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4	delays sulphide formation in the sediment and transfer to the bulk liquid in a model
5	shrimp pond
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7	Running title: Long-term inhibition of sulphate reduction through molybdate
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26 Abstract

Shrimp are commonly cultured in earthen aquaculture ponds where organic-rich uneaten feed 27 28 and faeces accumulate on and in the sediment to form anaerobic zones. Since the pond water is 29 rich in sulphate, these anaerobic conditions eventually lead to the production of sulphide. 30 Sulphides are toxic and even lethal to the shrimp that live on the pond sediment, but 31 physicochemical and microbial reactions that occur during the accumulation of organic waste 32 and the subsequent formation of sulphide in shrimp pond sediments remain unclear. Molybdate 33 treatment is a promising strategy to inhibit sulphate reduction, thus, preventing sulphide 34 accumulation. We used an experimental shrimp pond model to simulate the organic waste accumulation and sulphide formation in a long-term experiment (61 days) during the final 61 35 36 days of a full shrimp growth cycle. Sodium molybdate (5 and 25 mg/L Na₂MoO₄.2H₂O) was 37 applied as a preventive strategy to control sulphide production before oxygen depletion. Molybdate addition **partially** mitigated H₂S production in the sediment, and delayed its transfer 38 39 to the bulk liquid by pushing the higher sulphide concentration zone towards deeper sediment 40 layers. Molybdate treatment at 25 mg/L significantly impacted the overall microbial community 41 composition and treated samples (5 and 25 mg/L molybdate) had about 50% higher relative 42 abundance of sulphate reducing bacteria than the control (no molybdate) treatment. In 43 conclusion, molybdate worked has the potential to work as long-term mitigation strategy against sulphide accumulation in the sediment during shrimp growth by directly steering the 44 45 microbial community in a shrimp pond system. 46

47 Keywords: Aquaculture, molybdate, shrimp growth, sulphate reduction, sulphide toxicity

49 **1. Introduction**

50 The properties of pond bottom soil (sediment) and physicochemical and microbial interactions 51 on and in the sediment are crucial for the well-being and growth of the shrimp in aquaculture 52 ponds (Avnimelech and Ritvo, 2003; Burford et al., 1998). Sediments contain indigenous 53 nutrients and organic matter, derived directly from the environment, but also from uneaten and 54 digested feed of the numerous shrimp that dwell on the pond bottom, especially during semi-55 intensive and intensive stocking (50-300 shrimp/m³) (Avnimelech and Ritvo, 2003). This pond 56 bottom layer, *i.e.*, the interphase between the water and sediment, is an area that is densely 57 populated by microorganisms consuming the available organic matter. Due to the organic-rich 58 conditions on the pond bottom in combination with the typical temperatures of 25-30 °C in 59 shrimp ponds, the oxygen consumption by these microorganisms can cause a rapid drop in 60 dissolved oxygen in the sediment (Baxa et al., 2021; Dien et al., 2019). When oxygen 61 consumption exceeds the rate of oxygen transfer from the pond water phase to the sediment, 62 eventually sediment oxygen is depleted, and anaerobic conditions arise. Due to high sulphate 63 concentrations in the pond water, low redox conditions in the pond lead to production of 64 hydrogen sulphide (H₂S) from metabolic activity of sulphate reducing bacteria (SRB) (Avnimelech and Ritvo, 2003; Boyd, 1998). The H₂S formed creates a bad odour and black 65 colour in the sediment, and is also toxic to the shrimp that dwell at the pond bottom. Sulphide 66 67 toxicity to shrimp depends on both the H₂S concentration and pH (Thulasi et al., 2020; 68 Vismann, 1996), with lethal concentrations to kill 50% of the population (LC50) values ranging 69 between 0.0087 and 0.033 mg/L H₂S, depending on shrimp species and growth phase of the 70 shrimp (Chen, 1985; US-EPA, 2011). Exposure to sub-lethal concentrations of H₂S lowers 71 shrimp resistance to diseases and causes tissue corrosion (Suo et al., 2017). Overall, H_2S is 72 often the main cause for mortality or abnormal behaviour of shrimp, and may strongly impact 73 shrimp harvest (Panakorn, 2016).

74 Sulphide accumulation in shrimp ponds conventionally relies on labour-intensive, time-75 intensive and costly approaches, such as mechanical removal of reduced sediment or change of 76 culture water. An alternative approach is nitrate amendment, which has been shown to remove 77 the H₂S produced (Torun et al., 2020). However, as also demonstrated for sodium percarbonate, 78 the effect of nitrate towards sulphide removal was only transient, because when nitrate was 79 depleted, the H₂S production recovered (Schwermer et al., 2010; Torun et al., 2022; Torun et 80 al., 2020), requiring higher amounts of nitrate addition to compete with sulphate reduction. 81 Repeated and/or increased addition of nitrate is unwanted, because this may result in 82 cyanobacteria and algal blooms or the release of toxic metabolites, *e.g.*, nitrite or nitrous oxide. 83 A more targeted, preventive approach that achieves direct inhibition of the SRB, thus, 84 preventing sulphide production, is the application of molybdate (MoO₄). Because of its 85 stereochemical similarity to sulphate, molybdate inhibits the adenosine triphosphate 86 sulfurylase, which is the first enzyme in the sulphate reduction pathway (Peck, 1959; Stoeva 87 and Coates, 2019). Successful inhibition of sulphate reduction through the addition of 88 molybdate has been observed in studies on eutrophic lake sediments (Smith and Klug, 1981), 89 anaerobic digestion (Isa and Anderson, 2005; Ranade et al., 1999), and oil production systems 90 (Jesus et al., 2015; Kögler et al., 2021). Hence, its application in aquaculture systems also 91 warrants possibilities towards preventing H₂S formation in pond sediments. This was 92 demonstrated in a short-term experiment with a shrimp pond model in which molybdate 93 outperformed nitrate and sodium percarbonate in controlling H₂S formation, because of its 94 specificity and preventive mode of action (Torun et al., 2022). The applicability of molybdate 95 as a remediation strategy towards sulphide formation in aquaculture, however, strongly depends 96 on its long-term lasting effect during a 90-days shrimp growth cycle. 97 The objective of this study was to determine the long-term duration and magnitude of the

98 effect of molybdate towards H₂S mitigation in response to the gradual accumulation of organic

99 waste during a full shrimp growth cycle. A shift in the microbial community towards different 100 processes than sulphate reduction, in response to molybdate, could be beneficial to control 101 sulphide accumulation at the shrimp pond bottom. Because the accumulation of organic waste 102 was limited during the first 30 days of the shrimp growth cycle, *i.e.*, no O₂ depletion or H₂S 103 accumulation (Torun et al., 2023), only the final 61 days were considered in a lab-scale shrimp 104 pond bottom model.

105 **2. Material and methods**

106 2.1. Sampling and storage

107 2.1.1. Sediment sampling

Sandy clay and organic-rich sediments were obtained from the Ijzermonding Nature Reserve (Nieuwpoort, Belgium) from a creek (51°8′45″ N/2°44′38″E) that was regularly water-logged with tidal movement. Sampling were taken by scooping the top 5-10 cm of the sediment into a closed plastic container in which they were transported to the laboratory. The pH, conductivity, total solids (TS) and volatile solids (VS) of the fresh sediment were analysed directly upon arrival in the laboratory. A sample for sulphate and molybdate analysis was stored at 4°C until analysis, and a sample for DNA extraction was stored at -20°C.

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116 2.1.2. Feed and faeces collection and storage

Fresh shrimp faeces were collected from the flush outlet of shrimp tanks in which whiteleg shrimp (*Litopenaeus vannamei*) at post-larvae stage were fed with CreveTec Grower 2 (CreveTec, Ternat, Belgium) at the Aquaculture and *Artemia* Reference Center (ARC), Faculty of Bioscience and Engineering, Ghent University, Belgium. The faeces were stored at 4°C until use to avoid organic matter degradation during storage. The pH and conductivity of the faeces were measured directly after collection. A sample for DNA extraction was stored at -20°C.

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124 2.2. Experimental set-up and operation

A system mimicking organic matter accumulation in the shrimp ponds was designed and constructed, as described earlier (Torun et al., 2022), using 250 mL size glass beakers (outer diameter 70 mm) containing a 3.5 cm sediment layer and 5 cm overlaying artificial seawater (Instant Ocean, Aquarium Systems, Mentor, OH, USA). The salinity of the artificial seawater was adjusted to 20 g/L, representing a common salinity in shrimp ponds (15-25 g/L), and containing approximately 1.5 g/L sulphate. To avoid excessive water evaporation, the beakers were put in a transparent plastic box with a non-airtight lid in a temperature-controlled room at $28 \pm 1^{\circ}$ C without active aeration. No artificial or natural light was foreseen to avoid the growth of microalgae and keep a focused approach towards sulphide formation and

133 growth of increatigat and keep a focused approach towards supplied formation and 134 oxygen depletion.

135 The experiment was started with an initial cumulative waste of 30 days of shrimp culture (DOC 136 30) in the form of feed (CreveTec Grower 2 shrimp feed) and faeces, which was considered 137 day zero of the experiment. The cumulative waste for shrimp culture was calculated based on the 0.003848 m² bottom surface area of the beakers using commercial daily shrimp feeding 138 139 tables (Table S1). Feed and faeces were added based on semi-intensive stocking of 50 140 shrimp/m². About 25% of input feed was assumed to be accumulating in the pond bottom, with 141 15% considered digested feed (faeces), and 10% as uneaten feed. After adding the initial waste 142 of DOC 30, the respective amount of shrimp feed and faeces, based on daily uneaten feed and 143 faeces, were supplemented every 2-3 days (Table S2). The amounts of supplemented feed and 144 faeces were increased every 15 days to adapt to the growth of the shrimp.

145 Two different concentrations of 5 (M5) and 25 (M25) mg/L of sodium molybdate (Na₂Mo₄.2H₂O, Sigma Aldrich, St. Louis, Mo., US), were compared with a control treatment 146 147 (no molybdate addition) for a long-term experiment of the last 61 days of a shrimp growth 148 cycle. Molybdate was supplemented in a single dose on day 0 of the experiment. Each treatment 149 was carried out in 6 biological replicates. Measurements of dissolved oxygen (DO), H₂S and 150 pH in the bulk liquid were performed every 2-3 days. These measurements were taken from the 151 water column, about 1 cm above the sediment-water interface, since H₂S in the bulk liquid is 152 the major concern for the shrimp that dwell on the pond bottom. Apart from bulk liquid 153 measurements, microscale gradient depth profiles of DO, H₂S and pH at the water-sediment 154 interface and throughout the sediment were measured, using a microelectrode, on day 16, 30,

155 44 and 61 from three replicates of each treatment. After each depth profiling measurement (day 156 16, 30, 44), one replicate from each treatment was sacrificed for the measurement of molybdate 157 and sulphate concentrations in the bulk liquid. From these sacrificial beakers, sediment samples 158 were taken, and stored at -20°C for microbial community analysis. At the end of the experiment 159 (day 61), all remaining replicates (3 replicates) were sampled for sediment and liquid samples. 160 Sediment samples were taken from the upper 1 cm of the sediment layer after carefully decanting the liquid part. The liquid samples were filtered over a 0.20 µm Chromafil[®] Xtra filter 161 162 (Macherey-Nagel, PA, USA), and stored at 4 °C, prior to analysis of sulphate and molybdate 163 concentrations. During each liquid sampling, the degree of water evaporation was determined 164 by recording the water depth. The molybdate and sulphate measurements were corrected with 165 the evaporation factor.

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167 2.3. Microelectrode measurements

168 Microscale depth profiles of O₂, pH and H₂S were recorded using commercial microelectrodes 169 (Unisense A.S. Denmark, tip sizes pH: 200 µm, H₂S: 100 µm, O₂: 100 µm), operated with a 170 motorized micromanipulator (Unisense A. S., Denmark). Microscale measurements were 171 always performed before other samples were taken and before adding fresh waste to avoid 172 disturbance of the water column and sediment. The oxygen profiles were measured at 200 µm 173 resolution. The pH and H₂S were simultaneously recorded with the same resolution at 200 µm 174 in the water-sediment interphase, and at lower resolution deeper in the sediment. The sensors 175 were calibrated following standard calibration procedures, as described earlier (Malkin et al., 176 2014). The H₂S was calibrated with a 3-5 point standard curve using an acidified Na₂S standard 177 solution (pH 3.5-4.0). The O₂ sensor was calibrated with a 2 point standard curve, using 100% 178 in air bubbled seawater for the DO at saturation at 28°C and argon bubbled seawater for DO 179 zero. The pH sensor was calibrated with 2 point calibrations using commercial (Carl Roth GmbH & Co.KG, Karlsruhe, Germany) pH buffer solutions (4, and 7). Total sulphide concentrations were calculated as described earlier (Jeroschewski et al., 1996). For bulk liquid measurements, the same electrodes were used manually to take the measurements from approximately 1 cm above the sediment surface after ensuring that there was negligible variety in the duplicate measurements of the water column parameters.

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186 2.4. Analytical techniques

187 The TS and VS of the sediment were determined according to Standard Methods (Greenberg et 188 al., 1992). The pH of the overlaying water and sediment samples were measured with a pH 189 meter (Metrohm, Herisau, Switzerland), which was calibrated using pH buffer solutions at pH 190 4 and 7. The sulphate concentrations were measured through ion chromatography (930 191 Compact IC Flex, Metrohm, Herisau, Switzerland), equipped with a Metrosep A supp 5-192 150/4.0 anion column with conductivity detector, after diluting the samples 1:50 using ultra-193 pure water (Milli-Q, Millipore Corporation, Burlington, MA, USA). The detection range was 194 0.05 to 200 mg ion/L. Molybdate was measured using a commercial kit (Hach, Model Mo-2, 195 USA), based on the colorimetric determination of molybdenum using mercaptoacetic acid (Will 196 and Yoe, 1953). Standard solutions of 0, 5, 10, 25, and 50 mg/L Na₂MoO₄.2H₂O were prepared 197 to determine the standard curve at 425 nm using a UV-Vis Spectrophotometer (WPA Lightwave 198 II, Thermofisher, USA).

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200 2.5. Microbial community analysis

201 2.5.1. Amplicon sequencing

To analyse the changes in the bacterial community and SRB relative abundance, samples were taken from the upper 1 cm of the sediment from each sacrificial beakers in 3 replicates and frozen at -20 °C. The DNA was extracted directly from the frozen samples using a commercial 205 kit (DNeasy Power Soil Pro Kit, QIAGEN, Hilden, Germany), following the instructions of the 206 manufacturer. The quality of the DNA extracts was evaluated through agarose gel 207 electrophoresis and PCR analysis with the universal bacterial primers 341F (5'-208 CCTACGGGNGGCWGCAG) and 785Rmod (5'-GACTACHVGGGTATCTAAKCC) that 209 target the V3-V4 region of the 16S rRNA gene (Klindworth et al., 2013), following a PCR 210 protocol as described earlier (Boon et al., 2002). The samples were sent to LGC Genomics 211 GmbH (Berlin, Germany) for Illumina amplicon sequencing of the V3-V4 region of the 16S 212 rRNA gene of the bacterial community on the MiSeq platform with V3 chemistry. The 213 amplicon sequencing protocol and data processing are described in detail in the SI (S3).

214

215 2.5.2. Flow cytometry analysis

216 Absolute microbial cell counts in the sediment samples were determined using flow cytometry 217 (FCM). Prior to FCM analysis, sediment samples were defrosted, acclimated to room 218 temperature and diluted tenfold in sterile, 0.22 µm-filtered Instant Ocean® solution. To separate 219 the cells from sediment particles, samples were initially sonicated (O700 Sonicator, Osonica, 220 Newtown, CT, USA) for 3 minutes, followed by 3 minutes centrifugation at 500 g. The resulting 221 supernatant of the samples was stained with 1 vol% SYBR® Green I (SG, 100x concentrate in 222 0.22 µm-filtered DMSO, Invitrogen), and incubated in the dark at 37°C for 20 min. Immediately 223 after incubation, samples were analysed using a BD Accuri C6 Plus cytometer (BD Biosciences, 224 Erembodegem, Belgium), equipped with four fluorescence detectors (533/30 nm, 585/40 nm, 225 > 670528 nm and 675/25 nm), two scatter detectors and a 20-mW 488-nm laser. Samples were 226 analysed in fixed volume mode (30 µL). The flow cytometer was operated with Milli-Q water 227 (MerckMillipore, Belgium) as sheath fluid, and instrument performance was verified daily 228 using CS&T RUO Beads (BD Biosciences, Erembodegem, Belgium).

230 2.6. Statistical analysis

231 A table containing the relative abundances of the different OTUs (operational taxonomic units), 232 and their taxonomic assignment was created following amplicon data processing 233 (Supplementary Information File 2). All statistical analysis were carried out in R Studio version 234 4.03 (http://www.r-project.org) (R Development Core Team, 2013). A repeated measures 235 analysis of variance (ANOVA, *aov* function) was used to validate that the biological replicates 236 showed no significant (P < 0.05) differences in bacterial community composition. Next, 237 absolute singletons were removed, and the different samples were rescaled via the "common-238 scale" approach (McMurdie and Holmes, 2014) through which the proportions of all OTUs 239 were taken, multiplied with the minimum sample size, and rounded to the nearest integer. 240 Sampling depth of each sample was evaluated through rarefaction curves (Figure S1) (Hurlbert, 241 1971; Sanders, 1968). The packages vegan (Oksanen et al., 2016) and phyloseq (McMurdie and 242 Holmes, 2013) were used for microbial community analysis. A heatmap was created at the 243 phylum and family level (1% cut-off) with the *pheatmap* function (pheatmap package), and 244 biological replicates were collated according to the method described earlier (Connelly et al., 245 2017). The non-metric multidimensional scaling (NMDS) plots were constructed using the 246 Bray-Curtis (Bray and Curtis, 1957) distance measures. Significant differences between 247 treatments and timepoints were identified using pairwise permutational ANOVA 248 (PERMANOVA) analysis (9999 permutations) with Bonferroni correction, using the adonis 249 function (vegan).

250 **3. Results**

251 3.1. Impact of molybdate on oxygen depletion and sulphide production

252 3.1.1. Bulk liquid concentrations

253 The DO measurements in the bulk liquid showed that oxygen was completely depleted for the 254 first time on day 7 for all treatments (Figure 1a). In the following days, there was a fluctuation 255 in oxygen concentrations, *i.e.*, between 0 and 150 µM, for all treatments, with a regular oxygen 256 re-introduction in the bulk between day 7 to 35, keeping in mind that no active aeration was 257 applied. After day 35 of the experiment, a full oxygen depletion was observed for all treatments 258 with only limited oxygen re-introduction, resulting in oxygen concentrations only up to 50 µM. 259 For the entire experimental period, the DO concentration in the bulk did not show clear 260 difference between the different treatments, and also pH remained similar in the different 261 treatments (Figure S2).

262 No clear H₂S production was observed, *i.e.*, H₂S concentrations did not exceed 10 µM, in the 263 bulk liquid until day 35 of the experiment, coinciding with the time when oxygen depletion for 264 all incubations was recorded (Figure 1b). On day 35, a H₂S concentration of 64 ± 7 , 53 ± 11 , 265 and $39 \pm 3 \mu M$ was recorded in the bulk liquid for the control, M5, and M25 treatments, 266 respectively. This corresponded with a total bulk sulphide concentration that was 22 ± 1 % and 267 46 ± 1 % lower than the control for M5 and M25, respectively (Figure S3). Hence, there was 268 markedly lower H₂S production in the molybdate treatments compared to the control treatment, 269 especially in the M25 treatment. Also on day 56 of the experiment, H₂S concentration in the 270 bulk was clearly higher in the control treatment $(16 + 5 \mu M)$ compared to the M5 $(7 + 6 \mu M)$ 271 and M25 $(5 + 5 \mu M)$ treatments.

Residual molybdate measurements indicated that about 53 ± 1 % of the dosed molybdate disappeared from the bulk liquid for M25 at the end of the experiment, while this ranged between 5-15% for M5. (Table 1). Hence, in both molybdate treatments, residual molybdate 275 remained present. Residual sulphate concentration in the bulk gradually decreased throughout
276 the experiment, yet, no clear differences could be observed between the treatments (Table 2).

277

278 3.1.2. Sediment profiles

279 Microscale depth profiles of the DO in the sediment on day 16 revealed a 12 ± 4 % higher 280 concentration of oxygen in the M25 compared to the control treatment at the water-sediment 281 interface (Figure 2). Oxygen diffusion into the first 2 mm of the upper sediment layer was 282 observed for all treatments, with negligible differences between the treatments. On day 30, the 283 DO depth profile showed no apparent difference between the different treatments On day 44 284 and 61, no more oxygen was detected in the bulk liquid or sediment.

285 Microscale depth profiles of H₂S in the sediment were recorded on day 16, 44 and 61, while 286 day 30 gradient measurements of H₂S could not be obtained, due to technical problems with 287 the microelectrode (Figure 3). On day 16, although no H₂S could be observed in the bulk liquid, 288 H₂S gradient measurements showed a minor H₂S production in the control reaching a 289 concentration up to $5.8 \pm 0.1 \,\mu\text{M}$ at the sediment depth of 3.6 mm, while for the M5 and M25 290 treatments, no H₂S production was observed. On day 44, the control treated showed a maximum 291 H₂S concentration of $66.6 \pm 20.5 \,\mu$ M in the sediment deeper layers (sediment depth of 6.0 mm). 292 The M5 and M25 treatments showed a maximum H₂S concentration of only $22.7 \pm 4.2 \,\mu$ M and 293 $29.3 \pm 26.8 \,\mu\text{M}$ H₂S, respectively, both at 7.2 mm sediment depth, being 69 ± 11 and 60 ± 39 294 % lower than the maximum value in the control treatment, respectively. On day 61, the H_2S 295 concentration in the sediment was more similar between the different treatments, in contrast to 296 day 16 and 44. The M5 and M25 treatments showed a 17 ± 9 % and 26 ± 15 % lower maximum 297 H₂S concentration compared with the control, respectively. The H₂S production zone appeared 298 to be pushed to deeper sediment layers in the M5 and M25 treatments both for day 44 and 61 299 measurements, compared to the control. Total S microscale depth profiles showed a similar

300 pattern as the H₂S profiles (Figure S4), with the M5 and M25 treatments showing a markedly 301 lower total S concentration in the sediment in comparison with the control treatment. The pH 302 microscale depth profiles were similar between the different treatments, with limited variation 303 in function of time (Figure S5).

304

305 3.2. Microbial community analysis

Amplicon sequencing of the bacterial community resulted in an average of $23,917 \pm 9,397$ reads, which represented $2,922 \pm 876$ OTUs per sample (including singletons). Removal of absolute singletons and rescaling through the "common-scale" approach resulted in an average of $6,031 \pm 282$ reads and 681 ± 171 OTUs per sample. Repeated measures ANOVA revealed no significant differences (*P* < 0.0001) between the biological replicates.

311 Shrimp faeces (day 0) were dominated by Bacteroidota (35.9 \pm 7.8 %) and Fusobacteriota (35.1 312 \pm 5.7%) phyla, while the sediment used in the experiment was dominated by Actinobacteriota 313 $(22.3 \pm 8.8\%)$ and Proteobacteria $(33.5 \pm 1.6\%)$ phyla (Figure 4). The sediment samples (day 314 0) showed a relatively higher abundance of the phylum Desulfobacterota ($4.2 \pm 0.3\%$), which 315 contains several sulphate reducers, than the shrimp faeces $(0.2 \pm 0.0 \%)$ samples. Over time, 316 there was a clear shift in the bacterial communities, specifically for Proteobacteria, Bacteroidota 317 and Desulfobacterota relative abundances for all treatments. On day 16, the control, M5 and 318 M25 treatments showed markedly high relative abundances of Proteobacteria ($29.3 \pm 3.2 \%$, 27.2 ± 5.8 %, 30.7 ± 0.4 %, respectively). On day 30 and 44, the M5 and M25 treatment showed 319 320 an even further increase in the relative abundance of Proteobacteria (35.6 \pm 6.7 % and 34.7 \pm 321 6.2 % for day 30 and 34.2 ± 8.1 % and 38.0 ± 0.2 % for day 44, respectively), compared to the 322 control treatment (26.6 \pm 4.0 % for day 30, 22.8 \pm 3.1 % for day 44). This higher relative 323 abundance of Proteobacteria in the M5 and M25 treatments coincided with a more prominent 324 presence of the Rhodobacteraceae family (Figure S6) in the M5 ($16.1 \pm 4.5 \%$) and M25 (14.5

325 \pm 7.2 %) treatments, compared to the control (8.3 \pm 0.6%), on day 30 and later timepoints in 326 the experiment. The M5 (15.3 \pm 3.2 %) and M25 (16.1 \pm 4.1 %) treatment showed an overall 327 higher relative abundance of Flavobacteriaceae than the control treatment $(12.3 \pm 2.7 \%)$. The 328 high relative abundance of Bacteroidota (Figure 4) likely originated from the addition of faeces, 329 and reached values of 27.9 ± 1.2 %, 27.9 ± 0.8 % and 20.5 ± 1.8 % on day 16 for the control, 330 M5 and M25 treatments, respectively. However, in time, the relative abundance of Bacteroidota 331 decreased in all treatments $(22.3 \pm 2.4\%)$. There was no clear difference in Bacteroidota relative 332 abundance between the different treatments.

333 There was an apparent higher abundance of the Delsulfobacterota phylum, which contains 334 several sulphate reducers, in the M5 (15.3 \pm 3.1 %) and M25 (16.2 \pm 0.4%) treatments, 335 compared to the control treatment $(8.8 \pm 1.6\%)$ for all samples on day 16, 30, 44 and 61. The M25 treatment also showed a slightly higher abundance of this phyla compared to the M5 336 337 treatment. Family level analysis revealed that SRB species belonged to Desulfobulbaceae, Desulfolunaceae, 338 Desulfomonadaceae, Delsufovibrionaceae, Desulfosarcinaceae, 339 Delsulfobacteraceae and Desulfocapsaceae families (Figure S6). An absolute cell count 340 analysis of Delsulfobacterota phylum, by combining flow cytometry cell counts with amplicon 341 sequencing data, showed that all samples, including the samples treated with molybdate, 342 showed an increasing trend in time of absolute Delsulfobacterota cell counts. Molybdate treated 343 samples, especially M25, in general, showed even higher absolute cell counts for the 344 Desulfobacterota phylum compared to the control (Table 3).

The β -diversity analysis of the bacterial community, based on the Bray-Curtis distance measure, revealed that the M25 treatment showed an overall significantly different bacterial community composition than the control treatment (P = 0.0003) (Figure 5). However, none of the other treatments significantly differed (P > 0.05), and there was a limited impact of molybdate addition on the change of the bacterial community in function of time. The PERMANOVA

- analysis showed that there was a significant change in overall bacterial community composition
- 351 between day 16 and 30 (P = 0.0036), day 30 and 44 (P = 0.0174) and day 44 and 61 (P =
- 352 0.0066). On day 61, at the end of the experiment, the bacterial community composition for all
- 353 treatments showed a clear divergence.

354 **4. Discussion**

This study showed that molybdate addition, prior to H_2S formation, has a good potential to mitigate H_2S production in the sediment, and delay its transfer to the bulk liquid by pushing sulphide production zone in deeper layers of the sediment. Bacterial community analysis revealed a limited impact of molybdate addition on the change of the bacterial community in function of time. Molybdate treated samples did show a higher absolute abundance of the Desulfobacterota phylum compared to the control.

361

362 4.1. Molybdate effectively controls sulphide production and pushes higher sulphide363 concentration zones towards deeper sediment layers

364 The typical shrimp pond water with a salinity of 1.5-2.5 % contains about 1500 mg/L sulphate 365 (Torun et al., 2020). This high availability of sulphate and organic-rich conditions in the pond 366 bottom make the shrimp pond environment susceptible to the production of sulphides when 367 anaerobic conditions arise, due to the depletion of oxygen. The most effective method for 368 avoiding anaerobic conditions is to keep dissolved oxygen levels sufficiently high for the entire 369 depth of the pond water. However, mechanical aeration is usually applied on the water surface 370 (e.g., paddlewheel aerators). Apart from being costly and energy-consuming, these aerators 371 come with a risk of causing erosion in the pond bottom soil, when the water current is too 372 strong. Erosion degrades embankments, makes the harvest more difficult, and damages benthic 373 plants and animal communities, including the shrimp (Boyd, 1998). In a real pond system, 374 also the growth of microalgae could play a critical role, as they (1) enable in situ formation 375 of oxygen, and (2) by consuming CO₂, they could provoke an increase in pH, which could 376 reduce H₂S toxicity, but increase ammonia toxicity. They can even actively contribute to 377 an improved water quality (Huang et al., 2022). However, the direct involvement of microalgae in our model system would strongly add to the complexity of sulphide 378

formation, because of their multi-level impact on the shrimp pond nutrient dynamics, so we eliminated the possibility for photosynthetic growth from our model by not supplying natural or artificial light. Nitrate addition could serve as an alternative electron acceptor, in competition with sulphate. However, nitrate can only temporarily control sulphide production, and when nitrate is depleted, sulphide production recovers (Schwermer et al., 2010; Torun et al., 2022; Torun et al., 2020). These limitations substantiate the importance of a long-term lasting strategy to mitigate sulphide production in shrimp pond aquaculture systems.

386 In this study, 5 and 25 mg/L sodium molybdate clearly lower sulphide production in the 387 sediment, and pushed the higher H₂S concentration zone towards deeper sediment layers. Since 388 the transfer of H₂S from sediment to bulk liquid was delayed by this action, molybdate treated 389 samples had lower concentrations of H₂S in the bulk liquid on day 35 when peak concentrations 390 were observed. The H₂S concentration in the bulk liquid did fluctuate in function of time for all 391 treatments, which can be linked to the fact that the set-up used was an open system being 392 continuously exposed to the air inflow and disturbances created during the movement of 393 beakers for microelectrode measurements. The re-introduction of oxygen into the bulk liquid, 394 as confirmed by dissolved oxygen measurements, most likely re-oxidised a portion of the H₂S. 395 In addition, these external disturbances and the nature of the open system might have 396 accelerated H₂S to diffuse to the air from the bulk liquid. Alternatively, H₂S accumulated in the 397 bulk liquid might have reacted with ferrous iron to form iron sulphide that subsequently 398 precipitated in the sediment.

The inhibitory concentration of molybdate and a correlation between sulphate and molybdate concentrations were shown in several studies (Biswas et al., 2009; Chen et al., 1998; Jesus et al., 2015). In our previous study, we estimated that the inhibitory concentration for 1500 mg/L sulphate present in our experimental shrimp pond model should be approximately 15 mg/L of sodium molybdate for short-term prevention of H₂S production (Torun et al., 2022). In the 404 current study, the molybdate was only partially reduced, both in the M5 and M25 treatments, 405 and the production of H₂S in the sediment and its transfer to bulk liquid could not be fully 406 prevented in the long term experiment. This might be due to poor diffusion of the molybdate 407 in the deeper layer of the sediment, since in the upper layers of the sediment, there was markedly 408 lower H₂S concentration compared to the control treatment. The reason for molybdate having 409 potentially lower diffusion rates than the sulphate, might be related to adsorption of molybdate 410 on the sediment, as observed for pure quartz sand (Kögler et al., 2021). One can assume that 411 adsorbed molybdate could not inhibit microbial sulphate reduction.

412 In this study, residual sulphate concentrations did not show any apparent difference between 413 the control and molybdate treated samples, but these sulphate concentrations were measured in 414 the bulk liquid. The H₂S and total sulphide productions did show clear differences in the 415 sediment itself, with higher concentrations of sulphide in the control treatment, indicating the 416 effectiveness of molybdate to inhibit sulphate reduction. Residual sulphate in the bulk liquid 417 remained present in all treatments, so despite the high availability of organic matter, sulphate 418 reduction did not continue, as also observed in other studies on shrimp pond sediments (Torun 419 et al., 2022) and other anaerobic ecosystems, such as anaerobic digestion (Lippens and De 420 Vrieze, 2019). This apparent discrepancy was probably due to oxygen intrusion into the water 421 column, halting sulphate reduction in the bulk liquid. Hence, sulphate reduction might have 422 been locally interrupted in the bulk liquid, while it continued in the deeper layers of the 423 sediment.

Overall, it is clear that the biogeochemical sulphur cycle in such a pond system involves
various processes, i.e., sulphate reduction, sulphide/sulphur (re-)oxidation, precipitation
of metal sulphides, and production of polysulphides. Due to the nature of the open air
system (as is the case in real pond systems) in the current study, with the possibility of H₂S
escaping, it is not possible to make accurate sulphur mass balances.

430 4.2. Molybdate treatment changes the absolute abundance of sulphate reducing bacteria

431 When molybdate is provided in the presence of SRB, ATP sulfurylase uses molybdate (instead 432 of sulphate) and ATP to produce an unstable molecule equivalent to adenosine 5'-433 phosphosulfate (APS) that cannot be used as electron acceptor (Biswas et al., 2009). Under 434 molybdate excess, some studies indicated that SRB growth could be supressed altogether. 435 Kögler et al. (2021) showed that there were no SRB specific dsR genes isolated when molybdate 436 was continuously injected into sandpacks with residual oil in an oil reservoir. Nair et al. (2015) 437 reported that molybdate concentrations ranging between 50 and 150 µM increased the doubling 438 time of Desulfovibrio alaskensis G20, and 500 µM molybdate completely inhibited its cellular 439 growth. In the current study, molybdate was provided only once at lower concentrations than 440 the concentrations mentioned in the literature, but even such lower concentrations of molybdate 441 showed a promising impact towards decreasing sulphide concentrations in the bulk liquid.

442 A higher absolute abundance of the phylum Delsufobacterota, containing several SRB, was 443 detected in molybdate treated samples, despite the fact that mitigation of sulphide production 444 was observed. A similar trend was observed in an earlier study (Tenti et al., 2019), where SRB 445 counts in all samples from a lab-scale anaerobic digester, were similar with or without 446 molybdate, when molybdate concentration was lower than 1.2 mM. In this study, the detection 447 of SRB through 16S rRNA gene amplicon sequencing showed a relative increase of SRB, but 448 did not provide any information on their activity or absolute abundance. Hence, these relative 449 abundances were combined with the absolute cell counts, as obtained through flow cytometry 450 analysis to estimate absolute cell counts of the Desulfobacterota phylum. Such an approach has 451 been successfully applied in other ecosystems, and can be considered an established, reliable 452 way of quantifying microorganisms in environmental samples (Barr et al., 2021; Ou et al., 2017; 453 Props et al., 2017). An overall increasing trend in time was observed for the Desulfobacterota 454 phylum, including the samples treated with molybdate. The reason for the higher absolute 455 abundance of SRB, despite lower H₂S production, might be related to (partial) inactivation of 456 enzymes involved in sulphate reduction. In the study of Nair et al. (2015) on growth and 457 morphology of Desulfovibrio alaskensis G20, at least three important enzymes that play a 458 crucial role in energy production (alcohol dehydrogenase, pyruvate carboxylase, tungsten 459 formylmethanofuran dehydrogenase) showed downregulation or repression in the presence of 460 elevated molybdate concentrations. In the current study, molybdate treatment at 25 mg/L 461 showed an overall significantly different bacterial community composition compared to the 462 control without molybdate. The increase in SRB was unexpected, yet, next to sulphate 463 reduction, SRB can also carry out hydrogenic and/or acetogenic metabolisms. Hence, in the 464 absence of sulphate, many SRB can ferment organic acids or alcohol, producing hydrogen 465 acetate or carbon dioxide (Plugge et al., 2011; Zhang et al., 2022). The growth of Desulfovibrio 466 on lactate was reported in the absence of sulphate, in syntrophy with a methanogen (Bryant et 467 al., 1977), and the growth of the Delsufovibrionaceae family was also detected in this study. 468 Overall, a combination of reduced H₂S toxicity and the shift in the energy production 469 metabolism appeared to have increased the relative abundance of SRB in this study.

5. Conclusions

472	We showed that molybdate could be an effective mitigation agent against sulphide
473	accumulation in shrimp ponds as a long term strategy, since it can be applied in a single dose
474	and at relatively low concentrations. Although, sulphide production could not be avoided
475	completely, and only a temporal effect could be obtained, molybdate reduced H ₂ S productior
476	in the sediment, and delayed its transfer to the water column by pushing the sulphide production
477	zone towards deeper sediment layers. Molybdate induced a higher absolute abundance of
478	Desulfobacterota, but this was not reflected in increased sulphide formation. Overall
479	molybdate can has the potential to serve as a more environmentally friendly option compared
480	to other conventional strategies to mitigate sulphide production in shrimp pond systems.

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494

495 **Conflict of interest disclosure**

496 The authors declare they have no conflict of interest relating to the content of this article. Jo De497 Vrieze is a recommender for PCI Microbiology.

498

499 Data, script, code and supplementary material

500 The datasets generated and R scripts used during this research are included in this article, its 501 supplementary information, and were submitted to the Zenodo repository 502 (https://zenodo.org/doi/10.5281/zenodo.10149234). The raw fastq files that served as a basis 503 for the bacterial community analysis were deposited in the National Center for Biotechnology 504 Information (NCBI) database (Accession number SRP326102).

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617 **Tables:**

618 **Table 1** Molybdate concentrations of bulk liquid samples taken from the sacrificed replicates

on day 16, 30 and 44. At the end of the experiment (day 61), all three remaining replicates were

620 analysed, hence, the values for day 61 are average values and standard deviations of biological

621 replicates. M5 = Treatment with 5 mg/L molybdate addition. M25 = Treatment with 25 mg/L

622 molybdate addition.

	Molybdate concentration (mg/L)			
	Day 16	Day 30	Day 44	Day 61
Control	0.0	0.0	0.0	0.0 ± 0.0
M5	5.9	4.2	4.5	4.8 ± 4.3
M25	21.9	15.7	12.0	11.6 ± 0.7

Table 2 Sulphate concentrations of bulk liquid samples taken from the sacrificed replicates on day 16, 30 and 44. At the end of the experiment (day 61), all three remaining replicates were analysed, hence, the values for day 61 are average values and standard deviations of biological replicates. M5 = Treatment with 5 mg/L molybdate addition. M25 = Treatment with 25 mg/L molybdate addition.

	Sulphate concentration (mg/L)			
	Day 16	Day 30	Day 44	Day 61
Control	1396	1207	1283	1170 ± 206
M5	1334	1201	1261	1021 ± 75
M25	1389	1206	1351	1100 ± 78

Table 3 Absolute cell counts of the Desulfobacterota (10⁴ cells per mL), which contains several
sulphate reducing bacteria (SRB), as determined by combing flow cytometry cell counts with
amplicon sequencing data. The values are average values and standard deviations of biological
replicates. M5 = Treatment with 5 mg/L molybdate addition. M25 = Treatment with 25 mg/L
molybdate addition.

	Control	M5	M25
Day 16	10.4 ± 1.0	8.3 ± 0.9	19.1 ± 2.3
Day 30	15.2 ± 1.1	12.2 ± 1.9	20.5 ± 4.5
Day 44	21.9 ± 2.5	15.2 ± 0.4	23.7 ± 4.1
Day 61	24.3 ± 6.7	37.3 ± 6.8	29.1 ± 5.3



Figure 1 The bulk liquid concentrations of (a) O_2 and (b) H_2S in the control treatment, 640 molybdate treatment at 5 mg/L (M5) and molybdate treatment at 25 mg/L (M25). Values 641 represent averages of biological triplicates, and error bars represent the standard deviation.



Figure 2 The O₂ depth profiles for the (a) control treatment, (b) molybdate treatment at 5 mg/L
(M5) and (c) molybdate treatment at 25 mg/L (M25). Values represent averages of biological
triplicates, error bars are omitted to maintain the visibility of the graphs. Zero depth equals to
the sediment-water interface. On day 44 and 61, all O₂ values were below the detection limit.



Figure 3 The H₂S depth profiles for the (a) control treatment, (b) molybdate treatment at 5 mg/L (M5) and (c) molybdate treatment at 25 mg/L (M25). Values represent averages of biological triplicates, error bars are omitted to maintain the visibility of the graphs. Zero depth equals to the sediment-water interface. Because of technical problems with the microelectrode, data from day 30 are not included.



Figure 4 Heatmap showing the relative abundance of the bacterial community at the phylum
level in the faeces, the sediment and the different treatments on day 16, 30, 44 and 61. Weighted
average values of the biological replicates are presented. The colour scale ranges from 0 (white)
to 40% (red) relative abundance.





Figure 5 Non-metric multidimensional distance scaling (NMDS) analysis of the Bray-Curtis
distance measure of the bacterial community based on amplicon sequencing data at OTU level.
Different colours and symbols are used for different treatments and timepoints, respectively.