{}$
<u>3</u>	<u>k</u> L	2.86 (0.199)	<u>2.37</u>	<u>4.12</u>	<u>2.69</u>	<u>6.01</u>	1.34
<u>3</u>	<u>f</u> _	0.120 (0.00908)	0.126	0.0732	0.0281	0.214	<u>0.0496</u>
<u>3</u>	$\underline{\mathbf{k}}_{\mathrm{SL}}$	0.184 (0.0203)	0.138	0.124	0.185	0.137	<u>0.127</u>
<u>3</u>	<u>fsl</u>	0.881 (0.00906)	0.875	0.932	0.975	0.980	0.960
<u>4</u>	<u>k</u> L	2.65 (0.0255)	<u>2.58</u>	3.20	<u>2.03</u>	3.60	<u>2.73</u>
<u>4</u>	<u>fL</u>	0.195 (0.00698)	0.181	0.165	0.109	0.0711	0.152
4	$\underline{\mathbf{k}}_{\mathrm{SL}}$	0.176 (0.0126)	0.165	0.110	0.190	0.175	0.132
4	<u>fS</u> L	0.798 (0.00820)	0.814	0.833	0.887	0.928	0.844

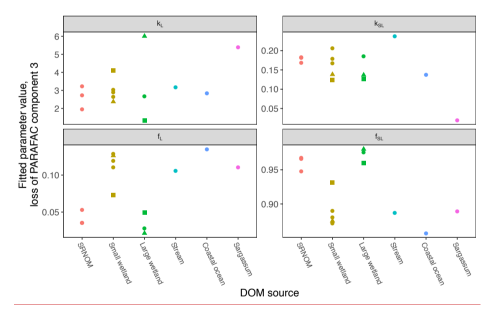


Figure A1. Fitted biexponential model parameters (Eq. 3) from the time series of PARAFAC component 3 (see Fig. $\underline{9}$ for data). f is unitless and k is \underline{m}^2 [mol photons] 1. For wetland samples, shapes represent different sampling dates (circles are 2017-10-04, triangles are 2017-12-20, and squares are 2018-04-01).

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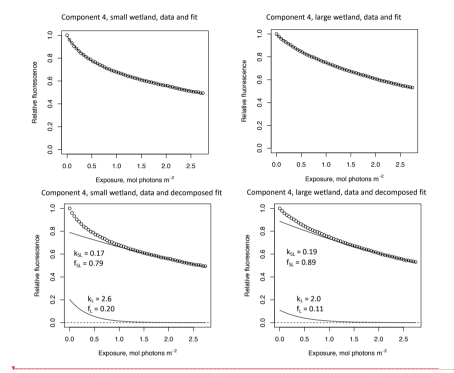
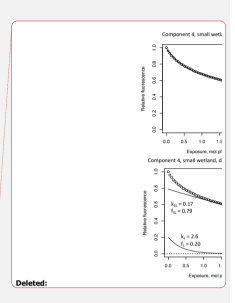


Figure A2. Data and model fit of PARAFAC component 4 loss in experiments with two wetland samples. Top panels show data and model fit (Eq. 3) while bottom panels decompose the fitted model into its two summed terms, $f_L e^{-k_L P}$ and $f_{SL} e^{-k_{SL} P}$, or labile and semi-labile terms.



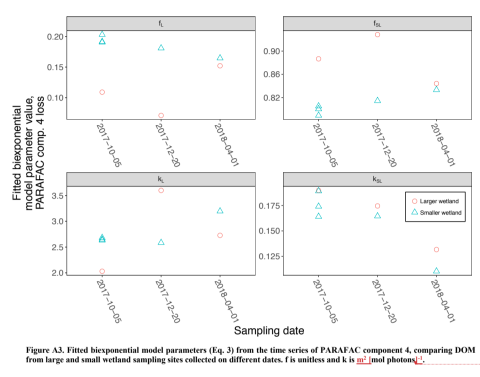


Figure A3. Fitted biexponential model parameters (Eq. 3) from the time series of PARAFAC component 4, comparing DOM from large and small wetland sampling sites collected on different dates. f is unitless and k is m^2 [mol photons] $^{-1}$.

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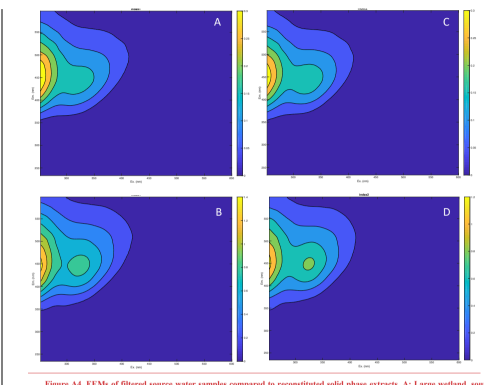


Figure A4. EEMs of filtered source water samples compared to reconstituted solid phase extracts. A: Large wetland, source water. B: Large wetland, solid phase extract. C: Small wetland, source water. D: Small wetland, solid phase extract. Figure shows samples originally collected 2017-10-05.

Data and code availability

670

Data and code used in this analysis are available from the Dryad repository at https://doi.org/10.5061/dryad.hmgqnk9d9 (Armstrong, 2020).

Author contribution

AA developed the method's applications for ecological inference, collected the data, analyzed the data, and drafted the manuscript. LP assisted in the method's conception, collected the data, assisted with data analysis, and edited the manuscript.

MG conceived the method, designed and optimized the instrument system, assisted with data analysis, and edited the manuscript.

Competing interests

The authors declare they have no conflict of interest.

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