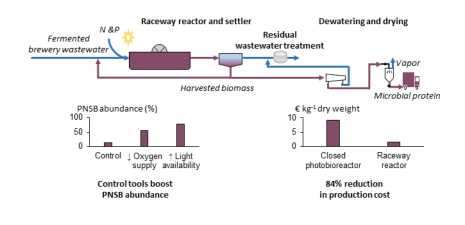
1	Control tools to selectively produce purple bacteria for microbial protein in raceway
2	reactors
3	
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# 15 Graphical abstract



17

# 18 Abstract

19 Purple non-sulfur bacteria (PNSB) show potential for microbial protein production on wastewater as animal feed. They offer good selectivity (i.e. uneven community with high 20 21 abundance of one species) when grown anaerobically in the light. However, the cost of a closed 22 anaerobic photobioreactor (PBR) is prohibitive for protein production. While open raceway 23 reactors are cheaper, their feasibility to selectively grow PNSB is thus far unexplored. This 24 study developed tools to boost PNSB abundance in the biomass of a raceway reactor fed with 25 volatile fatty acids as carbon source. For oxygen availability as tool, not stirring in the night 26 (i.e. reduced oxygen supply) elevated the PNSB abundance from 8% to 20%. For light availability as tool, a 24-h illumination increased the PNSB abundance from 8% to 31% 27 compared to a 12-h light/12-h dark regime. A reactor run at 2-d sludge retention time at the 28 highest surface-to-volume ratio (10 m<sup>2</sup> m<sup>-3</sup> increased light availability) showed productivities 29 up to 0.2 g protein  $L^{-1} d^{-1}$  and the highest PNSB abundance (78%). The estimated production 30 31 cost is €1.9 kg<sup>-1</sup> dry weight (vs. PBR €11.4 kg<sup>-1</sup> dry weight). This study pioneered in PNSBbased microbial protein production in raceways, yielding cost efficiency along with high 32 33 selectivity when avoiding the combined availability of oxygen, COD and darkness.

# 35 **1 Introduction**

36 Globally, only 4% of nitrogen and 17% of phosphorus fertilizers applied to the land, are eventually consumed.<sup>1,2</sup> These inefficiencies in the fertilizer-food chain severely distort the 37 38 carrying capacity of the Earth, surpassing the planetary boundaries (i.e. safe operating space for sustainability) beyond the zone of uncertainty.<sup>3</sup> Mitigation can be brought about by 39 40 upgrading wastewater resources to microbial protein or single-cell protein, which is the use of microorganisms as animal feed ingredient.<sup>4,5</sup> Resource recovery from food and beverage 41 42 wastewater is preferred. Brewery wastewater is a key target as it offers a relatively high chemical oxygen demand (COD) concentration (800-9000 mg COD L<sup>-1</sup>) and easiness to 43 prevent fecal contamination.<sup>6</sup> 44

Upgrading wastewater resources to microbial protein requires either chemo- or 45 photoheterotrophic microorganisms to convert the organic carbon as well as non-axenic 46 production conditions, as it is cost-wise redundant to sterilize vast amounts of water.<sup>4</sup> 47 Chemoheterotrophs, also known as aerobic heterotrophic bacteria (AHB; i.e. aerobic activated 48 sludge), make use of oxidation reactions for energy generation. These bacteria typically have 49 low biomass yields (0.6 g COD<sub>biomass</sub> g<sup>-1</sup> COD<sub>removed</sub>) and high growth rates (2-6 d<sup>-1</sup>).<sup>7</sup> To date, 50 AHB are pioneering in research, pilot, and full-scale implementations.<sup>8,9</sup> However, it is 51 challenging to produce an AHB product characterized by an uneven microbial community with 52 a high abundance of one dominant species (i.e. microbial selective production).<sup>8</sup> 53 54 Photoheterotrophic cultivation of PNSB may offer such potential, because of their unique ability to grow highly selectively under anaerobic conditions in the light.<sup>10-12</sup> PNSB are 55 characterized by high biomass yields (0.9-1.1 g COD<sub>biomass</sub> g<sup>-1</sup> COD<sub>removed</sub>) and have growth 56 rates between 0.6-3.7 d<sup>-1</sup>.<sup>13-15</sup> However, compared to AHB, there is a lack of full-scale PNSB 57

facilities for microbial protein production, probably due to costs (expensive photobioreactor;PBR).

To achieve selectivity with PNSB (i.e. uneven microbial community and high abundance 60 61 of one species), current research has focused on closed PBR such as anaerobic membrane bioreactors,<sup>16</sup> anaerobic tubular PBR<sup>13,17</sup> and illuminated anaerobic sequencing batch 62 reactors<sup>18</sup>. These closed PBR only allow the growth of phototrophs and anaerobic chemotrophs. 63 In the case of our previous study on synthetic wastewater, PNSB were able to be selectively 64 produced with an abundance of *Rhodobacter capsulatus* between 93-97% and a low diversity 65 index (exponent Shannon index) between 1.2-1.5 corresponding to an uneven microbial 66 community.<sup>19</sup> A current cost estimation for PNSB-based protein production on wastewater in 67 a closed PBR amounts to €22 kg<sup>-1</sup> protein,<sup>13</sup> which is roughly 11 times higher compared to the 68 price of fishmeal of  $\notin 2 \text{ kg}^{-1}$  protein.<sup>20</sup>. 69

An economically more interesting case can be made if PNSB were produced in open 70 raceway reactors conventionally used for microalgal processes.<sup>21</sup> These raceway reactors are 71 open systems with a reactor depth around 20 cm (vs. 6 cm diameter tubular PBR), a surface-72 to-volume ratio of 5 m<sup>2</sup> m<sup>-3</sup> (vs. 22 m<sup>2</sup> m<sup>-3</sup> tubular PBR) and are agitated through a paddlewheel 73 (vs. circulation pumps tubular PBR).<sup>21</sup> Investment costs of a raceway reactor approximate €56 74 m<sup>-3</sup> compared to €5,000 m<sup>-3</sup> for a closed PBR.<sup>22,23</sup> However, achieving selective PNSB 75 production is more challenging in these reactors, as air continuously enters the system, which 76 77 enables the proliferation of competing aerobic heterotrophs (i.e. non-PNSB). Moreover, the oxygen concentration in a raceway reactor is zero due to its direct use as electron acceptor, 78 making the growth of anaerobic chemotrophs also possible (e.g. acidogenic microorganisms 79 80 and sulfate-reducing bacteria; SRB). Currently, there is no published research available on PNSB production with raceway reactors, except for a trial focusing on polyhydroxyalkanoate 81 production.<sup>24</sup> However, it can be anticipated that the following control tools are essential to 82

maximize PNSB selectivity: (i) limiting the oxygen supply may decrease the growth of aerobic chemotrophs, (ii) increasing the light availability or the illumination period may aide PNSB in their competition for COD with aerobic chemotrophs and anaerobic chemotrophs such as acidogenic microorganisms and SRB, (iii) short sludge retention times (SRT) may washout slower-growing microorganisms such as microalgae, and (iv) limiting the COD-loading rate may decrease the competition with (an)aerobic chemotrophs during the dark period.

89 The hypothesis of this study was that PNSB can be selectively produced in a raceway 90 reactor provided a good combination of control tools. Batch experiments were first performed 91 to assess the phototrophic and (an)aerobic chemotrophic conversions of PNSB (axenic) and the (an)aerobic chemotrophic conversions of non-PNSB (non-axenic) to understand how they 92 93 individually would contribute in a raceway reactor. Afterwards, the effect of oxygen supply, 94 light availability, SRT and COD-loading rate was studied on PNSB abundance and community 95 diversity (non-axenic). A raceway reactor was then operated at a SRT of 2 d to understand whether PNSB can be selectively produced over sequential batches and explore the best 96 97 operational strategy for protein production and COD removal. All experiments were performed 98 using a synthetic medium composed of volatile fatty acids (VFA). The findings of the raceway 99 reactor were finally economically evaluated based on a production cost estimation, and 100 benchmarked with the production of a PNSB biomass in a closed PBR.

101 **2** 

Materials and methods

102 2.1 Inocula and medium

103 A *Rhodobacter capsulatus* strain, isolated in our previous study,<sup>13</sup> was used as model PNSB 104 for the axenic flask and non-axenic raceway reactor experiments. This species was selected 105 based on a prior evaluation made between five PNSB cultures, where it showed to have the 106 highest photoheterotrophic growth rate on synthetic wastewater. This species is able to grow photo- and chemoheterotrophically,<sup>10</sup> which enables examination of different PNSB growth
kinetics in raceway reactors. Aerobic activated sludge of a local brewery company (AB InBev,
Belgium, Leuven) was used as a proxy for a non-PNSB inoculum.

110 An adapted VFA-based medium as a proxy for fermented wastewater was used for all experiments, as we argued in a previous study that fermentation prior to protein production will 111 favor the microbial selectivity.<sup>4,13</sup> The COD concentration was increased to 3 g L<sup>-1</sup> and 112 contained a defined mixture of acetate, propionate and butyrate on a 1/1/1 COD basis. PNSB 113 grown photoheterotrophically on this VFA mixture have a biomass yield that approximate 1 g 114 COD<sub>biomass</sub> g<sup>-1</sup> COD<sub>removed</sub>.<sup>13,16</sup> This makes it easy to assess the chemoheterotrophic growth of 115 PNSB and of competing non-PNSB, as a lower biomass yield implies oxidation of COD to 116 117  $CO_2$ .

# 118 **2.2** Overview of the experiments

Five sets of flask and raceway reactor experiments were performed in this study to explore the conversions of PNSB and the effect of oxygen supply, light availability and SRT on PNSB selectivity (Table 1).

123 **Table 1** Objectives and experimental setup of five tests to grow a protein-rich PNSB biomass on brewery wastewater. *Rhodobacter capsulatus* was used as purple non-sulfur bacterium (PNSB)

124 inoculum and aerobic brewery sludge as non-PNSB inoculum. Experiments were performed at 28°C. Surface-to-volume (S/V) ratios were calculated based on the illuminated surface area. The

125 flasks were illuminated from the side and the raceway reactor from the top. OTR: oxygen transfer rate; SRT: sludge retention time

Objective	OTR (mg O <sub>2</sub> L <sup>-1</sup> d <sup>-1</sup> )	Stirring (on/off)	Illumination (light/dark)	S/V ratio (m <sup>2</sup> m <sup>-3</sup> )	SRT (d)	Inoculum	Cultivation	Reactor type
Assess conversions of PNSB	336		24 h	61	Batch*	PNSB	Axenic	Flask
and non-PNSB (section	0	24 h / 0 h	24 h			PNSB		
2.2.1)	336		0 h			PNSB		
	0		0 h			PNSB		
	336		0 h			Non-PNSB	Non-axenic	
	0		0 h			Non-PNSB	Non-axemic	
Influence of oxygen supply	72	24 h / 0 h	24 h		Batch* PN	DNGD	Axenic	Flask
on PNSB growth (Supporting Information S2-	336		24 h	61				
	72		0 h			PNSB		
3)	336		0 h					
Oxygen and light	204	24 h / 0 h	12 h / 12 h	5	Batch*	PNSB	Non-axenic	Raceway
availability as control tools to steer PNSB selectivity (section 2.2.2)		24 h / 0 h	24 h / 0 h	5				
		12 h / 12 h	12 h / 12 h	5				
		24 h / 0 h	12 h / 12 h	10				
Effect of SRT on PNSB	336				1.25		Non-axenic	Flask
growth (section 2.2.3)		24 h / 0 h	12 h / 12 h	61	2	PNSB		
					3			
PNSB selectivity over		24 h / 0 h	12 h / 12 h	5	2		Non-axenic	Raceway
sequential batches (section	204	12 h / 12 h		5	2	PNSB		
2.2.4)		24 h / 0 h		10	2			

126 \*Growth experiment between 50-150 h, stopped when stationary phase was reached

#### 127 2.2.1 Assess the conversions of PNSB and non-PNSB

Flask batch experiments were performed to explore the photoheterotrophic and (an)aerobic chemoheterotrophic conversions of PNSB along with the (an)aerobic chemoheterotrophic conversions of competing non-PNSB. These tests were conducted to understand how these conversions may individually contribute in a raceway reactor. The detailed cultivation conditions are described in Supporting Information S1

To explore the conversions of PNSB, four different conditions were tested under axenic conditions: (i) illumination with oxygen supply to study the combined photo- and chemotrophic growth (conditions prevalent in a raceway reactor); (ii) illumination without oxygen supply to study the phototrophic growth; (iii) no illumination with oxygen supply to study the aerobic chemotrophic growth and (iv) no illumination without oxygen supply to study the anaerobic chemotrophic growth (i.e. acidogenic metabolism).

An experiment was also performed to assess the effect of oxygen supply on the photoand chemoheterotrophic growth of PNSB. The methodology is explained in Supporting
Information S2.

#### 142 2.2.2 Light and oxygen availability as control tools to steer PNSB selectivity

These experiments were performed to explore the effects of light, oxygen supply, and the combination of light and oxygen on the biomass growth, biomass yield, biomass composition and PNSB selectivity.

A 100-L raceway reactor (MicroBio Engineering Inc., USA) was used to perform growth experiments in batch regime under non-axenic conditions. The stirring speed of the paddle was set at 30 rpm and the pH was controlled at 7 by sparging CO<sub>2</sub>. The maximum oxygen transfer rate (OTR) at this stirring speed was 204 mg  $O_2 L^{-1} d^{-1}$ . Temperature was controlled at 28°C with an external TetratecHT 300-W heater (Tetra, Germany). Approximately 5 L of tap water was added daily to compensate for evaporation. One halogen lamp was used to illuminate the

raceway reactor at an intensity of 54 W m<sup>-2</sup> (illumination spectrum see Supporting Information 152 S4). The reactor was filled with the VFA-based medium and *Rhodobacter capsulatus* was used 153 as inoculum at a total suspended solids (TSS) concertation of 0.02 g  $L^{-1}$ . 154

155 Four different conditions were tested in batch: (i) 12-h light/12-h dark with 24-h stirring at a surface-to-volume ratio of 5  $m^2 m^{-3}$  as benchmark (reactor filled up to 100 L); (ii) 24-h 156 light/0-h dark with 24-h stirring at a surface-to-volume ratio of 5  $m^2 m^{-3}$  to study the effect of 157 light; (iii) 12-h light/12-h dark with 12-h stirring (reduced oxygen supply vs. 24-h stirring) 158 during the light period at a surface-to-volume ratio of 5  $m^2 m^{-3}$  to study the effect oxygen 159 supply and (iv) 12-h light/12-h dark with 24-h stirring at a surface-to-volume ratio of 10 m<sup>2</sup> m<sup>-</sup> 160  $^{3}$  to study the effect of light (reactor filled up to 50 L). The absorbance of the biomass 161 162 suspension (660 nm) was monitored to determine the maximum specific growth rate. 163 Experiments were stopped when the stationary phase was reached. Samples were then taken 164 for further analysis.

165

#### 2.2.3 Effect of SRT on PNSB growth

These experiments were performed to explore the effect of SRT on the productivity, biomass 166 yield, biomass composition and PNSB selectivity. 167

Experiments were performed under combined photo- and chemotrophic conditions, 168 169 allowing the entry of oxygen along with illumination (i.e. conditions prevalent in raceway reactor). Flasks of 500 mL were used as reactors and illuminated through a natural 12-h 170 light/12-h dark regime with two halogen lamps at a light intensity of 30 W m<sup>-2</sup> (vs. previous 171 172 flask experiments section 2.2.1: 24-h light or 24-h dark). The flasks were filled with 200 mL of medium (section 2.1) corresponding to a maximum OTR of 336 mg  $O_2$  L<sup>-1</sup> d<sup>-1</sup>. The 173 experiment was performed non-axenically with Rhodobacter capsulatus as initial inoculum. 174 175 The tested SRT were 1.25 d, 2 d and 3 d. All tests were performed in biological duplicate. After 176 adding the medium and the inoculum, flasks were placed on a multipoint stirrer at 300 rpm.

Temperature during the light and dark period was respectively  $29.3 \pm 0.4$  °C (i.e. resulting from radiation heat) and  $24.2 \pm 0.7$  °C. The pH was  $7.4 \pm 0.2$ . Bottles were weighed daily to adjust for evaporation, remove a part of the broth and add fresh medium. Samples were taken daily to measure the absorbance (660 nm), pH, temperature and DO concentration. The moving average with a fixed subset size of three was determined. Steady state was reached when the daily absorbance overlapped with the moving average. Samples were then taken after the light and dark periods three days in a row to account for variability over time.

#### 184 **2.2.4** Operational strategies to steer PNSB selectivity and reactor performance

A final experiment was performed to demonstrate that PNSB can be maintained in a raceway reactor over multiple generations and determine the best operational strategy in terms of productivity and COD removal.

Temperature, pH and stirring were controlled as described in section 2.2.2. Three 188 operational strategies were tested: (i) 12-h light/12-h dark with 24-h stirring at a surface-to-189 volume ratio of 5 m<sup>2</sup> m<sup>-3</sup> as benchmark (reactor filled up to 100 L and depth 20 cm); (ii) 12-h 190 light/12-h dark with 12-h stirring (reduces oxygen supply vs. 24-h stirring) during the light 191 period at a surface-to-volume ratio of 5  $m^2 m^{-3}$  to study the effect oxygen supply and (iii) 12-h 192 light/12-h dark with 24-h stirring at a surface-to-volume ratio of 10 m<sup>2</sup> m<sup>-3</sup> (higher light 193 availability vs. 5 m<sup>2</sup> m<sup>-3</sup> reactor filled up to 50 L and depth 10 cm) to study the effect of light. 194 195 The SRT was chosen based on the maximal specific growth rate during the batch experiments (Figure 2). A value of 0.8 d<sup>-1</sup> was observed, which corresponds to a doubling time of 1.2 d. For 196 197 safety reason, a SRT of 2 d was chosen for the three conditions to prevent washout from the 198 reactor. Effluent was first removed and influent was then added before the start of the light 199 period. The absorbance was analyzed daily. After steady state, samples were taken by the end 200 of the light and dark periods three days in a row to account for variability over time. Samples 201 were stored at -20°C for further analysis.

#### 202 2.3 Analytical procedures

Standard methods were used to determine the TSS and volatile suspended solids 203 concentration.<sup>25</sup> The COD was measured using test kits (Macherey-Nagel, Germany) according 204 to the manufacturer's instructions. The volumetric mass transfer coefficient (K<sub>L</sub>a) of oxygen 205 was determined according to the sulfite oxidation method.<sup>26</sup> Protein concentration was 206 analyzed by Markwell, et al. 1978<sup>27</sup>, which is adapted from the Lowry procedure. The 207 bacteriochlorophyll a content was determined by an acetone/methanol solvent (7:2 v/v). 208 extraction.<sup>28</sup> Dissolved oxygen (DO) concentration (Hach, USA) and pH (Hanna Instruments, 209 USA) were determined with Handheld meters. 210

#### 211 **2.4 Microbial community analyses**

212 Genomic DNA was extracted from biomass samples collected (after steady state) across the reactor experiments using the DNeasy UltraClean microbial extraction kit (Oiagen, Venlo, the 213 214 Netherlands) according to the manufacturer's instructions. The DNA extracts were sent to a commercial company (Novogene, China) for amplicon sequencing analysis. In brief, the V3-215 V4 hypervariable region of the bacterial 16S rRNA gene pool of the DNA extracts was 216 amplified by PCR using the pair of 341f/806r primers prior to sequencing of PCR products 217 using a HiSeq 2500 sequencer (Illumina, USA). A detailed description of the wet-lab and dry-218 lab workflows can be found in Supporting Information S5. 219

# 220 2.5 Statistical analyses

Statistical analyses were performed in R (version 3.4.1) using RStudio (RStudio®, USA) for Windows.<sup>29</sup> Student's t-test were conducted to compare means. Normality of data residuals was tested using the Shapiro-Wilk normality test. The assumption of homoscedasticity was verified through a Levene's test. The non-parametric Kruskal-Wallis rank sum test was executed when normality was rejected. The Welch's t-test was used in case of heteroscedasticity. A significance level of p < 0.05 was chosen.

#### 227 **2.6 Production cost estimation**

228 The cost of four PNSB production scenarios was estimated and compared: (i) a closed tubular 229 PBR with 24-h stirring; (ii) a raceway reactor operated with 24-h stirring and a surface-tovolume ratio of 5 m<sup>2</sup> m<sup>-3</sup>; (iii) a raceway reactor operated with 24-h stirring and a surface-to-230 volume ratio of 10 m<sup>2</sup> m<sup>-3</sup> and (iv) a raceway reactor operated with 24-h stirring during the 231 light period and a surface-to-volume ratio of 5  $m^2 m^{-3}$ . Illumination for the four scenarios was 232 considered to be from sunlight according to a natural 12-h light/12-h dark regime For the 233 experiments, synthetic wastewater was used. This cost estimation was performed with brewery 234 wastewater, as a suitable model for food and beverage effluents where fecal contamination can 235 be avoided. 236

The goal of this model was to compare a closed PBR with a raceway reactor operated at three different strategies. It was not intended to determine an accurate production cost for PNSB. This cost estimation ought to be seen as a decision-making tool for research. Details and all input parameters are presented in Supporting Information S6.

#### 241 **3 Results and discussion**

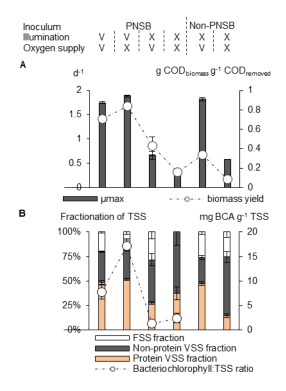
#### 242 **3.1** Assess the conversions of PNSB and non-PNSB

The results of the individual conversions of PNSB (Figure 1) indicated that during the 243 combined photo- and chemotrophic growth (i.e. light and oxygen supply), the phototrophic 244 245 metabolisms (i.e. under light) was dominant and not their chemotrophic metabolism (i.e. 246 oxygen supply). Biomass yields for their phototrophic and the combined photo- and chemotrophic metabolisms were similar (p > 0.05) and growth rates were almost equal. There 247 248 was, therefore, more photo-assimilation of COD than oxidation to CO2. The bacteriochlorophyll content was lower for the combined photo- and chemotrophic metabolism 249 than for the phototrophic metabolism, yet still 6 times higher than for the aerobic chemotrophic 250

growth (i.e. no illumination with oxygen supply). It should be noted that the chemotrophic metabolism of PNSB contributes more to growth when the oxygen supply increases (Supporting Information S3).

A similar experiment was performed by Sasaki, et al. 1998<sup>30</sup> with *Rhodobacter sphaeroides* on orange peel waste, showing a 1.6 times higher biomass yield for the combined metabolism compared to the chemotrophic metabolism. Our results were in line with this finding. This reveals that PNSB have the potential to use both their photo- and chemotrophic metabolism at once.

259 The DO concentration in a raceway reactor is zero due to the direct consumption of oxygen, allowing anaerobic fermentation of COD. The anaerobic chemotrophic growth of 260 261 PNSB was, therefore, tested with an organic more complex medium (vs. VFA-based medium 262 see Supporting Information S1). PNSB were able to anaerobically ferment organics (Figure 1), vet growth rates and biomass yields were relatively low  $(0.3 \pm 0.08 \text{ d}^{-1}; 0.16 \pm 0.09 \text{ g} \text{ COD}_{\text{biomass}})$ 263 g<sup>-1</sup> COD<sub>removed</sub>). A similar observation was made by Schultz Weaver 1982<sup>31</sup> with low anaerobic 264 growth rates and biomass yields for *Rhodobacter capsulatus* ( $\approx 0.08 \text{ d}^{-1}$ ; 0.09 g COD<sub>biomass</sub> g<sup>-1</sup> 265 COD<sub>removed</sub>) and *Rhodospirillum rubrum* ( $\approx 0.13 \text{ d}^{-1}$ ; 0.11 g COD<sub>biomass</sub> g<sup>-1</sup> COD<sub>removed</sub>). The 266 non-PNSB inoculum showed growth rates of  $0.58 \pm 0.03$  d<sup>-1</sup> or 2 times higher compared to 267 PNSB. Therefore, anaerobic fermentation will mainly be performed by competing non-PNSB. 268 Stronger competition during the light and dark period might arise from aerobic chemotrophic 269 270 non-PNSB since their growth rate was 2.8 times higher than for the aerobic chemotrophic 271 growth of PNSB and equal to the combined photo- and chemotrophic growth (p > 0.05).



272

Figure 1 (A) maximum specific growth rate and biomass yield for purple non-sulfur bacteria (PNSB) and non-PNSB along with (B) biomass fractionation and bacteriochlorophyll a (BCA) content. Tested conditions: combined photo- and chemoheterotrophic (illumination: V; oxygen supply: V), photoheterotrophic (V; X), aerobic chemoheterotrophic (X; V) and anaerobic chemoheterotrophic (X; X) growth. The oxygen transfer rate was 336 mg  $O_2 L^{-1} d^{-1}$ . Experiments were performed axenically with *Rhodobacter capsulatus* used as model PNSB. Non-PNSB were grown non-axenically. Averages with standard error. TSS: total suspended solids; VSS: volatile suspended solids; FSS: fixed suspended solids i.e. ash

#### 279 **3.2** Light and oxygen availability as control tools to steer PNSB selectivity

This experiment was set up to demonstrate that PNSB can be produced in a raceway reactor under non-axenic conditions and to investigate the effect of light, oxygen supply and the combination of both on PNSB selectivity.

The four tested conditions in Figure 2 showed an increase of the bacteriochlorophyll and carotenoid peaks during the batch growth experiments. Microbial community analysis confirmed these findings with PNSB abundances between 8-31% (Supporting Information S7). Hence, this research pioneers in demonstrating PNSB production in a raceway reactor without strict anaerobic conditions, thereby setting a precedent for future research. Lu, et al. 2019<sup>32</sup>

have also claimed to produce PNSB in a PBR at DO concentrations between 0.2-0.5 mg  $O_2 L^-$ 289 <sup>1</sup>, yet have not presented results of the microbial community composition.

For light availability as control tool, increasing the illumination time to 24-h light/0-h dark (vs. benchmark 12-h light/12-h dark) resulted in a biomass yield increase of 1.4 times (more photo-assimilation), an increase of the bacteriochlorophyll content by 3.2 times and increase of the PNSB abundance by 3.9 times (31% vs. benchmark 8%; Supporting Information S7). This was also the most effective control tool in terms of PNSB abundance.

295 Preventing oxygen supply during the dark phase (not stirring vs. benchmark 24-h stirring) 296 increased the biomass yield by 2.3 times, the bacteriochlorophyll content by 1.8 times and the PNSB abundance by 2.5 times (20%; Supporting Information S7). Hence, these results 297 298 reconfirm the findings of Supporting Information S3, where a lower oxygen supply resulted in 299 increased phototrophic growth. Increasing the surface-to-volume ratio had a dual effect. 300 Relatively to the benchmark, the biomass yield decreased by 1.5 times due to increased COD oxidation and the bacteriochlorophyll content increased by 2.4 due to an increased light 301 302 availability per reactor volume.

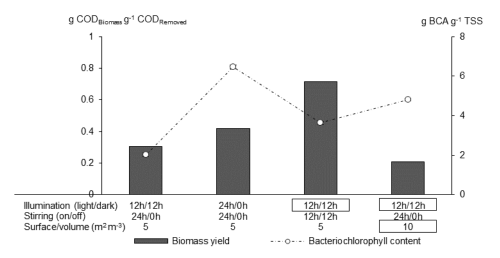


Figure 2 Batch raceway reactor experiment testing the effects of light (illumination), oxygen (stirring) and the combination
 of light (surface-to-volume ratio) on the biomass growth. Results show biomass yield (left y-axis) and bacteriochlorophyll a
 (BCA) content (right y-axis). Experiments were performed non-axenically with *Rhodobacter capsulatus* as initial inoculum.

Stirring (on/off) 12 h / 12 h implies stirring during the light period and not during the dark. Rectangular boxes show the change
 in reactor operation relative to the control. TSS: total suspended solids

#### 309 **3.3 Effect of SRT on PNSB growth**

310 This experiment was performed to study the effect of SRT on PNSB selectivity. Overall, PNSB 311 abundances did not show substantial differences between SRT (Figure 3). This is in line with 312 our previous observation where we tested the effect of SRT on PNSB abundance in a closed PBR.<sup>13</sup> Notably (Figure 3), the PNSB abundance at 6 g COD L<sup>-1</sup> for all SRT (50-67%) was 313 lower compared to 3 g COD L<sup>-1</sup> (88-94%). The exponent of the Shannon diversity index was 314 also lower at 3 g COD L<sup>-1</sup> (1.4-1.7) compared to 6 g COD L<sup>-1</sup> (2.7-3.0), which indicates a more 315 316 uneven community at lower loading rates. The abundance of the (an)aerobic 317 chemoheterotrophs Arcobacter was predominantly higher at higher loading rates (17-36%). Overall, it can be concluded that a higher loading decreased the PNSB selectivity. This 318 decreased PNSB selectivity was probably due to the higher COD availability especially during 319 the night period, leading to increased growth of competing chemotrophs. As such, COD 320 321 availability during the night will negatively influence the PNSB abundance.

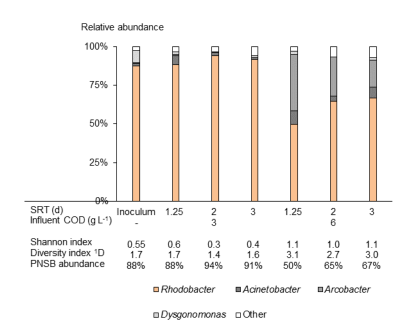


Figure 3 Effect of sludge retention time (SRT) on microbial community composition, Shannon's H index, exp(H') and purple
 non-sulfur bacteria (PNSB) abundance. Flasks were used as a reactor. The PNSB genera *Rhodobacter* and *Rhodopseudomonas* are all marked in orange. Samples obtained after 2-10 SRT.

#### 326 **3.4 Reactor performance and community dynamics over sequential batches**

This experiment was conducted to demonstrate that PNSB can be maintained in a raceway reactor over multiple generations and determine the best operational strategy in terms of productivity and COD removal.

The highest productivities (0.21 g protein  $L^{-1} d^{-1}$  corresponding with 0.43 g TSS  $L^{-1} d^{-1}$ ) 330 and removal rates (0.79 g COD  $L^{-1} d^{-1}$ ) were achieved when the reactor was operated with 24-331 h stirring and 12-h light/12-h dark at the highest surface-to-volume ratio of 10 m<sup>2</sup> m<sup>-3</sup> (Figure 332 4). A higher ratio of 10  $m^2$  m<sup>-3</sup> increased the light availability, resulting in higher biomass 333 concentrations (0.81  $\pm$  0.04 g TSS L<sup>-1</sup>) relative to the benchmark of 5 m<sup>2</sup> m<sup>-3</sup> (0.62  $\pm$  0.02 g 334 335 TSS L<sup>-1</sup>). For a closed PBR operated on the same medium at a SRT of 1 d (vs. 2 d for raceway reactor), we reached TSS productivities that were 1.5-2.6 times higher compared to the raceway 336 reactor.<sup>13</sup> This was probably due to the higher light availability in the PBR compared to a 337 raceway reactor (surface-to-volume ratio 33 vs. 5-10 m<sup>2</sup> m<sup>-3</sup>). For the microalga Chlorella 338 vulgaris cultivated in the same reactor (12-h light/12-h dark), the productivity was 0.009 g 339 protein L<sup>-1</sup> d<sup>-1</sup> or 22 times lower compared to PNSB.<sup>6</sup> This was probably due to the higher 340 growth rates of PNSB of 0.6-3.7 d<sup>-1</sup> compared to the ones of microalgae of 0.60-1.38 d<sup>-1</sup>.<sup>13,33,34</sup> 341 342 To a lesser extent, non-PNSB chemotrophs also contributed to biomass production in the 343 raceway reactor, thereby increasing the overall biomass productivity.

In terms of PNSB selectivity (Figure 5), preventing the combination of oxygen supply and darkness (not stirring) was an effective strategy in line with the results of section 3.2. The PNSB abundance was 56% (vs. 14% benchmark 24-h stirring) and the microbial community was more uneven showing a lower exponent of the Shannon diversity index (3.7 vs. 4.3 for benchmark). The decrease in PNSB abundance to 41% was notable during the dark period

along with the increase of the exponent of the Shannon diversity index from 3.7 to 7.2. Hence,
biomass harvesting should preferably be performed after the light period to assure a selective
microbial community.

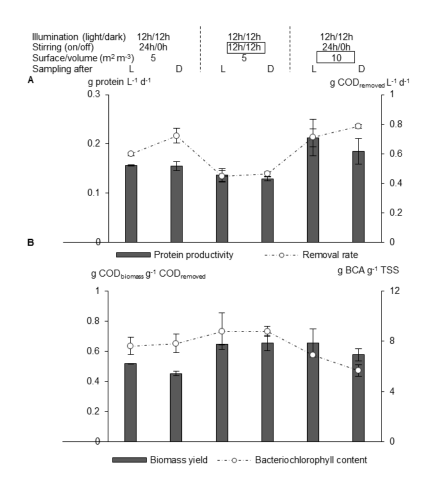
The reactor operated with 24-h stirring at a surface-to-volume ratio of  $10 \text{ m}^2 \text{ m}^{-3}$  was the 352 best strategy in terms of PNSB selectivity, showing a PNSB abundance of 78% or 5.6 times 353 354 higher compared to the benchmark and very comparable to the inoculum. The exponent of the Shannon diversity index was only 2.5, the lowest for all conditions and even lower than the 355 356 inoculum (3.5). This implies that light availability is key to boost PNSB selectivity in a raceway 357 reactor. The findings also show that a raceway reactor can approach the PNSB selectivity of a closed PBR. Potential higher PNSB abundances might even be achieved if oxygen supply is 358 359 prevented during the night along with high surface-to-volume ratios.

360 Other PNSB genera were also present in the system such as *Rhodopseudomonas* (2-3%), and *Blastochloris* (< 0.2%). The main competing genera were *Acinetobacter* (aerobic 361 chemoheterotroph), Dysgonomonas (anaerobic chemoheterotrophs), Arcobacter ((an)aerobic 362 363 chemoheterotrophs) and Alcaligenes (aerobic chemoheterotrophs) with an abundance of respectively 1-4%, 1-12%, 0-31% and 1-54%. Microalgae were not detected through the 364 365 absorbance spectra (no chlorophyll peaks) and no cyanobacteria were identified by amplicon sequencing. This was probably due to the short SRT (2 d), resulting in washout of slower-366 growing microalgae ( $\mu_{max}$  0.60-1.38 d<sup>-1</sup>).<sup>33</sup> Although the sulfate concentration in the medium 367 was 1.6 g L<sup>-1</sup>, there were no SRB detected (of note, primers not designed for archaea). SRB 368 require 0.7 g COD to remove 1 g of sulfate.<sup>35</sup> Therefore, they can contribute to COD removal, 369 yet biomass production will be negligible due to their low biomass yields of 0.015-0.033 g VSS 370 g<sup>-1</sup> SO<sub>4</sub><sup>2-</sup>.<sup>36</sup> 371

According to the authors' knowledge, this study is the first to publish results on PNSB production in a raceway reactor dedicated to microbial protein. Literature studying microbial

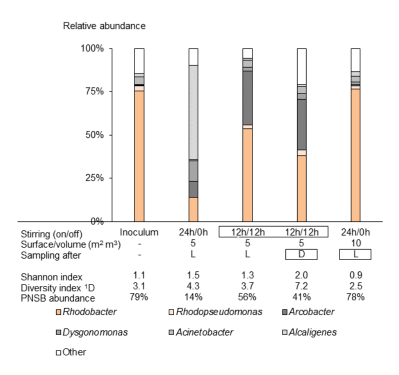
374 communities in waste lagoons are helpful for benchmarking, as these systems enable photosynthesis and are open to air. Do, et al. 2003<sup>37</sup> have investigated the correlation of 375 environmental factors on photosynthetic blooms, i.e. the spontaneous growth of purple bacteria 376 377 in waste lagoons. Their research showed that for a swine waste lagoon, up to 10% of the microbial community was made by a population of Rhodobacter. The authors observed a 378 379 positive correlation between *Rhodobacter* and the sulfate concentration. They claimed that it was due to competition between PNSB and SRB. More PNSB growth would result in a lack of 380 381 VFA for SRB and the accumulation of sulfate.

382 Our reactor was operated with artificial 12h-light/12h-dark conditions at constant temperature with a VFA-based medium. Hence, sunlight-driven pilot-scale validation should 383 384 still confirm these findings. As scale-up for raceways is mainly horizontal (depth is kept 385 constant), the main effects are expected from a different influent composition (more complex 386 COD mixture vs. synthetic), fluctuations in COD flow and temperature, day/night length and 387 light intensity. Another parameter that might be different for a for a full-scale raceway reactor 388 is the oxygen supply to the system. The oxygen supply rate during the experiment in this chapter was around 204 mg  $O_2 L^{-1} d^{-1}$ . However, for a full-scale system this would potentially 389 be lower due to higher reactor volume. The lower oxygen supply would positively influence 390 391 the PNSB growth, yet might enhance the competition with anaerobic heterotrophs such as SRB 392 and fermentative microorganisms.



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Figure 4 Production features of a raceway reactor operated at a sludge retention time of 2 d, testing the effect of oxygen (stirring) and light (surface-to-volume ratio). Sampling occurred after the light (L) or dark (D) period. Results show (A) the protein productivity and the volumetric removal rate along with (B) the biomass yield and bacteriochlorophyll (BCA) content. Experiments were performed non-axenically with *Rhodobacter capsulatus* as initial inoculum. Stirring (on/off) 12 h / 12 h implies stirring during the light period and not during the dark. Average values with standard error. 12 h / 0 h stirring occurred during the light period. Rectangular boxes show the change in reactor operation relative to the benchmark. TSS: total suspended solids



**Figure 5** Effect of operational strategy of a raceway reactor on microbial community composition, Shannon's H' index, exp(H') and purple non-sulfur bacteria (PNSB) abundance. Raceway reactor operated at a sludge retention time of 2 d, testing the effect of oxygen (stirring) and the combination of light (surface-to-volume ratio). Sampling occurred after the light (L) or dark (D) period. Stirring (on/off) 12 h / 12 h implies stirring during the light period and not during the dark. The PNSB genera *Rhodobacter* and *Rhodopseudomonas* are marked in orange. Rectangular boxes show the change in reactor operation relative to the benchmark.

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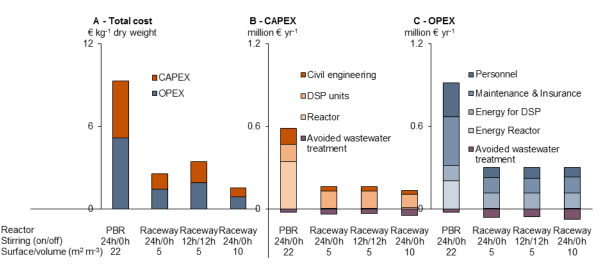
#### 409 **3.5** Production cost comparison of a closed PBR and an open raceway reactor

An anaerobic reactor such as a closed PBR is the best non-axenic technology to selectively produce PNSB on wastewater, yet production costs were estimated between 2.7-6.4 times higher compared to a raceway reactor (Figure 6). These high costs can mainly be contributed to the high investment costs for a tubular PBR (47% of capital expenditure; CAPEX) and the energy required for recirculation (23% of operational expenditure; OPEX).

415 PNSB cultivated in closed PBR have high biomass yields (1 g  $COD_{biomass}$  g<sup>-1</sup>  $COD_{removed}$ ) 416 due to photo-assimilation instead of oxidation of COD to CO<sub>2</sub>. This will result in lower COD 417 removal efficiencies compared to a raceway reactor. A raceway reactor can save around 418 €83,000-120,000 yr<sup>-1</sup> of the wastewater treatment cost or 1.7-2.4 times more compared to a PBR. Raceway reactors have the potential to remove 88% of the incoming COD. Therefore, wastewater treatment after PNSB production with a raceway reactor can be a simple aerobic activated sludge treatment process instead of a digester followed by aerobic treatment. On the contrary, PBR only remove 35% of the COD. Hence, a digester is still needed to lower the COD concentration.

The cost to produce PNSB in a raceway reactor was estimated between &1.3-3.0 kg<sup>-1</sup> dry weight (DW). Downstream processing was the most dominant cost factor contribution by 57-61% to the CAPEX and by 40-43% to the OPEX. A further cost reduction of 27-40% can be possible if ultrafiltration is replaced by gravitational settling (&0.9-2.4 kg<sup>-1</sup> DW). Currently, there is no published literature studying the settleability of PNSB except for Cerruti, et al. 2020 DOI: <u>https://doi.org/10.1101/2020.01.08.899062</u><sup>38</sup>. Further research should explore this closely, as it will be an important cost saver.

The operational strategy with 24-h stirring at a surface-to-volume ratio of  $10 \text{ m}^2 \text{ m}^{-3}$ (reactor depth 10 cm) resulted in the highest PNSB selectivity and was also the most costeffective. A regime of 24-h stirring will produce more biomass (more chemotrophic growth) and will remove more COD relatively to 12-h stirring. This will result in a cost that can be spread out over more products and additional savings for the subsequent aerobic activated sludge process.



**Figure 6** Cost comparison for a closed tubular photobioreactor (PBR) and an open raceway reactors using natural sunlight (12-h light/12-h dark). Results show (A) capital and operational expenditure (CAPEX, OPEX), (B) fractionation of CAPEX, and (C) fractionation of OPEX. The cost category "Avoided wastewater treatment" refers to activated sludge treatment costs that can be avoided due to COD removal by the microbial protein reactor. Stirring (on/off) 12 h / 12 h implies stirring during the light period and not during the dark. All scenarios were dimensioned based on brewery wastewater with a COD concentration of 2.1 kg m<sup>-3</sup> and a flow rate of 1150 m<sup>3</sup> d<sup>-1</sup>. DSP: Downstream processing i.e. harvesting, dewatering and drying.

# 444 **4 Conclusions**

- The goal of this research was to develop control tools to selectively produce PNSB in a racewayreactor. The main findings of this study are:
- 447 (i) This study pioneers in the selective production of PNSB in a raceway reactor with 448 productivities of up to 0.43 g TSS  $L^{-1} d^{-1}$  and COD removal rates of up to 0.79 g COD 449  $L^{-1} d^{-1}$ .
- 450 (ii) Avoiding oxygen supply during the dark phase and a higher surface-to-volume ratio
  451 were the best operational strategies to maximize the PNSB abundance (56-78%) and
  452 lower the diversity.
- 453 (iii) SRT does not show to have an impact on PNSB selectivity. However, COD availability
  454 should be avoided in the dark, as it decreases the PNSB abundance from 90% to 69%
  455 and increases the Shannon diversity from 0.45 to 1.1.
- 456 (iv) Flask and raceway experiments showed that PNSB competed mainly with aerobic
  457 chemotrophs and to a minor extent with anaerobic chemotrophs. Microalgae or SRB
  458 were not identified as major competitors.
- 459 (v) The combination of oxygen supply, higher COD load and darkness should be avoided.
- 460 (vi) Production costs for a raceway reactor amount to  $\notin 1.9 \text{ kg}^{-1}$  DW, which is six times 461 cheaper than a closed PBR.

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