

Interactive comment

Interactive comment on "New insights into mechanisms of sunlight-mediated high-temperature accelerated diurnal production-degradation of fluorescent DOM in lake waters" by Yijun Liu et al.

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Interactive comment on "New insights into mechanisms of sunlight-mediated high-temperature accelerated diurnal production-degradation of <code>incurecont DOM</code> in lake waters" by Yijun Liu et al. Dear Prof. Dr. Koji Suzuki, Associate Editor, Biogeosciences: Anonymous Referee 1 Received and published: 2 June 2020. We are very grateful to Review1 for the valuable and constructive comments on our manuscript. We are submitting the manuscript and Figures revised according to the Reviewer comments. We have considered duly all Reviewer comments, providing more examples, which could

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contribute to better understanding our FDOM research. Thank you.

Itemized responses (R) to Reviewer comments below: Note: all line numbers refer to the revised manuscript. Major comments The authors measured diurnal changes in CDOM components, DOC concentrations, and nutrients concentrations in two small lakes in Tianjin University, China to identify the biogeochemical processes controlling the diurnal DOM variation which is feasibly related to global warming. I think the research topics described in the manuscript would be of great interests to readers in Biogeosciences. However, I also think that the manuscript is not clearly written and difinAcult to follow, not technically sound and not appropriately discussed in the context of previous literature. Some of in Agures are not clear. Please see comments listed below (1) The authors ran PARAFAC modeling to determine in Cuorescent components for each time period and compare the components among time periods. First of all, since there is no description regarding how the authors determined the number of components and validated PARAFAC models. I cannot evaluate whether the conclusion derived from PARAFAC are scientiin Acally/technically sound or not. My opinion from our experiences of PARAFAC modeling is that it is not reasonable to apply PARAFAC for a dataset comprising the small number of samples (n < 20). It seems that the authors used small data set for PARAFAC modeling (lines197-200; I cannot understand the sentence though...). While, since the validity depends on the dataset for PARAFAC modeling, the authors should describe the validation method to identify the number of PARAFAC components and show the results of the validation. R-1. "Number of Samples". Our research group has provided ample previous evidence that only two EEM spectra from two samples are enough to identify the authentic PARAFAC components for freshwater and seawater samples (Mostofa et al., 2019; Mostofa et al., 2018-conference paper). In the 2019-EST paper and earlier studies, i.e. Mostofa et al. 2010; Mostofa et al. 2013, we have clearly assessed that "two freshwater samples, six inshore seawater samples and 12 offshore seawater samples are sufficient to run the PARAFAC model". My research group is doing FDOM research since 1999 and I have presented EEM data as a Keynote speaker in the Conference on Organic Matter

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Spectroscopy 2018 (WOMS18), held on 23-27 October 2018, in Carqueiranne City, France. Website: http://woms18.univ-tln.fr/moftofa-abstract/. In that presentation the number of samples issue, along with many other issues have been discussed providing substantial evidence. Three examples are provided below for further supporting the number of samples used in our PARAFAC model. (i) The PARAFAC model applied to upstream water (Namitaki, site-KR 1) of Kurose River was fitted by only one component (terrestrial humic-like substances) in three cases, i.e. all ten individual EEM data sets on samples from May 2002 to Feb 2003 (Fig. 1a), two EEM data sets on summer (May and July) samples (Fig. 1b) and two EEM data sets on winter (January and February) samples (Fig. 1c). Details about sampling can be found in Mostofa KMG et al. (2005) Geochemical J 39: 257-271.

(ii) The PARAFAC model applied to sewerage-impacted downstream water (Izumi, site-KR 5) of Kurose River was fitted two components in three cases, i.e. all 12 individual EEM data sets on samples from May 2002 to Feb 2003 (Fig. 2a-b), two EEM data sets on summer (May and July) samples (Fig. 2c-d), and two EEM data sets on winter (January and February) samples (Fig. 2e-f). Details about sampling can be found in Mostofa KMG et al. (2005) Geochemical J 39: 257-271. (iii) The PARAFAC model applied to inshore seawater of Seto Inland Sea was fitted by three components in two cases, i.e. all 12 individual EEM data sets (Fig. 3a-b-c) and six EEM data sets (Fig. 3d-e-f). Details about sampling can be found in Mostofa KMG et al. (2019) Environ Sci and Technol 53: 561-563.

R-1. 'Components and Model Validation'. First of all, we wish to escribe our knowledge of DOM and FDOM compositions in diverse surface waters. Without understanding of DOM or FDOM composition, without which we cannot dicuss about our views on "components and model validation'. FDOM are primarily originated from three key sources. First, the terrestrial source, i.e. FDOM derived from soil/land, which is predominantly detected in streams and rivers, and is largely decomposed in lakes/reservoirs/ponds/oceans (Coble, 1996; Mostofa et al. 2013; Mostofa et al.

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2019). It is well known that extracted soil humic substances are composed of three key components that include humic acids (HA), fulvic acids (FA) and protein-like fluorofores (PLF), with soil PLF entirely different from those detected in surface waters (Fig. 4: below- soil HA (a), FA (b) and PLF (c) of forest soil origin in Mohinuzzaman et al., 2020).

Second, the autochthonous source, i.e. FDOM originated from phytoplankton, which is mostly detected in stagnant waters such as lakes/reservoirs/ponds (Zhang et al., 2009; Parlanti et al. 2000; Mostofa et al. 2013; Yijun et al.- this study). However, some rivers/streams in winter dry season can present stagnant water areas where phytoplankton can grow and produce autochthonous FDOM. As phytoplankton cannot grow in streams or upstream rivers where water is only released from groundwater/terrestrial land, these waters do not contain autochthonous source-FDOM. It is well known that the production of microorganisms/phytoplankton needs time (several days) and stagnant waters for their growth under sunlight conditions. Therefore, stream/upstream river FDOM can only be of terrestrial source and not autochthonous source. Third, an anthropogenic source or sewerage-derived FDOM (e.g. detergent-like components), is detected only in untreated sewerage-impacted rivers in specific site (Baker et al. 2001; Mostofa et al. 2005; Mostofa et al. 2010). Fourth, another important issue is that seawater FDOM is strongly affected by salinity, pH and photochemical degradation, which are completely different from those of freshwater FDOM. For example, salinity can cause the peak C of terrestrial humic-like substances to shift at longer wavelength region (red-shifted phenomena), together with other changes of other FDOM components with respect to river freshwaters (Coble 1996; Yamashita and Zaffé, 2008; Yamashita et al. 2010-Deep-Sea Res-II, 57, 1478-1485; Mostofa et al. 2010; Mostofa et al. 2013; Mostofa et al. 2019). Fifth, in this study (Yijun et al. 2020; see Figure 5) and other studies (Zhang et al., 2009; Mostofa et al. 2013; Parlanti et al. 2000). Terrestrial humic acid and terrestrial humic-like substances (C type) show fluorescence properties (peak positions and EEM images) similar to those of autochthonous humiclike substances (C type) of phytoplankton origin The two autochthonous and terrestrial

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sources of FDOM components cannot be distinguished by PARAFAC analysis when all diverse samples are mixed together. This is the reason why one cannot distinguish between these two autochthonous sources.

Sixth, in this and other studies (Yijun et al. 2020; see Figure 6; Zhang et al., 2009; Mostofa et al. 2013; Shammi et al. 2017) terrestrial fulvic acid and terrestrial humic-like substances (M type) (Mostofa et al. 2019; Mostofa et al. 2013; Mohinuzzaman et al. 2020) show fluorescence properties (peak positions and EEM images) similar to autochthonous humic-like substances (M type) of phytoplankton origin). The two autochthonous (Fig. 6a-d) and terrestrial (Fig. 6e-g) sources of FDOM components cannot be distinguished by PARAFAC analysis when all diverse samples are mixed together. This is the reason why one cannot distinguish between these two autochthonous sources.

Seventh, untreated sewerage-affected river could transport detergent-like substances and standard household detergents into river waters (see Fig. 7a-c; Mostofa et al. 2013; Mostofa et al. 2005; Baker et al. 2001).

Based on all seven different types of FDOM sources, it is very important to choose "Selective Characteristic Samples" that can represent authentic FDOM components (Mostofa et al. 2019), whereas mixing all seven types of FDOM sources together in a PARAFAC model cannot evaluate authentic FDOM. Stedmon et al. (2003) firstly used the PARAFAC model on a total of 90 samples covering all major terrestrial sources of DOM, streams and estuary DOM mixed with seawater from the adjacent Kattegat, which resulted in a five component model. Since the study of Stedmon et al. (2003), the use of mixed samples in the PARAFAC model has been repeated in several other studies to date (Murphy et al., 2013; Kulkarni et al., 2017; Wu et al., 2018; Yue et al., 2019). However, artifact due to mixing. Mostofa et al. (2019) provided evidence that these components were artifacts due to mixing. "Model Validation". Stedmon et al. (2003) and Murphy et al. (2013) have used the split-half analysis method for validation of the PARAFAC components. This method consists in dividing the EEM data set into

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orescent components does not correspond to the respective peak position appearing

outside the usual DOM fluorescence (i.e. Rayleigh and water Raman peak positions), then the model should be rechecked for 3 components. Differently, if all components are verified with those of standard substances or specific field/experimental observations, the model and the respective fluorescent components are validated. Finally, the fluorescent components obtained from the model can be characterized and classified (Yamashita and Tanoue, 2003; Sheng and Yu, 2006; Zhang et al. 2009; Mostofa et al., 2013; Shammi et al. 2017a, 2017b, 2017c; Mostofa et al., 2019)." in lines 206-225. We believe that it is important to find out the various sources of FDOM which can be related

to biogeochemical facts and their significances. To clarify the principles on which the PARAFAC model is based, we have also added an additional paragraph "The parallel factor (PARAFAC) analysis is a three-way multivariate method that can be applied to EEM data of DOM to decompose them into trilinear components (Harshman. 1970; Caroll and Chang, 1970). For any fluorophore of FDOM, the emission intensity, xjk, at a specific wavelength, i, which corresponds to excitation at the wavelength, k, can be

expressed by the following equation (Harshman. 1970): wherea is the concentration (in arbitrary units, a.u.) of the analyte (fluorophore or FDOM), bj is the relative emission at the wavelength j, and ck is the relative amount of light absorbed at the excitation wavelength k. For any number of analytes and samples, the PARAFAC model can be expressed as a set of trilinear terms and a residual array

as (Stedmon et al., 2003):

where xijk is the fluorescence intensity of the ith sample at the emission wavelength j and excitation wavelength k, aif is directly proportional to the concentration (in a.u.) of the analyte f in sample i. bif is directly proportional to the quantum efficiency of fluorescence of the analyte f at the emission wavelength j, ckf is linearly related to the specific absorption coefficient at the excitation wavelength k, F is the number of components in the model, and ε ijk is the residual matrix that indicates the variability not accounted for by the model." in lines 182-198.

(2) The authors concluded that some components were disappeared over a 24-h diur-

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33.5°C), but not in lower temperature months). (3) The authors described temporal

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tochemical and microbial processes into low molecular weight (LMW)-non-fluorescent

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discussed this issue based on the reviewer2 comments in the new addition "4.4. High-

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temperature production of FDOM from phytoplankton and its sequential entire degradation at 24-h diurnal scale The complete degradation of FDOM after its formation from phytoplankton can be ascribed to the gradual increase in SI (from 1.36 to 2.97 MJ/m2 for a total of 15.26 MJ/m2) at the highest WT (30.2-33.7 °C) and AT (35.4-41.8 °C) (Fig. 2). These environmental conditions are directly responsible of the rapid production of FDOM from phytoplankton and its corresponding rapid degradation by day-time sunlight-induced and night-time microbially-induced processes. Such entire sequential diurnal degradation of FDOM was not observed in October, May and June samples (Figs. 3-5) when solar intensity, AT and WT were relatively low compared to July samples (Fig. 2). As discussed previously, the production and degradation of FDOM are related to simultaneous significant fluctuations in the contents of NO3-, NH4+, NO2-, DON, PO43-, DSi and DOC (Fig. 6; Table S2). Photochemical and microbial processes can transform EPS into various forms of FDOM (Shammi et al., 2017a, 2017b) that are related to field observations, which show that different components of EPS vary over different timescales and temperatures (Shammi et al., 2017c; Sheng and Yu. 2006). Many studies provided evidence that photochemical and microbial degradation processes increase with increasing temperature and light intensity (Matsumoto et al., 2007; Weston and Joye, 2005; Malinverno and Martinez, 2015; Whelan and Rhew, 2015; McKay and Rosario-Ortiz, 2015; Grannas et al., 2006; Farias et al., 2007). In turn, these results would indicate that high SI, WT and AT would accelerate the complete transformation of autochthonous FDOM into LMW DOM and other mineralization end-products on a 24-h diurnal cycle, which could be further increased by the influence of future GW. Such changes can to be reasonably expected to occur in the future on the basis of increasing water temperature, extended summer season and increased water stratification in response to the predicted GW from 1.5 to 2.0°C (Huisman et al., 2006; Watanabe et al., 2011; Rogelj et al., 2019)."

(5) It seems that the authors generally consider/discuss degradation of EPS to explain daytime changes in PARAFAC components. Why don't the authors consider DOM production including the PARAFAC components with primary production during daytime?

R-5. DOM is well known to comprise many types of organic substances, which include EPS that can gradually convert into various FDOM components, non-fluorescent photoproducts that can also be produced from FDOM, non-fluorescent microbial products, non-fluorescent LMW substances (aldehydes, ketones), etc. Besides terrestrial sources, autochthonous DOM is a key fraction of DOM compositions in lakes and reservoirs, but it is still uncertain how it is released from phytoplankton. Our research group has gained a wide experience on EPS extraction from large volume of surface waters, and its photo-microbial products, which is the key to better understand the relation between FDOM and DOM. We believe that results of this study will contribute to un-

derstand the sequential release of FDOM from EPS and their subsequent degradation (Figs. 8-9; this study). As commonly reported in all other studies, in this manuscript

DOM is discussed as DOC changes.

Other comments I listed some other comments below. Similar errors with some of the comments were found throughout the manuscript. Lines 69-72: What is "key DOM components"? It's not clear. R. The sentence has been revised according to this comment with addition of "Key DOM components include terrestrial humic substances (fulvic and humic acids), EPS, and aromatic amino acids (Coble, 1996; Yamashina and Tanoue, 2003; Yamashina and Tanoue, 2004; Shammi et al. 2017a, 2017b, 2017c; Zhang et al., 2009). Lines 99-101: I could not understand the message of the sentence. The authors would like to mention "diurnal degradation processes depended on the photosynthetic activity of primary producers"? R: Yes. The sentence has been rearranged according to this comment as "Diurnal day-time (sunlight) and night-time (microbial) degradation processes are a natural phenomenon that depends on the photosynthetic activity of primary producers in surface waters which is ultimately related to daily biogeochemical changes of C, N and P cycling".

Lines 102-105: "a signiīň Acant decrease of its ĩň Cuorescence intensity with increasing water depth" is due to photoinduced degradation? I don't agree with it. R. Yes, we agree. The sentence has been revised properly in lines 108-109. Line 105: Which

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occur during the gradual increase of SI". R. All references in line 261 have been deleted

from the mentioned sentences "FDOM production by degradation of EPS would occur during the gradual increase of SI (from 1.36 to 2.97 MJ/m2 for a total = 15.26 MJ/m2) at WT of 30.2-33.7 $^{\circ}$ C and AT of 35.4-41.8 $^{\circ}$ C (Fig. 2)."

Lines 245-247: I could not follow how the authors estimate the abundance of EPS quantitatively. R. The text has been revised as "..... In particular, the decrease in fluorescence intensity of EPS in this time period was approximately 3Line 254: The citation of Ma and Green (2004) is not appropriate for the sentence, because their work was carried out Lake Superior. R. Reference deleted in line 288-289 and revised according to our own results with addition of "(b) night-time extended microbial degradation, which was also supported by the net decrease of DOC along with nutrients at night period, which will be comprehensively discussed in the next section.". Line 415: What is "photoinduced respiration of phytoplankton"? In addition, it seems that the authors discuss the photosynthesis rather than respiration at the rest of this paragraph. R. The text has been revised as "Further evidence of photosynthetic activities of phytoplankton was provided by the significant shifting of the δ 15N value of NO3-." . Lines 439-465: I think the authors do not discuss "A global view of production and degradation pathways of FDOM in lake waters" in this paragraph. R. For reviewer's kind information. Our group published previously on EPS extraction at large scales using 40 liters of surface waters and on the subsequent photo-microbial experiments that can sequentially release FDOM from EPS (Shammi et al. 2017a, 2017b, 2017c). The results of our current study, as well as some unpublished data, appear to confirm all previous results on FDOM, EPS and their byproducts. Although it is till now unclear how autochthonous FDOM originates from phytoplankton, the results of this study demonstrated that EPS are produced in the morning and then convert into many autochthonous FDOM components that are ubiquitously detected in surface waters, as previously reported in experimental observations (Zhang et al., 2009). The paragraph in question attempts to provide a conceptual model based on our research findings by describing the sequential release of FDOM, which ultimately describes dynamics of DOM that originates from phytoplankton. This global view contributes to further under-

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standing autochthonous FDOM and its degradation processes. Line 435: I could not in all in al

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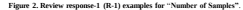
Figure 1. Review response-1 (R-1) examples for "Number of Samples".

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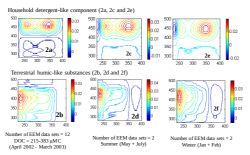
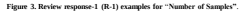


Fig. 2. Figure 2. Review response-1 (R-1) examples for "Number of Samples".

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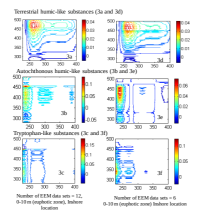


Fig. 3. Figure 3. Review response-1 (R-1) examples for "Number of Samples".

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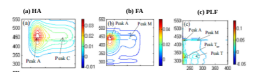
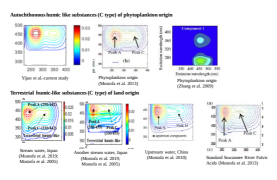


Fig. 4. Figure 4. Author response (R-1) for 'Components and Model Validation'.

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Fig. 5. Figure 5. Author response (R-1) for 'Components and Model Validation'.

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Figure 6. Author response (R-1) for 'Components and Model Validation'. 250 300 350 400 Origin from EPS under sunlight conditions Origin from lake phytoplankton (Oct, Jingya lake) (Shammi et al. 2017-Scientific Report) after resuspenssions in Milli-Q incubation (Zhang et al. 2009) water under dark incubation (Mostofa et al. 2013) Terrestrial humic-like substances (M type) of land origin (e-g) Upstream water, site KR-2, Kurose River, Japan (Mostofa et al. 2019; Soil fulvic acid after water extraction from forest soil (Mohinuzzaman et al. 2020) from Yellow River, China (Mostofa et al. 2013)

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Mostofa et al. 2005)

Figure 7. Author response (R-1) for 'Components and Model Validation'.

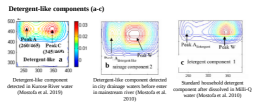
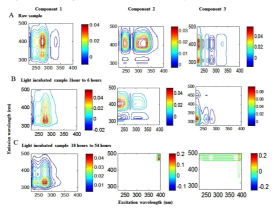


Fig. 7. Figure 7. Author response (R-1) for 'Components and Model Validation'.

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Fig. 8. Figure 8. Author response (R-2) for review comments by Reviewer#1

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Figure 9. Author response for specific comment by reviewer#1



Fig. 9. Figure 9. Author response for specific comment by reviewer#1

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Figure 10. Author response for specific comment by reviewer#1

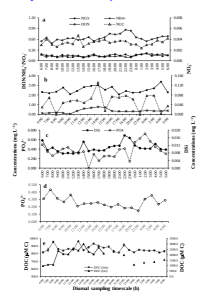


Fig. 10. Figure 10. Author response for specific comment by reviewer#1

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