We would like to thank Dr. Dileep Kumar for looking over this work and providing valuable critiques to our paper. The comments are thoughtful and bring up many important points, which we addressed individually below.

## Abstract

Line 9: 'Bubbles adsorb and transport particulate matter both in industrial and marine systems' – include lakes systems here.

We agree that this opening sentence falls short of encompassing the full scope of the impact of bubbles. Industrial and marine systems have been the primary focus of the research, but this phenomenon applies more broadly, and certainly applies to lakes as we show in the manuscript. We adjusted the opening sentence to provide a broader significance to the work.

## "Bubbles adsorb and transport particulate matter in a variety of natural and engineered settings, including industrial, freshwater, and marine systems."

Line 9-12: 'methane-containing bubbles emitted from anoxic sediments are found extensively in aquatic ecosystems' – the word "extensively" is inappropriate for marine systems since methane- containing bubbles can only be found in a few select coastal ecosystems. However, this issue assumes greater and global significance in vertical transportation of dissolved and particulate materials scavenged across a few meters below to sea surface by the rising wind-induced bubbles, particularly in shallow marginal systems.

Thank you for this comment. We changed the sentence to reflect that methane containing bubbles would be a particular concern in freshwater systems such as lakes, reservoirs, and wetlands ("are found widely in freshwater ecosystems"), and also broadened the first sentence to bring in more of the global significance of bubblemediated transport (above).

## Introduction

Lines 34-35: 'Metals can be mobilized from sediments via solubilization by oxidation reduction reactions, and by sediment resuspension or bioturbation' – mobilization can also occur through acidification of lakes.

Thank you for bringing this omission to our attention. We have added acidification to the list of mechanisms.

"mobilized from sediments via solubilization by oxidation-reduction reactions, and by sediment resuspension, acidification, or bioturbation (Calmano et al., 1993;Eggleton and Thomas, 2004;Schaller, 2014;Schindler et al., 1980)."

Line 35-36: 'transport to surface waters of contaminants mobilized from the sediment is affected by lake hydrodynamic conditions, notably stratification' – an interesting question to the current investigation be how does stratification influence

methane bubble rising to surface during minimal wind induced turbulent conditions? A strongly stratified upper water column will inhibit (slow down speed of rising) or even prevent particularly small sized but proportionately with large surface areas from rising across the strong pycnocline. This is possible if vertical profiling is done with close intervals of sampling to find density gradients across the pycnocline and assessing the bubble rise rates in hypo- and epilimnion layers.

The impact of stratification on bubble rise is an interesting question to both methaneemission from lakes and bacterial or chemical transport that should be addressed in future work. If stratification does prevent small bubbles from penetrating, bubbles may be an additional mechanism concentrating organisms or chemicals at these interfaces, resulting in the thin layers of organisms that can often be seen within the water column. However, the changes in density across naturally occurring pycnoclines might be gradual enough and the bubbles buoyant enough pass through it without bursting or accumulating. The proposed vertical profiling experiment would make a very good follow-up study.

## We have also added a discussion of these questions

"However, many questions remain regarding bubble-mediated transport in natural systems, including how the change in water density at the thermocline affects bubble rise and associated chemical and biological material."

Lines 40-41: 'Verspagen et. al. (2005) showed that recruitment from sediments of the potentially toxic cyanobacterium Microcystis was a major driver of the summer bloom Verspagen et al., 2005)' – referenced twice in the same sentence!

*Thank you for pointing out this redundancy. It has been changed to "Previous research showed .."* 

Lines 66-67: 'the full extent of bubble particle flotation in aquatic systems remains unknown.' – even the present manuscript cannot make it 'full', which requires many attempts by many investigators!

We agree we cannot hope to determine the full extent of bubble particle flotation with this study, and have removed full from this sentence.

Line 70: Fig. S1 should show pictures before and after the bubble event to highlighting the emergence of particles following the bubbling.

We have changed Figure S1 to show the water surface near the beginning of a bubble triggering event, as well as at the end. This highlights the visible accumulation of particulate matter on the water surface.



**Figure S1.** Picture of the lake surface near the beginning (A) and end (B) of a triggered bubble event at 15 m depth showing an accumulation of particulate matter (visible as light specks on the water surface).

Lines 78-80: 'Given the expected importance of both bubble size and total bubble volume, we used a bubble size sensor (Delwiche et al., 2015; Delwiche and Hemond, 2017) to measure bubble diameter distributions both in the lake and in the laboratory.'– adsorption or scavenging of particles by bubbles is expected to be proportional to the surface area of the bubble (similar to metal adsorption on to a particle) and therefore representing bubble characteristic in terms of 'surface area' than its 'size' or 'volume' would be preferable.

The surface area is an important bubble characteristic for transport along with other key characteristics, but typically the bubble diameter is provided as a key metric of bubble characteristics. We have rephrased this as. "Given the expected importance of bubble size on key characteristics (e.g. surface area, buoyancy, diffusion of gas), we used a bubble size sensor (Delwiche et al., 2015;Delwiche and Hemond, 2017) to measure bubble diameter distribution both in the lake and in the laboratory."

Methods Lines 101 and 250: 'another lake' – please name the lakes.

# The lakes referenced as "other lakes" in the text are Lake Scharmützelsee and Lake Limmaren, which have been explicitly stated in the text.

Lines 111-112: 'All bubbles rising through the bubble size sensor or collection funnel entered the flexible tubing and rose into the sample cup.' – as the particles and the associated substances are adsorptive in nature it is likely that some of the rising bubble attached particles are adsorbed in the flexible tubing etc. before they reached sample cup. Authors may include a statement on this possible loss of particles during sample processing.

The reviewer makes a valid point, transported particles were indeed adsorptive and some stuck to the sample tubing. We have added a statement to introduce this possible sample processing artifact. "The interaction of bubbles with the flexible tubing resulted in visible particle attachment to the tubing, making our estimates of particle mass transport a lower bound"

Line 117: Word 'approx.' may not be necessary as the coordinates are specified to third decimal.

This was removed.

Lines 119-120: 'preventing mixing from of the sediment to the surface.' – requires rephrasing.

We have rephrased as "preventing mixing of sediment to the surface"

Lines 121-124: Good strategy.

We found sediment contamination within the collection cups that were deployed for much longer periods capturing natural bubble events, but the possibility of contamination and subsequent growth, death or decay of transported cells made it impossible to have reliable estimates from this method. While there are some issues with this approach, it made the measurements of particle transport feasible.

Line 250: Please correct the flux units 'cells m-2' to cells m-2 d-1.

This error was corrected.

**Results and Discussion** 

Lines 262-263: 'demonstrate that bubbles transport particles from depths of at least 15 m to the lake surface.' – It may be revised as "demonstrate that bubbles transport particles from depths to the lake surface" since bubbles if formed even in deeper waters can transport materials to surface.

This statement is confusing, and was revised to "Both field and bubble column experiments demonstrate that bubbles can transport particles from the sediment to the lake surface."

Line 307: Lines 134-135 mention 'On 26 June 2018 we sampled for cyanobacteria bubble transport using similar procedures, except we used a simple inverted funnel instead of a custom bubble size sensor to intercept rising bubbles' whereas Fig. S7 caption shows "Frequency distribution numbers are approximate because the bubble size sensor is unable to measure fast bubble flux or very small bubbles" – It is important to check the compatibility between these statements, particularly for data of 26 June 2018 if used.

The data from Fig. S8 is only from the column experiments, not collected during the 26 June 2018 sampling. Fig S7 shows the frequency of bubble diameter naturally occurring (from previous work) and triggered (this work, not for the 26 June 2018 date) only collected during the October 2017 sampling event, where cyanobacteria were not measured.

To clarify the difference between Fig. S7 and Fig. S8, and to further clarify the figures themselves, we have updated the figure captions to read:

**Figure S7**. Frequency of occurrence of bubble diameter during triggered (purple) and natural (gray) events in Upper Mystic Lake. Mean (black lines) and standard deviation (shaded regions) for each event type. Bubbles were triggered by dropping an anchor multiple times during the October 2017 sampling event, while natural bubble size distribution are based on continuous measurements from the summer of 2015 and 2016 (Delwiche and Hemond, 2017). Frequency distribution numbers for the triggered bubbles are approximate because the bubble size sensor is unable to measure the rapid bubble flux that sometimes occurred with anchor-triggered bubble events.

**Figure S8.** Frequency of bubble diameter observed across multiple trials (a-k) during the cyanobacteria experiment in the laboratory bubble column. Some trials had a bimodal diameter distribution. Panels f and k represent trials where air was bubbled directly above the sediment, and remaining panels represent trials where air was bubbled into the sediment. Note the different y-axis scales.

Line 317: replace ug with \_g. *This mistake was corrected.* 

Lines 342-343: Besides 'a significant fraction of the arsenic input to epilimnetic waters can be attributed to inflow from the Aberjona River (Hemond, 1995)' aerial transport of dust associated arsenic/metals should be invoked here to be among the unknown inputs.

We have not properly accounted for all other forms of input of metals to the lake by restricting the next sentence to just surface water input, so we have adjusted that to include atmospheric deposition with "However, bubble-mediated fluxes of arsenic or other sediment-borne metals may represent a larger fraction of epilimnetic input in other lakes having lower influx rates from surface water inflow or other external sources, such as atmospheric deposition (Csavina et al., 2012)."

Lines 361-362: 'Bubble-transported particulate matter contained cells at a rate of approximately 30 cells mL-1 gas, indicating that bubbles are capable of transporting cyanobacteria through' – May be revised as "Bubble-transported particulate matter contained cells at approximately 30 cells mL-1 gas, indicating that bubbles are capable of transporting cyanobacteria through". A 'rate' is expected to be material transferred during a specific duration (time). \*\*\*

Thank you for identifying this error. It has been changed as suggested.

We would like to thank the referee for looking over this work and providing valuable critiques to our paper. The comments are thoughtful and bring up many important points, which we addressed individually below.

## Anonymous Referee #1

Authors state that the particles associated with the bubbles, almost entirely originated from the sediments, rather than from the water. Will this statement hold true in case of turbid waters? Please clarify.

We do not actually know whether the sediment particles have been scavenged from the plume of sediment in the water column, or the sediment directly. We have evidence to suggest that only a small portion (~10%) seem to originate in the water column from the column experiments, but the concentration of particles in the water column could have been different between experimental conditions in the column and field. As such, more turbid waters could result in larger concentrations originating from the water column as compared to the sediment, but further work is needed to understand this difference.

We were also not clear about the water column conditions when we conducted our tests for particle scavenging in the experimental column. Because these tests were done after tests where bubbles were emitted from the sediment bed, the water column was visibly turbid and contained many suspended particles. We have added two sentences to clarify this point:

(In Methods) "Scavenging tests were conducted after particle transport tests, so the water column above the sediment bed was turbid and contained a plume of sediment particles. "

(In Results) "We conducted the scavenging tests when the water column was visibly turbid and contained a plume of suspended particles from previous tests."

## We also add this as a possible mechanism in section 3.1:

"These particle loadings on bubbles, and any ecosystem-wide flux estimates derived from them, must be qualified by the fact that neither triggered bubbles nor bubbles in the bubble column fully replicate natural bubbling. In particular, the triggering of bubbles with an anchor may have raised plumes of suspended sediment through which some fraction of produced bubbles had to rise, and within which the possibility of scavenging should be considered."

Add the details of dissolved oxygen concentration, temperature and total suspended matter in the water column at the lake sampling station.

We have added a figure (Fig. S3) showing the temperature profile taken during the June 26, 2018 sampling event. Previous work on Upper Mystic Lake has shown that

dissolved oxygen tracks closely with temperature (Delwiche and Hemond, 2017). We do not have a total suspended matter profile.



*Figure S3. Water temperature profile taken during June 16, 2018 sampling event on Upper Mystic Lake.* 

Delwiche, K. B., and Hemond, H. F.: Methane Bubble Size Distributions, Flux, and Dissolution in a Freshwater Lake, Environ Sci Technol, 51, 13733-13739, https://doi.org/10.1021/acs.est.7b04243, 2017.

Did you observe any bubble breakup during the transport through the flexible tubing? If yes, does it affect the final bubble size count and volume transported?

The bubble size sensor was placed below the sample cup set-up, which contained the flexible tubing, so any breakup within the tubing (which did occur) did not affect the measured size distribution. However, the size distribution could have been affected by rapid bubble flux, which can cause bubbles to coalesce within the funnel constriction leading to the bubble size sensor (as described in Delwiche et al, 2017). To address this fact, we have modified the text:

Anchor-triggered bubbles were significantly smaller (average diameter 5.6 mm) than those measured for natural bubbling events (average diameter 6.4 mm) during a 2016 field campaign [Fig. S7, (Delwiche and Hemond, 2017)], even though relatively high bubble flux events (such as those triggered by anchor dropping) can lead to some bubble coalescence within the funnel constriction in the bubble size sensor [ (as described previously (Delwiche and Hemond, 2017)].

Line 114, please add the grade of HNO3 used for rinsing.

We used reagent grade  $HNO_3$  for all acid washing, and have amended the text to reflect this:

"All sample cups were soaked in 5-10% reagent grade HNO3 for 24 hours..."

Authors dropped a cinderblock to trigger bubble release. Please state the difference in bubble volume during natural release and forced release.

This information is presented in section 3.2 Triggered bubbles are smaller than natural bubbles, but both are larger than 1 mm where differences between sizes decrease, making it unlikely that their difference in size should substantially change transport.

The impact of cinderblock on the lake floor would have re-suspended a significant amount of sediments. Does the forced release, thus suggest a much larger than natural bubble release mediated particle transport?

We agree with the reviewer that triggering a bubble release with an anchor drop suspends a significant amount of sediments. We also wondered if this suspended sediment would artificially raise the measured rates of bubble particle transport. To address this question, we conducted the particle scavenging experiments in the bubble column, as described in section 2.3. The scavenging tests were done when the water column had significant amounts of suspended sediment from previous trials. Bubbles passing through this sediment cloud had only around 10% of the particle mass from bubbles emitted from the sediment, indicating that while particle scavenging does occur, it is relatively minor. However, we agree that anchor dropping could still influence bubble mediated particle transport, and future research is needed to assess the particle transport rates for naturally occurring bubbles.

The collection of sediment by dredge and subsequent transport in bucket, would have resulted in the release of a significant amount of gas from the sediments. Can the authors provide the difference in the gas content of in-situ sediments and those collected by dredge and brought to the lab in a bucket?

The gas content of the sediment was not measured, but would certainly be lower once removed from the environment by the dredge and placed into the bucket. However, the gas content of the sediment was not critical to the development of bubbles in the experimental bubble chamber. We used a syringe pump to inject gas into the sediment bed. For this reason, we did not find it critical to measure the gas content of the sediments collected in the environment.

What was the percentage of bubbles breaking up, when striking the inverted funnel and releasing the cyanobacteria?

As the reviewer points out, there are a number of potential experimental artifacts that could decrease the measured amount of sediment and cyanobacteria transport (including particles adhering to the sampling apparatus, as discussed earlier and now included in the manuscript). However, we have not found that bubbles break up when encountering an inverted funnel. Previous work looking at potential bubble break-up when bubbles reach the bubble sensor funnel found instead that bubble coalescence can occur when bubble flux is high enough. This coalescence relates to the reviewer's previous comment on how bubbles break up could affect size measurements, so we encourage the reviewer to see that response.

Authors used air, instead of methane in the laboratory experiment. Will there be a difference in the particle transport by an air bubble as compared to methane bubble? Please discus in the text.

The composition of the air in the bubble was dramatically different between the experimental column and the field, given the origins of both gases. If the experiment was conducted at high pressure, such as in the deep ocean, this difference in gas composition in the bubble could reach a critical point where it could affect the bubbles and particle transport. However, at the pressures found within our system (both lake and column), the composition of gas is unlikely to influence bubble properties or particle transport.

In support of the conclusion above, using either air in the column or gas from the sediment resulted in a similar amount of particle transport per ml gas (" $0.01 \pm 0.006$  mg/mL in the bubble column, compared to  $0.01 \pm 0.01$  mg/mL on June 2018 in the field"). However, the differences between those amounts and the amounts measured in the field in October 2017 ( $0.09 \pm 0.07$  mg/mL) are substantial, so we do not fully understand all of the factors (potentially gas composition) that influence particle transport.

We also added some general caveats to this approach, which would include gas composition (e.g.):

" There remains the possibility that our measured bubble particle transport rates differ significantly from those from naturally emitted bubbles, and this remains an important area for future research."

"While this variability in cell transport between column measurements and estimates of potential field transport highlights the need for continued research, it is useful to estimate the potential range of cyanobacterial transport."

How did the authors decide the rate of injection of air into the sediments? What happened to the gases already present in the sediments when authors injected the air?

We have added the following text to the manuscript to clarify these points:

"The bubbling rate was calibrated to achieve a relatively steady release of bubbles without substantial wait time in between. While we expect that much of the gas naturally existing within the sediment was released during sediment collection and as it was transferred to the sample bed (indeed we did not observe natural bubble release from the sediment bed prior to experimental trials), remaining gas could have been incorporated in to rising bubbles."

Line 266, authors did not estimate the gas reserve in the sediments. How can they infer that the lower gas volume did not indicate a smaller gas reserve?

As you point out, we did not measure the gas reserve in the sediment, so we cannot speculate as to the cause of the lower gas volume in June 2018. We have re-framed the section to focus on the observations and avoid undue speculation: "Both field and bubble column experiments demonstrate that bubbles can transport particles from the sediment to the lake surface. A positive correlation (p < 0.05 level for October 2017 ( $r^2 = 0.76$ ), p=0.15 ( $r^2=0.38$ ) for June 2018 ) was found between total particle mass and gas volume in bubble traps for both field sampling campaigns (Fig. 1). The general magnitudes of particle loadings on bubbles in column experiments and on bubbles observed in triggered experiments in the field were of similar magnitude;  $0.01 \pm 0.006$  mg mL<sup>-1</sup> in the column vs  $0.09 \pm 0.07$  mg mL<sup>-1</sup> on October 2017 and  $0.01 \pm$ 0.01 mg mL<sup>-1</sup> on June 2018 in the field."

If the positing of boat influenced the bubble release, then how can they quantify the bubble volume and associated particle transport?

It was indeed a challenge to position the boat above the sample plume, particularly when winds blew us off course between anchor drop and bubbles reaching the surface. However, since we were interested in particle transport **per gas volume**, our results should not be affected by whether we captured all gas from a particular bubbling event.

We note that this sentence is now re-written in response to other comments, as mentioned above.

Line 273, I do not agree with the comparison of experimental column release with that from the natural lake environment. As stated above the conditions in the lab were completely different than that in the lake, and thus any comparison between the two is superfluous.

As any controlled environment will have many differences from the natural environment, we hope that you will agree that the experimental columns were within the range observed in the field, thus can be used to verify that cyanobacteria can move quickly on these bubbles. The bubble column work was necessary to test the importance of particle shedding and scavenging (something we could not test in the field), and the fact that bubble column particle transport was of similar magnitude to field results ( $0.01 \pm 0.006 \text{ mg/mL}$  in the bubble column versus  $0.09 \pm 0.07 \text{ mg/mL}$  and  $0.01 \pm 0.01 \text{ mg/mL}$  in the field) indicated that the bubble column results could inform field processes. However, to acknowledge the necessary differences between the controlled and natural environments, we have added the following text:

"Although this is significantly higher than the measurements made in the bubble column, the conditions in the column are substantially different from the conditions in the field and the sediments used in column had been stored for 8 months, so the cyanobacteria cell concentration was 10 times less than fresh sediments. While this variability in cell transport between column measurements and estimates of potential field transport highlights the need for continued research, it is useful to estimate the potential range of cyanobacterial transport."

Authors state a large difference in the size of natural and forced release of bubbles. Then what is the reliability of the volume and particle transport estimated by the authors?

There is a large amount of uncertainty in amount of particle mass transported per ml of bubble volume in our measurements, which was not properly emphasized before in the manuscript. The differences in bubble size could be one aspect of this uncertainty. In response to this comment and other referee comments, we have emphasized the uncertainty in the text and removed amounts of cells or arsenic transported from the abstract. Even with these large uncertainties, we can still put our results into context by saying that we expect that this type of transport might be small compared to other inputs for arsenic, but that bubble-mediated cell transport could be a substantial part of the life cycle of cyanobacteria in this lake. This provides contexts for what should be pursued in future experiments while still emphasizing the uncertainty in our measurements. We hope that this provides better insight into the reliability of these measurements.

Line 25, change 'Concentrations' to 'Concentration' Line 27, change 'concentrations' to 'concentration'

We have also changed the "A concentration of 10<sup>5</sup> cyanobacteria cells mL<sup>-1</sup> is considered to present a risk of both acute and chronic health effects (Backer, 2002), and many states, including Massachusetts, issue public health warnings for recreational water bodies when the cyanobacteria cell concentration exceeds this value."

Line 40, modify 'et. al.' with 'et. al.'

According to other referee comments, we have changed this sentence to "*Previous* research showed that recruitment..."

Line 48, insert space after 2008;

It seems that many of the references required spaces to separate them. This has been addressed here and in many other instances in the text.

Line 71, change 'volumes' to 'volume'

This has been changed.

Line 74, change 'greatest' to 'a considerable'

We agree that removing greatest is advisable, but tried to improve the sentence structure with the following "This potential transport pathway could be relatively more important for metal and cyanobacteria transport in eutrophic, deep, stratified lakes, such as UML."

Line 79, change 'distribution' to 'distribution'

This "s" has been removed from "distribution".

Line 119, change 'mixing from of the' to 'mixing from the'

This has been changed to "preventing mixing of sediment to the surface"

Line 123, change 'an' to 'a'

This has been changed.

Line 148, change 'column is comprised' to 'column comprised'

This has been changed to "The column is composed of four section..."

Line 176, change 'um' to '\_m'

This has been changed.

Line 180, change 'metals analysis on bulk sediment' to 'metal analysis in bulk sediment'

This has been changed.

Line 185, change 'which use' to 'with use'

This part of the sentence has been removed.

Line 186, change 'analysis on' to 'analysis of'

This was changed.

Line 188, 5 \_mol filter? Is it correct?

umol was not correct and we changed to 5 um.

We would like to thank the referee for looking over this work and providing valuable critiques to our paper. The comments are thoughtful and bring up many important points, which we addressed individually below.

## Anonymous Referee #4

The issues with sample collection make me call into question the quantitative results and budget. Please see my specific comments below for further details. Ultimately, the data need to be published, but the manuscript needs major revisions to remove the budgets which are likely inaccurate, given the sample collection procedure. Please refocus the manuscript to state the observations and cast your results in light of how the samples were collected.

We agree that the quantitative results and budget analysis are highly speculative, so the suggestion of removing the budget analysis would certainly be one way of addressing this issue. However, we propose keeping the budget calculations in the text, but making sure to emphasize the proper uncertainty associated with these budget estimates and to replace any specific estimates highlighted in the abstract or conclusions with a statement that more work is needed to calculate a proper budget for this mechanism. We hope that this approach would provide some context for the observations while remaining realistic about the fact that the information isn't at the level it needs to be for estimating a proper budget. We hope that our revisions have captured the spirit of this comment, while still providing some context to interpret our observations and to inspire future research.

## Specific Comments:

L 23-24: Define "problematic". What does this mean for cyanobacteria? Be more specific.

This statement was clarified as "In a 2012 national assessment, 15.2% of surveyed lakes in the U.S. were categorized as Most Disturbed due to the concentration of cyanobacteria, a significant increase in lakes with this categorization (8.3%, 95% confidence intervals 4.0-12.5%) over the 2007 assessment (U.S. Environmental Protection Agency, 2016)."

L 29-30: What about the "improved understanding"? What type of understanding? Be specific.

We have changed this to be more specific as "Identifying the sources and mechanisms of transport of these substances within lake ecosystems can help predict the fate of contaminants and aid remediation efforts."

L 110-111: How do you know the bubble transported biology and chemistry is no adhered to the inner walls of sampling equipment? Do your measurements represent an underestimate?

This is a point that was also brought up by a previous referee, so we have added a comment about this potential sampling artifact, which would underestimate transport:

"The interaction of bubbles with the flexible tubing resulted in visible particle attachment to the tubing, making our estimates of particle mass transport a lower bound."

L 172-173: Are these filter measurements meant to be volumetric? If so, do you know how much water passed through each filter before clogging?

For these filter measurements, we recorded the total volume filtered and the total mass accumulated, whether or not this was distributed over more than one filter because of clogging. Thus, we do not know the volumes passed through individual filters, only the total volume of water associated with a total particle mass.

## We have amended our text to read:

"Due to filter clogging, we typically used multiple filters for each sample, and total particulate transport per sample was calculated by summing the particle mass on each filter and dividing by the total gas volume associated with the sample. "

L181: I don't know how this relates to the accuracy and precision of your measurements? How do counts per second relate to concentration?

The relative standard deviation of the ICP-MS counts relates to the uncertainty in the measurements. The uncertainty for the sediment digests is quite low, and while it is higher in the less concentrated bubble transported particle samples, this uncertainty is still low relative to the experimental uncertainty. We have added the following line to the text:

"These relatively low RSD values indicate that analytical uncertainty is low, especially compared experimental uncertainty."

L 266: This is an excellent study and I think your experiments and testing shows bubbles play a role in lakes that has not been considering from a biological perspective. This study needs to be published, but I can't get over the anchor drop issue. I have thrown many anchors overboard in lakes and the plume of sediment is always significant. I have a hard time decoupling this disturbance with your results. There needs to be a paragraph describing how the laboratory results follow the lake results and the anchor had minimal impact on the lake results. Although, your laboratory results show sediment disturbance impact the bubble transported particles. How can you decouple these methodological problems with your results? What if you shift the focus of your manuscript to documenting that bubbles DO transport chemistry and biology, but stop short of the full budgets, as I think those are biased due to the methodological problems. We agree with the reviewer that triggering bubbles with an anchor drop leads to substantially different conditions than naturally ebullition. We wish we could have collected samples from natural ebullition alone, but this would have resulted in long wait times and probable changes in the cyanobacteria population prior to sample analysis. We attempted to alleviate some of this concern by using the laboratory bubble column experiments to demonstrate that particle scavenging when bubbles rise through a plume of sediment is still a relatively minor contribution to total particle transport. However, we agree that this experiment alone cannot account for all potential effects of the anchor drop. We feel this is an excellent area for future research, either in systems with much higher ebullition rates such that natural bubbles could be used, or potentially with updated experimental apparatus that can utilize natural bubbles.

To address these concerns, we have re-worded the text in numerous areas to highlight the uncertainty while still providing context for whether these observations could substantially impact chemical cycling or cyanobacterial life cycle. Some examples include:

Abstract- "Although more work is needed to reduce uncertainty in budget estimates, bubble-facilitated cyanobacterial transport has the potential to contribute substantially to the cyanobacteria cell recruitment to the surface of this lake and may thus be of particular importance in large, deep, stratified lakes."

**Results-** "These particle loadings on bubbles, and any ecosystem-wide flux estimates derived from them, must be qualified by the fact that neither triggered bubbles nor bubbles in the bubble column fully replicate natural bubbling. In particular, the triggering of bubbles with an anchor may have raised plumes of suspended sediment through which some fraction of produced bubbles had to rise, and within which the possibility of scavenging should be considered."

" However, many questions remain regarding bubble-mediated transport in natural systems, including how the change in water density at the thermocline affects bubble rise and associated chemical and biological material."

"There remains the possibility that our measured bubble particle transport rates differ significantly from those from naturally emitted bubbles, and this remains an important area for future research. However, despite this uncertainty, broad-scale estimates of arsenic and cyanobacteria cycling can provide important context as to whether these processes may be significant in UML."

" These calculations demonstrate that bubble transported cyanobacteria could negatively impact water quality, though more research is warranted to improve these estimates."

" Using the maximum observed recruitment rate of 2.3 x 10<sup>5</sup> cells m<sup>-2</sup> day<sup>-1</sup> (Brunberg and Blomqvist, 2003) from sediments for the area of the lake above 12 meters, we

estimate that bubbling could contribute 14 % of cyanobacterial recruitment in the lake, but 95% confidence intervals range from less than 0 to 46% of overall recruitment. While we cannot rule out the possibility that this is an insignificant source of cells given the large uncertainty in these measurements, the potential for bubble-mediated transport to contribute substantially to the source of cyanobacteria cells at the lake surface warrants further investigation."

## Conclusions-

"Bubble mediated transport of cyanobacteria cells may contribute substantially to cellular recruitment from the sediment, but the uncertainties in our measurements make these estimates speculative."

L 268-270: This observation is baseless since you caused the ebullition.

The reviewer makes a good point that natural variation in ebullition has nothing to do with the variation in mass transport observed in our triggered bubbling events. We have removed this sentence.

L 277-280: This is analogous to dropping an anchor on the lake sediments. How do you reconcile these laboratory experiments with what you did in the field? Again, this is evidence the focus of the manuscript should be focused to an observation that bubbles do transport chemistry and biology, but do not calculate budgets because the evidence shows they are not accurate.

Two observations from the columns with recently disturbed sediments (similar to the anchor drop, as mentioned in the comment) are similar to those with "normal" sediment, so the impact of these disturbances creates a complicated relationship with particle transport that we can not fully understand. The combination of both measurements ("normal" and "recently disturbed") resulted in transport that were similar to one field collection date, so it is at least in a similar range to what is occurring in the field.

This comment again highlights the uncertainty in our measurements. We agree with this comment and address it by making the uncertainty in our calculations more prominent, downplaying numbers in the abstract and conclusions, but keeping the budgets for context. We have re-written the text in numerous locations to highlight sources of uncertainty (mentioned above). However, we do still see value in budget calculations, however uncertain they may be. For example, the rough budget calculations for arsenic show a several order of magnitude gap between potential bubble arsenic transport rates and other transport rates within UML, indicating that even if our estimates are biased low, they are unlikely to be high enough to matter in UML. Conversely, the upper threshold for cyanobacteria transport in UML does fall within the realm of an important flux, which is a justification for further research in this area. We therefore think these estimates give a useful perspective, but we emphasize the large uncertainty that exists in these measurements and that the budgets are a best guess. L 283-285: Were there particles to scavenge? This was tap water, right?

This reviewer and one other have helpfully pointed out that we were not clear about the water column conditions when we conducted our tests for particle scavenging. As discussed previously in this response, scavenging tests were done after tests where bubbles were emitted from the sediment bed, so the water column was visibly turbid and contained many suspended particles.

We have added a sentence to clarify this point: "We conducted the scavenging tests when the water column was visibly turbid and contained a plume of suspended particles from previous tests."

Section 3.3 header: Again, I have a hard time reconcile the topic of this section that particles originated in the sediment after traveling through a plume of sediment. Maybe scavenging is a more active process and makes up a larger percentage of the particles when not passed through a plume of sediment.

We agree that bubble scavenging of particles within the water column could contribute to the particle burden, and thus not all particles originate in the sediment. Indeed, our scavenging tests shows that approximately 10% of the particles transported to the surface could be picked up within the relatively turbid water column. This indicates that within our experiments, a substantial fraction of the particles appear to come from the sediment bed itself. However, as pointed out previously, the artificial conditions for bubble release in both our laboratory and field experiment could influence our results. To acknowledge this uncertainty, we have changed the section title to:

# "3.3 Bubble-transported particles have chemical and biological characteristics similar to sediment

The data on bubble particle mass transport clearly shows that bubbles are capable of transporting particles from relatively deep depths, and minimal rates of particle shedding and scavenging in the water column suggests that these particles originate primarily in the sediment. "

L 325-326: Observations like this are the reason this manuscript needs to be published.

We appreciate your support for the publication of this work. To highlight the finding of potential ephippia in the particles, we have added a reference to the specific panel in Figure S10 that may show ephippia (Fig. S10-B).

L 353-354: This is a major finding of this study and should be a highlight.

We appreciate the reviewer's enthusiasm for the content. The referenced sentence in L 353-354 speculates that since cyanobacteria overwinter in the lake sediments, bubble-

mediated transport could be a mechanism of inoculating the upper water column with these cells. We believe we have highlighted this possibility with the mass budget calculations that compare potential bubble cell transport to other methods of cell recruitment. However, as discussed previously in responses to this reviewer, there remains a high degree of uncertainty around our estimated cell flux.

L 374: What does it mean to have a negative rate of transport? Are bubbles actually sequestering cells from the surface waters? This is another reason why I think the budgets need to be removed and the focus placed on the observations and laboratory experiments.

A negative transport rate is not meaningful, but is another aspect of the variability of our measurements that add uncertainty to the budgets. As discussed earlier, we agree with the reviewer that more attention should be given to the uncertain nature of our budget calculations, and have re-written portions of our text accordingly. Furthermore, we now conclude with the statement that:

"Using the maximum observed recruitment rate of  $2.3 \times 10^5$  cells m<sup>-2</sup> day<sup>-1</sup> (Brunberg and Blomqvist, 2003) from sediments for the area of the lake above 12 meters, we estimate that bubbling could contribute 14 % of cyanobacterial recruitment in the lake, but 95% confidence intervals range from less than 0 to 46% of overall recruitment. While we cannot rule out the possibility that this is an insignificant source of cells given the large uncertainty in these measurements, the potential for bubble-mediated transport to contribute substantially to the source of cyanobacteria cells at the lake surface warrants further investigation."

L 400: Given the large errors in your bubble transport of cells, I have a hard time following how the error now is so small. The error propagation is not well explained.

This is an error, and the range of values reported comes from using both 9 meters and 12 meters as the cut-off for where cyanobacteria would be able to recruit to the surface without bubbles. We agree that this does suggest a smaller uncertainty in the final budget than is warranted from the data.

To address this and the comment from above, we propose to still include the budgets in the presentation of the data for perspective, but to better emphasize the speculative nature of these budget results and the uncertainty associated with it. This provides context for the results and motivates additional research in the future on this topic, while still being realistic about whether these transport rates are well constrained. Even with the large uncertainty in particle transport values, the arsenic transport is unlikely to be a substantial part of arsenic found in the lake surface, but bubbles could still be an important part of cyanobacteria transport.

Since there are a number of uncertainties associated with cyanobacteria transport, we can emphasize that bubble-mediated transport has the potential to be a significant source of cell recruitment, especially in deep, eutrophic lakes. However, more work is

needed to better constrain these values to determine the actual contribution.

Technical Comments: L 22: Delete "are". Thank you for finding this glaring error in our first sentence, we have deleted the "are".

## L 22-23: First sentence needs a citation.

We have added two references that provide an overview of how water quality is a wide-spread phenomenon that will be likely exacerbated with increases in urbanization and climate change. "Deterioration of water quality is wide-spread and expected to become more acute with increased urbanization and climate-change (Zhang, 2016; Paerl et al., 2011)."

L 32-34: First sentence of the paragraph, poor sentence structure, please rewrite.

We have clarified this sentence to read:

"Because sediments are typically major repositories of contaminants (Nriagu et al., 1996; Pan and Wang, 2012; Taylor and Owens, 2009), it is important to understand the processes leading to contaminant mobilization."

L 35-37: "However, transport to surface: ::" Poor sentence structure, please rewrite.

We agree that this sentence was poorly worded. We have restructured the whole paragraph to improve readability:

"Because sediments are typically major repositories of contaminants (Nriagu et al., 1996; Pan and Wang, 2012; Taylor and Owens, 2009), it is important to understand the processes leading to sediment mobilization. Metals can be mobilized from sediments via solubilization by oxidation-reduction reactions, and by sediment resuspension, acidification or bioturbation (Calmano et al., 1993; Eggleton and Thomas, 2004; Schaller, 2014; Schindler et al., 1980). Likewise, over-wintering cyanobacteria and algae concentrated in the sediments are mobilized through germination, wind-induced resuspension, or bioturbation (Ramm et al., 2017; *Verspagen et al., 2004; Stahl-Delbanco and Hansson, 2002).* In some cases, the number of resting cells in sediment can be predictive of the severity of subsequent bloom events (Anderson et al., 2005). Previous research showed that recruitment from sediments of the potentially toxic cyanobacterium Microcystis was a major driver of the summer bloom (Verspagen et al., 2005). Cyanobacterial recruitment to surface waters from deep sediments is expected to be inhibited by stratification, low oxygen concentration, and low light levels (Ramm et al., 2017). Metals mobilized from sediment under stratified water columns will also be inhibited from reaching surface waters due to stratification (Wetzel, 2001)."

45-46: "Bubbling from anoxic sediment: : :" Sentence missing numerous citations. Thank you for bringing this to our attention, we have added the following two citations showing substantial contribution of methane bubbling to total freshwater emissions:

Bastviken, D.; Tranvik, L. J.; Downing, J. A.; Crill, P. M.; EnrichPrast, A. Enrich- prast, A. Freshwater methane emissions offset the continental carbon sink. Science 2011, 331, 50–50.

Deemer, B.; Harrison, J.;Li, S.; Beaulieu, J.;DelSontro, T.; Barros, N.; Bezerra-Neto, J.; Powers, S.; Dos Santos, M.; Vonk, J. Greenhouse gas emissions from reservoir water surfaces: A new global synthesis. BioScience 2016, 66 (11), 949–964.

Citations for the ability of bubbles to transport particles are already provided in subsequent sentences detailing this process in industry and marine systems.

L 50-

53: "Bubble-mediated particle: : :" Poor sentence structure, confusing, please rewrite.

We agree this sentence was quite poorly written, and have changed it to:

"Bubble-mediated particle transport also occurs in the open ocean where bubbles are injected into the water by breaking waves, scavenge surface-active particles as they rise, and then deposit these particles on the ocean surface (Aller et al., 2005; Blanchard, 1975; Wallace et al., 1972; Liss, 1975)."

L184-185: "We filtered bubble: : :" I did not understand this sentence. *We agree this sentence is confusing, and have shortened it to say:* 

"We filtered bubble column samples using pre-weighed 5.0  $\mu$ m and 0.2  $\mu$ m Whatman Nuclepore membrane filters (47mm diameter)."

L 187: How much lower are the blanks? Actual numbers would be better. Two orders of magnitude can range from 110-fold lower to 900-fold lower. These are very different blanks.

To clarify the blank question, we have calculated that the Whatman filters contained less than a nanogram of arsenic contamination, far below the sample concentrations. For the Nucleopore membranes, the 5  $\mu$ m filters had arsenic levels below the ICP-MS detection limit, and the 0.2  $\mu$ m filters had 0.003  $\pm$  0.002  $\mu$ g per filter for the 0.2  $\mu$ m filter (less than 1% of the arsenic found in the least concentrated sample). We have added the following text:

"Duplicate analysis of clean Nuclepore membranes (blank) was used to determine arsenic contamination of the filters and was below the detection limit for the 5  $\mu$ m

filters and 0.003  $\pm$  0.002 µg per filter for the 0.2 µm filter (less than 1% of the arsenic found in the least concentrated sample). "

L 249: mL-1 gas volume or mL gas volume-1? *This was changed to "mL gas volume-1"* 

L 250: Estimate – estimated (past tense). *Thank you, we have made this change.* 

L 258: Bring eq. 1 up so that the reader knows the equation before getting the variables.

We have Equation 1 to the top of the paragraph, along with a summary description of each variable to aid in readability.

Rewrite the part about the depth interval for germination. I was lost.

We have improved the readability of this section as: "We conservatively assumed that germination could occur to a depth of 12 meters based on typical light, temperature, and oxygen levels observed in UML (Varadharajan, 2009). The fraction ( $F_g$ ) of the surface area (SA = 580,000 m<sup>2</sup>) of lake above 12 meters that could support cyanobacterial recruitment through germination is approximately 0.50 (Varadharajan, 2009)."

L 362: This is a concentration, not a rate. *Thank you, we have eliminated "a rate of".* 

L 365: Keep units consistent. Use slash or exponent throughout.

Thank you for noticing this inconsistency. We have used exponents throughout.

#### Vertical transport sediment-associated metals of and

## cyanobacteria by ebullition in a stratified lake

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Abstract. Bubbles adsorb and transport particulate matter in a variety of natural and engineered settings, including 10 industrial, freshwater, and marine systems. While methane-containing bubbles emitted from anoxic sediments are found widely in freshwater ecosystems, relatively little attention has been paid to the possibility that these bubbles transport particle-associated chemical or biological material from sediments to surface waters of freshwater lakes. We triggered ebullition and quantified transport of particulate material from sediments to the surface by bubbles in Upper Mystic Lake, MA and in a 15 m tall experimental column. Particle transport was positively correlated with the volume of gas bubbles released from the sediment, and particles transported by bubbles appear to originate almost entirely in the sediment, rather 15 than being scavenged from the water column. Concentrations of arsenic, chromium, lead, and cyanobacterial cells in bubbletransported particulate material were similar to those of bulk sediment, and particles were transported from depths exceeding 15 m, implying the potential for daily average fluxes as large as 0.18  $\mu$ g of arsenic m<sup>-2</sup> and 2 x 10<sup>4</sup> cyanobacterial cells m<sup>-2</sup> in the strongly stratified Upper Mystic Lake. Bubble-facilitated arsenic transport currently appears to be a modest component 20 of total arsenic cycling in this lake. Although more work is needed to reduce uncertainty in budget estimates, bubblefacilitated cyanobacterial transport has the potential to contribute substantially to the cyanobacteria cell recruitment to the

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surface of this lake and may thus be of particular importance in large, deep, stratified lakes.

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#### **1** Introduction

35 Deterioration of water quality is wide-spread and expected to become more acute with increased urbanization and climatechange, (Zhang, 2016; Paerl et al., 2011). In a 2012 national assessment, 15.2% of surveyed lakes in the U.S. were categorized as Most Disturbed due to the concentration of cyanobacteria, a significant increase in lakes with this categorization (8.3%, 95% confidence intervals 4.0-12.5%) over the 2007 assessment (U.S. Environmental Protection Agency, 2016). A concentration of 10<sup>5</sup> cyanobacteria cells mL<sup>-1</sup> is considered to present a risk of both acute and chronic health effects (Backer, 2002), and many states, including Massachusetts, issue public health warnings for recreational water bodies when the cyanobacteria cell concentration exceeds this value. Metals are also important contaminants in freshwater systems because of their persistence and toxicity (Bronmark and Hansson, 2002). In 2004, 1.5 million lake-acres in the U.S. were impaired by metals, such as lead, chromium and arsenic (Environmental Protection Agency, 2004). Jdentifying the

sources and mechanisms of transport of these substances within lake ecosystems can help predict the fate of contaminants 45 and aid remediation efforts.

- Because sediments are typically major repositories of contaminants (Nriagu et al., 1996; Pan and Wang, 2012; <u>Taylor and Owens, 2009), it is important to understand the</u> processes leading to <u>contaminant</u> mobilization, Metals can be mobilized from sediments via solubilization by oxidation-reduction reactions, and by sediment resuspension, <u>acidification, or</u> <u>bioturbation</u> (Calmano et al., 1993; Eggleton and Thomas, 2004; Schaller, 2014; Schindler et al., 1980). Likewise, over-
- 50 wintering cyanobacteria and algae concentrated in the sediments are mobilized through germination, wind-induced resuspension, or bioturbation (Ramm et al., 2017; Verspagen et al., 2004; Stahl-Delbanco and Hansson, 2002). In some cases, the number of resting cells in sediment can be predictive of the severity of subsequent bloom events (Anderson et al., 2005). Previous research showed that recruitment from sediments of the potentially toxic cyanobacterium *Microcystis* was a major driver of the summer bloom (Verspagen et al., 2005). Cyanobacterial recruitment to surface waters from deep sediments is, inhibited by stratification, low oxygen concentration, and low light levels (Ramm et al., 2017). Metals mobilized from sediment under stratified water columns will also be inhibited from reaching surface waters due to

mobilized from sediment under stratified wat stratification (Wetzel, 2001).

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An alternative mechanism for vertical transport of metals and cells from sediment to surface water could be bubble-

facilitated transport. Bubbling from anoxic sediments, driven by methanogenesis, is widespread in freshwater systems
(Bastviken et al., 2011; Deemer et al., 2016), and bubbles are known to be effective particle transporters. Bubble particle
flotation, a process by which amphiphilic particles attach to a bubble's gas-water interface and are transported upwards
during bubble rise, is used extensively in industry for applications such as separating valuable minerals from gangue (Min et al., 2008; Rodrigues and Rubio, 2007), removing ink during paper recycling (Vashisth et al., 2011), recovering desirable
proteins and microorganisms from industrial bioreactors (Schugerl, 2000), and treating wastewaters (Aldrich and Feng, 2000; Lin and Lo, 1996; Rubio et al., 2002). Bubble-mediated particle transport also occurs in the open ocean where bubbles are injected into the water by breaking waves, scavenge surface-active particles as they rise, and then deposit these particles on the ocean surface (Aller et al., 2005; Blanchard, 1975; Wallace et al., 1972; Liss, 1975).

Despite this previous work, little is known about the importance of particle transport by bubbles in freshwater systems. Bubbles produced by methanogenesis in anoxic sediments are prevalent in freshwater systems, and bubbles are released to the surface during drops in hydrostatic pressure, sediment disturbance, or upon sufficient gas accumulation

(Chanton et al., 1989; Joyce and Jewell, 2003; Scandella et al., 2011; Liu et al., 2016; Maeck et al., 2014; Varadharajan and Hemond, 2012). Bubble flotation could thus potentially provide a chemical and biological link from deep water to surface waters that would otherwise not occur through advective or eddy-diffusive transport alone. Additionally, the relatively rapid

rise time of bubbles limits the time available for oxidation reactions, and suggests that particulate matter from the

- 105 hypolimnion could reach the lake surface in a reduced state, with possible consequences for both toxicity and reactivity. Some evidence does suggest that bubbles can transport polycyclic aromatic hydrocarbons (Viana et al., 2012) and manufactured gas plant tar from sediments (McLinn and Stolzenburg, 2009). Additional work has shown that bubblemediated transport of microorganisms including methane oxidizing bacteria (MOB) is an important mechanism connecting benthic and pelagic populations at 10 m water depth (Schmale et al., 2015). However, researchers in the previous study were
- 110 unable to quantify the importance of bubble-mediated transport to overall recruitment of pelagic MOB populations, and the extent of bubble particle flotation in aquatic systems remains unknown.

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Microsoft Office User 2/17/20 12:37 PM Deleted: (Aldrich and Feng, 2000;Lin and Lo, 1996;Rubio et al., 2002). Bubble-mediated particle transport also occurs in the open ocean and contributes to an accumulation of surface-active particles at the surface of the ocean (Aller et al., 2005;Blanchard, 1975;Wallace et al., 1972), as bubbles are injected into the ocean surface by breaking waves, and their particle burden is obtained by scavenging of particles as the bubbles rise (Liss, 1975).

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Microsoft Office User 2/17/20 12:31 PM Deleted: full The present study is motivated by authors' observations of particle accumulations associated with bubbling events at Upper Mystic Lake (UML), where bursting bubbles often left black particles distributed on the water surface in a ring pattern (Fig. S1). Particles were also observed at the air-water interface in bubble traps during long-term deployments (data not shown). The significant volume of gas observed to bubble from UML during previous studies (Delwiche and Hemond,

135 2017; Varadharajan and Hemond, 2012), together with strong thermal stratification suppressing other mechanisms of sediment transport to the surface, led to the hypothesis that bubbles could serve as a relatively important mode of particle
 transport from the sediment to the water surface. This potential transport pathway could be relatively more important for metal and cyanobacteria transport in eutrophic, deep, stratified lakes, such as UML.

In the present study, we quantified particle transport by bubbles in UML, an urban lake with a history of sediment contamination. We also used a 15 m tall bubble column to study bubble-mediated particle transport under controlled lab conditions. Given the expected importance of bubble size on key characteristics (e.g. surface area, buoyancy, diffusion of gas), we used a bubble size sensor (Delwiche et al., 2015; Delwiche and Hemond, 2017) to measure bubble diameter distribution both in the lake and in the laboratory. We address the following questions:

1. How much sediment is transported to the surface through ebullition?

145 2. How does bubble-mediated sediment transport contribute to metal cycling?

3. How does bubble-mediated sediment transport contribute to cyanobacteria recruitment to the upper water column?

### 2 Methods

### 2.1 Upper Mystic Lake field site history

UML in Arlington, MA is an urban, dimictic kettle lake with an average depth of 15 m, a maximum depth of 24 m, and a surface area of 0.58 km<sup>2</sup>. The lake is used extensively for recreational and scientific purposes, and previous studies have characterized several aspects of methane ebullition (Delwiche and Hemond, 2017; Scandella et al., 2016; Varadharajan and Hemond, 2012) and microbial community structure and function (Preheim et al., 2016; Arora-Williams et al., 2018). Microsoft Office User 2/17/20 12:31 PM Deleted: volumes Microsoft Office User 2/17/20 12:31 PM Deleted: (Delwiche and Hemond, 2017;Varadharajan and Hemond, 2012)

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Chemical manufacturing and leather tanning industries during the late 1800s and 1900s released toxic metals such as arsenic, chromium, and lead that flowed into the lake and were deposited in the lake sediments. Sediment cores reveal a distinct layered pattern with peak metal/metalloid concentrations traceable to years of peak manufacturing or subsequent earth-moving (Spliethoff and Hemond, 1996). Additionally, high nutrient loading promotes the growth of algal and cyanobacterial blooms. A public health advisory was issued for UML as recently as July 2017 for cyanobacteria cell concentrations

>70,000 cells mL<sup>-1</sup> (https://www.arlingtonma.gov/Home/Components/News/News/4965/16 accessed on 06/05/2019).

Years of field observations at UML have provided a thorough picture of the typical hydrological conditions in the lake. Significant volumes of gas are produced from the sediments, which escape to the surface via ebullition, resulting in an average release rate of 22 ml of bubble volume m<sup>-2</sup> d<sup>-1</sup>(Varadharajan, 2009). From June - Oct., the oxycline and thermocline are typically found between 6-12 m and 3-9 m, respectively (Varadharajan, 2009; Delwiche and Hemond, 2017). The Secchi depth in the lake is typically 2-3 m during the same period (Varadharajan, 2009). Light was sufficient for germination down to 12 m in <u>Lake Scharmützelsee</u> with a similar average Secchi depth (Ramm et al., 2017), thus we assume light does not limit cyanobacteria germination down to a depth of at least 12 m when estimating the impact of bubbling on cyanobacteria recruitment.

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#### 2.2 Field sampling

Bubble-transported particles were collected from both the laboratory column and the lake in 350 mL plastic sampling cups affixed either to the top of a custom bubble size sensor [sensor described previously (Delwiche et al., 2015; Delwiche and Hemond, 2017)], or to the top of a collection funnel (the bubble sensor was used in 2017 sampling; the funnel alone was used for sampling in 2018). The plastic sampling cup lid contained a barbed bulkhead fitting connected via flexible plastic tubing to an on-off valve and a quick-release adapter (Fig. S2). The sampling cup, valve, and adapter were connected to the custom bubble size sensor or collection funnel with flexible tubing. All bubbles rising through the bubble size sensor or collection funnel entered the flexible tubing and rose into the sample cup. The interaction of bubbles with the flexible tubing resulted in visible particle attachment to the tubing, making our estimates of particle mass transport a lower bound. The

sample cup lid contained a secondary valve to release water upon bubble entry. All sample cups were soaked in 5-10%

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Deleted: (Delwiche et al., 2015;Delwiche and Hemond, 2017) 200 <u>reagent grade HNO<sub>3</sub></u> for 24 hours and rinsed and filled with Milli-Q water prior to use. Gas and associated particles accumulated in the collection cup during sampling, and were then transported back to the lab for analysis.

On 17 October 2017 we sampled for bubble-mediated sediment mass fluxes and associated particulate metal fluxes in an area of the lake previously found to have relatively high ebullition rates [42.432 latitude, -71.151 longitude, and 16 m deep; (Delwiche and Hemond, 2017)]. This previous work showed that sediment ebullition rates from this location remain

- high from July to November, yet the water column remains stratified, preventing mixing  $\rho f$  sediment to the surface. Previous work at this particular location within the lake indicated that natural bubble fluxes were around 45 mL m<sup>-2</sup> day<sup>-1</sup> with high spatial and temporal variability (Delwiche and Hemond, 2017). Given the need to collect samples as soon after bubbling as possible to minimize potential changes in cyanobacteria population, and the difficulty with predicting flux from natural bubble events, we chose to trigger ebullition manually by dropping <u>a</u> 20 cm x 20 cm x 20 cm cinderblock anchor into the
- 210 sediment. This procedure enabled us to collect multiple samples during a single field trip, with minimal time for samples to change after collection in the sampling cup. Since anchor triggering was expected to release a plume of sediment, we used laboratory experiments to explore whether bubbles rising through suspended sediment would scavenge particles (more details below).

After bubble triggering, the bubble size sensor was positioned above the bubble plume and 1 m below the water 215 surface. Bubbles exiting the sensor, together with any particles adhered to the bubble/water interface, were collected in the sample cup described previously. Several anchor drops within an area of approximately 10 m by 10 m were required to intercept a sufficient number of bubbles for mass quantification per sample, and we intentionally collected samples with different total gas volumes. We collected blank water samples to correct for background contributions of particulate matter, arsenic, lead, and chromium. Bubbling resulted in the visual accumulation of particles at the surface (Fig. S1) and in the 220 sampling cup. During a separate field visit in November 2017 we used an <u>Ekman</u> dredge to collect sediment and stored the

sediment in a 5 gallon bucket below 4 C° until use in February 2018 for bubble column experiments.

On 26 June 2018 we sampled for cyanobacteria bubble transport using similar procedures, except we used a simple inverted funnel instead of a custom bubble size sensor to intercept rising bubbles. The sampling funnel was placed 10 m below the water surface, where cyanobacteria concentrations were expected to be lower than the surface based on previous

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Microsoft Office User 2/17/20 12:31 PM Deleted: Eckman 230 observations (Preheim et al., 2016) to reduce sample contamination with cyanobacteria from the surrounding water column. Water temperature measurements taken using a Hydrolab sonde (Hach Co.) confirmed that the thermocline depth was above
10 m in this location during sampling, (Fig. S3). We collected 30-40 mL of water samples at 15m, 11m, 10m, and 1m depths for background cell concentration counts, and gathered sediment grab samples with an Ekman dredge. All sample cups were sterilized prior to use by rinsing with 10% bleach followed by 70% ethanol and deionized, sterile water, and cups were filled

with sterile water prior to sample collection. Samples were stored in a dark cooler on ice and were refrigerated upon return
 to the lab. On 26 June 2018 we also used an <u>Ekman</u> dredge to collect a bulk sediment sample, which was kept in a dark
 refrigerator at 4 C° until use in February 2019 for cyanobacteria transport in the experimental bubble column.

#### 2.3 Large laboratory column design and sampling

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To study bubble particle shedding and scavenging, we built a 15 m tall bubble column in the laboratory stairwell. The column is <u>composed</u> of four sections of 6-inch (15.3 cm) nominal diameter transparent polyvinyl chloride (PVC) pipe joined by threaded unions with O-ring seals. The base of the column is a reducing tee fitting with a removable spigot for drainage, and the column was filled from the top with tap water. We built a sediment container connected to 1/8 inch (3.1 mm) outer diameter copper tubing that could be lowered into the column and secured at any depth. The container was filled

with sediment originally collected with an Ekman dredge from the same place in UML used for field sampling. We used a syringe pump to push air into the sediment through the tubing at a controlled rate, resulting in bubble release from the sediment. The bubbling rate was calibrated to achieve a relatively steady release of bubbles without substantial wait time in between. While we expect that much of the gas naturally existing within the sediment was released during sediment collection and as it was transferred to the sample bed (indeed we did not observe natural bubble release from the sediment to be prior to experimental trials), remaining gas could have been incorporated into rising bubbles.

We conducted one set of column experiments in February 2018 to quantify shedding, scavenging, and metals transport, and another set of column experiments in February 2019 to quantify cyanobacterial transport. For the shedding, and metals transport experimental runs, we filled the sediment bed with sediment collected from the same site as ebullition experiments during our November 2017 field visit, and we injected 50 mL of air at 0.7 mL min<sup>-1</sup> into the sediment bed.

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- 260 Prior to the start of each run we collected water samples to correct for background contributions to particulate matter and arsenic concentrations in bubble-transported particle data. Three experimental runs were each conducted at each of three depths: 5m, 10m, and 15m, with the mobile sediment bed being repositioned between runs. To quantify particle scavenging rates, we also conducted trials in which we injected air into the water column several centimetres above the sediment surface. Scavenging tests were conducted after particle transport tests, so the water column above the sediment bed was turbid and contained a plume of sediment particles. For the cyanobacterial transport experiments, we used sediment from the
- June 2018 field visit and injected variable volumes of air into the sediment bed. We ran four experiments each at 6 m and 13 m depth, with the sediment being replenished between the 6 m and 13 m runs. Six surface water grab samples were collected at multiple times throughout the experiment to quantify background cell concentrations, and at each depth one trial was run where air was bubbled into the water directly below sensor. For both sets of experiments, bubbles passed through
- 270 the same customized bubble size sensor (Delwiche et al., 2015; Delwiche and Hemond, 2017) and sample cup apparatus used in the field setting.

#### 2.4 Sample processing for particle mass and heavy metals analysis

We filtered the field samples collected from UML for metals analysis within 24 hours of sampling with preweighed Whatman Grade 41 quantitative cotton filters (nominal pore size 20 µm, 25 mm diameter). Due to filter clogging, we typically used multiple filters for each sample, and total particulate transport per sample was calculated by summing the particle mass on each filter and dividing by the total gas volume associated with the sample. After filtering we air-dried the filters, weighed them, transferred each to microwave digestion vessels, and added 10 mL of nitric acid from Fisher Scientific (Optima grade for ultra-trace elemental analysis). Samples were digested in a MARS6 microwave oven, diluted with 30 mL of Milli-Q water, and then filtered with a 0.2 µm polyethersulfone membrane syringe filter. For analysis, we diluted samples to 2% nitric acid, added a rhodium internal standard, and analyzed the samples using an Agilent 7900 inductively coupled plasma mass spectrometer (ICP-MS) with a 5<sup>x</sup> point calibration curve from 0.05 - 10 ppb. Blank analysis to determine background arsenic concentrations in the Whatman cotton filter paper found levels of Jess than a nanogram of arsenic. For metal analysis in bulk sediment sample we added 100 mg of dried sediment to 10mL of nitric acid and digested as described Microsoft Office User 2/17/20 12:31 PM Deleted: (Delwiche et al., 2015;Delwiche and Hemond, 2017)

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above. The relative standard deviation (RSD) values for counts-per-second from the ICP data were on average  $5.2\% \pm 2.8\%$  for the bubble-transported sediment particles, and were  $1.1\% \pm 0.6\%$  for the bottom sediment digests (which contained more particle mass per digest). These relatively low RSD values indicate that analytical uncertainty is low, especially compared experimental uncertainty.

We filtered bubble column samples using pre-weighed 5.0  $\mu$ m and 0.2  $\mu$ m Whatman Nuclepore membrane filters 300 (47mm diameter). Filters were dried, weighed, digested, diluted, and analyzed as described above. Duplicate analysis of clean Nuclepore membranes (blank) was used to determine arsenic contamination of the filters and was below the detection limit for the 5  $\mu$ m filters and 0.003  $\pm$  0.002  $\mu$ g per filter for the 0.2  $\mu$ m filter (less than 1% of the arsenic found in the least concentrated sample).

#### 305 2.5 Sample processing for cyanobacteria analysis

For both the field and bubble column cyanobacterial transport experiments, we filtered a subset of the samples within 24 hours with 0.2 µm pore size filters held in autoclaved Swinnex filter holders (25 mm diameter). Filters were then removed from the filter holders and transferred to PowerWater bead beating tubes (Qiagen, Inc.). Approximately 8-9 <u>mL</u> of remaining liquid for each sample was preserved with 1-2 mL of formamide (10% final concentration volume/volume) for

310 microscopic cell counts. Lastly, the remaining sample volume was filtered on pre-weighed Whatman Nuclepore membrane filters (0.2 μm pore size, 47mm diameter), air dried, and re-weighed to estimate bulk mass transport.

For qPCR analysis on the June 2018 bulk sediment samples, 0.13 g of wet sediment was suspended in 15 mL of sterile water and then filtered as described above. For microscopy cell counts, 0.14 g wet sediment were preserved in 2% by volume paraformaldehyde. Water column samples from the June 2018 field campaign were also preserved in 2% by volume

315 paraformaldehyde for cell counts. For qPCR analysis of the June 2018 sediment samples before use in the bubble column, we filtered 0.7 g of wet sediment (0.007 g dry sediment). For microscopy cell counts of the June 2018 sediment samples before use in the experimental bubble columns, we placed 0.8 and 2.0 mg of wet sediment (0.08 and 0.18 mg dry weight, respectively) in to 10 mLs of 10% formalin.

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#### 2.6 Cyanobacteria cell quantification

Cyanobacteria cell counts were assessed through quantitative polymerase chain reaction (qPCR) and microscopy.

These two methods estimate cyanobacteria cells numbers by targeting different features of cyanobacterial cells. qPCR targets

- 335 the unique genetic signatures in the 16S ribosomal RNA (rRNA) gene of cyanobacteria (Nubel et al., 1997) to estimate cell number from gene copy numbers. Microscopy takes advantage of the unique fluorescence spectra of cyanobacterial photosynthetic pigments to identify cells (Salonen et al., 1999). Positive control Microcystis aeruginosa UTEX LB 2386 and negative control Pseudomonas aeruginosa samples were used to optimize amplification conditions to ensure specificity for cyanobacteria qPCR. Microcystis and Pseudomonas cultures were grown overnight (12 h) under fluorescent lights at 25
- 340 °C in BG11 and Luria Broth media, respectively. Microcystis stock culture was serially diluted in phosphate buffered saline to make a standard curve, filtered onto 0.22 µm polyethersulfone membrane filters (Millipore Sigma, Inc.) and frozen at - 80 °C until DNA extraction. Additionally, serial dilutions of Microcystis cultures were fixed with 1% formalin (final concentration, volume/volume) for microscopy. While Microcystis cells were used as a positive control to test the method, qPCR primers targeted all cyanobacteria cells (not limited to Microcystis).
- 345 To estimate the total number of cells in the Microcystis stock culture and samples with microscopy, between 4.6 mL to 10.4 mL of fixed water samples or 1000 µL fixed Microcystis stock culture were filtered onto 0.22 µm polyethersulfone membrane filters (Millipore Sigma, Inc.). Cells were visualized under a Zeiss AxioObserver Epifluorescence SIM microscope [excitation: 545 nm; emission: 572 nm (Salonen et al., 1999)]. The total number of autofluorescent cells per filter was estimated from twenty to forty random fields of view spanning the entire area of each filter. Cells were identified from images with ImageJ (Schneider et al., 2012). First, background noise was reduced by excluding low intensity pixels, 350 with threshold values ranging between 14-162 (pixel intensities ranged from 0-255 for 8-bit gray-scale images). Next, only particles within the size range of  $0.1 \ \mu m^2 - 29.4 \ \mu m^2$  were counted as cells. A dilution series of *Microcystis* fixed culture was created by diluting cultures 2-fold in 1% formalin to test the variance and accuracy of this counting method (Fig. 54). We

did not test the quantification below 20 cells per field of view and all the experimental samples (not controls) had an average 355

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of less than 20 cells per field of view, so microscopy measurements were only used for detection, not quantification.

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For cyanobacteria cell quantification with (qPCR), DNA was extracted using PowerWater kits (Qiagen, Inc)

- 360 following the manufacturer's protocol, with the addition of 20 μl proteinase K and incubation at 65 °C for 10 min before bead beating as an alternative lysis step. Primers were used to amplify *Cyanobacteria* 16S rRNA genes as previously described (Nubel et al., 1997), with CYA359F (5'- GGG GAA TYT TCC GCA ATG GG) and an equal mixture of CYA781R(a) (5'- GAC TAC TGG GGT ATC TAA TCC CAT T) and CYA781R(b) (5'- GAC TAC AGG GGT ATC TAA TCC CTT T). qPCR reactions contained 10 μl of SsoAdvanced Universal SYBR Green Supermix (BioRad Laboratories,
- 365 Inc.), 1.6 μl DNA template, 2 μl forward primer (10 mM), 2 μl reverse primer (10 mM), and 4.4 μl deionized, reagent grade sterile water. The following cycling conditions were used: denaturation at 98 °C for 30 seconds, annealing at 68 °C for 30 seconds, and elongation at 72 °C for 30 seconds followed by visualization step for 40 cycles. A dilution series of *Microcystis* was created by diluting cells 10-fold in PBS before filtration and DNA extraction. Cell numbers for environmental samples were determined from a linear regression of threshold cycle number (Cq) values of *Microcystis* and the number of cells
- 370 calculated for each dilution, (e.g. Fig. \$5) and different batches were calibrated with internal standards of *Microcystis* culture. Inhibition was determined for a subset of samples by spiking known concentrations of *Microcystis* DNA into environmental DNA extracts and measuring the resulting threshold cycle number (Fig. \$6). In all cases tested, inhibition was negligible. The limit of quantification is 5 cells per filter, based on a signal to noise ratio (SNR) 2-3 x the average cell concentration of the blanks (2.76 SNR).

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#### 2.7 Cyanobacterial recruitment estimates for cells from Upper Mystic Lake

The contribution (%) of ebullition to cyanobacteria recruitment ( $P_e$ ) was calculated as:

 $P_e = 100 \times \frac{c_e \times SA}{(c_e \times SA) + (c_g \times F_g \times SA)} \tag{1}$ 

Where  $C_e$  is the average cell flux from ebullition to the lake surface,  $C_g$  is the recruitment rate due to germination, SA is lake380surface area, and  $F_g$  is the fraction of the lake surface area that could support recruitment through germination. We estimated<br/> $C_g$  using our measured range of potential particle transport and the concentration of cells in the lake sediment. Values for<br/>recruitment estimates were calculated assuming ebullition occurs at an average rate of 22 mL m<sup>-2</sup> d<sup>-1</sup> for the entire summer<br/>from all areas of the lake equally based on previous lake-wide ebullition surveys, (Varadharajan, 2009), We used the average

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	cyanobacteria cell concentration from this study of 880 cells $m_{\pm}$ gas volume— to calculate the average flux of cells to the	
	surface via ebullition (C <sub>e</sub> ) of 2 x $10^4$ cells m <sup>-2</sup> <u>d<sup>-1</sup></u> . We <u>estimated</u> the recruitment rate due to resuspension and germination	
390	$(C_g)$ as the maximum observed rate from a previous experiment in Lake Limmaren of 2.3 x 10 <sup>5</sup> cell m <sup>-2</sup> d <sup>-1</sup> (Brunberg and	
I	Blomqvist, 2003), and applied this recruitment rate to areas of the lake suitable for germination, although this is likely an	
	overestimation. We conservatively assumed that germination could occur to a depth of 12 meters based on typical light,	
	temperature, and oxygen levels observed in UML (Varadharajan, 2009). The fraction $(F_g)$ of the surface area (SA = 580,000	
	$m^2$ ) of lake above 12 meters that could support cyanobacterial recruitment through germination is approximately 0.50	
395	(Varadharajan, 2009).	

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but only at the p=0.15 level ( $r^2$ =0.38) for the June 2018 data. Total gas collected per sample was a

function of the number of anchor drops per sample and our ability to position the boat above the bubble plume, so the lower gas volumes on June 2018 do

not necessarily indicate a smaller reserve of gas in the sediment. Samples show variable mass transport

rates both between and within sampling dates. Such variability is not unexpected given the previously documented spatial and temporal variability of

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bubble particle transport from 15 m depth of  $0.01 \pm 0.006$  mg/mL in the bubble column, compared to  $0.09 \pm 0.07$  mg/mL on October 2017 and  $0.01 \pm 0.01$ 

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ebullition rates in this lake (Varadharajan 2009;Varadharajan and Hemond, 2012).

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mass transport data showed a

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**Microsoft Office User 2/17/20 12:31 PM Deleted:** The bounds for germination suitability from the sediments, found using previous measurements of light, temperature, and oxygen at UML (Varadharajan, 2009), were assumed to occur 9 to 12 meters deep in the lake. The fraction ( $F_{s}$ ) of the surface area ( $SA = 580,000 \text{ m}^2$ ) of lake above 9 and 12 meters that could support cyanobacterial recruitment through germination is 0.40 to 0.50,

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#### **3 Results and Discussion**

#### 3.1 Rate of bubble-particle transport

Both field and bubble column experiments demonstrate that bubbles <u>can</u> transport particles from the sediment to the lake surface. A positive correlation (p < 0.05 level for October 2017 ( $r^2 = 0.76$ ), p=0.15 ( $r^2=0.38$ ) for June 2018 ) was found between total particle mass and gas volume in bubble traps for both field sampling campaigns (Fig. 1). The general magnitudes of particle loadings on bubbles in column experiments and on bubbles observed in triggered experiments in the field were of similar magnitude;  $0.01 \pm 0.006$  mg mL<sup>-1</sup> in the column vs.  $0.09 \pm 0.07$  mg mL<sup>-1</sup> on October 2017 and  $0.01 \pm 0.01$  mg mL<sup>-1</sup> on June 2018 in the field.

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These particle loadings on bubbles, and any ecosystem-wide flux estimates derived from them, must be qualified by the fact that neither triggered bubbles nor bubbles in the bubble column fully replicate natural bubbling. In particular, the triggering of bubbles with an anchor may have raised plumes of suspended sediment through which some fraction of produced bubbles had to rise, and within which the possibility of scavenging should be considered. Likewise, bubbles could shed particles part-way up the water column during rise. To estimate the significance of particle shedding, we used the bubble column to compare transport rates from bubbles released at 5 m, 10 m, and 15 m depths (Fig. 2). We found no

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significant difference in transport rates from any depths, suggesting that net particle shedding was not a major process. We did however note that the first bubble column test conducted after repositioning the sediment source yielded a higher particle

transport rate than those found in subsequent tests (Fig. 2), <u>consistent with the intuitively reasonable possibility that</u>
mechanical sediment disturbance <u>can affect particle loading on bubbles</u>. We also note that while bubbles do dissolve as they
rise, bubbles in the size range seen during this study remain relatively constant <u>in volume during their rise through 15 m of</u>
water column because dissolution is partially compensated by bubble expansion during rise (Delwiche and Hemond, 2017),
and we therefore do not expect bubble dissolution to substantively impact particle shedding.

To <u>observe</u> the <u>possible extent</u> of bubble scavenging of particles from the water column, we compared data from 5 m and 10 m column experiments to samples gathered when gas was bubbled from several centimetres above sediment, thus allowing maximum opportunity for scavenging to occur. We conducted the scavenging tests when the water column was visibly turbid and contained a plume of suspended particles from previous tests. Particle mass scavenging represented only around 10% of the mean particle <u>loading</u> for <u>bubbles in the</u> 5 m and 10 m experiments (grey diamonds in Fig, 2), indicating that while scavenging rates were non-zero, the large majority of the particulate matter transported to the top of the water column originated in the sediment. <u>Taken together</u>, the minimal particle shedding and particle scavenging in column

experiments suggests that particles observed on bubbles in the field, even when bubble release was triggered, mainly originated in the sediment.

While bubbles transported sediment directly from the bottom of the laboratory column to the water surface, a vertical distance of 15 m, there appears to be no reason that transport of particles from significantly larger depths cannot occur. Such transport provides a direct chemical and biological link between sediment and surface waters, and this could be the dominant link between deep sediments and the surface water during months of stratification. However, many questions remain regarding bubble-mediated transport in natural systems, including how the change in water density at the thermocline affects bubble rise and associated chemical and biological material.

#### 485 3.2 Bubble size distribution similar between field triggered and natural bubbles

Bubble volume has been found to significantly affect particle flotation rates in industrial processes (Yoon and Luttrell, 1989), and therefore it is important to compare the anchor-triggered bubble sizes to naturally-occurring bubble sizes to understand how our measured transport rates may reflect naturally occurring transport rates. Anchor-triggered bubbles were significantly smaller (average diameter 5.6 mm) than those measured for natural bubbling events (average diameter 6.4 mm)

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scavenging indicates that the large majority of

particles deposited on the surface by bubbles

originate in the sediment.

during a 2016 field campaign [Fig. §7, (Delwiche and Hemond, 2017)], even though relatively high bubble flux events (such as those triggered by anchor dropping) can lead to some bubble coalescence within the funnel constriction in the bubble size sensor [as described previously (Delwiche and Hemond, 2017)]

510 However, both natural and triggered bubbles were still very large compared to bubbles used in traditional flotation chambers (Yoon and Luttrell, 1989; Rubio et al., 2002). While research on particle flotation for large bubbles is limited, several previous studies have found that differences in particle transport rates decrease for bubbles above 1 mm diameter (Dai et al., 1998; Koh and Schwarz, 2008), indicating that particle transport rates should be similar between natural and triggered bubbles. Bubble sizes measured in the cyanobacteria transport experiment displayed a bimodal distribution (Fig. 515 S8) that was not observed in other bubble experiments. This bimodal distribution could be a result of artificially pumping gas in to the sediment, but the impact of this on particle transport is unknown.

#### 3.3 Bubble-transported particles have chemical and biological characteristics similar to sediment

The data on bubble particle mass transport clearly shows that bubbles are capable of transporting particles from 520 relatively deep depths, and minimal rates of particle shedding and scavenging in the water column suggests that these particles originate primarily in the sediment. Concentrations of arsenic, chromium, and lead in the bubble-transported particulate matter collected during field experiments were similar to concentrations in the sediment (Fig. 3). Bubbletransported particles contain arsenic, chromium, and lead at average ratios of 100 µg kg<sup>-1</sup> 120 µg kg<sup>-1</sup> and 240 µg kg<sup>-1</sup> (respectively, excluding outlier in chromium data, see Fig 3), compared to 136 ug kg<sup>-1</sup>, 160 ug kg<sup>-1</sup>, and 330 ug kg<sup>-1</sup> 525 (respectively) in bulk sediment samples. In the bubble column, arsenic and chromium levels are similar to the bulk sediment in the column experiments (Fig. §9), although lead levels appear to be higher. Overall, this similarity supports our conclusion that bubbles are primarily transporting sediment matter to the lake surface with only modest amounts of scavenging or particle shedding, despite the relatively deep water column.

In addition to the heavy metal results indicating that the transported particles are from the sediment, biological evidence also suggests a sedimentary origin. All particle samples transported by bubbles contained an abundance of 530 biological structures (Fig. <u>\$10</u>), such as the apparent head shields and carapaces of Bosmina spp., which have also been found extensively in other freshwater lake sediments (Kerfoot, 1995). These particle samples also contained structures that

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appear to be ephippia, (Fig. S10-B), the protective cases enclosing diapausing eggs produced by zooplankton such as *Daphnia*. Ephippia can overwinter in lake sediments or survive periods of desiccation, providing a seed bank to recolonize

565 the water column when <u>favourable</u> conditions return <u>(Caceres and Tessier, 2003; Hairston, 1996)</u>. These biological findings further support the finding that bubbles are transporting sediment particles through the profundal water column. <u>There</u> remains the possibility that our measured bubble particle transport rates differ significantly from those from naturally emitted <u>bubbles</u>, and this remains an important area for future research. However, despite this uncertainty, broad-scale estimates of arsenic and cyanobacteria cycling can provide important context as to whether these processes may be significant in UML.

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#### 3.4 Implications for arsenic and heavy metal cycling

The presence of arsenic and other heavy metals in the bubble-transported particles could have significant implications for chemical cycling in aquatic ecosystems. Measured rates of arsenic flotation in field samples were about  $8 \pm 4 \mu g$  arsenic L<sup>-1</sup> of gas bubbled (Fig. 4a). Typical natural bubble flux for UML was estimated as  $0.02 \pm 0.02$  L m<sup>-2</sup> day<sup>-1</sup> during previous ebullition studies (Varadharajan, 2009), which corresponds to a potential arsenic flux of  $0.2 \pm 0.2 \mu g$  m<sup>-2</sup> day<sup>-1</sup> from the sediment to the lake surface. This flux would be highly episodic given the spatial and temporal heterogeneity of methane bubbling in UML (Varadharajan, 2009; Scandella et al., 2016).

This estimate of daily arsenic flux can be compared with historical measurements showing significant arsenic accumulation within the epilimnion at rates exceeding 30  $\mu$ g m<sup>-2</sup> day<sup>-1</sup> (Knauer et al., 2000). This flux is two orders of magnitude larger than our estimate for bubble transported arsenic of 0.2  $\mu$ g m<sup>-2</sup> day<sup>-1</sup>, indicating that bubble-arsenic transport may be of relatively low importance in UML where a significant fraction of the arsenic input to epilimnetic waters can be attributed to inflow from the Aberjona River (Hemond, 1995). However, bubble-mediated fluxes of arsenic or other sediment-borne metals may represent a larger fraction of epilimnetic input in other lakes having lower influx rates from surface water inflow<sub>w</sub> or other external sources, such as atmospheric deposition (Csavina et al., 2012). In addition, particles

585 transported by bubbles are deposited directly at the water surface, potentially positioning them for easy ingestion by recreational swimmers.

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<b>Deleted:</b> related to arsenic cycling in UML (UML has not been extensively studied for chromium and lead cycling). In 2000, for example, the
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Although bubble-facilitated transport does not appear to dominate arsenic transport in UML, much higher ebullition

610 rates have been reported elsewhere in the world (Deemer et al., 2016). For example, a mid-latitude reservoir in Switzerland was reported to have an order of magnitude higher ebullition flux [0.225 L m<sup>-2</sup> day<sup>-1</sup> (Delsontro et al., 2010)]. Co-occurrence of high ebullition rates and contaminated sediment could lead to significant bubble-facilitated contaminant cycling.

#### 3.5 Implications for cyanobacteria transport and possible bloom initiation

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Since cyanobacteria are known to overwinter in lake sediments, bubble-mediated transport could be one mechanism by which resting cells inoculate the upper water column. <u>Bubble column experiments showed that bubble-transported</u> <u>particulate matter contained cells at</u> approximately 30 cells mL<sup>-1</sup> gas, indicating that bubbles are capable of transporting cyanobacteria through deep water columns. We also measured cyanobacteria transport in the field with bubble traps, but our measurements were contaminated by cyanobacteria in the surrounding water column (see SI for results and discussion).

- 620 While we could not directly measure bubble cyanobacteria transport in the field, we can estimate it using a combination of measured bubble particle flotation rates, and the average cyanobacteria cell concentration in lake sediment. Estimated cell transport using this method is  $880 \pm 1140$  cyanobacteria cells mL<sup>-1</sup> of bubble volume. Although this is significantly higher than the measurements made in the bubble column, the conditions in the column are substantially different from the conditions in the field and the sediments used in column had been stored for 8 months, so the cyanobacteria cell
- 625 concentration was 10 times less than fresh sediments. While this variability in cell transport between column measurements and estimates of potential field transport highlights the need for continued research, it is useful to estimate the potential range of cyanobacterial transport.

To assess the likelihood that bubble-mediated cell transport could significantly inoculate surface waters, we use the upper transport estimate of 880 ± 1140 cyanobacteria cells mL<sup>-1</sup> of bubble volume and the bubble flux estimate mentioned previously of 22 ± 20 mL m<sup>-2</sup> day<sup>-1</sup> to estimate a daily transport of  $2 \times 10^4 \pm 3 \times 10^4$  cells m<sup>-2</sup> day<sup>-1</sup>. If cyanobacteria cells concentrate within the upper 1 meter of the lake, <u>outcompeting other phytoplankton species for sunlight (Xiao et al., 2018)</u>, this results in an increase in concentration of 20 ± 30 cells L<sup>-1</sup> day<sup>-1</sup>. While this concentration is not a human health concern,

such a concentration from the average rate of bubbling could represent a significant inoculum. At a maximum growth rate of

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Deleted: We estimated the concentration of cvanobacteria cells in sediments and associated with bubble-transported particles with quantitative polymerase chain reaction (qPCR), and then used bubble column experiments to demonstrate cyanobacteria cell transport via bubble flotation. The column was filled with tap water to minimize cyanobacteria contamination from the surrounding water column. Bubbles were emitted directly under the bubble trap as a control, or within a sediment bed placed at 6 or 13 m below the surface. Prior to bubbling, the water column cyanobacteria cell concentration was less than 1 cell mL-1 of water (Fig. 5). After bubbling air through the sediment at the bottom of the column, the cyanobacteria concentration in the column increased to an average of approximately 7 cells mL<sup>-1</sup>, but still remained lower than the concentration of cells in the trap (Fig 5). Bubble-transported particulate matter contained cells at a rate of

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Deleted: Since bubble particle transport rates varied across experiments (from  $0.01 \pm 0.006$ mg/mL in the bubble column to  $0.09 \pm 0.07$  mg/mL on October 2017 and  $0.01 \pm 0.01$  mg/mL on June 2018 in the field), and cell flotation rates are likely tied to total particle transport rates, we conclude that the best estimates of transport of cyanobacteria cells in the field should be based on average estimates of particle transport per L gas across all column and field experiments and the average cyanobacteria cell concentration in the sediment. From an average sediment transport rate across all field and column experiments and an average sediment concentration, we can extrapolate that approximately 8.8 x 105 ± 11.4 x 105 cyanobacteria cells are transported per L bubble volume in the lake. Bubble-mediated transport could be a significant mechanism of

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**Deleted:**  $1.9 \times 10^4 \pm 3.1 \times 10^4$  cells m<sup>-2</sup> day<sup>-1</sup>. Harmful cyanobacterial species, such as *Microcystis*, often aggregate in the surface layers to outcompete other algae for sunlight (Xiao et al., 2018).

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Microsoft Office User 2/17/20 12:31 PM Deleted: <sup>1</sup> approximately 1 day<sup>-1</sup> (Robarts and Zohary, 1987) and absent significant losses, this cell concentration alone would result in 680 a cell density greater than the Massachusetts Health limit of 7 x 10<sup>4</sup> cells mL<sup>-1</sup> in about three weeks. Larger bubbling events [e.g. (Deemer et al., 2016)] could result in the same cyanobacteria cell concentration within approximately 15 days. The estimated growth of bubble-transported cyanobacterial cells is dependent on the cells being viable. Incubating cells to assess viability will be an important step for future studies. These calculations demonstrate that bubble transported cyanobacteria could negatively impact water quality, though more research is warranted to improve these estimates.

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Given the potential impact on bloom formation, we compared this source of cells to other pathways of cell recruitment to the lake surface, especially in deep, stratified lakes like UML. Cyanobacteria are thought to largely be recruited to surface waters from shallow areas due to a combination of higher light, temperature, and oxygen levels that promote germination, and increased wind-driven sediment resuspension (Ramm et al., 2017). While sediment cyanobacteria concentrations are higher in deeper areas of the lake, cells are not able to germinate because of the dark, anoxic conditions in

- 690 deep, eutrophic lakes (Ramm et al., 2017). Bubble-mediated transport is a mechanism by which this large reservoir of "lost" cells in deep sediments could contribute to overall recruitment to the surface waters. To determine the potential contribution of bubble mediated transport to cyanobacteria recruitment to the surface, we assume that germination does not occur significantly past the oxycline (12 m) in UML between June and Oct., as low oxygen concentrations and low light levels prevent germination, and wind-driven mixing cannot resuspend sediments across the shallow thermocline (Varadharajan,
- 695 2009). We also assume that cells resuspended in the spring overturn in March would have germinated, settled, lysed or have been consumed by grazers by June [e.g. (Tijdens et al., 2008; Verspagen et al., 2005)]. Furthermore, we do not include external inputs of cyanobacteria to the lake, such as from the river [e.g. (Bouma-Gregson et al., 2019)] or air (Seiffried et al., 2015; Lewandowska et al., 2017; Evans et al., 2019). Since literature estimates of recruitment rates for these sources are lacking, we assume these inputs are small compared to shallow sediment and bubble-mediated recruitment. Using the

700 maximum observed recruitment rate of 2.3 x 10<sup>5</sup> cells m<sup>-2</sup> day<sup>-1</sup> (Brunberg and Blomqvist, 2003) from sediments for the area of the lake above 12 meters, we estimate that bubbling could contribute 14% of cyanobacterial recruitment in the lake, but 95% confidence intervals range from less than 0 to 46% of overall recruitment. While we cannot rule out the possibility that Microsoft Office User 2/17/20 12:31 PM **Deleted:** (Deemer et al., 2016)

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#### 4 Conclusions

Bubble-particle transport between the sediment and surface of UML is a novel transport pathway capable of moving particulate matter upwards through a stratified water column, over depths of 15 m or greater, without shedding a major

- 725 fraction of their particle burden or accumulating large amounts of additional particles as they rise. Bubble-facilitated metal transport in present-day UML appears minor compared to surface inflows, but lakes with higher ebullition flux or more contaminated surficial sediments may experience more significant chemical transport from contaminated sediments. Bubble mediated transport of cyanobacteria cells may contribute <u>substantially</u> to cellular recruitment from the sediment, <u>but the uncertainties in our measurements make these estimates speculative</u>. Bubble transport is expected to be particularly
- 730 important in deep, eutrophic lakes in which alternative mechanisms of sediment regeneration to surface waters are limited. Further work is warranted to more thoroughly quantify this ebullitive transport pathway, and its implications for chemical and biological cycling. In addition, future work should include alternative methods of bubble triggering as well as the quantification of particle transport rates on naturally-occurring bubbles.

#### 735 Code/Data availability

All data necessary to validate the research findings are available on JHU Dataverse, doi.org/10.7281/T1/7WXPIN.

#### Author contributions

KD, HH and SP designed the experiments and KD and JG carried out experimental work. KD, JG and SP analyzed the data.

KD prepared the manuscript with contributions from all co-authors.

#### 740 Competing interests

The authors declare that they have no conflict of interest.

### Acknowledgements

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Microsoft Office User 2/17/20 12:31 PM Deleted: thereby inoculating epilimnetic waters of stratified water bodies with cyanobacterial cells, at rates capable of triggering significant blooms in a few weeks 750 Funding for part of this work was supported by Johns Hopkins Whiting School of Engineering. Funding was also provided

by the Singapore-MIT Alliance for Research and Technology, MIT Center for Environmental Health Science, the MIT

Martin Family Fellowship to K. Delwiche, the W. E. Leonhard 1941 professorship to H. Hemond, and the National Science

Foundation under grant number EAR-1045193. This work was also supported in part by the NIEHS Superfund Basic

Research Program, NIH, P42 ES027707.

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Figures and Legends



Figure 1: Total particle mass in mg associated with the bubbles captured during each field campaign with bubble triggering events in Oct. 2017 (filled circles) and June 2018 (open circles). Triggering events yielded different bubble volumes (given 925 in mL).



**Figure 2:** Transported particle mass per L of gas bubbled in the large bubble column, as a function of bubble release depth. Solid circles represent samples where bubbles were emitted from the sediment bed, diamonds represent samples where gas was bubbled directly above the sediment bed. Hollow circles around solid circles denote samples with recently-disturbed sediments.

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935 Figure 3: Comparison between mass of arsenic, chromium, and lead per kg of sediment (open triangles) and bubbletransported particulate matter (solid circles). Standard deviation scale similar to point size and therefore omitted for figure clarity.



Figure 4: Chemical amounts observed in bubble traps associated with bubble-mediated transport of sediment particles. (a)
Arsenic mass, (b) chromium mass, (c) lead mass (in µg) transported versus the bubble volume of each sample (in mL, as
measured at the lake surface a-c). Standard deviation is added to each measurement but the scale similar to point size for most measurements.



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Figure 5: The concentration of cyanobacteria cells (as measured by quantitative PCR) increases in the experimental water column and bubble traps after initiating bubbling within sediments. The background concentration of cyanobacteria cells in water in the column was initially very low (Before bubbling) but increased after bubbling. The concentration of cells in the bubble trap increased because of the contamination from the surrounding water column, even without bubbling within sediment (Just air). However, the highest concentration of cyanobacteria in the bubble trap was observed when initiating bubbling from within the sediment (Bubble transport). The increase in cell concentration in both the water column and the bubble trap after bubbling within sediment is evidence for cyanobacteria transport via bubble floatation. Error bars show standard deviation across measurements.

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Bubble column experiments show bubble particle transport from 15 m depth of  $0.01 \pm 0.006$  mg/mL in the

bubble column, compared to  $0.09 \pm 0.07$  mg/mL on October 2017 and  $0.01 \pm 0.01$  mg/mL on June 2018 in

the field. This rate reflects transport to the lake surface, but total bubble-particle transport rates would also

include particles that were transported part-way up the bubble column and then shed from the bubble.

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