

Delft University of Technology

#### Harnessing the metabolic versatility of purple non-sulfur bacteria

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DOI 10.4233/uuid:c98876c1-aa83-4c8d-9816-191daf5cc423

Publication date 2022

**Document Version** Final published version

Citation (APA)

Cerruti, M. (2022). Harnessing the metabolic versatility of purple non-sulfur bacteria. [Dissertation (TU Delft), Delft University of Technology]. https://doi.org/10.4233/uuid:c98876c1-aa83-4c8d-9816-191daf5cc423

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Marta Cerruti

Harnessing the metabolic versatility of purple non-sulfur bacteria

# Harnessing the metabolic versatility of purple non-sulfur bacteria

### Harnessing the metabolic versatility of purple non-sulfur bacteria

Dissertation for the purpose of obtaining the degree of doctor at Delft University of Technology by the authority of the Rector Magnificus, Prof.dr.ir. T.H.J.J. van der Hagen, chair of the Board for Doctorates To be defended publicly on 17.02.2022, 12:30 o'clock

by

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ISBN 978-94-6384-303-4

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### Propositions

- 1. A univocal quantitative description of the metabolism of hyperversatile purple phototrophic bacteria is still a far-away goal. *(This proposition pertains to this dissertation)*
- 2. A mixed community is always more robust than a pure culture. *(This proposition pertains to this dissertation)*
- 3. Without a clear understanding of the mechanisms inside the reactor black box, process control is a blind operation. *(This proposition pertains to this dissertation)*
- 4. Knowledge development, and not self-celebration, should be the main driver in research.
- 5. Tools to process data are as important as concepts.
- 6. It's not science *per se* that should be questioned, but methods and interpretations.
- 7. Coaching spans from micro-managing to advising. Only the latter can be beneficial for both mentor and trainee.
- 8. Collaborations are the science counterpart of animal mating rituals.
- 9. Bacterial communities teach us that, with the appropriate environmental conditions, each individual can thrive.
- 10. 'You might have working hours. Bacteria don't.'

These propositions are regarded as opposable and defendable, and have been approved as such by the promotor Prof.dr.ir. M.C.M. van Loosdrecht and the copromotor Dr.ir. D.G.Weissbrodt

Dai diamanti non nasce niente dal letame nascono i fior

Fabrizio de Andrè

### Summary

A transition to a circular economy is necessary to mitigate the negative effects on the environment of the exploitation and disposal of materials and to achieve society-wide benefits. The current produce-waste-dispose model is slowly changing toward a more sustainable produce-userecycle-upcycle model. In this context, bio-based processes using microbial mixed cultures are crucial to develop waste-to-resource valorization processes.

Purple phototrophic bacteria (PPB) form a guild of hyper-versatile organisms found in almost all aqueous environments, thriving on infrared light energy, capturing organics by photoorganoheterotrophy, and even recycling  $CO_2$  by photolithoautotrophy. Due to their outstanding metabolic versatility, their organic and nutrient capture ability, and their biomass yields over substrate approaching 1 g CODx g<sup>-1</sup> CODs, PPB are dedicated organisms to study and use for the development of water resource recovery applications. Despite already 80 years of research on PPB, their physiology still needs to get deciphered, and their environmental biotechnological exploitation is at its infancy.

The aim of this thesis was to study and harness the metabolic versatility of PPB at different levels, from the elucidation of light-driven physiologies in pure cultures to the management of selection phenomena, population dynamics, and distributed metabolic functionalities in mixed cultures. The findings were aggregated to derive to mixed-culture bioprocess application perspectives for capturing organics and nutrients from municipal sewage and agri-food wastewater and producing valuable products, as bioplastics, biohydrogen or photopigments. In this thesis, a comprehensive overview of the potential of PPB for water resource recovery is given. The molecular principles and ecological dynamics governing the PPB metabolism were elucidated with the goal to demonstrate the potential of PPB-based biotechnologies.

In **Chapter 1**, the General Introduction delivers a targeted overview of the purple bacteria, their key metabolic properties, and their implementation potential for water resource recovery. The current European regulations for resource recycling and upgrading are reviewed, with a focus on water and wastewater bodies. A description of the complex metabolism of PPB, along with their potential industrial applications is given. Furthermore, the research questions and research objectives of this thesis are presented.

In Chapter 2, Rhodopseudomonas palustris was used as model organism to study the physiological responses to light/dark diel energy variations and associated variations in electron sinks. The aim was to elucidate the physiological allocation of electrons derived from organic substrate under light / dark alternation periods. Light is the primary energy source for PPB. In natural environments, it is provided in cycles, alternating with dark periods. This implies a discontinuous energy supply, that can impact the electron distribution in the cells. Redox balance is important for the physiology of PPB, but it has not been addressed in relation to Under dynamic illumination patterns. growth conditions. Rhodopseudomonas palustris redirects the electrons in excess toward different routes. Aside biomass, products such as dihydrogen (H<sub>2</sub>) and poly-β-hydroxyalkanoates (PHAs) are the main electron sinks. The impact of three different light patterns (24 h continuous infrared illumination, 16h light/8h dark, and 8h light/16h dark) was evaluated in a continuous culture fed with balanced nutrients medium. Under continuous illumination, a stationary state was attained, with a constant growth rate equal to 0.04 h<sup>-1</sup>. Under dark phases, the biomass was not produced, and acetate, the substrate, accumulated in the reactor. Under light/dark diel cycles, substrate uptake and growth were unbalanced, and Rhodopseudomonas palustris first directed the excess of carbon and electrons toward PHA production. Only secondarily, the electron excess was redirected toward H<sub>2</sub> production. A numerical model was used to describe the process dynamics and stoichiometric and kinetic parameters for the conversions. Elucidating the preferential carbon and electron redistribution allows a selective control of the bioproduct formation under dynamic energy conditions.

Due to their high growth yield on substrates, mixed cultures of PPB raise interest for their ability to directly capture organics and nutrients from used aqueous streams by assimilation in their biomass. Prior investigations have been made in diluted cell suspensions with low settling rate, requiring energy-expensive mechanical downstream processing (*e.g.*, membrane filtration or centrifugation) for biomass recovery. The aim of **Chapter 3** was to enrich and produce a concentrated, fast-settling PPB biomass with high nutrient removal capacity in a sequencing batch reactor (SBR). After seeding with a conventional activated sludge inoculum, the microbial community was rapidly enriched with PPB within the first 24 h of operation under infrared light illumination and feeding of an acetate-based synthetic wastewater, and PPB reached up to 90% of the 16S rRNA gene amplicon sequencing dataset. Managing the SBR operational conditions, namely decreasing the hydraulic retention time to 16 h, and increasing the organic loading rates to 1.3 kg COD  $d^{-1} m^{-3}$  stimulated biomass aggregation, settling, and accumulation in the system. Biomass concentration was as high as 3.8 g VSS L<sup>-1</sup>, 10 times higher than previously reported in literature. Carbon, nitrogen and phosphorous were removed up to 96, 77 and 73 % respectively, reaching the European nutrient discharge limits. The here-examined microbial ecology insights and the aggregation propensity of PPB mixed-cultures can help developing photobiobased processes for water treatment and recovery.

Water resource recovery involves different types of aqueous waste streams. To develop robust PPB-based mixed-culture biorefinery processes, it is necessary to evaluate the impacts of organic streams on the symbiotic / competitive interactions of PPB with other microbial guilds in the engineered community. The aforementioned results highlighted that the supply of infrared light together with acetate as sole carbon source and electron donor preferentially selects for PPB. In carbohydrate-rich wastewaters, fermentative chemoorganoheterotrophic bacteria (FCB) play a key role in sugar dissimilation. The interaction between FCB and PPB has been previously studied in axenic pure cultures or co-cultures. Little is known about their metabolic interactions in open mixed cultures. Chapter 4 addressed this knowledge gap by assessing the competitive and syntrophic interactions between PPB and FCB. The effects of reactor regimes (batch or continuous) and illumination mode (continuous light or light/dark diel cycles) were studied on glucose conversions and process ecology. Under batch regime, PPB were outcompeted by FCB which displayed a faster maximum growth rate and scavenged the glucose resource, consequently decreasing the pH to 4 in the batch cultures. Under continuous culturing, three main FCB families were enriched to up to 70% (Enterobacteriaceae, Lachnospiraceae and Clostridiaceae) while the PPB family of Rhodobacteraceae accounted for the other 30% of the community. The analysis of the glucose degradation products allowed for the assessment of the syntrophic relationship between FCB and PPB. FCB first fermented glucose into volatile fatty acids and alcohols, and PPB grew on fermentation products. The integration of the effects of reactor regimes and light patterns is crucial in the perspective of designing an efficient anaerobic mixed-culture processes for carbohydrate biorefinery based on associations of PPB and fermentative bacteria.

Light supply is one of the major costs in phototrophic-based technologies for water treatment and bioproduct formation, therefore its effects on microbial physiology need to be carefully estimated. From pure culture studies, it is known that PPB physiology is linked to the light intensity provided to the cells, and the growth rates increase with the light intensity up to a saturation threshold. Nonetheless, little is known about light intensity effects on PPB mixed cultures. In **Chapter 5** this knowledge gap is addressed, and 9 different light intensities (from 350 to 0 W m<sup>-2</sup>) were tested on a PPB enrichment under SBR conditions. Under all conditions, PPB were the dominant guild, accounting for more than 70% of the community. Light correlated with a logistic regression to the biomass growth rates, and the half saturation constant was estimated at 10  $\mu$ mol<sub>photons</sub> s<sup>-1</sup> g<sub>DW</sub><sup>-1</sup>. It was found that the mass fraction of the photopigments responsible for light harvesting was *ca* 5 times higher at lower light intensities compared to high light intensities, potentially compensating for the lower energy supply. As light is a crucial parameter for phototrophic growth, the assessment of the effects of light intensities on the physiology and microbial ecology of a PPB enrichment paves the way for the development of PPB-based biofactories.

At the end of this book, the main findings of the thesis are summarized in the **Chapter 6**. A perspective on current challenges and future research is given in **Chapter 7**, including applicability of the PPB-based biotechnologies for the treatment different aqueous waste streams, the limitations of light supply and the recovery of high-value products like carotenoids. Chapter 1

General introduction

#### Purple bacteria for wastewater resource recovery

#### 1. Wastewater treatment and resource recovery: drivers of the circular economy

#### 1.1 Circular economy principles in the European context

In a linear model of industrial production, the resources are extracted, processed, used, and discarded. To reduce the environmental impact, circular economy principles aim to promote a closed loop for resource exploitation. In this context, the value of the material, resources, products and services is maintained as long as possible, and the waste production is minimized (EEA, 2020).

In 2015, the European Union (EU) adopted the first circular economy action plan, which defined 54 measures to regulate production, consumption, and waste management (European Commission, 2015). The actions aimed to accelerate Europe's transition toward a closed loop of product lifecycle, implementing the recycle and re-use principles. In 2020, the European Union adopted a second action plan, to complete the ongoing tasks and transition toward a regenerative growth model, including waste reduction targets and waste prevention (European Commission, 2020).

The European Green Deal (2019) has the goal to achieve a sustainable EU growth by 2050, setting goals to achieve zero-pollution and protect the natural environment (European Commission, 2019). It describes a set of guidelines to move toward a clean circular economy, reduce the environmental impact and limit ecosystems pollution by improving nutrient management and reducing emissions. The 'Zero Pollution Action Plan for Air, Water and Soil' initiative, included in the European Green Deal, (European Commission, 2021) aims to take measures so that "Air, water and soil pollution is reduced to levels no longer considered harmful to health and natural ecosystems and that respect the boundaries our planet can cope with, thus creating a toxic-free environment."

#### 1.2 Water safeguard: a priority for the European Union

Water is one of the most important resources on Earth. It is crucial for the living world, and it sustains public health, development, and economic growth (European Commission, 2010).

Water resources, also in the EU, are increasingly under risk of scarcity and quality deterioration (The European Parliament and the Council, 2020). The safeguard of water resources is one of the European priorities (European Commission, 2018). Along with protective actions of natural water bodies, a wider re-use of treated wastewater helps reducing this pressure.

In the 2015 action plan for circular economy, the European Commission committed to take actions to promote the use of treated wastewaters, including the development of legislative guidelines for water reuse. Reclaimed treated waters from urban wastewater treatment plants (WWTPs) form a reliable water supply for different purposes, including agricultural applications. The use of treated wastewaters has generally a lower environmental impact compared to other processes such as desalination. It extends the water life cycle and helps preserving water resources (European Commission, 2018).

#### 1.3 Water and wastewater management in the European Union

Clean unpolluted water is crucial for human health and the ecosystem. Agricultural activities and industrial discharges are the principal pollution sources, adding to the domestic sewage emissions. Hazardous substances released in the environment comprehend micropollutants, pesticides, fecal microbes, excess nutrients, and oxygen-consuming substances (organic matter). An excess of nutrients such as carbon, nitrogen, and phosphorus in the water bodies is linked to the eutrophication process. It causes an accelerated growth of algae or plants that disturbs the ecological balance in the waters and the water quality itself. When decaying, algal and plant materials are consumed along with oxygen by microorganisms, therefore depleting the oxygen levels in the water body and hampering aquatic life (Abdel-Raouf et al., 2012; Bartram et al., 2002).

From 1991 onwards, the EU is regulating the wastewater treatment and discharge (European Council, 1991). The water quality objectives set the limits to cumulative pollution. The emission limit values focus on the concentration and loads of pollutants that can be discharged from different sources in the environment. The urban wastewater treatment (UWWT) directive aims to protect waters from adverse waste discharge. To meet the imposed limits, water coming from industrial or municipal sources requires treatment before discharge in the environment (European Council, 1991).

Wastewater treatment incudes different steps to reintroduce clean, unpolluted water in the natural environments. At the forefront of the WWTP, mechanical pre-treatments physically remove large objects from the wastewaters. The primary clarification treatment must reduce by at least 20% the biochemical oxygen demand (BOD) and 50% the total suspended solids (TSS) (Pescod, 1992).

The UWWT directive stipulates that in agglomerates over 2000 persons equivalents (p.e.), water must be subjected to secondary (or biological) treatment, to substantially reduce soluble organic matter, nitrogen and phosophorus pollution, and further remove suspended solids. In some specific cases, a tertiary, more stringent, treatment can be advised to further remove C, N, P compounds and avoid eutrophication.

Historically, sanitation technologies have been developed to safeguard public health by channeling excreta and removing pathogens. Progressively, with the evolution of environmental protection targets, WWTPs were developed to remove organic matter (C), nitrogen (N) and phosphorus (P). Most efforts made in industrialized regions over the last 50 years have been mainly addressed to design technologies for the removal of these C-N-P macropollutants.

Biological wastewater treatment (BWWT) technologies like activated sludge processes benefit from 100 years of development (Jenkins & Wanner, 2014; Sheik et al., 2014). BWWT takes advantage of the ability of microorganisms to uptake nutrients from the wastewaters and convert them into biomass (Hedaoo et al., 2012).

Control and regulation of biological processes allowed the design of efficient BWWT (Pell & Wörman, 2011). Conventional processes are designed for biological nutrient removal (BNR) in series of tanks where different redox conditions are carefully imposed (e.g., anaerobic-anoxicaerobic sequences in the A2/O process) to select in the sludge for guilds of microorganisms with metabolisms specialized for the removal of organic matter, nitrogen and phosphorus (Oleszkiewicz & Barnard, 2006). While physicochemical technologies are also used, all nutrients can be removed biologically. Organic matter is typically oxidized by aerobic- or anaerobic-respiring chemoorganoheterotrophic organisms. Nitrogen is removed by association of nitrifiers (that oxidize ammonium into nitrite and nitrate, aerobically) and denitrifiers (that reduce nitrate and nitrite to nitrogen oxides and inert dinitrogen gas using organic matter as electron acceptor) (Fenchel et al., 2012). Phosphorus is taken up in the biomass by polyphosphate-accumulating organisms (PAOs) by alternation of anaerobic and aerobic conditions who simultaneously remove organic matter: in the anaerobic tank they store volatile fatty acids (VFAs) as intracellular poly-β-hydroxyalkanoates (PHAs); in the anoxic or aeration tanks they grow on PHAs using either nitrite/nitrate or dissolved oxygen as terminal electron acceptors, respectively, and store orthophosphates as intracellular polyphosphate; phosphorus therefore ends up in waste sludge (Weissbrodt et al., 2017). These microbial processes can be assembled in large systems using flocculent activated sludge or intensified with biofilms and granular sludge depending on treatment targets.

The waste sludge derived from these processes was initially simply discharged or incinerated. In more recent times, thanks also to the implementation of circular economy principles, a particular attention has been placed in the valorization of the biomass. Beside the main objectives for public health sanitation and environmental protection, valuable resources can be recovered from used water streams. Anaerobic digestion is already well established since the 1980's for the production of biogas from waste biomass (Lettinga, 1995). Technologies are currently developed to upgrade organics, nutrients and biomass into, *e.g.*, biofuels, biomaterials, fertilizers or food / feed ingredients. WWTPs therefore transition into water resource factories.

## 2. Phototrophs and their metabolic opportunities for resource protection and recycling

#### 2.1 Phototrophic plasticity: the key for a bio-based circular economy

In parallel to the conventional chemotrophic processes involved in WWTPs, phototrophic systems raise interest in the tertiary WWT (Abdel-Raouf et al., 2012). Phototrophic organisms, like (micro)algae and bacteria, harness the light energy and numerous carbon sources and electron donors / acceptors for biomass growth and bioproducts formation. They use a broad spectrum of inorganic ( $CO_2$ ) and organic carbon sources, along with numerous nitrogen and phosphorous compounds, promoting a carbon neutral bio-economy (Renuka et al., 2021).

Due to their high nutrient removal abilities, they are used as tertiary WWT. In some cases, their interaction with chemotrophic bacteria have been exploited to improve the WWT performances and reduce the aeration costs (Abdel-Raouf et al., 2012).

Phototrophic systems have been studied for water sanitation since the 1950s (Chu & Phang, 2019). Microalgal systems are primarily used for biofuel production, due to the high lipid content coupled to high growth rates and low production costs (Shin et al., 2018). In WWT, the use of phototrophs has the advantage of removing pathogens and pollutants, nutrient recovery, and energy savings. Microalgal systems have been explored and are currently exploited for simultaneous WWT and biofuel production (Chia et al., 2018). Algal-bacteria consortia have been proposed as efficient processes. Along with established technologies, PPB mixed cultures have gained wide interest for WWT only in the past 5-10 years.

The ability to convert light energy into chemical energy is an evolutionary advantage for phototrophs, that can compete for substrates with heterotrophs under nutrient-limited conditions (Nowicka & Kruk, 2016). In fact, at least part of the energy required for biomass growth is derived from light, and therefore nutrients can be allocated into anabolic reactions to produce biomass and cellular components.

Phototrophs display numerous metabolic types: they are able to use organic and inorganic carbon sources, different electron donors and acceptors, and photons at different wavelengths.

Phototrophs can potentially cover all the possible metabolic modes (Tang et al., 2011). Mixotrophy is common in green microalgae and cyanobacteria, whereas purple bacteria can adapt their metabolism to the environmental conditions. The plasticity of the phototrophic metabolism is a key parameter for the utilization of the resources present in both natural and anthropogenic environments. From an industrial point of view, the phototrophs versatility makes them suitable for a wide range of application. Their applications span from WWT to biofactories for the production of lipids, proteins, biofuels and high-value molecules, such as carotenoids.

#### 2.2 Phototrophy: a metabolism widespread across life domains

Phototrophic metabolisms are widespread along the different life domains, from eukaryotic superior plants and algae to prokaryotic bacteria and archaea. All phototrophs share the ability to capture light photons as energy source to convert it into chemical energy and reducing power through membrane bound complexes (Peretó, 2011). Besides this common trait, phototrophs are diverse organisms with peculiar metabolisms that have found their own niche in diverse environments, both natural and anthropogenic (Stomp et al., 2007).

Three different types of phototrophic metabolism have originated early during the formation of the biosphere. They are known as oxygenic (*i.e.*, cyanobacteria, algae, plants), anoxygenic (e.g., phototrophic green bacteria, purple bacteria and heliobacteria), and rhodopsin-based (e.g., archaea and some bacteria) phototrophs.

The exact evolution of the phototrophic organisms is still uncertain, as microbial phototrophy is dispersed among the three life domains (Eukaryotes, Bacteria and Archaea), with more than 20 diverse and unrelated phyla (Battistuzzi et al., 2004; Heimann & Huerlimann, 2015). The evolutionary evidences displayed by chemical markers, stromatolites fossils and microfossils support different possible origins (Olson et al., 2004). The primordial environment, poor in oxygen, might have selected

for the evolution of anoxygenic phototrophs. Chemosynthetic processes used for carbon fixation were dependent on specific substrates and ran with low efficiencies (Nowicka & Kruk, 2016). Photosynthesis allowed for the retention of energy from the sun, whereas reduced compounds potentially used as electron donors, such as  $H_2$  or  $H_2S$ , were available in the oceans (Hamilton et al., 2017).

#### 2.2.1 Rhodopsin-based phototrophic metabolism, a primordial way to harvest energy from light

The simplest phototrophic metabolism is used by archaea and some prokaryotes, utilizing photoactivated rhodopsins. Rhodopsins are a class of membrane-bound, light-driven pumps that harvest light to convert it into ATP, found in prokaryotic organisms and in the visual system of animals (Spudich & Jung, 2005). The prokaryotic rhodopsin pumps ions outside the membranes after light-induced isomerization of the retinal, creating an energy potential used by the ATP-synthase to produce ATP (Bamann et al., 2014; Bryant & Frigaard, 2006). With the advance of new-generation sequencing technologies like metagenomics, rhodopsins have been related to a wide taxonomic distribution. Many of the planktonic prokaryotes present in the photic zones of water surfaces that do not possess a photosynthetic reaction center possess rhodopsin-like systems (Bryant & Frigaard, 2006).

#### 2.2.2 Early anoxygenic phototrophs and the first photosynthesis

The first phototrophs developed around 3.5 billion years ago in an almost oxygen-free environment (Olson, 2006). Reduced compounds as  $H_2$ ,  $H_2S$ , sulfide, were greatly available in the primordial atmosphere. Anoxygenic phototrophs, using light and these reduced compounds as electron donors, synthesized complex organic molecules by reducing CO<sub>2</sub> or simple organic molecules. Anoxygenic phototrophs are unable to use water as an electron donor, and therefore no O<sub>2</sub> is produced as side product (Hanada, 2016).

The wide phylogenetic distribution reflects the environmental adaptability and the hyper-versatile metabolisms (Dubbs & Tabita, 2004) of these organisms, found in saline and non-saline environments across oxygen, sulfide and light gradients (Hallenbeck, 2017b). Members of the anoxygenic phototrophs guild can be found in three prokaryotic groups of green bacteria, purple bacteria and heliobacteria.

#### 2.2.3 Cyanobacteria and the premises of oxygenic photosynthesis

Around 3 to 2.4 billion years ago, cyanobacteria developed the ability to oxidize  $H_2O$ , with  $O_2$  formation as side product (Blankenship, 1992). They couple the fixation of  $CO_2$  with the production of  $O_2$ , in the in the

widely known process of photosynthesis. It is hypothesized that this event favored the accumulation of oxygen in the atmosphere, contributing to its development (Dismukes et al., 2001). The increasing  $O_2$  concentration both in aquatic and soil environments potentially allowed for the development of increasingly complex organisms, including algae and plants (Dismukes et al., 2001; Sharif et al., 2017). The development of eukaryotes is linked to endosymbiotic events between chemotrophic bacteria and oxygenic phototrophs. The presence of chloroplasts in algae and superior plants proofs this endosymbiosis (Whatley, 1993).

#### 2.3 Niche partitioning along light wavelengths and electron donors

Light, as the main energy source, is a key parameter for niche partitioning and selection mechanisms among phototrophic microorganisms. The light spectrum is distributed unevenly in waters, being influenced by the water depth (Stomp et al., 2007). Underwater, light intensity is diminished due to absorption and scattering phenomena (Gallegos & Moore, 2000), generally described by the Lambert-Beer equations (Abdelrhman, 2017).

$$I_L = I_0 \cdot e^{(-\varepsilon L)} \tag{1}$$

With  $I_L$  the irradiance at distance L from the source,  $I_0$  the incident irradiance and  $\varepsilon$  the extinction coefficient.

Light penetration in water columns is also dependent on the wavelengths (Stomp et al., 2007).

Short wavelengths, like blue light, penetrate deeply in pure water (> 30 m), whereas light with long wavelengths, like infrared light, can only penetrate less than 1 m (Chiang et al., 2011). Microbial photopigments absorb light at different wavelengths, from short wavelengths (*i.e.* carotenoids) to far infrared light (*i.e.* bacteriochlorophylls). This differential light absorption allows niche partitioning across the light spectrum (Cardona et al., 2018).

Photopigments are membrane-bound molecules, that absorb light and start an electrons cascade that ultimately reaches the production of ATP. Primary photopigments absorbing light are (bacterio-) chlorophylls, in the reaction centers of the photosynthetic units. Peripheral to those, carotenoids are accessory photopigments, that have the double function of absorbing light at lower wavelengths and prevent photooxidation (Madigan et al., 2015). Photopigments absorption peaks range between 300 and 1100 nm (Hou, 2014), allowing for a spatial distribution of colorful niches of phototrophs in the aquatic environments (Stomp et al., 2007). In aqueous environments irradiated by infrared light, purple bacteria find their niche.



Figure 1 Whole-cell wavelength spectrum of three PPB species. The absorption of light at different wavelengths allows niche partitioning across the light spectrum. Data from wavelength scan of Rhodobacter, Rhodopseudomonas and Rhodospirillum pure cultures isolated from an in-house enrichment.

#### 3. Purple phototrophic bacteria: whole communities in single cells

Purple phototrophic bacteria (PPB) form a guild of hyper-versatile organisms that thrive in ever-changing environments. The first observations of PPB date back to the 1920's, when van Niel, in Delft, discovered them. He was interested by these colorful bacteria (yellow, brown, red, and purple), and their interaction with light. Although representative lineages such as *Rhodobacter*, *Rhodopseudomonas* and *Rhodospirillum* are regrouped under the term PPB, they are not all / always purple *per se*. For instance, *Rhodobacter* relatives can reveal both purple and yellow colors depending on species, phototrophic niche and redox state of the environment in which they grow (Figure 2A). The diverse coloration is given by the photopigments present in the intracellular membranes, in particular carotenoids. The expression of different carotenoids induces a change in bacterial color appearance, which is as well linked to concentration and oxidation state of the carotenoids themselves (Aklujkar & Beatty, 2005).



Figure 2 PPB pure cultures.

A: Pure cultures of Rhodobacter blasticus grown aerobically (in pink) and anaerobically (in yellow). The color is linked to the oxidation state of the carotenoids present in the membranes; **B:** PPB show different coloration depending on the species. This is due to the presence of different types of photopigments, and, in particular, to a different expression of the carotenoid series.

PPB are a major phototrophic guild, that inhabit soils and prevalently aquatic environments. Stimulated by light, they participate in the carbon cycle by both fixing  $CO_2$  and oxidizing organic compounds. Anoxygenic phototrophs, PPB are among the first organisms to have developed on Earth, thanks to their ability to use inorganic compounds as electron donors (Yurkov & Beatty, 1998). Their photosystem is simpler compared to the oxygenic phototrophs, but PPB evolved complex metabolic strategies to find their niche. In nature, they can be found in aquatic environments exposed to light. They are present in sewage, lagoons, waste lagoons and lake or marine waters. Furthermore, they can be found in extremophilic environments, including hypersaline areas, at very low or very high temperature and pH conditions (Madigan, 2003).

More than 50 genera are now classified as PPB, but a univocal definition of the term PPB is not present (Hunter et al., 2009). Traditionally, PPB have been divided into purple sulfur bacteria (PSB) and purple non-sulfur bacteria (PNSB), based on their tolerance to sulfur compounds (van Niel 1944). This classification has later been replaced by molecular studies: 16S rRNA gene sequencing showed that PSB belong to the Gamma proteobacteria phylum, whereas PNSB to the Alpha- and Betaproteobacteria (Imhoff et al., 2005). Metabolically, PSB are primarily photoautotrophs, whereas PNSB are the most abundant photoorganoheterotrophic organisms in nature (Hunter et al., 2009).

Due also to their wide phylogenetic distribution, PPB present metabolic differences, and are probably the most versatile class of organisms know so far. The metabolic versatility has been reported not only between genera, but also in single species. PPB adapt their metabolism based on the environmental conditions, being able to growth phototrophically but also chemotrophically by aerobic and anaerobic respirations and fermentation in the dark, using numerous carbon sources, electron donors and acceptors present in their environment (Schultz & Weaver, 1982; Siefert & Pfennig, 1979). PPB exhibit a variety of ecophysiologies and phenotypes as across cell morphology, motility, salt tolerance, internal membrane structure, lipids content, and photopigments type and disposition. The only common trait PPB share is the cyclic electron transfer process in the photosynthetic membranes.

#### 3.1 Phototrophic machinery and cyclic photophosphorylation in PPB

From his studies on PPB, van Niel discovered that photosynthesis is a lightdependent reaction reducing  $CO_2$  to cellular material (Barker & Hungate, 1990). He generalized the reaction as

$$2H_2A + nCO_2 + light \to 2A + (CH_2O)_n + H_2O$$
(2)

With A as electron donor.

To capture the light, phototrophs use light-harvesting complexes, placed in a complex of intracellular membranes (Ruivo et al., 2014). Contrary to oxygenic phototrophs, which utilize a linear electron flow to produce ATP and reducing power, PPB convert light energy to chemical energy through a cyclic electron flow (McEwan, 1994).

Light is captured by antenna pigments in the reaction center, embedded in the photosynthetic membranes. It is used to excite the bacteriochlorophylls (Bchl), that becomes a strong electron donor. The excited electrons are transferred through an electron transport chain from low electronegative carriers to high electronegative carriers. They pass from the bacteriochlorophyll to the bacteriopheophytin, to the quinone pool, to the ferredoxin, to the cytochrome c, eventually returning to the reaction center. The electron flow generates a proton motive force. The reduced quinones are oxidized by protons derived from the tricarboxylic acid (TCA) cycle. The protons are transferred to the periplasm. Here, they are used to activate the ATP-synthase and produce ATP. The reducing power (NADH) required for growth is often achieved through a reverse electron transport. In this process, the electrons from the quinone pool are forced against the electrochemical gradient to reduce NAD<sup>+</sup> to NADH through the NADH-dehydrogenase (Madigan et al., 2015).



Figure 3 Schematic representation of the photosynthetic complexes in the membranes and cyclic electron flow in PPB. Photons captured by the bacteriochlorophylls (Bchl) in the Light Harvesting complexes (LH I and II) and used to excite electrons in the Reaction Center (RC). From here, electrons are transferred through the membrane-bound complexes, creating a proton motive force, that activates the ATP synthase.

#### 3.2 PPB: champions in metabolic versatility

As the name says, PPB are phototrophs, harvesting light from the sun spectrum and converting it to chemical energy. Under anaerobic conditions, and with light as energy source, they can use both organic and inorganic carbon sources. PPB can use several organic compounds as carbon sources, electron donors and electron acceptors, depending on the oxidation state of the electron donor. They aerobically as chemotrophs can grow both or anaerobically as photoheterotrophs. Being able to adapt to several environmental conditions, they are considered champions in metabolic versatility (Nelson & Fraser, 2004). A schematic overview of the possible metabolic modes of PPB is presented in Chapter 3 (Table 1).



Figure 4 Schematic of the possible metabolic modes of PPB. Depending on the environmental conditions, PPB can show a photoorganoheterotrophic metabolism, photolithoautotrophic, chemoorganotrophic metabolism, dark and light fermentation.

#### 3.2.1 Photolithoautotrophic growth

Photolithoautotrophic growth was one of the first to develop, with the use of reduced compounds, such as  $H_2$  or  $H_2S$  as electron donors, and biomass as electron acceptor (Nowicka & Kruk, 2016). Opposite to what happen in oxygenic phototrophs, oxygen is not used by PPB as electron acceptor during phototrophic growth. Indeed, phototrophic growth under aerobic conditions is limited, as oxygen inhibits the formation of the photosynthetic complexes (Donohue et al., 1988).

Photosynthetic organisms can concentrate and fix inorganic carbon supplied by dissolved carbon dioxide ( $CO_2$ ) and bicarbonate ( $HCO_3^-$ ) in the well-known process called photosynthesis. All phototrophic organisms can grow with photolithoautotrophic metabolism, utilizing different routes for  $CO_2$  fixation. Photolithoautotrophic growth is divided into two steps. In the first light is harvested as previously described to produce ATP and reducing power.

During the second step, in the so-called dark reactions, the inorganic carbon is assimilated and converted / reduced into carbohydrates and complex organic molecules, that can be used to build the cell materials (Madigan et al., 2015).

Five known pathways enable  $CO_2$  fixation, namely the Calvin-Benson cycle (used by eukaryotic and prokaryotic microalgae, heliobacteria, and PPB) (Blankenship et al., 1995), the reductive or reverse tricarboxylic acid (RTCA) cycle (used by green sulfur bacteria) (Tang & Blankenship, 2010), the 3-hydroxypropionate bi-cycle (used by green non-sulfur bacteria) (Shih et al.,

2017), the reductive acetyl-CoA pathway (Tang et al., 2011), and the 3-hydroxypropionate/4-hydroxybutyrate cycle (Thauer, 2007).

The most widely distributed pathway for  $CO_2$  fixation is the Calvin-Benson cycle, used also by PPB. The Calvin-Benson cycle utilizes reducing power and ATP produced in the light reactions to fix  $CO_2$  to carbohydrates, through four major phases: *i*) the carboxylation phase:  $CO_2$  is condensed to a molecule of ribulose bisphosphate (5-C molecule) by the enzyme ribulose bisphosphate carboxylase to form two molecules of 3-phosphogliceric acid (PGA); *ii*) the reduction phase: the PGA is phosphorylated and reduced to glyceraldehyde-3-phosphate; *iii*) in the regeneration phase, the 5-C sugar is reformed, by fixating another molecule of  $CO_2$  to 3 to 7 sugar phosphates; and *iv*) the production phase, where carbohydrates, fatty acids, amino acids and organic acids are synthetized (Richmond, 2003).

In PPB, photolithoautotrophic growth is secondary to photoorganoheterotrophic growth, and can only occur in presence to electron acceptor different from  $O_2$ . However, in presence of an excess of electrons,  $CO_2$  fixation can be used as electron sink during photoheterotrophic growth (McKinlay & Harwood, 2010).

#### 3.2.2 Photoorganoheterotrophic growth

The photoorganoheterotrophic metabolism is the preferred growth mode of PPB, allowing them to thrive on carbon and electrons derived from organic compounds. Besides being less widely described than photolithoautotrophy, photoorganoheterotrophy is relatively widespread in natural environments.

Contrary to chemoorganoheterotrophic organisms that need to catabolize a substantial part of the organics to produce ATP, photoorganoheterotrophs derive their energy directly from light. They can therefore grow with high biomass yields over organic substrates, sometimes close to 1 g COD<sub>x</sub> g<sup>-1</sup> COD<sub>s</sub> (Alloul et al., 2019). In the photoorganoheterotrophic growth, PPB use organics as electron donors, and, in balanced conditions, biomass as electron acceptor. The electron and carbon sources span from volatile fatty acids (VFAs), to alcohols, to carbohydrates. The terminal electron acceptor is, when an electron imbalance is present, redirected toward other routes (see Chapter 2). In the light, PPB are also able to activate a photofermentative metabolism. In this case, sugar degradation is not coupled with growth, but it is only used as dissipative mechanism with electron redirection toward H<sub>2</sub> formation (Basak et al., 2014).

On top of the photolithoautotrophic growth using  $CO_2$ , PPB can grow on other C1 compounds, like methanol, formate and CO. In this case, C1 compounds are oxidized to  $CO_2$ , which is then fixed via the Calvin-Benson Cycle (Quayle & Pfennig, 1975; Sahm et al., 1976). The RubisCO enzyme is here used both for carbon fixation and for redox balance, as it allows to use  $CO_2$  as electron acceptor.

Acetate is a volatile fatty acid, often used as a model compound since it can potentially be assimilated by all PPB. Acetate is metabolized through different pathways, depending on the species. For example, acetate can be used by PPB both as electron donor and electron acceptor (Butow & Dan, 1991) as well as carbon source. As carbon source, it is assimilated by species related to *Rhodopseudomonas* through the glyoxylate cycles (Petushkova et al., 2021), whereas *Rhodobacter* species uses the ethyl-malonyl pathway (Laguna et al., 2011). Alternatively, in species lacking the glyoxylate cycle, the citramalate pathway has been proposed as an alternative pathway (Berg & Ivanovsky, 2009). Other organics, like VFAs or ethanol, are first metabolized to acetyl-CoA and then incorporated into the cellular metabolism through the TCA cycle (Hunter et al., 2009). In PPB metabolism, the TCA cycle is active independently from the carbon source present in the medium, and serves both to catabolic and anabolic reactions. It catalyzes the full oxidation of acetyl-CoA to CO<sub>2</sub>, providing reducing equivalents for ATP formation during dark respiratory processes, and at the same time it supplies the biosynthetic pathways with metabolites and reducing power (Petushkova et al., 2019).

#### 3.2.3 Chemotrophic growth

PPB are also capable to grow as chemotrophs when light is absent. The presence of oxygen inhibits at gene-regulation level the formation of the photosynthetic complexes. Under these conditions, PPB can still grow as aerobic chemoorganoheterotrophs, even though with lower growth rates compared to phototrophic metabolism (Madigan et al., 1980). Under dark conditions PPB can ferment sugars (both carbon and electron donors) for biomass formation and biohydrogen production. Growth in the dark is limited, and strictly dependent to the presence of an external source of energy and reducing power (Madigan & Gest, 1979; Petushkova et al., 2021)

## 4. Environmental biotechnologies fostered by purple phototrophic bacteria

Microorganisms have been used as living factories for centuries, as exemplified by the production of beer and bread. In more recent times, microbial cell factories have been developed growing interest, as workhorses for industrial bioprocesses. The extreme metabolic variability of phototrophs is reflected on the spectrum of compounds that they produce as secondary metabolites under different environmental constrains. Used as electron excess sinks, nutrient storage compounds, or for self-defense, bioproducts from phototrophs attract interests for technology. The main fields of applications include biogas (methane and hydrogen), feedstock additives (with single cell protein production), biopolymers formation (as bioplastics like PHAs) and pharmaceutically interesting compounds (antioxidants and quinones).



Figure 5: Understanding the complex metabolism of PPB, from molecular to process scale, allows us to harness its potential, from wastewater treatment to high-value produce production.

#### 4.1 Biohydrogen

Hydrogen (H<sub>2</sub>) is a renewable and clean source of energy. Four major pathways can be harnessed for biological H<sub>2</sub> production: *i*) direct biophotolysis, the simplest and most effective process to produce hydrogen, directly transferring electrons from water to hydrogen-evolving enzymes. It is accompanied by the simultaneous evolution of O<sub>2</sub> and H<sub>2</sub> (Rastegari et al., 2019). *ii*) Indirect biophotolysis, linked to aerobic photosynthesis reactions (Hallenbeck, 2017a). *iii*) Dark fermentation, where H<sub>2</sub> is a side product of carbohydrates fermentation (Abo-Hashesh & Hallenbeck, 2012). *iv*) Photofermentation, performed by PPB under anaerobic light conditions. Short-chain organic acids are used as electron donors, and H<sub>2</sub> is used as electron sink (Androga et al., 2014). Opposite to the dark fermentation, this process does not lead to biomass growth. Two enzymes are responsible for H<sub>2</sub> production in PPB, namely the nitrogenase and hydrogenase enzymes. Nitrogenase is a nitrogen fixing enzyme active under anaerobic conditions (Basak et al., 2014). In presence of nitrogen gas, H<sub>2</sub> production is only secondary to nitrogen fixation, according to the Equation 3

$$N_2 + 8H^+ + 8e^- + 16ATP \rightarrow 2NH_3 + H_2 + 16(ADP + Pi)$$
 (3)

In absence of  $N_2$ , the nitrogenase catalizes the production of hydrogen from protons through an irreversible reaction, as in Equation 4

$$2H^+ + 2e^- + 4ATP \rightarrow H_2 + 4(ADP + Pi) \tag{4}$$

The hydrogenase, instead, catalizes a simple and reversible redox reaction, as in Equation 5

#### $2H^+ + 2e^- \leftrightarrow H_2$

(5)

#### 4.2 Poly-β-hydroxyalkanoates (PHA)

Polyhydroxyalkanoates (PHA) is a group of polyesters produced by microorganisms, stored intracellularly in granules as carbon and reducing equivalents sinks (Madison & Huisman, 1999). They have material properties similar to polypropylene, and find application as petrochemical plastic substitutes (Bugnicourt et al., 2014). PHA have the advantage of being recyclable and biodegradable. Biologically, PHA are produced as reducing power and carbon storage in presence of carbon excess. In PHA producing organisms, PHA can be stored up to 90% of the weight of the cells. In a similar way, PPB can store up to 90% of the dry weight as PHA in pure cultures (Kranz et al., 1997; Sangkharak & Prasertsan, 2007), but the accumulation is dependent on the strain and on the carbon source used. In mixed cultures, it was possible to achieve a maximum of 20% of the dry weight (Luongo et al., 2017).

#### 4.3 Single-cell proteins

Microbial (single-cell) proteins (SCP) are an alternative to increase nutrient content in the animal feed. PPB are an interesting sources of SCP, due to their high protein content, vitamins and photopigments. The use of PPB in shrimp feed has increased the survival and healthiness of the animals. Further applications explore the use of PPB for human consumption in space missions (Chumpol et al., 2018).

#### 4.4 Photopigments

Photopigments are crucial in PPB for light harvest and following energy production. The bacteriochlorophylls are the main light harvesting pigments. They have been proposed as photosensitizers for photodynamic therapy of cancer (Liu et al., 2012; Pantiushenko et al., 2015). Carotenoids, have broad and known utilization ad dyes in food, and as food and feed additives, thanks to their antioxidant properties (Barreiro & Barredo, 2018; Cheng et al., 2020). PPB tune their photopigment content based on light conditions (Liu et al., 2019). PPB utilization as photopigment producers has been explored as animal feed, mainly in combination to SCP production (Chumpol et al., 2018).

## 5. Closing the loop with purple bacteria: from nutrient removal to production of high-value products from wastewater

Wastewaters can present fluctuating characteristics that influence the substrate consumption abilities of the biological sludges used, including temperature and nutrient (C, N, P) load. Despite the plasticity of the microbial communities, the varying conditions can severely affect their composition and activity, factually reducing the removal of pollutants. PPB, instead, thrive in ever-changing environment. Their ability grow on numerous substrates and electron sources rises interest for high-strength wastewater treatment. Coupled with the possibility to produce industrially interesting compounds, PPB are a promising guild for bio factories.

Wastewater treatment with green phototrophs has been proposed as an efficient process for nutrient capture and biovalorization, in example for lipids and biofuels production. Compared to green phototroph systems, PPB present some advantages: i) an hyperversatile metabolism, adaptable to several environmental conditions. When in presence of infrared light, PPB become the dominant community. Laboratory scale experiments showed a stable enrichment above 70% (Hülsen et al., 2016). *ii*) High quantum and carbon yields. The efficiency of photosynthetic conversion of PPB is close to 1 (Sener et al., 2016): this implies that almost all the energy harvested from the photosynthetic complexes can theoretically be converted into chemical energy. Compared to microalgae, in example, PPB show photosynthetic conversion efficiencies 2 times higher (Perrine et al., 2012). (Perrine et al., 2012) iii) Similarly, PPB have high biomass yields over their carbon sources, with a theoretical maximum close to  $1 \text{ gCOD}_x \text{ gCOD}_s^{-1}$ <sup>1</sup>(Alloul et al., 2019). The production of ATP through light-mediated reactions allows the cells to utilize the carbon sources entirely for biomass formation, without having to dissipate a part for energy formation. A higher biomass yield is advisable in biofactories, where the product recovery is dependent on the cell concentration.

#### Knowledge gaps

#### 1. Electron reallocation in pure cultures

Loads of information is available on the fundamental molecular mechanisms governing the PPB metabolism. Their fascinating complexity has been studied in detail in pure culture systems, and metabolic models have been constructed to clarify and simplify their complexity. Pure culture studies provide a simplification of the genomic and functional potential of the systems. They have the advantage to allow a detailed examination of the metabolism and the responses of the cultures to environmental factors, in example. Despite the numerous studies on the effects of light cycles on PPB, the internal electron allocation under intermittent energy supply remains puzzling, and the mechanisms that govern the preferential electron flow have not been explained.

#### 2. Light: a central parameter often overlooked

Light, being the energy source for PPB, is of foremost importance. Physiological responses of PPB have primarily been studied under continuous illumination conditions. However, in nature, daily light / dark cycles occur, implying a discontinuous supply of energy. Its effects on the PPB metabolism have been overlooked, and only few studies provide observations on the responses to light / dark cycles. The importance of light to PPB cultures is proven by pure culture studies, as in the case of *Rhodopseudomonas*, where it was possible to correlate growth rates and light intensity (Kuo et al., 2012). However, the effect on mixed cultures is yet to be elucidated. The control of light supply, and its relationship with PPB kinetics, is crucial to design PPB-based WWT processes.

#### 3. Microbial ecology of the PPB black box

The applicability of mixed cultures PPB to WWT has been demonstrated, but often the microbial ecology has been neglected. Mixed-culture systems rely on the metabolic performance of microbial communities, that compete and interact for resources in an ecosystem. However, in environmental biotechnology, microbial communities are often considered as a black box, where kinetics and stoichiometry are used as predominant parameters to define the performances of a process. Understanding the community dynamics, with deeper insights into the microbiological black box, is crucial for a systematic understanding of the observed phenomena. Recent studies focus on the overall performances of PPBbased WWT, but the microbial ecology is overlooked. The principles for an efficient selection and aggregation have not been described. Different carbon and electron inflow streams affect the selection principles and therefore the applicability of PPB to numerous waste streams. Harnessing microbiomes relies on managing the metabolisms of microorganisms selected to perform the conversions of interest. The effects of environmental factors, as nutrient type and load, irradiation intensity and patterns, have been studied in pure cultures. Nonetheless, a microbial community responds differently compared to a pure culture to the same environmental factors. Inter-species interactions, such as cooperation and competitions phenomena, are crucial to understand the microbial complexity and applicability in biobased processes.
### Research question

The core research question of this thesis will focus on the fundamental investigation of the metabolism of PNSB in the context of water resource recovery, and is formulated as such:

### How can we harness the microbial ecology and metabolic versatility of purple non-sulfur phototrophic bacteria towards water resource recovery and biorefinery?

The thesis will develop around four main research questions:

- 1. How do PPB respond to redox imbalance and allocate carbon and electrons during light / dark diel cycles?
- 2. How can we enrich and aggregate PPB in mixed cultures for an intensified capture/removal of nutrients?
- 3. How are PPB differently selected on fermented (*i.e.*, VFAs) *vs.* fermentable substrates (*i.e.*, carbohydrates/sugars)?
- 4. What are the effects of decreasing light intensities on the physiology/growth of PPB, community composition, and conversions?

### Objectives

With this thesis I aimed to shed light on the microbial ecology principles governing PPB enrichments for wastewater recovery.

In Chapter 2, we evaluated the effects of diel light/ dark cycles in the model organisms Rhodopseudomonas palustris. We aimed to identify the metabolic drivers for the selective production of PHA or H<sub>2</sub>, two major redox sinks for PPB with commercial applications. In Chapter 3, a method for PPB enrichment and aggregation was proposed. We coupled a high PPB enrichment to high nutrient removal, so that to comply to the current European law. Depending on the wastewater, the type of carbon and electron source variates. In example, agricultural / food wastes are prevalently composed of sugars. In Chapter 4 we evaluated the interactions of PPB with fermentative bacteria in glucose-based environment, in regard also to the reactor regimes. PPB convert light energy into chemical energy, factually being able to assimilate carbon sources for biomass growth. In Chapter 5, we evaluated the metabolic requirements of a PPB community subjected to a decreased light intensity, to establish a minimum irradiation limit for PPB biomass growth and nutrient removal. The general conclusions of this thesis are given in Chapter 6. Chapter 7 presents an outlook on future research, with a particular focus on the application of PPB mixed communities for wastewater resource recovery.

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Effects of light/dark diel cycles on electron allocation to PHAs and H<sub>2</sub> in the photoorganoheterotrophic metabolism of *Rhodopseudomonas palustris* 

This chapter is deposited in a modified version as preprint with contributions of: Marta Cerruti, Heleen T. Ouboter, Viktor Chasna, Mark C. M. van Loosdrecht, Cristian Picioreanu, David G. Weissbrodt (2020) Effects of light / dark diel cycles on the photoorganoheterotrophic metabolism of *Rhodopseudomonas palustris* for differential electron allocation to PHAs and H<sub>2</sub>, *bioRxiv*, preprint, 2020.08.19.258533. https://doi.org/10.1101/2020.08.19.258533

### Abstract

Purple non-sulfur bacteria (PNSB) are hyperversatile photoorganoheterotrophs that foster organic aqueous waste biorefinery from pure-culture to mixed-culture biotechnologies. Light/dark diel cycles can impact the electron distribution in their metabolism and resulting product formation. For survival and growth, PNSB have developed different mechanisms to allocate excessive electrons under dynamic growth conditions. Products such as dihydrogen  $(H_2)$  and intracellular poly-B-hydroxyalkanoates (PHAs) can form alternative electron sinks to biomass formation. Coupling between illumination patterns and redox balancing in the PNSB physiology remained to be uncovered. A controlled continuous culture of Rhodopseudomonas palustris, a model PNSB isolated from an in-house enrichment, was fed with an acetate-based balanced nutritive medium and exposed to three irradiance conditions: 24 h continuous infrared illumination, 16 h light/8 h dark, and 8 h light/16 h dark alternations. Durations of light and dark phases over a cycle determined the light energy availability level and attainment of a metabolic stationary state. Under long dark phases, acetate accumulated to levels (ca. 8 C-mM L<sup>-1</sup>) that were depleted by growth in the light. Under short dark phases, acetate was rapidly consumed in the light with most of phototrophic growth occurring under acetate-limiting conditions. Diel cycles resulted in an imbalance between substrate uptake and biomass growth. Rhodopseudomonas shunted the excess of carbon and electron flows first toward PHA production. Only secondarily when PHA storage got saturated, the remaining electron excess was redirected toward H<sub>2</sub>. A numerical model was developed to describe dynamics of biomass, nutrients, off-gas compounds (CO<sub>2</sub>, H<sub>2</sub>) and PHAs during light/dark cycles. Simulations allowed for extracting stoichiometric and kinetic parameters exhibited by *Rhodopseudomonas*. Insights on ecophysiological dynamics of PNSB induced by light/dark diel cycles can help understand, predict and control their functioning in biorefinery processes.

### 1. Introduction

Purple non-sulfur bacteria (PNSB) form a guild of hyper-versatile anoxygenic phototrophs which establish their niche in the infrared (IR) light wavelength domain (Blankenship et al., 1995). PNSB grow on various organic and inorganic substrates (Alloul et al., 2019), capture nutrients (Cerruti, Stevens, et al., 2020a), and produce biocompounds of interest for the industry and circular economy like fertilizing nutrient-rich biomasses, single-cell proteins for food ingredients, carotenoids antioxidants, intracellular poly- $\beta$ -hydroxyalkanoates (PHAs) for biodegradable plastics, and dihydrogen (H<sub>2</sub>) gas for biofuel (Hülsen et al., 2018; Hustede et al., 1993; Vincenzini et al., 1997; Wang et al., 2017). PNSB foster biorefinery opportunities from pure-culture to co-culture and mixed-culture biotechnologies. *Rhodopseudomonas palustris* is one model PNSB (Madigan & Jung, 2009), widely described and used biotechnologically.

Under light and in presence of organic substrates as carbon and electron sources, PNSB grow photoorganoheterotrophically. In the dark, they can grow chemoorganoheterotrophically on sugars by fermentation (Madigan et al., 1980), or by aerobic or anaerobic respiration when exogenous electron acceptors are supplied (as dissolved oxygen, nitrate, trimethylamine-N-oxide or dimethyl sulfoxide) (Ho Lee et al., 2002; Imhoff, 1995; Madigan et al., 1982).

Light is crucial for phototrophy by providing energy to cells. PNSB capture the photonic energy and couple the light-driven oxidation of the photopigments to an electron transfer through membrane bound-enzymes. The trans-membrane gradient of hydrogen protons ( $H^+$ ) supports ATP synthesis through cyclic photophosphorylation (Madigan et al., 2015). NADH is mainly produced during catabolic processes. In some cases, a reverse electron flow takes places, and the NADH-dehydrogenase catalyzes the proton transfer from the ubiquinone pool to NAD<sup>+</sup>. The control of the redox balance is crucial for cell survival and growth. The ratio between NADH/NAD<sup>+</sup> is important for intracellular redox homeostasis (Green & Paget, 2004). PNSB generate NADH in three different ways (Figure 1), namely: 1) catabolic processes (Laguna et al., 2011); 2) light-driven reactions through the quinone pool (Blankenship et al., 1995); and 3) reverse electron transfer (Dupuis et al., 1998).



Figure 1. Schematic representation of the reducing power allocation in Rhodopseudomonas. <u>Red lines</u>: electron flow; <u>Black dashed lines</u>: proton transfer; <u>Blue dotted lines</u>: NADH is normally produced in the catabolic processes, whereas NADPH is used in the anabolic processes, such as poly-hydroxyalkanoate (PHA) formation. Biomass is the primary electron sink in PNSB. PHAs and H<sub>2</sub> are two of the other possible electron sinks.

To cope with possible redox imbalances, PNSB redistribute the electrons toward different routes (Spero et al., 2016). In Rhodopseudomonas, the first and most important electron sink is the biomass itself, that receives the intermediates of the tricarboxylic acid cycle (TCA) and the NADPH produced there (McKinlay & Harwood, 2011). Secondly, the Calvin-Benson-Bassham (CBB) cycle is constitutively active also under photoorganoheterotrophy as central recycling mechanism of redox cofactors coming from the anabolic process (Gordon & McKinlay, 2014; McKinlay & Harwood, 2011). Under photoorganoheterotrophic anaerobic conditions, a surplus of electrons can be disposed through PHA or H<sub>2</sub> production, besides CO<sub>2</sub> fixation (Harwood, 2008). Both these electron sinks are normally produced under nitrogen limitation. H<sub>2</sub> production by PNSB is substantially mediated by nitrogenase (Blankenship et al., 1995). H<sub>2</sub> is a side-product of the dinitrogen fixation process, that can only happen when preferred nitrogen sources like ammonium are depleted (Basak et al., 2014; Kim et al., 1980; Oda et al., 2005). Hydrogenase-mediated H<sub>2</sub> production has been reported in presence of an oxidized organic compound (*i.e.* malate) (Willison et al., 1983). In the environment, numerous bacterial species accumulate PHAs as a means of carbon and electrons storage, conferring them a competitive advantage (Kutralam-Muniasamy et al., 2017). In lab-scale experiments and industrial

processes, to achieve an overproduction, these storage polymers are synthetized when acetyl-CoA and NADPH are in excess but nutrients like nitrogen, sulfur or phosphorus are limiting (Montano-Herrera et al., 2017). Similarly, PNSB can produce PHAs (Hustede et al., 1992) as a means of carbon and electron balance, utilizing the reducing power to build the storage polymers (De Philippis et al., 1992; Haywood et al., 1988). PHA production in PNSB occurs under balanced growth, and it increases under nutrient limitations (Higuchi-Takeuchi et al., 2016). The two processes of  $H_2$  and PHA productions are considered competitive electron dissipation pathways (Hustede et al., 1993; Padovani et al., 2016). However, due to the hyper-versatility of PNSB metabolisms, a univocal response to redox imbalances has not yet been identified.

In natural environments, phototrophs are subjected to diel (*i.e.*, 24-h period) light/dark cycles. Since light is responsible for the production of energy and reducing power, the irradiation patterns impact the cellular physiology. The light and dark cycles impact redox and ATP balance of the cells, subjecting them to metabolic switches. The impact of diel cycles on the metabolic response of PNSB and internal electron allocation patterns remains puzzling. Allocation of reducing power toward the aforementioned routes is known, but mechanisms that govern the preferential electron flow have not been explained yet.

Here, we investigated the effects of diel light/dark cycles on the physiology and electron allocation in PNSB. We used *Rhodopseudomonas palustris* as representative organism isolated from an in-house PNSB enrichment culture implemented for nutrient capture from wastewater (Cerruti, Stevens, et al., 2020). The light/dark dynamic conditions imposed on the acetate-fed continuous culture were hypothesized to influence the organismal physiology by the intermittent supply of the IR light energy source. The process mimicked natural day/night cycles that generate a redox imbalance. Using wet-lab experiments and mathematical modelling, we elucidated the preferential redistribution of carbon and electron flows toward biomass, PHAs and  $H_2$  in function of the lengths of the light and dark periods and of the underlying dynamics in substrate conversions.

### 2. Material and Methods

### 2.1 Strain

The Rhodopseudomonas strain was isolated by agar colony selection from an inhouse PNSB enrichment culture run for nutrient removal from synthetic wastewater (Cerruti, Stevens, et al., 2020a). The isolate was characterized by full-length 16S rRNA gene sequencing. The genomic DNA was extracted using UltraClean® Microbial DNA Isolation Kit (MO BIO Laboratories, Inc., USA), following manufacturer's instructions. The full 16S rRNA gene was amplified polymerase forward by chain reaction using GM3R (5'-AGAGTTTGATCMTGGC-3') GM4F (5'and reverse TACCTTGTTACGACTT-3') primers (Muyzer et al., 1995), and sequenced for phylogenetic identification by Sanger sequencing (Baseclear, NL). The nucleotide sequence was aligned over the NCBI BLAST database (Altschul et al., 1990), and resulted in a 98.1% identity and query cover with Rhodopseudomonas palustris. The amplified sequences are deposited under the accession number (see Supplementary information SUB10460307 rps chem4E GM3 OK350353-4).

### 2.2 Medium

The inflow medium of the continuous cultures was adapted from Cerruti et al. (2020). It consisted of (per liter):  $0.914 \text{ g CH}_3\text{COONa}\cdot3\text{H}_2\text{O}$  (13.5 C-mmol L<sup>-1</sup> and 54 mmol electrons L<sup>-1</sup> when expressed via degree of reduction or 432 mg COD L<sup>-1</sup> when expressed as chemical oxygen demand), 0.229 g NH<sub>4</sub>Cl (*i.e.*, 4.281 mmol N L<sup>-1</sup> or 60 mg N-NH<sub>4</sub><sup>+</sup> L<sup>-1</sup>), 0.014 g KH<sub>2</sub>PO<sub>4</sub> and 0.021 g K<sub>2</sub>HPO<sub>4</sub> (*i.e.*, 0.223 mmol P L<sup>-1</sup> or 7 mg P L<sup>-1</sup>), 0.200 g MgSO<sub>4</sub>·7H<sub>2</sub>O, 0.200 g NaCl, 0.050 g CaCl<sub>2</sub>·2H<sub>2</sub>O, 1 mL solution of vitamins, 1 mL solution of trace elements, and 4.7 g 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) used as pH buffer. The medium was autoclaved to assure sterility.

The solution of vitamins was composed of (per liter): 200 mg thiamine–HCl, 500 mg niacin, 300 mg  $\rho$ -amino-benzoic acid, 100 mg pyridoxine–HCl, 50 mg biotin and 50 mg vitamin B12.

The solution of trace elements was composed of (per liter): 1100 mg Na EDTA $\cdot$ 2H<sub>2</sub>O, 2000 mg FeCl<sub>3</sub>·6H2O, 100 mg ZnCl<sub>2</sub>, 64 mg MnSO<sub>4</sub>·H<sub>2</sub>O, 100 mg H<sub>3</sub>BO<sub>3</sub>, 100 mg CoCl<sub>2</sub>·6H<sub>2</sub>O, 24 mg Na<sub>2</sub>MoO<sub>4</sub>·2H<sub>2</sub>O, 16 mg CuSO<sub>4</sub>·5H<sub>2</sub>O, 10 mg NiCl<sub>2</sub>·6H<sub>2</sub>O and 5 mg NaSeO<sub>3</sub>.

The stock solutions of trace elements and vitamins were filtered on 0.22  $\mu m$  filters (Whatman, USA).

### 2.3 Reactor setup

Continuous cultures were run to evaluate the influence of the light/dark diel cycles on the physiology of *Rhodopseudomonas palustris*. A 1.5-L continuous-flow stirred-tank reactor (CSTR) with 1.2-L working volume and cylindrical geometry was connected to a programmable logic controller (In-Control and Power unit, Applikon, NL) and operated with stirring at 350 rpm, at pH 7.0  $\pm$  0.5, and a temperature of 30 $\pm$ 1 °C during illumination phases and 20°C during dark phases. This temperature pattern resulted from the intermittent switches of the IR light supply. Argon gas (Linde, NL, >99% purity) was sparged continuously in the bulk liquid phase at 120 mL min<sup>-1</sup> to maintain anaerobic conditions. The use of N<sub>2</sub> gas was avoided since interfering with nitrogenase-mediated H<sub>2</sub> production pathways (Harwood, 2008). The continuous flow rate was set at 0.048 L h<sup>-1</sup>. It corresponded to a dilution rate of 0.04 h<sup>-1</sup> chosen based on the growth rate of *Rhodopseudomonas* pure cultures previously measured at 0.11 h<sup>-1</sup> (data not shown). The biomass was maintained in the CSTR at a low concentration of 0.26  $\pm$  0.05 g dry weight L<sup>-1</sup> to minimize light shading effects.

The reactor was placed in a shaded hood to tightly control the irradiation patterns. Two halogen floodlight lamps (Handson, NL) were positioned at opposite sides of the reactor diameter. The incident white light spectrum was filtered to supply infrared (IR) light ( $\lambda > 700$  nm) using two Black 962 Infrared Transmitting Perspex Acrylic Sheets of 70 x 70 cm (Black Perspex 962, Plasticstockist, UK). The IR light intensity measured at the reactor surface with a pyranometer (CMP3; Kipp & Zonen, NL) was 300 W m<sup>-2</sup>, enabling non-light-limited conditions in the reactor (Cerruti et al., in prep.). An automatic timer device (Grundig, NL) was used to switch on/off the light at required time sets. Three irradiance conditions were tested: 1) continuous illumination (*i.e.*, 24 h light), 2) 16 h light and 8 h dark cycles, and 3) 8 h light and 16 h dark cycles.

### 2.4 Analytical methods

The CO<sub>2</sub> and H<sub>2</sub> in the offgas were measured using an online mass spectrometer (Thermofisher, Prima BT Benchtop MS) connected to the bioreactor. The production rates of these components were calculated using the argon gas inflow rate (120 mL  $h^{-1}$ ).

The biomass concentration was measured from the mixed liquor by absorbance at 660 nm (A<sub>660</sub>) using a spectrophotometer (Biochrom, Libra S11, USA). A calibration curve was established to correlate A<sub>660</sub> to dry weight (DW) concentration of biomass:  $c_X$  (g DW L<sup>-1</sup>) = 0.64 A<sub>660</sub> – 0.06. Biomass dry weight was obtained by taking samples from the mixed liquor, filtering them using 0.45 µm filters (Whatman, USA), and drying them in a 70 °C stove for 72 h (adapted from Lip et al. 2020). Acetate was measured from the filtrates with a high-performance liquid chromatograph (HPLC) (Waters, 2707, NL) equipped with an Aminex HPX-87H column (BioRad, USA). Analytes were eluted using  $H_3PO_4$  (1.5 mmol L<sup>-1</sup>, flowrate of 0.6 mL min<sup>-1</sup>, temperature of 60°C) prior to detections by refraction index (Waters 2414) and UV (210 nm, Waters 484) spectrophotometry.

Concentration of ammonium (as  $N-NH_4^+$ ) and orthophosphate (as  $P-PO_4^{3-}$ ) were measured from the filtrates with a discrete analyser (Thermoscientific Gallery, NL) using manufacturer's reagents.

PHA measurements were performed with fluorimetry for high sensitivity on low concentrations of biomass maintained in the reactor. The low biomass concentration did not allow for traditional extractions of PHA and gas chromatographic measurements of monomers. PHAs were stained in the biological samples with Nile red (CAS n. 7385-67-3, Sigma Aldricht), as in (Zuriani et al., 2013). Fluorescence was measured with a microplate reader (1000M pro, Tecan), with excitation at 535 nm and emission at 605 nm.

A calibration curve was used to correlate fluorescent counts to mol fractions of PHAs expressed as hydroxybutyrate (HA) monomer equivalents (C-mmol<sub>HB</sub> C-mmol<sub>X</sub><sup>-1</sup>) (Supplementary information SI-1). PHA quantifications for the calibration line were performed as in Vermeer et al., (2021).

### 2.5 Mathematical model

A mathematical model was developed to characterize the behavior of the continuous pure culture of *Rhodopseudomonas* exposed to light/dark cycles. The following assumptions were adopted to build the model structure: (*i*) photoorganoheterotrophic growth was assumed during the light period; (*ii*) under dark, a completely ceased metabolic activity was considered since acetate was the sole carbon source and cannot be metabolized further in absence of light and external electron acceptors. The light-driven growth stoichiometry was defined as in Table 1, with the biomass composition adapted from McKinlay & Harwood (2010).

The growth process was described by a system of balance equations for the relevant substances present in the mixed liquor (Table 2, equations 1-7), namely biomass ( $C_X$ ), acetate ( $C_S$ ), ammonium (NH<sub>4</sub><sup>+</sup>,  $C_N$ ), phosphate (PO<sub>4</sub><sup>3-</sup>,  $C_P$ ), carbon dioxide (CO<sub>2</sub>,  $C_C$ ), and PHAs as hydroxybutyrate monomer equivalents per mass fraction (HB,  $C_{PHA}$ ). NH<sub>4</sub><sup>+</sup>, PO<sub>4</sub><sup>3-</sup>, were dissolved in the liquid phase. CO<sub>2</sub> was dissolved in the liquid and the gas phases. Biomass and PHAs were present in the solid phase.

Components i →	Acetate	Ammonium	Phosphate	Biomass	Carbon dioxide	PHAs	Water	Protons
Abbreviation →	(S)	(N)	(P)	(X)	(C)	(HB)	(M)	(H)
Molecular formula →	сн <sub>3</sub> соон	$\mathrm{NH}_4^+$	$PO_3^{4-}$	$\begin{array}{c} C_1 H_{1.8} O_{0.} \\ {}_{38} N_{0.18} P_{0.0} \end{array}$	$CO_2$	$C_4H_8O_3$	$\rm H_2O$	$\mathrm{H}^{\dagger}$
Units →	C-mol	mol	mol	14 C-mol	C-mol	C-mol	mol	mol
↓ Microbial processes j								
<ol> <li>PPB growth on acetate Acetate → Biomass</li> </ol>	-0.57125 $(-Y_{SX,I} = -1/Y_{XS,1})$	$-0.18$ (- $Y_{N/X,1}$ )	-0.014 $(-Y_{P/X,1})$	1 fixed	$0.1425$ $(Y_{CX,1})$	0	$0.5335$ $(Y_{WX,1})$	$0.138$ $(Y_{HX,1})$
2) Acetate storage as PHAs Acetate → PHA	-2.25 (- $Y_{S/HB,2}$ = -1/ $Y_{HB,S,2}$ )	0	0	0	$0.5$ $(Y_{C/HB,2})$	1 fixed	$\begin{array}{c} 0.5 \\ (Y_{W\!/\!H\!B,2}) \end{array}$	$0$ $(Y_{H/HB,2})$
3) PPB growth on PHAs PHA → Biomass	0	-0.18 $(-Y_{NX,3})$	$-0.014$ ( $-Y_{PX,3}$ )	1 fixed	$0.0157$ $(Y_{CX,3})$	-0.254 (- $Y_{HB/X,3}$ )	$\begin{array}{c} 0.406 \\ (Y_{WX,3}) \end{array}$	$\begin{array}{c} 0.138 \\ (Y_{H(X,3)} \end{array}$

Table 1. Stoichiometric matrix of the mathematical model developed to characterize photoorganoheterotrophic growth and PHA storage by Rhodopseudomonas palustris under light/dark diel cycles. According to experimental data, growth was assumed to only occur under periods of illumination with infrared light. Stoichiometric coefficients ( $v_{ii}$ ) are expressed as yields normalized per unit of biomass for growth processes and per unit of PHAs (expressed as equivalents of hydroxybutyrate monomers, HB) for the intracellular storage processes.

Table 2. Mole balance equations comprising the transport and biokinetic conversion terms of the mathematical model for the photoorganoheterotrophic growth of *Rhodopseudomonas palustris* and intracellular storage.

Compounds	Mole balance equations	
Biomass	$\frac{dC_X}{dt} = -D \cdot C_X + (\mu_{ace} + \mu_{PHA}) \cdot C_X$	(1)
Acetate	$\frac{dC_S}{dt} = D \cdot (C_{S,in} - C_S) - (q_{S,X} + q_{S,PHA}) \cdot C_X$	(2)
Ammonia	$\frac{dC_N}{dt} = D \cdot (C_{N,in} - C_N) - (q_{N,S} + q_{N,PHA}) \cdot C_X$	(3)
Phosphate	$\frac{dC_P}{dt} = D \cdot (C_{P,in} - C_P) - (q_{P,} + q_{P,PHA}) \cdot C_X$	(4)
CO <sub>2</sub> (aq.)	$\frac{dC_{CO_2}}{dt} = -D \cdot C_{C,L} + q_C \cdot C_X + k_L a_C \cdot (C^*_{C,L} - C_{C,L})$	(5)
CO <sub>2</sub> (gas)	$\frac{dC_{CO_2,gas}}{dt} = -\frac{F_g}{V_g} \cdot C_{C,g} - k_L a \cdot (C^*_{C,L} - C_{C,L}) \cdot \frac{V_L}{V_g}$	(6)
PHAs (HB)	$\frac{dC_{PHA}}{dt} = -D \cdot C_{PHA} + (q_{PHA,S} - q_{PHA,X}) \cdot C_X$	(7)

With  $C_X$  the state-variable concentration of biomass in the mixed liquor (C-mmol L<sup>-1</sup>); *t* the process time (h); *D* the dilution rate (h<sup>-1</sup>);  $C_{S,in}$ ,  $C_{P,in}$  the constant concentrations of dissolved acetate, ammonium and phosphate in the influent ;  $C_S$ ,  $C_N$ ,  $C_P$  the state-variable concentrations of dissolved acetate, ammonium and phosphate in the bulk liquid;  $C_{CL}$  the state-variable concentration of dissolved CO<sub>2</sub> in the bulk liquid;  $C_{PHA}$  the state-variable concentration of PHAs (as hydroxybutyrate monomer equivalents);  $\mu$  the total growth rate split as growth rates on acetate  $\mu_{ace}$ (process 1 in Table 1) and on PHAs  $\mu_{PHA}$  (process 3 in Table 1);  $q_S$  the biomass-specific rate of acetate uptake split as acetate uptake for growth ( $q_{SX}$ ) and for PHA storage ( $q_{SPHA}$ ).

A mole balance on the gas phase was integrated for CO<sub>2</sub> (Eq. 6): with  $C_{CL}^*$  (mmol CO<sub>2</sub> L<sup>-1</sup> liquid) the maximum saturation concentration of CO<sub>2</sub> dissolved in the liquid phase, and  $C_C$  (mmol CO<sub>2</sub> L<sup>-1</sup> liquid) the actual concentration of CO<sub>2</sub> dissolved in the liquid phase. The volumetric mass transfer coefficient of CO<sub>2</sub> between the gas and liquid phases ( $k_La$ , h<sup>-1</sup>) was calculated in function of the power input, liquid volume, and gas velocity (see SI-2.1 in Supplementary information). The maximum dissolved CO<sub>2</sub> concentration ( $C_{CL}^*$ ) was calculated in function of the measured CO<sub>2</sub> concentration in the gas phase ( $C_{Cg}$  mmol CO<sub>2</sub> L<sup>-1</sup> gas), the Henry's coefficient ( $H_C$ , mol L<sup>-1</sup>atm<sup>-1</sup>), the respective temperatures (T) of 20°C in dark and 30°C under light, and pressure (1 atm) (SI-2.2).

Parameters and units are described in Table 3.

The liquid flow rate F (L h<sup>-1</sup>), liquid volume V (L), gas flow rate  $F_g$  (L h<sup>-1</sup>), gas volume  $V_g$  (L), and the concentrations of dissolved chemical compounds in the influent ( $C_{i,in}$ , mmol L<sup>-1</sup>) were all fixed in the experiments.

Parameter name	Symbol	Value Units		Source	
Stoichiometric parameters					
Biomass yield on acetate	Y <sub>X/S</sub>	1.75	C-mol X · C-mol <sup>-1</sup>	Fitted	
Biomass yield on PHA (HB)	Y <sub>X/HB</sub>	3.94	C-mol X $\cdot$ C-mol <sup>-1</sup> HB	Stoichiometry	
PHA (HB) yield on acetate	$Y_{HB/S}$	0.44	C-mol HB · C-mol <sup>-1</sup>	Stoichiometry	
Ammonium yield over biomass	$Y_{N/X}$	0.07	s mol N · C-mol⁻¹ X	Fitted	
Phosphate yield over biomass	$Y_{P/X}$	0.007	$mol \ P \cdot C\text{-}mol^{-1} \ X$	Fitted	
CO <sub>2</sub> yield on acetate	$Y_{C/S}$	0.24	C-mol $CO_2 \cdot C$ -mol	Fitted	
CO <sub>2</sub> yield on HB	$Y_{C/HB}$	0.06	$C-mol CO_2 \cdot C-mol^{-1} HB$	Stoichiometry	
Biokinetic parameters					
Maximum biomass-specific	$\mu_{max}$	0.15	C-mol X $\cdot$ h <sup>-1</sup> $\cdot$ C-	Fitted	
Half-saturation coefficient for acetate	$K_S$	0.1	C-mmol S $\cdot$ L <sup>-1</sup>	Lower than in Kaewsuk et al.	
Half-saturation coefficient for acetate (growth on HB)	$K_{S2}$	0.003	C-mmol S $\cdot$ L <sup>-1</sup>	(2010) (0.3 mM) Arbitrary	
Half-saturation coefficient for PHAs (HB)	$K_{HB}$	0.001	C-mmol HB $\cdot$ L <sup>-1</sup>	(De Kreuk et al., 2007)	
Half-saturation coefficient	$K_N$	0.001	mmol N $\cdot$ L <sup>-1</sup>	(Wolf et al., 2007)	
Half-saturation coefficient	$K_P$	0.003	mmol P $\cdot$ L <sup>-1</sup>	(Puyol et al.,	
for phosphate Half-saturation coefficient for light	$K_I$	10	$W \cdot m^{-2}$	2017) (Prachanurak et al., 2019)	
Reactor, transport and light					
Reactor radius	R	0.055	М	Measured	
Liquid volume	V	1.2	L	Measured	
Liquid flow rate	F	0.048	$L \cdot h^{-1}$	Measured	
Dilution rate	D	0.04	h-1	= F/V	
Gas volume	$V_g$	0.3	L	Measured	
Gas flow rate	$F_g$	7.14	$L \cdot h^{-1}$	Measured	
Inflow acetate concentration	$C_{S,in}$	7	C-mmol S $\cdot$ L <sup>-1</sup>	Measured	
Inflow ammonium concentration	$C_{N,in}$	4.28	mmol N $\cdot$ L <sup>-1</sup>	Measured	
Inflow phosphate concentration	$C_{P,in}$	0.22	mmol P $\cdot$ L <sup>-1</sup>	Measured	
Volumetric mass transfer coefficient	<i>k</i> <sub>L</sub> a	32	$h^{-1}$	Calculated (SI-1)	
of CO <sub>2</sub> Henry coefficient for CO <sub>2</sub>	Hc	0.03 (at 30°C) 0.04	$C$ -mol $\cdot L^{-1} \cdot atm^{-1}$	(Sander, 2015)	
T :- 1 4 :- 4 : 4 4 11-	T	(at 20°C)	<b>W</b> 2	Maaaaad	
Light intensity at reactor walls	10	300	w · m <sup>-2</sup>	Measured	
Lignt extinction coefficient per biomass concentration	ε	0.1	m <sup>2</sup> · g <sup>-1</sup>	(Kim et al., 1980)	

Table 3. Stoichiometric and kinetic constants used in the mathematical model describing the photoorganoheterotrophic growth of *Rhodopseudomonas palustris* and intracellular storage. The yields and maximum biomass-specific growth rate were fitted to experimental data. The other parameters were measured during the experiments, calculated or obtained from literature.

The biomass specific growth rate ( $\mu$ , h<sup>-1</sup>) was calculated both in function of acetate ( $\mu_{ace}$ ) and PHAs ( $\mu_{PHA}$ ) (Eq 8-9). They were initially assumed to be limited by the concentrations of multiple chemical compounds and by incident light intensity (I, W m<sup>-2</sup>). According to liquid phase measurements, acetate and light were the main limiting state variables for biomass growth. Ammonium and phosphate were supplied in excess.

$$\mu_{ace} = \mu_{max} \cdot \frac{c_S}{c_S + K_S} \cdot \frac{c_N}{c_N + K_N} \cdot \frac{c_P}{c_P + K_P} \cdot \frac{I}{I + K_I}$$
(8)

$$\mu_{PHA} = \mu_{max} \cdot \frac{C_{HB}}{C_{HB} + K_{HB}} \cdot \frac{C_N}{C_N + K_N} \cdot \frac{C_P}{C_P + K_P} \cdot \frac{I}{I + K_I} \cdot \frac{K_{S,2}}{C_S + K_{S,2}}$$
(9)

With  $\mu_{max}$  defined as the biomass-specific maximum growth rate for *Rhodop-seudomonas palustris*.

The volumetric rates of acetate  $(r_s)$ , ammonium  $(r_N)$  and phosphate  $(r_P)$  uptake and of CO<sub>2</sub> production (in the gas phase)  $(r_C)$  (all in mmol L<sup>-1</sup> h<sup>-1</sup>) follow from the overall growth stoichiometric coefficients (*i.e.*, yields) and the biomass growth rate ( $\mu$ ):

$$r_S = -Y_{S/X} \cdot r_X = -\frac{1}{Y_{X/S}} \cdot r_X \tag{10}$$

$$r_N = -Y_{N/X} \cdot r_X \tag{11}$$

$$r_P = -Y_{P/X} \cdot r_X \tag{12}$$

$$r_C = Y_{C/X} \cdot r_X \tag{13}$$

The biomass-specific rates were calculated as:

$$q_{S,X} = \mu_{ace} \cdot \frac{1}{Y_{X/S}} \tag{14}$$

$$q_{S,PHA} = q_{S,max} \cdot \frac{c_S}{c_S + K_{S,2}} \cdot \frac{l}{l + K_I}$$
(15)

$$q_{N,S} = \mu_{ace} \cdot Y_{N,S/x} \tag{16}$$

$$q_{N,PHA} = \mu_{PHA} \cdot Y_{N,HB/X} \tag{17}$$

$$q_{P,S} = \mu_S \cdot Y_{P,S/X} \tag{18}$$

$$q_{P,PHA} = \mu_{PHA} \cdot Y_{P,HB/X} \tag{19}$$

$$q_{PHA,X} = \mu_{PHA} \cdot \frac{1}{Y_{X/HB}} \tag{20}$$

$$q_{PHA,S} = Y_{HB/S} \cdot q_{S,HB} \tag{21}$$

$$q_{CO_2} = Y_{1,CO_2/S} \cdot q_{S,HB} + Y_{2,CO_2/S} \cdot q_{S,X} + q_{PHA,S} \cdot Y_{CO_2/HB}$$
(22)

The yields of biomass growth on acetate  $(Y_{X/S})$  and  $CO_2$  production over biomass production  $(Y_{C/X})$  and the biomass-specific maximum growth rate  $(\mu_{max})$ were computed by fitting the model outputs to the measurements from the experiment performed with 16 h light / 8 h dark. Additional estimations of  $\mu_{max}$ were made based on data collected from preliminary batch cultures of the *Rhodopseudomonas* strain incubated at 30°C under IR light with the same cultivation medium as for the reactor, as well as from the CSTR experiments run under continuous illumination and 8 h light / 16 h dark. Overall, an average  $\mu_{max}$  value of 0.15 ± 0.05 h<sup>-1</sup> was obtained for this organism under photoorganoheterotrophy. The yields of ammonium consumption ( $Y_{N/X}$ ) and phosphate consumption ( $Y_{P/X}$ ) during biomass growth, and consequently the N and P compositions of the biomass ( $n_N$  and  $n_P$ , respectively) were determined from the 8 h light / 16 h dark experiment.

Two light sources with resulting IR irradiation intensities  $I_0$  measured at the reactor wall (300 W m<sup>-2</sup>) were placed at opposite sides next to the reactor. The light intensity decrease away from the reactor wall was assumed to follow the Beer-Lambert attenuation law. Therefore, the total light intensity  $I_t$  at a radial position x (m) across the reactor diameter was obtained by summing up the light intensities coming from both sides as following:

$$I_t = I_0 \cdot \left( e^{-\varepsilon \cdot c_X \cdot x} + e^{-\varepsilon \cdot c_X \cdot (2 \cdot R - x)} \right)$$
(23)

with *R* (m) the reactor radius,  $\varepsilon$  (m<sup>2</sup> g<sup>-1</sup>) the biomass-specific light attenuation coefficient, and  $C_X$  (g m<sup>-3</sup>) the biomass concentration. We considered that the liquid mixing enabled the cells to be exposed to an average light intensity *I* (W m<sup>-2</sup>) computed by integrating eq. 23 over the reactor diameter:

$$I = I_0 \cdot \frac{1 - e^{-2 \cdot \varepsilon \cdot c_X \cdot R}}{\varepsilon \cdot c_X \cdot R}$$
(24)

The model was solved in MATLAB (R2018b, Mathworks, Natick, MA, <u>www.mathworks.com</u>) using a stiff solver for the ordinary differential equations system. The initial concentrations were taken to coincide with the measurements. The parameter estimation was performed by a constrained optimization routine minimizing the sum of squares of relative errors between model and experimental data. The fixed model parameters and the fitted parameters are listed in Table 3.

### 3. Results

The tree IR irradiance regimes were tested in the acetate-fed, anaerobic CSTR targeted and enabled to elucidate the physiology of the PNSB *Rhodopseudomonas palustris*, namely: (i) continuous illumination, (ii) cyclic 16 h light / 8 h dark, and (iii) cyclic 8 h light / 16 h dark. A specific focus was given to carbon and electron allocation to biomass, H<sub>2</sub> and PHA productions. The mathematical model calibrated on the 16 h light / 8 h dark experiment enabled to extract key stoichiometric and kinetic parameters like the biomass production yield on acetate ( $Y_{XS}$ ), the CO<sub>2</sub> production yield during growth ( $Y_{CX}$ ), and the maximum biomass-specific growth rate ( $\mu_{max}$ ) (Table 3). The calibrated model was used to predict the behavior of the 8 h light / 16 h dark cycles.

### 3.1 Biomass production of Rhodopseudomonas palustris

Under continuous illumination, the biomass reached a constant concentration of  $12.10 \pm 0.44$  C-mmol L<sup>-1</sup>. The outflow of biomass from the reactor corresponded to 88% of the carbon and 99% of the reduction equivalents (COD) provided with the influent. Around 11% of the supplied carbon was recovered as CO<sub>2</sub> (0.112 C-mmol L<sup>-1</sup>) in the off-gas. HPLC results showed no residual acetate in the bulk liquid at steady state.

Under cyclic operations (Figure 2), the system reached a pseudo-steady-state conditions with similar dynamics in concentrations per light/dark cycle.

In the dark, the biomass did not grow and was washed out of the reactor with the effluent. The biomass wash-out rate  $(F / V \cdot C_X)$  was very well represented by the numerical model (Figure 2A,B). The corresponding increases in acetate, ammonium and phosphate concentrations during the dark phases matched with the expected change based on the feeding rate. No conversion occurred in the dark.

Under 16-h light periods, the biomass accumulation exceeded the dilution rate (D = F / V). An initial growth rate  $(\mu)$  of 0.15 h<sup>-1</sup> (assumed to be equivalent to  $\mu_{max}$  since the reactor experiences a high residual concentration of acetate right after the dark phase) was observed. It reached a steady value  $(\mu = D)$  of 0.04 h<sup>-1</sup> after 2 h of exposure to IR light (Figure 2A). The residual acetate concentration

reached a minimal steady value  $(0.7 \pm 0.6 \text{ C-mmol } \text{L}^{-1})$  when the biomass growth rate achieved the steady state.

Under shorter 8-h irradiance periods, the biomass growth rate did not reach a stationary state in a cycle, being around 0.1 h<sup>-1</sup>. Acetate was not depleted at the end of the light phase (Figure 2B). The increase in biomass concentration from 11 C-mmol L<sup>-1</sup> on the first cycle to 5 C-mmol L<sup>-1</sup> on the fourth cycle. The growth rate  $\mu$  during the 8-h light period was approximately 0.12 h<sup>-1</sup> thus lower than the aforementioned  $\mu_{max}$ .

The estimated biomass yield on acetate  $(Y_{X/S})$  during the photoorganoheterotrophic growth of *Rhodopseudomonas palustris* was 0.88 C-mol X C-mol<sup>-1</sup> acetate under both irradiance periods of 16 h and 8 h, and matched with the yield measured under 24-h continuous light regime.



Figure 2. Dynamics of biomass, acetate, ammonium and phosphate concentrations during light-dark diel cycles in continuous-flow cultures of Rhodopseudomonas palustris: (A and C) 16-h light / 8-h dark regime; (B and D) 8-h light / 16-h dark regime. Gray areas represent the dark periods, white areas the light periods. The measured concentrations of biomass (black circles) and acetate (red open squares) are displayed in panels A and B. The lines display the model simulations. The model was calibrated with the dataset of the 16-h light / 8-h dark and used to predict evolutions under the 8-h light / 16-h dark regime. In the first case, the biomass concentration achieved a stationary state, while acetate reached limiting residual concentrations 2 h after the start of the 16-h light periods. In the second case, the biomass increased without reaching a steady state during the shorter 8-h light periods, while acetate was not fully consumed. The measured concentrations of ammonium (blue diamonds) and phosphate (green triangles) are given in panels C and D, together with the model simulations. Concentrations of N and P were not limiting in any illumination phase.

### 3.2 Carbon, nitrogen, and phosphorus assimilation

Under continuous illumination, no residual acetate was detected in the bulk liquid. Under 16 h of light, acetate was almost fully depleted, down to a concentration around 1 C-mmol L<sup>-1</sup>. This was very well represented by the numerical model. Under 8 h of light, acetate was still present in the bulk liquid in concentrations ranging from 4 to 1.5 C-mmol L<sup>-1</sup>. In this case, the model overestimated the rate of acetate consumption, *i.e.* predicting acetate depletion at the end of the light phase.

Nitrogen (N) and phosphorus (P) sources were provided in excess, and never became limiting. Under light, ammonium and phosphate were consumed, resulting in a decrease in their concentrations (Figure 2C,D). Under dark, the fed N and P accumulated in the liquid phase since not assimilated by the nongrowing biomass. At the end of the dark phase,  $3.6 \pm 0.6 \text{ mmol N L}^{-1}$  and  $0.14 \pm 0.01 \text{ mmol P L}^{-1}$  remained the bulk liquid. N and P dynamics showed irregular behavior under 16 h light (Figure 2C). Therefore, the yields of ammonium ( $Y_{N/X}$ ) and phosphate ( $Y_{P/X}$ ) consumptions during growth were fitted on the 8-h light experiment (Figure 2D). The computed yields ( $Y_{N/X} = 0.18 \text{ mol N C-mol}^{-1} X$ ,  $Y_{P/X} = 0.009 \text{ mol P C-mol}^{-1} X$ ) were very close to the theoretical yields resulting from the elemental composition of *Rhodopseudomonas* biomass ( $Y_{N/X} = 0.18 \text{ mol N C-mol}^{-1} X$ ,  $Y_{P/X} = 0.014 \text{ mol P C-mol}^{-1} X$ ) that has been measured by McKinlay & Harwood (2010). With these yields, the N and P dynamics during the 8-h light cycles were well represented by the model: accumulation under dark and consumption by assimilation in the biomass growing under light.

### *3.3 CO*<sub>2</sub> production follows the biomass growth patterns

The CO<sub>2</sub> production rate was constant at  $0.075 \pm 0.001$  mmol h<sup>-1</sup> during continuous illumination (Figure 3A): the chemostat achieved a stationary operation.

Under 16 h of light, CO<sub>2</sub> emission increased rapidly within the first hour, reaching 0.25 mmol  $h^{-1}$ , following the biomass growth rate with acetate uptake (Figure 3B). Once the acetate reached the minimal level, the CO<sub>2</sub> production also decreased to a stable level of 0.07 mmol  $h^{-1}$  after the second hour of illumination, reflecting the steady-state operation. The numerical model reproduced the observed trends during light phase, but the residual CO<sub>2</sub> production in the dark phase could not be explained numerically.

Under the 8-h light phases the CO<sub>2</sub> production did not achieve a steady state. In the light phase, the CO<sub>2</sub> production increased with 0.05 to 0.08 mmol h<sup>-1</sup> during the cycles, reaching a peak after 4 h (0.88  $\pm$  0.013 mmol h<sup>-1</sup>), which was described qualitatively by the model (Figure 3C). The reason for the discrepancy between model and experimental data is the

larger computed rate of acetate consumption, which leads to a higher  $\rm CO_2$  formation rate in the model results.

In both illumination regimes, the  $CO_2$  production dropped to the same stable value of 0.033 mmol h<sup>-1</sup> during the dark phase. This residual baseline value of  $CO_2$  in the off-gas under dark periods may relate to the detection limit of the online mass spectrometer (MS) instrument.

### $3.4 H_2$ is produced only during the short light periods

A very low  $H_2$  production was recorded during the continuous illumination with rates of around 0.004 mmol h<sup>-1</sup> (Figure 3D). Occasionally, unexplained  $H_2$  spikes were recorded, possibly relating to the background noise of the online MS.

During the 16-h light/8-h dark experiment, the emission of hydrogen in the dark was negligible and no specific production pattern was measured in the light (Figure 3E). The very low residual  $H_2$  concentration may relate to an instrumental offset as well.

In contrast, H<sub>2</sub> was produced constantly during the 8-h light periods (Figure 3F), with an increase of  $0.02 \pm 0.01 \text{ mmol h}^{-1}$  per cycle and reaching a maximum of 0.05 mmol h<sup>-1</sup>. The biomass-specific rate of H<sub>2</sub> production under 8 h of light was 10 times higher (0.156 mmol H<sub>2</sub> h<sup>-1</sup> g<sup>-1</sup> DW) than under all other conditions (0.014 mmol H<sub>2</sub> h<sup>-1</sup> g<sup>-1</sup> DW). Under continuous illumination, H<sub>2</sub> production corresponded to about 0.008 mmol electrons L<sup>-1</sup>, *i.e.*, 0.3% of the electrons supplied with the influent.

Under the 16-h light / 8-h dark periods, H<sub>2</sub> production accounted for  $0.43 \pm 0.28$  % (in the light) and  $0.33 \pm 0.13$  % (in the dark) of the electron balances. Under the 16-h dark periods, H<sub>2</sub> contribution made  $0.70 \pm 0.45$  % of the electron balance. Under the 8-h light phases, H<sub>2</sub> made a more important contribution to the electron balance, reaching 2%, *i.e.*, higher than in all other phases (Table 4).

	balance balance	ol h <sup>-1</sup> ) (% C) (% e)		0.004 98% 99%	.32 115±2% 118±16%	.54 103±9% 120±10%	.21 158±8%* 154±17%*	.29 130±25%* 158±23%*	
out		C-mmol h <sup>-1</sup> ) (e <sup>-</sup> -mm		65 2.662±	75±0.01 3.06±0	68±0.06 2.79±0	97±0.05 4.03±1	79±0.15 4.12±1	
п2 Σ	Off-gas	) (e-mmol h <sup>-1</sup> ) (C		0.008±0.004 0.	0.008±0.006 0.	0.007±0.002 0.	0.04±0.02 0.	0.02±0.02 0.	
(as HB)	Effluent	(e <sup>-</sup> -mmol h <sup>-1</sup>		<lod<sup>1</lod<sup>	0.22±0.06	$0.16 \pm 0.01$	$0.28 \pm 0.09$	$0.31 {\pm} 0.11$	
	Off-gas	(C-mmol h <sup>-1</sup> )		$0.070 \pm 0.001$	0.08±0.01	$0.07 \pm 0.02$	0.07±0.01	$0.03 \pm 0.002$	
		(e-mmol h <sup>-1</sup> )		2.654	2.45±0.29	2.42±0.21	2.44±0.60	2.84±0.72	
Biomass	Effluent	(C-mmol h <sup>-1</sup> )		0.58	$0.58 \pm 0.03$	0.50±0.06	$0.57 \pm 0.14$	$0.57 \pm 0.16$	
		(e <sup>-</sup> -mmol h <sup>-1</sup> )		<lod<sup>1</lod<sup>	$0.52 \pm 0.50$	$0.64 \pm 0.41$	$1.26 \pm 0.60$	$0.96 \pm 0.46$	
Acetate	Effluent	(C-mmol h <sup>-1</sup> )		<lod<sup>1</lod<sup>	$0.02 \pm 0.01$	$0.04 \pm 0.02$	$0.28 \pm 0.03$	$0.19 \pm 0.07$	
	.9	(e-mmol h <sup>-1</sup> )		2.69±0.40	2.69±0.40		2.44±0.05		
Acetate	Influent = $\Sigma$	(C-mmol h <sup>-1</sup> )		$0.66 \pm 0.05$	0 00 00	CO.0±00.0	0.61+0.006	000.0-10.0	
Component	Phase	Units	Illumination regime	) 24-h continuous light	) 16-h light period	8-h dark period	) 8-h light period	16-h dark period	
		I		I ⊖	ାଳି		6		

# Table 4. Carbon (C-mmol h<sup>-1</sup>; grey columns) and electron (e-mmol h<sup>-1</sup>; white columns) balances under the different illumination regimes.

# <sup>1</sup> Limit of detection (LOD)

biomass concentration. The absorbance measurements of biomass might have been influenced by the presence of PHAs inside the cells. \* The surpassing carbon and electron balances for the third illumination regime can be due to an overestimation of the

# 3.5 *PHA production follows CO*<sub>2</sub> *formation under 16 h of light but not under 8 h of light*

PHAs were not detectable under continuous illumination.

A PHB peak was detectable in the 16-h light phases, 2 h after the light switch  $(0.114 \pm 0.013 \text{ C-mmol HB C-mmol}^{-1} \text{ X})$  (Figure 3G), concomitantly to the peak of CO<sub>2</sub> production. The PHAs decreased constantly after the peak, reaching a baseline at  $0.063 \pm 0.001 \text{ C-mmol HB C-mmol}^{-1} \text{ X}$ . Under the 16-h light / 8-h dark cycles, PHA corresponded on average to 7% of the electron balance, regardless the illumination conditions. The numerical model caught well the PHAs accumulation during the first period of the 16-h light phases, followed by a decrease both under light and under dark conditions.

Under 8-h light / 16-h dark cycles (Figure 3H), PHA production did not present peaks, with an average production of  $0.119 \pm 0.015$  C-mmol HB C-mmol<sup>-1</sup> X, corresponding to 12% of the electron balance. About 1.7 times more PHAs were detectable in the dark phase of the 8-h light / 16-h dark regime than of the 16-h light / 8-h dark one. The numerical model indicated an oscillation in the PHA production that was not confirmed with experimental data (Figure 3H).

The diel light/dark regimes induced PHA formation. The longer light periods (16 h) generated a typical feast-famine behavior (Montiel Corona et al., 2017). The shorter light periods (8 h) resulted in a higher and more stable PHA content in the cells (Figure 3H).



Figure 3. Dynamics of CO<sub>2</sub>, H<sub>2</sub> and PHA during light-dark cycles: (A and D) continuous illumination; (B, E and G) 16-h light / 8-h dark regime; (C, F and H) 8-h light / 16-h dark regime. Gray areas represent the dark periods, white areas the light periods. The measured (red circles) CO<sub>2</sub> production rates are displayed in panels A-C, together with the model simulations for the light/dark cycles (red lines). CO2 production was constant during continuous illumination, but peaks appeared during light-dark cycles, correlating with the acetate uptake. The measured H<sub>2</sub> production rates (blue circles) are given in panels D-F, but were not modelled since H<sub>2</sub> contributed only to minor fractions of the electron balances. H<sub>2</sub> was produced at a constant low level during continuous illumination and 16-h light cycles. H<sub>2</sub> production significantly increased out of the instrumental noise of the online mass spectrometer during each light period of the 8-h light cycles. PHAs (green diamonds) were indirectly quantified in the light/dark experiments through sensitive fluorimetry measurements calibrated to PHA mol fractions (expressed as hydroxybutyrate, HB) in biomass, and modelled (green lines) (G-H). A PHA peak was measured at the start of each 16-h light phase. Under 8 h of light, PHA contents remained constantly at a higher level during light periods.

### 4. Discussion

### *4.1 Acetate rather than light drives the metabolic responses of Rhodopseudomonas palustris*

Light energy source is central for the catabolic processes in PNSB. Organic substrates are used as C-source for biomass synthesis and as electron donors. When no external electron acceptor is present, the difference in degree of reduction between substrate and biomass has to be balanced with internal processes of electron reallocation. The dynamics of the change in N and P concentrations were congruent with assimilation fore biomass formation and their measured yields were close to the elemental composition of *Rhodopseudomonas palustris* biomass ( $C_1H_{1.8}O_{0.38}N_{0.18}P_{0.014}$ ) reported by McKinlay & Harwood (2010). Carbon balances closed for the continuous illumination experiment and for the 16-h light / 8-h dark experiment. This was not the case for the 8-h light / 16-h dark cycles where the over-assessed C-balances seemed to indicate a net production (Table 4). PHA granules that were always present in the cells under 8 h light / 16 h dark may have led to an absorbance overestimation of the biomass present in the reactor. The overestimation of the biomass consequently affected the electron balances as well.

Light/dark cycles occur daily in natural environments. Phototrophic organisms have adapted to cope with imbalances in energy supply. PNSB have higher biomass production under light/dark cycles (Koku et al., 2003; Zhi et al., 2019). H<sub>2</sub> production is increased under discontinuous illumination, following the biomass trend (Li et al., 2011; Wakayama et al., 2000).

Here, the light cycles are important not only for the energy that light provides, but most importantly for the illumination time available for the cultures to metabolize the nutrients. The light intensity at the surface of the reactor was 300 W m<sup>-2</sup>. Considering the attenuation due to the biomass concentration and the reactor depth, the minimum calculated light available in the reactor was 180 W m<sup>-2</sup> (Figure SI-3.1 in Supplementary information). Under any given condition, the available light intensity was not limiting the microbial metabolism since always exceeding the half-saturation coefficient for light ( $K_I$ ) arbitrarily set at 10 W m<sup>-2</sup> (Prachanurak et al., 2019). PNSB adapt the number and type of photosynthetic unit components based on the different light intensities (Brotosudarmo et al., 2011). However, Imam et al. (2015) have reported that light may be saturating for PNSB growth already from 100 W m<sup>-2</sup> onward. We can therefore assume that the irradiance intensity used in our study during the light periods did not deeply affect the metabolic state of the cells.

The continuous cultivation system was initially set-up based on the stoichiometric and kinetic parameters derived from anaerobic,

photoorganoheterotrophic batches of the *Rhodopseudomonas palustris* isolate cultivated under the same medium, infrared irradiance and temperature conditions as the CSTR.

In the continuously illuminated chemostat, the biomass reached a steady-state, acetate was limiting (*i.e.*, low residual concentration), and the biomass growth rate equaled to the dilution rate (0.04  $h^{-1}$ ). Once the light/dark cycles were applied, a disturbance of the steady state was immediately observed.

Under long (16 h) illumination, the biomass consumed the acetate for growth and after an initial increase of the growth rate the steady-state conditions were restored for the remaining two thirds of the light cycles (with a specific growth rate again equal to the dilution rate).

Longer (16 h) dark periods led to an increased accumulation of acetate in the bulk liquid (twice as high as in the 8 h dark), since acetate was fed twice longer and there was no consumption in the dark. Due to the shorter light (8 h light) periods, acetate was not fully consumed at the end of the light phases. Therefore, acetate was no more growth-rate-limiting and the light period was too short to reach a steady state. The growth rate in the light phase exceeded the value of the imposed dilution rate, leading to a transient accumulation of biomass (Figure 2B).

The modelled biomass dynamics agreed with the measurements, fitting quantitatively the values for the 16-h light experiment, where the exact steadystate values were obtained. When using the same model parameters for the short (8 h) illumination experiment, the calculated amount of biomass formed during the light periods was underestimated. The disagreement may be related to the carbon imbalance reported in Table 4. A longer illumination phase would allow the biomass to reach exactly the same steady state as in the longer light period experiment (Figure SI-3 in Supplementary information) after about 10 h of light and acetate attained the minimum.

The average volumetric rate of acetate consumption was 0.08 C-mol h<sup>-1</sup> L<sup>-1</sup> under 16 h of light and 0.15 C-mol h<sup>-1</sup> L<sup>-1</sup> under 8 h of light (Figures SI-3.2 in Supplementary information). Initially, the biomass grew at the maximum substrate uptake rate. If the light phase was too short (as in the 8-h light periods), acetate was not fully consumed (residual of  $2.5 \pm 1.1$  C-mmol L<sup>-1</sup>) and the biomass grew at its maximum growth rate (0.15 h<sup>-1</sup>).

# 4.2 The PHA synthesis patterns reflect the metabolic state of the Rhodopseudomonas cells

PHAs can be formed under dynamic conditions or because of imbalance in the degree of reduction between the biomass and carbon source. PHAs constitute carbon, electron and energy stocks for microorganisms surviving dynamic environments (van Aalst-van Leeuwen et al., 1997; Van Loosdrecht et al., 1997). In 24-h continuously irradiated chemostat conditions, carbon was

continuously fed to the cultures and cells did not store PHAs. PHAs synthesis in PNSB has primarily been reported under nitrogen limitation, as an intrinsic mechanism for the redistribution of carbon excess. PHAs accumulation is conventionally reported for PNSB under growth-limiting conditions (Braunegg et al., 1998; Narancic et al., 2016) when the carbon/electron sources are available but the nutrients (such as N, P or S) needed to produce cellular components are limited, resulting in the accumulation of acetyl coenzyme A (acetyl-CoA) in the cells.

During the anabolic reactions, *Rhodopseudomonas* incorporates acetate in the form of acetyl-CoA and releases at the same time the coenzyme A (CoA). In highly active cells, the levels of acetyl-CoA are low, but the levels of CoA are high. In contrast, in non-growing cells, acetyl-CoA is not utilized in anabolic processes and accumulates in the cells (McKinlay et al., 2014). A high acetyl-CoA to CoA ratio and the presence of NADPH are required for the initiation of the PHA pathway (Satoh et al., 2003).

The ratio between carbon sources and other nutrients is important to maintain the balance between substrate uptake rate and growth rate. The composition of the inflow of the CSTR was defined based on the biomass composition of PNSB reported by Puyol et al. (2017): a C:N ratio of 5.68:1 mol C-acetate mol<sup>-1</sup> N-NH<sub>4</sub><sup>+</sup> was used in the medium. Nitrogen was provided in excess preventing N-limitation imbalance. Consequently, under continuous illumination, no production of PHAs was detected.

After a period of darkness, the biomass was transiently subjected to an excess of light and nutrients (acetate, ammonium, phosphate) resulting in an increased substrate uptake rate. For the first 2 h after switching on the 16-h light phase, the growth rate was close to the previously measured  $\mu_{max}$  of 0.15 h<sup>-1</sup>. After 3 h, when acetate became limiting, the growth rate stabilized again at the dilution rate value of 0.04 h<sup>-1</sup>. When shorter (8 h) light periods were applied, the initial growth rate was around 0.1 h<sup>-1</sup>. Under both conditions, cells exhibited growth rates close to  $\mu_{max}$ . This resulted in the production of reducing power, which became in excess, and had to be reallocated. As reported by Kanno et al. (2018), the photosynthetic units are not disassembled in the dark, even under starvation. These are readily available, once the light conditions are restored, to produce ATP and NADH for the biosynthetic processes. This, linked to the prompt availability of the enzymes for PHA formation that are constitutively expressed (Kranz et al., 1997), leads to the immediate production of PHAs. Under 16 h light, a peak of PHB was observed 2 h after the light switch, but it decreased to baseline steady-state levels once the growth rate stabilized again at the dilution rate of 0.04 h<sup>-1</sup>. The PHB decreased and no H<sub>2</sub> production was observed. Cells reached a maximal capacity of substrate uptake, though not coupled to a maximal growth rate, similarly to what is described in Krishna & Van Loosdrecht (1999).

Under 8-h light / 16-h dark cycles, acetate was not completely consumed during illumination and increasingly accumulated in the bulk liquid during the dark phases. This resulted in a further imbalance in the redox state of the cells. The redox imbalance generated by the continuous presence of acetate, rather than the illumination conditions alone, led to a more constant production of PHAs. The cells were highly active, with a growth rate close to the maximal growth rate, leading to a low availability of CoA that resulted in PHA formation.

### 4.3 $H_2$ production is a secondary pathway of electron dissipation

The H<sub>2</sub> production rates here reported were low compared to other studies that have exposed PNSB to similar light intensities (Imam et al., 2015). However, our experimental conditions were not designed to stimulate  $H_2$  production.  $H_2$ can be produced either via the nitrogenase system, either via the ferredoxinhydrogenase system. Ammonium is known to inhibit N<sub>2</sub> fixation in photosynthetic bacteria. It also effectively prevents photoproduction of H<sub>2</sub>, due to inhibition and inactivation of nitrogenase (Basak et al., 2014). The presence of ammonium in non-limiting amounts in the medium indicated that potentially  $H_2$  production was driven by the hydrogenase rather than by the nitrogenase. The molar C:N ratio (5.7) was 7 times lower than in most other studies on H<sub>2</sub> production (average C:N ratio 40) (Koku et al., 2002). Nonetheless, the H<sub>2</sub> production rate per gram biomass during the 8-h light phases was around 11 times higher than in all other conditions tested here (0.156 vs. 0.014 mmol  $h^{-1}$  g<sup>-1</sup> <sup>1</sup> DW). H<sub>2</sub> contribution to the electron balance was therefore 3 to 7 times higher in the 8-h light phases than in all other conditions, indicating that H<sub>2</sub> production acts as further electron dissipation pathway. A similar H<sub>2</sub> production pattern under alternating irradiation in PNSB cultures has been found (Adessi et al., 2012), with  $H_2$  production only during the light phases.  $H_2$  production was not integrated in the numerical model, since H<sub>2</sub> did not contribute significantly to the stoichiometric balance and kinetics  $(0.87 \pm 0.78 \% \text{ of the electron balance})$ .

The absence of relevant H<sub>2</sub> production during 16 h of light  $(0.0040 \pm 0.0004 \text{ mmol } \text{H}_2 \text{ h}^{-1} \text{ or } 0.75 \pm 0.01 \text{ mmol electrons } \text{h}^{-1})$  and the low but constant H<sub>2</sub> production during 8 h of light  $(0.025 \pm 0.009 \text{ mmol } \text{H}_2 \text{ h}^{-1} \text{ or } 0.97 \pm 0.05 \text{ mmol electrons } \text{h}^{-1})$  indicates that the H<sub>2</sub> sink did not play a major role in the electron redistribution patterns in these experiments.

### 5. Conclusions

Overall, PNSB are one of the most versatile guilds of microorganisms, whose metabolisms are impacted by dynamics in environmental conditions such as light/dark diel cycles. We elucidated the mechanisms of electron allocation using quantitative biotechnology by involving a continuous culture of the representative PNSB *Rhodopseudomonas palustris* isolated from an in-house enrichment and metabolic modelling.

It can be concluded from this work that:

- 1. Under light-saturating conditions and continuous-flow reactor regime, durations of light and dark phases in a 24-h diel cycle set the availability of achievement substrate and the of а steady state in the photoorganoheterotrophic metabolism of *Rhodopseudomonas*. Longer dark phases (16 h) result in an excess of substrate available at the start of the light phase for biomass growth. Longer light phases (16 h) lead to substrate limitation and steady conditions.
- 2. Even in actively growing *Rhodopseudomonas* cells, carbon allocation is in place, namely toward PHA production.
- 3. In growing cells,  $H_2$  production during illumination was a minor electron sink and secondary to PHA production, under the experimental conditions tested.
- 4. The numerical mathematical model caught the evolution trends in biomass, nutrient and PHAs dynamics measured during the light/dark cycles of different durations (16/8 vs. 8/16 h), as well as the CO<sub>2</sub> production and peaks emitted in the 16-h light periods. It allowed for determining stoichiometric and kinetic parameters of *Rhodopseudomonas palustris*.

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### Supplementary material

Appendix 1: Calibration line translating the PHA contents measured as fluorescent counts per C-mmol biomass into mol fractions of PHAs in the *Rhodopseudomonas* biomass (C-mmol<sub>HB</sub> C-mmol<sub>X</sub><sup>-1</sup>).



Figure SI-1: regression between fluorescent counts and C-mol<sub>HB</sub> C-mol<sub>X</sub><sup>-1</sup>.

To determine the HB content, *Rhodopseudomonas* was inoculated in the media and cultivated as in Materials and Methods. After one week of incubation, half of the biomass was centrifuged for 15 min at 10000 x g and separated from the supernatant. The biomass was re-incubated for one week in the same media but deprived of the nitrogen sources to promote HB accumulation. The other half of the biomass was centrifuged and stored at -20°C until further manipulation after addition of formaldehyde to stop all metabolic activity. To measure the PHA in the cells, the biomass grown in the complete medium was mixed with the one grown under N-limiting conditions in the following ratio:

1:0, 1:1, 1:4, 1:9, 1:18, 0:1 N-limited : non-limited.

For each point of the dilution series, 1 mL of sample was used for Nile-red staining as in the Material and Methods.

All the remaining biomass was freeze-dried and PHAs were quantified according to Vermeer et al., 2021.

The conversion factor between the fluorescent counts  $g^{-1}$  DW and the mass fraction mg HB  $g^{-1}$  DW was calculated from the regression slope here presented.
## Appendix 2: CO<sub>2</sub> solubility

SI-2.1 Estimation of the kLa of gas compounds in the process

The calculation of the volumetric mass transfer coefficient for  $CO_2$  was performed considering the equation for salt solutions (Platas Barradas et al., 2011):

$$k_L a (s^{-1}) = 0.002 \cdot (\frac{P}{V_{liquid}})^{0.7} \cdot u_{gas}^{0.4}$$

where *P* is the agitation power input (W),  $V_{liquid}$  is the bulk liquid volume (m<sup>3</sup>), and  $u_{gas}$  is the superficial gas velocity (m·s<sup>-1</sup>).

The agitation power input *P* was calculated through the following equation:  $P = \rho \cdot N_p \cdot N_i^3 \cdot D_i^5$ 

where  $\rho$  is the density (kg m<sup>-3</sup>) of the solution,  $N_p$  is the impeller power number,  $N_i$  is the impeller rotating speed (s<sup>-1</sup>), and  $D_i$  is the impeller diameter (m).

The power number of the impeller  $N_p$  is a function of the impeller type and the Reynolds number Re of the impeller. Re was calculated through the following equation:

 $Re = \frac{\rho N_i D_i^2}{\mu}$ where  $\mu$  is the viscosity of the solvent (Pa·s).

Considering the density  $\rho$  (997 kg m<sup>-3</sup>) and viscosity  $\mu$  of water at 25 °C (0.00089 Pa·s), the Reynolds number *Re* of the impeller was slightly higher than 10<sup>5</sup> (-). For that *Re* value and for a 6-blade Rushton turbine impeller, the power number *Ni* was around 5~6 (-). For the calculations, a power number of 6 was assumed. The diameter of the impeller was around 4 cm and the rotating speed 350 rpm (5.83 s<sup>-1</sup>). Using those values the power input of the impeller was found equal to 0.101 W.

The superficial gas velocity was calculated by dividing the gas flowrate F (7.14 L·h<sup>-1</sup>) with the surface area A (=  $\pi \cdot R^2$ ). The reactor radius R was 5.5 cm, *i.e.*, 0.055 m. The calculated superficial gas velocity  $V_g$  was 2.09 ·10<sup>-5</sup> m s<sup>-1</sup>.

Using the calculated values of the superficial gas velocity, power input and liquid volume  $(1.2 \cdot 10^{-3} \text{ m}^3)$  the k<sub>L</sub>a of carbon dioxide was calculated equal to 32 (h<sup>-1</sup>) through the following equation. The diffusion coefficient used are  $1.92 \cdot 10^{-5} \text{ cm}^2 \text{ s}^{-1}$ .

$$kLa_{x} = \frac{kLa_{CO_{2}}}{\sqrt{\frac{D_{CO_{2}}}{D_{x}}}}$$

### SI-2.2

The maximum dissolved CO<sub>2</sub> concentration (was calculated in function of the measured CO<sub>2</sub> concentration in the gas phase ( $C_{C,g}$ , mmol CO<sub>2</sub> L<sup>-1</sup> gas), the Henry's coefficient ( $H_c$ , mol L<sup>-1</sup>atm<sup>-1</sup>), the respective temperatures (T) of 20°C in dark and 30°C under light, and pressure (1 atm) (SI-2.2).

$$C_{C,L}^* = H_C \cdot 1000 \cdot C_{C,g} \cdot T \cdot R \cdot \frac{1}{101325} \pmod{\text{m}^{-3}}$$
  
With *T* the temperature (°K), *R* the gas constant (8.314 J mol<sup>-1</sup> K<sup>-1</sup>) and  $H_C$  the

Henry's constant at higher temperature, calculated as

$$H_C = H^{\circ} \cdot \exp\left[\frac{\Delta H}{R} \cdot \left(\frac{1}{T} - \frac{1}{T_0}\right)\right]$$

With *H* the Henry's constant for CO<sub>2</sub> at 298 K (0.0340 mol L<sup>-1</sup> atm<sup>-1</sup>),  $\Delta H$  the enthalpy of dissolution, and *T* and *T*<sub>0</sub> the temperatures.

## **Appendix 3: Mathematical model simulations**



Figure SI-3.1. Data and simulation with an additional extended light phase after the 8-h light / 16-h dark cycles. (A) **Biomass (black circles)** and acetate (red open squares) measured concentrations, with lines being the model results. (B) Measured (red circles) and model prediction (line) of the CO<sub>2</sub> production rate A steady state is reached once the concentration of acetate is at its minimum.



Figure SI-3.2. Model results for production and uptake rate in the 16 h light / 8 h dark cycles. (A) specific growth rate. (B specific acetate uptake rate. (C) specific ammonium uptake rate. (D) specific phosphate uptake rate. (E) specific  $CO_2$  production rate.



Figure SI-3.3. Model results for the light intensity inside the reactor. (A) 16 h light / 8 h dark cycles. (B) 8 h light / 16 h dark cycles. Under any given condition light became a limiting factor.



Figure SI-3.4. Model results for production and uptake rate in the 8 h light / 16 h dark cycles. (A) specific growth rate. (B specific acetate uptake rate. (C) specific ammonium uptake rate. (D) specific phosphate uptake rate. (E) specific  $CO_2$  production rate.

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Chapter 3

# Enrichment and aggregation

of purple non-sulfur bacteria in a mixed-culture sequencing-batch photobioreactor for biological nutrient removal from wastewater

This chapter is published in a modified version as original research article with contributions of: Marta Cerruti, Berber Stevens' Sirous Ebrahimi, Abbas Alloul, Siegfried E. Vlaeminck, David G. Weissbrodt (2020) Enriching and aggregating purple non-sulfur bacteria in an anaerobic sequencing-batch photobioreactor for nutrient capture from wastewater. *Frontiers in Bioengineering and Biotechnology*, 8, 557234. https://doi.org/10.3389/fbioe.2020.557234

## Abstract

Mixed-culture biotechnologies are widely used to capture nutrients from wastewater. Purple non-sulfur bacteria (PNSB), a guild of anoxygenic photomixotrophic organisms, raise interest for their ability to directly assimilate nutrients in the biomass. One challenge targets the aggregation and accumulation of PNSB biomass to separate it from the treated water. Our aim was to enrich and produce a concentrated, fast-settling PNSB biomass with high nutrient removal capacity in a 1.5-L, stirred-tank, acetate-fed, anaerobic sequencingbatch photobioreactor (SBR). PNSB were rapidly enriched after inoculation with activated sludge at 0.1 gVSS  $L^{-1}$  in a first batch of 24 h under continuous irradiance of infrared (IR) light (>700 nm) at 375 W m<sup>-2</sup>, with Rhodobacter reaching 54% of amplicon sequencing read counts. SBR operations with decreasing hydraulic retention times (48 to 16 h, *i.e.*, 1 to 3 cycles  $d^{-1}$ ) and increasing volumetric organic loading rates (0.2 to 1.3 kg COD  $d^{-1}$  m<sup>-3</sup>) stimulated biomass aggregation, settling, and accumulation in the system. reaching as high as 3.8 g VSS L<sup>-1</sup>. The sludge retention time (SRT) increased freely from 2.5 to 11 d. Acetate, ammonium, and orthophosphate were removed up to 96% at a rate of 1.1 kg COD  $d^{-1}$  m<sup>-3</sup>, 77% at 113 g N  $d^{-1}$  m<sup>-3</sup>, and 73% at 15 g P d<sup>-1</sup> m<sup>-3</sup>, respectively, with COD:N:P assimilation ratio of 100:6.7:0.9 m/m/m. SBR regime shifts sequentially selected for *Rhodobacter* (90%) under shorter SRT and non-limiting concentration of acetate during reaction phases, for *Rhodopseudomonas* (70%) under longer SRT and acetate limitation during reaction, and Blastochloris (10%) under higher biomass concentrations, underlying competition for substrate and photons in the PNSB guild. With SBR operations we produced a fast-settling PNSB biomass for nutrient capture from wastewater. This provided insights into the microbial ecology of PNSB-based processes for water resource recovery.



## 1. Introduction

Biological nutrient removal (BNR) is one of the main goals of wastewater treatment to safeguard aquatic ecosystems from anoxia and eutrophication. Water quality regulations become stricter on the limits of nutrient discharge and removal. European quality criteria target the following residual concentrations and removal of organic matter (125 mg  $COD_{Tot} L^{-1}$  and 75% removal; 25 mg  $BOD_5 L^{-1}$  and 70-90% removal), nitrogen (10-15 mg  $N_{Tot} L^{-1}$ , 70-80% removal), and phosphorus (1-2 mg  $P_{Tot} L^{-1}$ , 80% removal) (EUR-Lex, 1991; Guimarães et al., 2018). Besides conventional activated sludge systems, research and innovation target the use of novel microbial processes for water resource recovery (Alloul et al., 2018; Guest et al., 2009; Verstraete & Vlaeminck, 2011) on top of pollution control.

In the resource recovery context, purple non-sulfur bacteria (PNSB) can propel a sustainable treatment by capturing nutrient resources from used water (Daniel Puvol et al., 2017; Verstraete et al., 2016), valorizing waste into biomass, bioenergy, bulk chemicals, and biomaterials. PNSB form an attractive guild of phototrophic organisms with a facultative anaerobic and hyperversatile metabolism that allows them to grow under ever-changing environmental conditions (J.F. Imhoff, 2017; van Niel, 1944). They populate the surface of aquatic environments by absorbing the infrared (IR) spectrum of the sunlight (700-1200 nm) using carotenoids and bacteriochlorophylls, with a competitive advantage in a mixed-culture microbial ecosystem. PNSB can switch between photoorganoheterotrophy, photolithoautotrophy, respiratory or fermentative chemoorgano-heterotrophy, respiratory chemolithoautotrophy, and nitrogen fixation depending on the composition of electron donors and acceptors present in their surroundings (Madigan & Jung, 2009). This enables them to thrive on different pools of electron donors, recycle electrons, achieve redox homeostasis, and grow under alternation of light and dark (McEwan, 1994). PNSB ferment reduced organics into carboxylates in the dark, photoferment them into dihydrogen, or accumulate and condense them as intracellular storage polymers like biopolyesters (e.g., poly-\beta-hydroxyalkanoates, PHAs) as electron sinks under nutrient limitations (Eilert Hustede et al., 1993). Rediscovering PNSB for ecotechnologies and nutrient capture goes via basic study of their metabolism and selection features from pure to mixed cultures, and eco-design to develop robust, non-axenic, and economically appealing processes (Bryant & Frigaard, 2006).

The potential of PNSB for converting diverse carbon sources such as volatile fatty acids (acetate, malate, butyrate and propionate), sugars or alcohols, has been screened with isolates (Alloul *et al.*, 2019; Madigan & Gest, 1979a; Stoppani *et al.*, 1954), underlying the potential of populations of this guild for water treatment. PNSB can assimilate carbon (C), nitrogen (N), and phosphorus (P) from wastewater at COD:N:P ratio of 100:7:2 versus 100:5:1 m/m/m for

activated sludge, with an elemental formula for purple phototrophic biomass given as  $C_1H_{1,8}O_{0,38}N_{0,18}$  (degree of reduction of 4.5 mol e<sup>-</sup> C-mol<sup>-1</sup> X<sub>PPB</sub>) (D. Puvol et al., 2017). Their photon-capturing and energy-recycling physiology lead PNSB to achieve rapid biomass specific maximum growth rates  $(\mu_{max})$  of 1.7 to 5.3 d<sup>-1</sup> and biomass yields ( $Y_{X/COD}$ ) on organic substrates (expressed as chemical oxygen demand, COD) of 0.6 to 1.2 g  $COD_x$  g<sup>-1</sup> COD<sub>s</sub> from mixed to pure cultures (Eroglu et al., 1999; Hülsen et al., 2014), involving additional electron sources from the bulk liquid phase. New-generation biological wastewater treatment processes aim to decrease sludge production and handling, by making use of slow-growing and low-yield microorganisms such as polyphosphate-accumulating and anammox bacteria. In contrast, the use of organisms with a high biomass yield such as PNSB is of definite interest to remove, capture and concentrate carbon, nitrogen and phosphorus nutrient resources out of the wastewater by assimilation into the biomass. The biomass can be then valorized to generate energy through methanization and to produce, e.g., single-cell proteins (*i.e.*, source of microbial proteins), bioplastics via PHAs, and biohydrogen on concentrated streams (Honda et al., 2006; Daniel Puyol et al., 2017).

Technically, one important challenge of photobiotechnologies resides in the limitation of photon supply across the reactor bulk (Pulz, 2001), therefore, many PNSB-based processes have been operated at concentrations below 1 g VSS  $L^{-1}$ . Light limitation is often considered *a priori* as a killing factor for the process performance and economics, while such low biomass concentration can remain a drawback for the intensification of volumetric conversions.

Mixed-culture processes are actively investigated to harness the ability of PNSB to treat wastewater (Hülsen et al., 2016; Nakajima et al., 1997; Verstraete et al., 2016), starting from stabilization ponds (Almasi & Pescod, 1996; Freedman et al., 1983). Process configurations involved continuous up-flow system (Driessens et al., 1987), continuous-flow stirred tank reactor (Alloul et al., 2019), tubular reactor (Carlozzi et al., 2006), sequencing batch reactor (SBR) (Chitapornpan et al., 2012; Fradinho et al., 2013), membrane bioreactor (MBR) (Hülsen et al., 2016), and membrane sequencing batch reactor (MSBR) (Kaewsuk et al., 2010). One challenge in the application of PNSB organisms is considered to remain in the solid-liquid (S/L) separation of the biomass from the aqueous stream. Decoupling the hydraulic (HRT) and solid (SRT) retention times is crucial to retain the biomass in the process.

The use of membrane filtration has been recommended because PNSB have been hypothesized to primarily grow in suspension for catching photons and to settle slowly (Chitapornpan et al., 2012). However, membranes are intended to separate biomass from the treated wastewater, but do not foster the formation of a good settling sludge. In the lab, MBRs are used to maintain biomass in suspension (van der Star et al., 2008). A centrifugation step is still needed after the membrane filtration to efficiently concentrate and harvest the PNSB biomass downstream. Alternatives to MBRs can lead to capital and operational savings, since membrane filtration and fouling relate to substantial pumping energy and maintenance costs besides the use of plastic materials.

Intensification of PNSB-based environmental biotechnologies should be targeted by enhancing the bioaggregation and biofilm-forming ability of the biomass. Although previous works have not tailored SBR regimes to this end (Chitapornpan et al., 2012; Fradinho et al., 2013), the application of substrate gradients via SBR operation can be efficient to stimulate microbial aggregation and biomass accumulation. Granulation of activated sludge biomasses in SBR systems has been the trigger of BNR process intensification (Aqeel et al., 2019; Pronk et al., 2015; M. K. H. Winkler et al., 2018). This should lead to an efficient S/L separation, resulting in lowering costs for downstream processing by potentially reducing the need for ultrafiltration and centrifugation to concentrate the biomass. A SBR design also offers operational flexibility (Wilderer, 1998) to manipulate reactor cycles and loading rates. Although offering less surface-to-volume ratio, the use of simple stirred-tank designs in SBR application can in addition lead to simpler scale-up than flat-sheet, tubular, or membrane-based processes.

Here, we investigated the possibility to develop a mixed-culture biotechnology process based on the enrichment of a concentrated and well-settling PNSB biomass out of activated sludge in a stirred-tank photobioreactor operated under SBR regime and continuously irradiated with IR light. Conditions to enrich and maintain a PNSB mixed culture were elucidated at bench scale, along with microbial competition in the PNSB guild. Biomass growth, aggregation, and composition were analysed along with volumetric rates of C-N-P removal. The here-examined microbial ecology insights and aggregation propensity of the PNSB guild can sustain the development of bioengineering strategies for mixed-culture process development in simple SBR design for wastewater treatment and resource recovery from aqueous nutrient streams.

## 2. Material and Methods

## 2.1 Cultivation medium

The composition of the cultivation medium was calculated based on stoichiometric requirements to sustain PNSB growth and complemented with other minerals. It was adapted from Kaewsuk et al. (2010) to meet with C-N-P anabolic requirements of PNSB. The stock solution consisted of (per liter): 0.914 g of CH<sub>3</sub>COONa·3H<sub>2</sub>O, 0.014 g of KH<sub>2</sub>PO<sub>4</sub>, 0.021 g of K<sub>2</sub>HPO<sub>4</sub>, 0.229 g of NH<sub>4</sub>Cl, 0.200 g of MgSO<sub>4</sub>·7H<sub>2</sub>O, 0.200 g of NaCl, 0.050 g of CaCl<sub>2</sub>·2H<sub>2</sub>O, 0.100 g of yeast extract, 1 mL of vitamin solution, and 1 mL of trace metal solution. The vitamin solution contained (per liter) 200 mg of thiamine–HCl, 500 mg of niacin, 300 mg of  $\rho$ -amino-benzoic acid, 100 mg of pyridoxine–HCl, 50 mg of biotin, and 50 mg of vitamin B12. The trace metal solution contained (per liter) 1100 mg of EDTA–2Na·2H<sub>2</sub>O, 2000 mg of FeCl3·6H<sub>2</sub>O, 100 mg of ZnCl2, 64 mg of MnSO<sub>4</sub>·H<sub>2</sub>O, 100 mg of H<sub>3</sub>BO<sub>3</sub>, 100 mg of CoCl<sub>2</sub>·6H<sub>2</sub>O, 24 mg of Na<sub>2</sub>MoO<sub>4</sub>·2H<sub>2</sub>O, 16 mg of CuSO<sub>4</sub>·5H<sub>2</sub>O, 10 mg of NiCl<sub>2</sub>·6H<sub>2</sub>O, and 5 mg of NaSeO<sub>3</sub>. Carbon sources (acetate) were separated from nitrogen and phosphate sources to avoid contaminations.

### 2.2 Anaerobic, sequencing-batch, photobioreactor setup

The PNSB enrichment was performed in a 1.5-L cylindrical, single-wall, glass, stirred-tank reactor (Applikon Biotechnology, Netherlands) (Figure 1.A). The reactor was inoculated at 0.1 g VSS L<sup>-1</sup> of flocculent activated sludge taken from the BNR WWTP Harnaschpolder (the Netherlands) after washing the sludge with the cultivation medium 3 times. The reactor operated for 6 months under SBR regime at controlled temperature of  $30 \pm 1$  °C and pH of 7.0  $\pm 1.0$  on the cultivation medium.

To select for purple phototrophs and to avoid the proliferation of green phototrophs, the reactor was placed in a dark fume hood providing only IR light. A white light source was beamed with two halogen lamps (120 W, Breedstraler, GAMMA, Netherlands) placed at the side of the reactor and filtered for IR wavelengths (>700 nm) with two filter sheets (Black Perspex 962, Plasticstockist, UK) placed in front of the lamps. Irradiance was measured at the reactor surface with a pyranometer (CMP3, Kipp & Zonen, Netherlands) and set at a relatively high value of 375 W m<sup>-2</sup> to promote PNSB enrichment and biomass growth.



Figure 1. Stirred-tank photobioreactor enriched in PNSB. The reactor was inoculated at 0.1 g VSS L<sup>-1</sup> with BNR activated sludge. One initial batch of 40 h was used to activate the biomass and test the enrichment of PNSB prior to switching to SBR operation over 5 months.

### 2.3 Anaerobic, sequencing-batch, photobioreactor operation

After a first 40-h period under batch regime to check for the selection of PNSB, the system was switched to an SBR regime, consisting of discharge, idle, feed and settling phases (Figure 1.B). Different cycle timings and HRT, reaction phase length, and COD loading rates were tested as followed in three operational modes. In SBR1, 24 cycles of 24 h each were applied (i.e., 24 days of experiment), consisting of: biomass settling (3 h) effluent withdrawal (5 min), influent feeding (5 min), and reaction (20.75 h). In SBR2, the total length of the cycles was decreased 3-fold and set at 8 h. The reaction phase was shortened to 4.75 h, while all the other phases were maintained. The reactor was operated over 205 cycles. In SBR3, the cycle composition was maintained as in SBR2, while the COD concentration was doubled from 430 to 860 mg  $COD_{Ac} L^{-1}$  in the influent to prevent COD-limitations along the reaction phase. All SBRs were operated at a volume exchange ratio of 50%. The stepwise adaptation of the SBR operations from 1 to 3 cycles day<sup>-1</sup> resulted in HRTs from 48 h (SBR1) to 16 h (SBR2 and SBR3) and in volumetric organic loading rates (OLRs) of 0.215 (SBR1) to 0.645 (SBR2) and 1.290 (SBR3) kg COD  $d^{-1}$  m<sup>-3</sup> (Table 1). The sludge retention time (SRT) was let freely evolve across the SBR operations without controlled purge of the biomass, with median values ranging from 1.5 to 11 d as calculated from eq. 2 in Supplementary material 1, as a result of biomass accumulation.

The pH of the mixed liquor was controlled at  $7.0 \pm 1.0$  by automatic addition of HCl or NaOH at 1 mol L<sup>-1</sup> each. The bulk liquid was sparged with argon gas (quality 99.999%) to maintain anaerobic conditions, while continuously stirring at 378 rpm (potentiostat ADI 1012, Applikon, the Netherlands) during the reaction phase.

Table 1. Operational conditions of the three SBR regimes across the experimental period. SBR1 was run with a 24-h cycle composed of 20.75 h of reaction time and with  $257 \pm 54$  mg COD L<sup>-1</sup> in the bulk liquid phase after feeding. In SBR2, the total cycle length was shortened to 8 h, with a reaction time of 4.75 h. In SBR3, the initial COD concentration was doubled compared to the first two SBRs. All SBRs were run at a volume exchange ratio of 50%.

Process parameters	Units	SBR1	SBR2	SBR3		
SBR cycles						
Number of SBR cycles per day	(-)	1	3	3		
Reaction phase length per cycle	(h)	20.75	4.75	4.75		
Discharge, idle, feeding phases lengths	(min)	5	5	5		
Settling phase length	(h)	3	3	3		
Retention times						
Hydraulic retention time (HRT)	(h)	48	16	16		
Sludge retention time (SRT) <sup>1</sup>	(d)	1.5	7.2	10.6		
Loadings						
Volumetric organic loading rate (OLR)	$(g \text{ COD } d^{-1} L_r^{-1})$	0.215	0.645	1.29		
C:N:P ratio in the influent	(m / m / m)	100 : 35.8 : 3.8	100 : 35.8 : 4.2	100 : 11.2 : 1.7		
Measured initial concentrations in the bulk liquid phase at the beginning of reaction phases						
Acetate	(mg COD L <sup>-1</sup> )	$257\pm54$	$232\pm18$	$443\pm76$		
Ammonium	$(mg N-NH_4^+ L^{-1})$	$92 \pm 21$	$83 \pm 16$	$49\pm17$		
Orthophosphate	$(mg P-PO_4^{3+} L^{-1})$	9.7 ± 1.3	$9.9\pm3.3$	$7.7 \pm 1.1$		

<sup>1</sup> The sludge retention time (SRT) was let freely evolve over the experimental period. The reactor was operated without purge of biomass. The SRT increased as a result of biomass accumulation in the reactor. The median value over each SBR period is provided. The distributions of SRT are displayed in Figure 2 and detailed evolutions in Supplementatry material 3.

## 2.4 Analytical methods to measure growth and nutrient consumptions

### 2.4.1 Measurements of biomass growth and nutrient concentrations

Biomass growth was monitored spectrophotometrically by absorbance at a wavelength of 660 nm (DR3900, Hach, Germany) 4-5 times a week (Supple-mentary material 2), and gravimetrically by quantifying the concentration of volatile suspended solids (VSS) as described in experimental methods for wastewater treatment (van Loosdrecht et al., 2016). For the 40-batch and SBR1, absorbance measurements were adequate since the biomass was low in concentration and in suspension.

For SBR2 and SBR3, the biomass aggregated and VSS measurements were much more accurate.

The consumption of the dissolved nutrients was monitored by sampling the mixed liquor at the beginning and end of the reaction phase, after centrifugation (5 min, 17000 x g) and filtration of the supernatant on 0.45-µm filters (Millex-HV, PVDF, Germany). The concentrations of COD, ammonium (as N-NH<sub>4</sub><sup>+</sup>) and orthophosphate (as P-PO<sub>4</sub><sup>3-</sup>) were measured by colorimetric assays (LCK kits no. 114/614/COD, 302/303/ammonium, 348/350/phosphate; Hach-Lange, Dusseldorf, Germany) followed by spectrophotometry (DR3900, Hach, Germany). The COD colorimetric method measured all oxidizable substances (here notably acetate, yeast extract, and EDTA from the trace element solution). As technical control, samples were measured in triplicates and the relative standard deviation was 0.5 - 1.9%.

### 2.4.2 Computations of microbial conversions and extraction of growth parameters

All symbols and equations used to compute microbial conversions and extraction of growth parameters are available in Supplementary material 1. In short, the average percentage of removal ( $\eta_s$ , %), total rate of nutrient removal ( $R_s$ , kg S d<sup>-1</sup>), apparent volumetric rate of removal of nutrients ( $r_s$ , kg S d<sup>-1</sup>), and apparent growth rate ( $\mu_{max}$ , d<sup>-1</sup>) were calculated using mass balances over the C-N-P nutrients and biomass at a volumetric exchange ratio (VER) of 50%. Measurements of nutrients were performed at the beginning and end of the batch reaction phases of the SBR. Influent concentrations were back-calculated using the VER. The concentrations of nutrients in the effluent were assumed identical as at the end of the reaction phase.

Basic kinetic and stoichiometric parameters for microbial conversions and growth were assessed from nutrient consumptions and biomass production profiles using Aquasim (Reichert, 1994). A mathematical model was constructed using mass balances for substrate consumption and biomass production, and fitted to the experimental data. The maximum biomass-specific rate of acetate consumption ( $q_{S,max}$ , kg S d<sup>-1</sup> kg X), maximum yield of biomass production on substrate ( $Y_{X/S,max}$ , kg X kg<sup>-1</sup> S), and maintenance rate on substrate ( $m_S$ , kg S d<sup>-1</sup> kg X) were derived by parameter fit from the Herbert-Pirt relation of substrate allocation for growth and maintenance. The biomass-specific maximum rate of growth ( $\mu_{max}$ , d<sup>-1</sup>) was computed from the relation between  $q_{S,max}$  and  $Y_{X/S,max}$ , assuming the maintenance rate negligible compared to the maximum growth rate during the exponential phase of the batch reaction period.

## 2.5 Analysis of biomass and microbial community compositions

2.5.1 Light microscopy analysis of microbial morphotypes and bioaggregates Microbial morphotypes present in the enrichment were visually observed by phase contrast microscopy (Axioplan 2, Zeiss, Germany).

## 2.5.2 Wavelength scan analysis of pigment content in the PNSB-enriched biomass

The evolution of the biomass contents in bacteriochlorophyll a (BChl a) and carotenoids in the biomass was used as a proxy for tracking the PNSB enrichment in the mixed liquor. Measurements were performed by wavelength scan over the visible and near-infrared spectrum from 400 to 1000 nm (DR3900, Hach, Germany). A focus was attributed to absorbance peaks between 800-900 nm (BChl a) and 400-600 nm (carotenoids).

# 2.5.3 V3-V4 16S rRNA gene amplicon sequencing of bacterial community compositions

Genomic DNA was extracted from biomass samples throughout the duration of the experiment, using UltraClean Microbial Isolation kits (MOBIO laboratories, Inc., USA) following manufacturer's instructions, and stored at -20 °C. The concentrations and qualities of the DNA extracts were measured by Qbit3 fluorimeter (Thermofisher Scientific, USA), according to manufacturer's instructions. The DNA extracts were sent to Novogene (China) for amplicon sequencing. The V3-V4 regions of the 16S rRNA gene were amplified by polymerase chain reaction (PCR) using the set of forward V3-V4 forward 341f (5'-CCTACGGGAGGCAGCAG

-3') and reverse 806r (5'- GGACTACHVGGGTWTCTAAT-3') primers (Takahashi *et al.*, 2014). The amplicon sequencing libraries were pooled and sequenced in an Illumina paired-end platform. After sequencing, the raw read were quality filtered, chimeric sequences were removed and OTUs were generated on the base of  $\geq$  97% identity. Subsequently, microbial community analysis was performed by Novogene using Mothur & Qiime software (V1.7.0). For phylogenetical determination the most recent SSURef database from SILVA (<u>http://www.arb-silva.de/</u>) was used. Relative abundances of OTUs were reported as % total sequencing reads count.

## 3. Results

3.1 High and simultaneous removal of C-N-P nutrients was achieved in the PNSB-enriched, mixed-culture, stirred-tank SBR

The nutrient removal performances achieved by the PNSB-based process from SBR1 to SBR2 and SBR3 regimes are displayed in Figure 2 and Table 2. The detailed dynamics in nutrient and biomass concentrations and compositions are provided in Supplementary material 3.

#### *3.1.2 Nutrient removing activities were detected during the initial batch*

During the first 40 h of batch used to activate the biomass, nutrients were removed at 98 % COD (as acetate), 52 % N-NH<sub>4</sub><sup>+</sup>, and 60% P-PO<sub>4</sub><sup>3-</sup> (Figure 2.A). These related to apparent volumetric removal rates of 0.190  $\pm$  0.048 kg COD d<sup>-1</sup> m<sup>-3</sup>, 21.5  $\pm$  8.6 g N d<sup>-1</sup> m<sup>-3</sup>, and 2.5  $\pm$  0.5 mg P d<sup>-1</sup> m<sup>3</sup> (Figure 2.B). The COD:N:P consumption ratio was 100:7.5:0.12 (m/m/m) in this batch.

# 3.1.3 A complete removal of acetate was achieved across all SBR operation modes

During SBR1, the average percentage of removal of the biodegradable COD was 96%, with an average volumetric consumption rate of  $0.220 \pm 0.060$  kg COD d<sup>-1</sup> m<sup>-3</sup>. During SBR2, 96% of the COD was removed as well at a 4-fold higher rate of  $0.891 \pm 0.235$  kg COD d<sup>-1</sup> m<sup>-3</sup>. The carbon was fully removed over the first hour of the reaction phase, resulting in remaining 3.75 h of substrate limitation. During SBR3, the COD load in the influent was doubled, and as a result, no nutrient limitation occurred during the reaction phase. COD remained highly removed at 91%, with a volumetric removal rate of  $1.08 \pm 0.32$  kg d<sup>-1</sup> m<sup>-3</sup>.

# *3.1.4 A maximum of 85% of ammonium and 74% of phosphate was removed from the inflow*

The ammonium removal rates increased from  $26 \pm 13$  g N-NH<sub>4</sub><sup>+</sup> d<sup>-1</sup> m<sup>-3</sup> in SBR1 to  $83.4 \pm 35$  g N d<sup>-1</sup> m<sup>-3</sup> during SBR2, and  $113.3 \pm 62$  g N d<sup>-1</sup> m<sup>-3</sup> during SBR3. Average N-removal percentages evolved from 53% to 44% and 77% of the ammonium load across the three SBRs, respectively. Removal rates of or-thophosphate increased from  $3.0 \pm 0.7$  g P-PO<sub>4</sub><sup>3-</sup> d<sup>-1</sup> m<sup>-3</sup> of SBR1 to  $10.7 \pm 4.5$  g P d<sup>-1</sup> m<sup>-3</sup> in SBR2 and  $15.2 \pm 4.6$  g P d<sup>-1</sup> m<sup>-3</sup> in SBR3, with average P-removal percentages of 57, 45 and 73% per cycle, respectively. Under the non-limiting COD conditions of SBR3, the acetate, ammonium, and orthophosphate were released at median concentrations of 43 (min = 17; 1<sup>st</sup>-3<sup>rd</sup> quartile = 28-62) mg COD L<sub>Eff</sub><sup>-1</sup>, 10 (2; 8-13) mg N-NH<sub>4</sub><sup>+</sup> L<sub>Eff</sub><sup>-1</sup>, and 2.0 (0.4; 1.7-2.9) mg P-PO<sub>4</sub><sup>3-</sup> L<sub>Eff</sub><sup>-1</sup>, *i.e.*, close to European discharge criteria.

Thus, the average apparent COD:N:P assimilation ratio evolved from 100:7.5:0.12 in the batch to 100:9.2:1.2 (SBR1-2) under COD-limitation and 100:6.7:0.9 (SBR3) under non-COD-limitation. Across and beyond the experimental period, the intrinsic kinetics and stoichiometry of the PNSB-enriched biomass ranged with a biomass specific maximum growth rate ( $\mu_{max}$ ) of 0.96-2.16 d<sup>-1</sup> and a maximum yield of biomass production on substrate ( $Y_{X/COD,max}$ ) of 0.21-0.74 g VSS g<sup>-1</sup> CODs, respectively. This related to a yield value of 0.34-1.19 g CODx g<sup>-1</sup> CODs when using a theoretical elemental composition of C<sub>1</sub>H<sub>1.8</sub>O<sub>0.38</sub>N<sub>0.18</sub> (1.607 g CODx g<sup>-1</sup> VSS) for purple phototrophic bacteria (D. Puyol et al., 2017). The maximum biomass specific consumption of acetate

 $(q_{COD,max})$  ranged from 0.03-0.78 kg CODs h<sup>-1</sup> kg<sup>-1</sup> VSS. These measurements were performed directly during SBR cycles at the actual concentration of the biomass present in the system. More accurate measurements and derivation of these physiological parameters can be performed at diluted initial concentrations of PNSB biomass to prevent nutrient and light limitations during batch tests.



Figure 2. Nutrient removal and biomass characteristics across SBR operations in the mixed-culture PNSB photobioreactor, A: Increases in COD, ammonium, and orthophosphate nutrient removal percentages from SBR1 to SBR3. On average, 95% of the COD was removed during all operational states. N-NH<sub>4</sub><sup>+</sup> and P-PO<sub>4</sub><sup>3</sup> reached 77% and 73% of removal from the synthetic influent. B: Gradual increases in volumetric rates of C-N-P nutrient removals from SBR1 to SBR3. C: Increase in the fraction of mixed-liquor biomass that settled in the bioreactor along SBR1 (days 3-30), SBR2 (days 30-100), and SBR3 (days 100-135) after inoculation with BNR activated sludge and a first batch of 40 h. D: Accumulation of biomass in the photobioreactor from SBR1 to SBR2 and SBR3. The grey triangles relate to cleanings and resuspensions of the wall biofilm in the biosystem. They indicate the total amount of biomass that accumulated in the reactor. E: Distributions of biomass concentrations in the reactor at the end of the reaction phase and in the effluent after settling. The settling ability of the biomass increased steadily during SBR operations. The residual biomass concentration in suspension at the end of the settling phase in SBR3 was 10 times lower than the concentration in the mixed liquor during reaction time, displaying the well-settling property of the PNSB-enriched biomass. F: The SRT was let freely evolve in the reactor, increasing from median values of 2 d (SBR1) to 7.5 d (SBR2) and 11 d (SBR3) along with biomass accumulation.

Table 2. Nutrient removal by the PNSB-enriched biomass in SBR1, SBR2, and SBR3 presented as averages and maximal values of removal rates and removal percentages. Residual concentration and removal percentages for COD met with European legislation limits for all SBRs (averages above 90% removal, with residues close to 60 mg COD L<sup>-1</sup>). Ammonium and orthophosphate were removed up to 77% and 73%, respectively, with residual concentrations reaching 11 mg N-NH<sub>4</sub><sup>+</sup> L<sup>-1</sup> and 2 mg P-PO<sub>4</sub><sup>3-</sup> L<sup>-1</sup>.

Chemical parameter	Units	SBR1	SBR2	SBR3
Organic matter (as COD)				
Volumetric removal rates	(kg d <sup>-1</sup> m <sup>-3</sup> )	$0.220\pm0.056$	$0.891\pm0.235$	$1.019\pm0.318$
Removal percentage	(%)	$96 \pm 7$	$96 \pm 11$	$91 \pm 11$
Maximal volumetric removal rates	$(\text{kg d}^{-1} \text{ m}^{-3})$	0.387	1.488	2.437
Maximal removal percentage	(%)	100	100	98
Average residual concentrations	(mg L <sup>-1</sup> )	$67 \pm 25$	$62 \pm 32$	$61 \pm 6$
Ammonium (as N-NH <sub>4</sub> <sup>+</sup> )				
Volumetric removal rates	$(g d^{-1} m^{-3})$	$26 \pm 13$	$83\pm35$	$113 \pm 62$
Removal percentage	(%)	$53 \pm 18$	$44 \pm 10$	$77 \pm 21$
Maximal volumetric removal rates	$(g d^{-1} m^{-3})$	52	159	65
Maximal removal percentage	(%)	83	60	94
Average residual concentrations	(mg L <sup>-1</sup> )	$39 \pm 16$	$39 \pm 6$	$11 \pm 7$
Orthophosphate (as P-PO <sub>4</sub> <sup>3-</sup> )				
Volumetric removal rates	$(g d^{-1} m^{-3})$	$3 \pm 1$	$11 \pm 5$	$15 \pm 5$
Removal percentage	(%)	$60 \pm 11$	$45 \pm 10$	$73 \pm 14$
Maximal volumetric removal rates	$(g d^{-1} m^{-3})$	5	18	25
Maximal removal percentage	(%)	91	59	95
Average residual concentrations	$(mg L^{-1})$	$4 \pm 1$	$5 \pm 1$	$2 \pm 1$

### 3.1.5 Kinetic and stoichiometric parameters of microbial growth

The maximum biomass specific rate of substrate consumption  $(q_{S,max})$ , the yield of biomass growth of substrate consumption  $(Y_{X/S})$  and the maximum biomass specific growth rate  $(\mu_{max})$  were obtained by parameter fit to batch evolutions of acetate and biomass during selected reaction phases of the SBRs (Table 3). Under the conditions of the initial batch and of SBR1 (i.e., long HRT of 48 h, low OLR of 0.215 kg COD d<sup>-1</sup> m<sup>-3</sup>, very low biomass concentration of 0.1 g VSS L<sup>-1</sup>, and low SRT of 1.5 d), the highly-enriched PNSB biomass displayed a high  $q_{S,max}$  between 5.8-13.7 g COD<sub>S</sub> d<sup>-1</sup>  $g^{-1}$  VSS on which it maximized its growth rate:  $\mu_{max}$  (2.16-3.36 d<sup>-1</sup>) was substantial, ranging between values reported in literature for mixed cultures and pure cultures of PNSB. The biomass thus developed at a relatively low yield  $Y_{X/S, max}$  (0.23-0.39 g VSS  $g^{-1}$  COD<sub>S</sub>). The maintenance rate (m<sub>s</sub>) was estimated to 0.72 g COD<sub>S</sub> d<sup>-1</sup> g<sup>-1</sup> VSS. Under the conditions of SBR2 and SBR3 (i.e., 3-times lower HRTs, 3 to 6times higher OLRs, 16 to 30-times higher biomass concentrations, and 5 to 7-times longer SRTs), the biomass consumed acetate at a 3 to 8-fold lower  $q_{Smax}$  (1.8-2.2 g  $COD_s d^{-1} g^{-1} VSS$ ). The maximum growth rate and yield values could not be extracted from the data collected from the reactions phases at high biomass concentration (low sensitivity of absorbance and VSS measurements to detect changes). More accurate estimates are obtained with batch tests conducted with a diluted concentration of biomass.

Table 3. Observed physiological parameters of the biomass of the PNSB mixed culture extracted from the reaction periods of the initial batch and the three SBR periods, and comparison with literature data obtained from pure-culture and mixed-culture PNSB systems. The values are given based on measured COD units for the acetate substrate and absorbance-calibrated VSS units for the biomass. The elemental formula for purple phototrophic bacteria C1H1.800.38N0.18 (4.5 mol e- C-mol-1, 36 g COD C-mol-1, 22.4 g VSS C-mol-1, 1.607 g CODX g-1 VSS) (D. Puyol et al., 2017) may be used for conversion of VSS in to COD units.

System	q <sub>S,max</sub>	Y <sub>X/S,max</sub>	$\mu_{max}$
	$(g COD_S d^{-1} g^{-1} VSS)$	$(g VSS g^{-1} COD_S)$	$(d^{-1})$
PNSB pure cultures	n.a.	0.98-1.23 <sup>a</sup>	5.28 <sup>a</sup>
PNSB mixed culture	n.a.	0.23-0.63 <sup>b</sup>	0.72-1.68 <sup>b</sup>
Initial batch	5.76	0.39	2.16
SBR1 <sup>c</sup>	$13.68 \pm 5.04$	$0.23\pm0.02$	$3.36 \pm 1.4$
SBR2 <sup>c</sup>	$1.76\pm0.91$	n.a. <sup>d</sup>	n.a. <sup>d</sup>
SBR3 <sup>c</sup>	$2.22\pm0.79$	n.a. <sup>d</sup>	n.a. <sup>d</sup>

<sup>a</sup> Taken from literature (Eroglu et al., 1999; Jih, 1998)

<sup>b</sup> Taken from literature (Hülsen et al., 2016; Hülsen et al., 2014; Kaewsuk et al., 2010; Puyol et al., 2017)

<sup>c</sup> Average values collected over different cycles monitored over SBR1 (cycles 11, 17), SBR2 (cycles 21, 145, 166), and SBR3 (cycles 10, 31, 49, 91).

<sup>d</sup> High biomass concentrations in the system. VSS and absorbance measurement are no sensitive enough to detect growth over reaction period. External batches with diluted biomass should be performed to this end.

## 3.2 SBR operations enhanced the settling ability and accumulation of the PNSB biomass

The settling ability of the PNSB biomass increased across enrichment SBR operations, leading to substantial accumulation of biomass in the system from 0.1 (SBR1) to 1.6 (SBR2) to 3.0 (SBR3) g VSS L<sup>-1</sup> as median values (Figure 2.C-D). The enhancement of the settling ability was measured by comparing these biomass concentrations present in the mixed liquor at the end of the reaction phase with the concentrations in the effluent after the settling phase which was for all SBRs as low as 0.13-0.15 g VSS  $L_{Eff}^{-1}$  (Figure 2.C-D). The fraction of settled biomass increased across SBR1 from 12% to 53% of the VSS present in the mixed liquors at the end of reaction phases, reached 96% by end of SBR2. and remained high at 97±3% over SBR3 (Figure 2.C). The total rates of biomass accumulation calculated over the full settling period of 3 h increased from 0.02±0.01 (SBR1) to 0.69±0.46 (SBR2) and 1.30±0.45 (SBR3) g VSS h<sup>-1</sup>, or from  $0.02\pm0.01$  to  $0.46\pm0.31$  and  $0.87\pm0.30$  kg VSS h<sup>-1</sup> m<sup>-3</sup>, respectively, when translated into volumetric rates. At the beginning of SBR1, the full 3 h period was required to settle the suspended biomass. At the end of SBR3, most of the 5.9 g VSS of biomass that aggregated and accumulated in the system settled in about 10 min (*i.e.*, 35 gVSS h<sup>-1</sup> or 24 kg VSS h<sup>-1</sup> m<sup>-3</sup> effectively). This high settling rate obtained on SBR3 corresponds to a sedimentation G-flux of solids of 4.7 kg h<sup>-1</sup> m<sup>-2</sup>. This displayed the well-settling property of the aggregated PNSB biomass. It underlined potential for considerably shortening the settling phase and SBR cycle length in order to increase the daily loading of the system.

The fraction of VSS in the TSS remained relatively high with 85% (SBR1) to 93% (SBR2) to 80% (SBR3) as median values, *i.e.*, corresponding to a fraction of inorganic suspended solids (ISS) between 7-20%. During SBR3 a period at lower VSS fraction with values below 60% and higher ISS fraction (>40%) was detected between days 97-117, underlying potential accumulation of inorganics, *e.g.*, as intracellular polyphosphate (not measured), during nutrient assimilation in the biomass.

The SRT was let to increase freely, without controlled purge of biomass, as a result of the enhancement of settling properties of the biomass: it rose from 2 d in SBR1 to 7 d in SBR2 and 11 d in SBR3 as median values (Figure 2.D). Strategies can be tested to control the SRT at specific values on the range between, *e.g.*, 3-10 days, depending on nutrient capture and biomass production targets.

The PNSB enrichment process could be easily be tracked visually with the gradual increase in the purple color intensity in the bioreactor (Figure 3.A-D).

3.3 Microscopy images showed an increasing size in microbial aggregates After inoculation with flocculent activated sludge, phase-contrast microscopy imaging revealed the presence of dense aggregates already in SBR1 formed by the PNSB biomass (Figure 3.E-G). Some cells clustered in flower-shaped aggregates, in a way comparable to the typical morphotype of *Rhodopseudomonas*. Other rod-shaped cells were present, putatively belonging to *Rhodobacter* and *Blastochloris* genera. The size of the aggregates increased from 50 to 150  $\mu$ m during the operational time along with the better settling abilities of the biomass.

## 3.4 Wavelength scans highlighted the enrichment of carotenoid and bacteriochlorophyll pigments in the biomass and a shift in predominant populations in the PNSB guild

Carotenoids and bacteriochlorophylls, and their increase along the enrichment of the PNSB biomass, were detected by the presence of absorbance peaks at wavelengths between 450-500 nm and between 800-900 nm. The wavelength scan data presented in Figure 3.H are normalized by the biomass concentrations, expressed as absorbance units at 660 nm. Peaks at 800 nm and 850 nm were already present at the end of the initial batch phase, and persisted during SBR1. At the end of SBR2, the absorbance peaks shifted to higher wavelengths of 805 nm and 865 nm. During SBR3, another peak was detected at 1000 nm that is characteristic for the genus *Blastochloris*. This suggested a shift in predominant microbial populations harbouring different types of pigments in the PNSB guild across the mixed-culture enrichment process.



Figure 3. Evolution of the pigmentation and aggregative characteristics of the PNSBenriched biomass. A: Phase-contrast microscopy images of the aggregates present in SBR1 to SBR3. The size of the aggregates increased during time along with increased settling abilities of the biomass. B: Wavelength scans of intact cultures, normalized for the biomass content (at 660 nm). The presence of PNSB was tracked at peaks around 800-900 nm (Bchl a) and 400-500 nm (carotenoids). After the initial batch phase of 40 h, the peaks typical for PNSB pigments were present, and persisted in the biomass until the end of SBR3.

## 3.5 Amplicon sequencing revealed selection shifts from Rhodobacter to Rhodopseudomonas and Blastochloris genera within the guild of PNSB

The composition of the bacterial community of the mixed culture and underlying shifts in predominant populations were analysed by V3-V4 16S rRNA gene amplicon sequencing. The times series of PNSB populations are displayed in Figure 4. The detailed times series of the full set of identified genera across the sequencing dataset is given in Supplementary material 5.

The BNR activated sludge inoculum presented a diversity of genera, with *Rhodobacter* as the main PNSB detected at 4% of the sequencing read counts. The typical populations of the BNR sludge like ammonium oxidizer (*Nitrosomonas*), nitrite oxidizer (*Nitrospira*), denitrifier (*Zoogloea*), polyphosphate- ("*Candidatus* Accumulibacter") and glycogen-accumulating ("*Ca.* Competibacter") organisms got rapidely outcompeted right after start-up of the first batch under PNSB-selective conditions (Supplementary material 5).

At the end of the 40-h batch phase, *Rhodobacter* reached a relative abundance of 52%. At the end of the first cycle of SBR1, a high-grade enrichment of 90% of *Rhodobacter* was obtained. Around the 10<sup>th</sup> cycle of SBR1 (10 days), the genus *Rhodopseudomonas* got enriched at 15%, and reached 50% at the end of the 23<sup>rd</sup> cycle (23 days after inoculum). The compositions of the communities of the mixed liquor and of the biofilm that developed on the walls of the reactor during the 13<sup>th</sup> cycle revealed that *Rhodopseudomonas* (55%) was outcompeting *Rhodobacter* (5%) in the biofilm, while *Rhodobacter* (60%) was more enriched than *Rhodopseudomonas* (10%) in the mixed liquor. Then, *Rhodobacter* decreased constantly from cycle to cycle, while *Rhodopseudomonas* progressively took the lead in the flocculent biomass as well.

After 18 cycles of SBR2 (*i.e.*, 4.5 and 22.5 days from the starts of SBR2 and SBR1, respectively), *Rhodopseudomonas* became dominant (70%), outcompeting *Rhodobacter* (17%) in the enrichment culture. Interestingly, after 20 days of SBR2, the genus *Blastochloris*, also an affiliate of the PNSB guild, got selected, while the relative abundance of *Rhodopseudomonas* decreased to 60% at the end of SBR2. In SBR3, *Blastochloris* reached 10% of the bacterial community dataset.



x-axes = Time since SBR start-ups (d)

Figure 4. Time series of V3-V4 16S rRNA gene amplicon sequencing of bacterial community compositions in the PNSB-enriched mixed-culture process along SBR regime shifts. After inoculating the reactor with BNR activated sludge ("BNR"), a first PNSB genus Rhodobacter was initially enriched during the first 40-h batch ("40 h") and early SBR1 period. The second PNSB genus Rhodopseudomonas was predominantly selected across operations of SBR2 and SBR3. The third PNSB genus Blastochloris popped up by end of SBR2 and SBR3. The PNSB guild remained predominant in the biomass across the process with an average total relative abundance of sequencing reads affiliated to known PNSB above 60% of the total community dataset (median = 81%; min-max = 60-93%). In SBR1, both the mixed liquor and the wall biofilm were sampled on day 22 and sequenced. In SBR3, both the settled biomass and s were sampled on day 33 after settling, and sequenced. The full set of genera is given in Supplementary material 5.

## 4. Discussion

# 4.1 A high-grade enrichment of a concentrated, well-settling PNSB biomass can be obtained under SBR regime

The enrichment of PNSB has often been successful, while most PNSB mixed cultures reported so far have mainly been in membrane systems (Hülsen et al., 2016). Here, we successfully enriched a mixed culture of PNSB out of activated sludge under traditional SBR regime in a stirred-tank system without the use of a membrane module to separate the biomass and the bulk liquid phase. This went by using the natural propensity of PNSB to form biofilms and bioaggregates. SBR regimes result in substrate gradients across reactor operation from high concentrations at the beginning of the cycle to low residual concentrations at the end. Such substrate gradients are known to promote the bioaggregation of microorganisms (Aqeel et al., 2019; Pronk et al., 2015; M. K. H. Winkler et al., 2018).

Promotion of bioaggregation of PNSB is key for a good S/L separation and accumulation of biomass in the system. One important outcome of this study highlighted that aggregation of PNSB can be stimulated under SBR regime to intensify the volumetric conversions and to facilitate downstream processing. After inoculation at 0.1 g VSS L<sup>-1</sup>, a high concentration of a PNSB-enriched biomass of up to a maximum of 4.0 g VSS  $L^{-1}$  was obtained in SBR3. The good settling ability of the PNSB biomass obtained under this regime resulted in the emission of less than 5% of the mixed liquor biomass in the effluent of SBR3. as low as 0.1 g VSS L<sup>-1</sup>. Interestingly, Driessens et al. (Driessens et al., 1987) have early reported the flocculation and good sedimentation (G-flux of 7-9 kg h m<sup>-2</sup> comparable to well-flocculated activated sludge) of *Rhodobacter* capsulatus in an upflow continuous photobioreactor operated under loading rates of 2.5-5.0 kg C d<sup>-1</sup> m<sup>-3</sup> (as calcium lactate; *i.e.*, 6.7-13.3 kg COD d<sup>-1</sup> m<sup>-3</sup>) and 0.5-1.0 kg N d<sup>-1</sup> m<sup>-3</sup> (as ammonium) with 87% C and N assimilation in the biomass (3.3-4.2 g VSS L<sup>-1</sup>). The PNSB-enriched biomass during SBR3 displayed a high sedimentation G-flux of 4.7 kg h<sup>-1</sup> m<sup>-2</sup> relatively close to the values reached by Driessens et al. (Driessens et al., 1987) under highly concentrated loading rates 5 to 10-fold higher than used here (max. 1.3 kg COD  $d^{-1}$  m<sup>-3</sup> in SBR3). Collectively, this comparison sustains that PNSB can be aggregated for a higher accumulation and retention of biomass to intensify nutrient conversions.

In the PNSB mixed culture, the HRT was initially set high to 48 h (*i.e.*, 1 cycle  $d^{-1}$  at a volume exchange ratio of 50%) to maintain biomass during start-up, prior to decreasing it to 16 h (3 cycles  $d^{-1}$ ) from SBR2 onward. This value was in the range of the HRTs of 8-24 h that have been used in the operation of continuous photo anaerobic membrane bioreactor (PAnMBR) to enrich for purple phototrophic bacteria (PPB) at bench (Hülsen et al., 2016). It was also in

the range of traditional SBRs operated with conventional activated sludge (Mace & Mata-Alvarez, 2002). An operation at 4 cycles  $d^{-1}$  may be foreseen. Decreasing the settling phase length would lead to selectively retain the biomass fraction with higher settling property, with granulation potentialities. This is a typical approach to form a granular sludge out of flocculent activated sludge (Merle K. de Kreuk & van Loosdrecht, 2006; Lochmatter & Holliger, 2014; M. K. H. Winkler et al., 2018).

The settling ability increased along the SBR operation, with a settled biomass fraction raising from 12% (SBR1) to 97% (SBR2-3). Amplicon sequencing revealed that the settled biomass accounted for a 3-fold higher relative abundance of PNSB (80% as sum of *Rhodobacter*, *Rhodopseudomonas*, and *Blastochloris*) than the non-settled biomass (25%) (Figure 4). Together with the phase-contrast microscopy measurements, this highlighted that PNSB are capable of forming bioaggregates with good settling properties. Such increased settling ability links to a more efficient separation of the PNSB biomass from the treated bulk liquid, thus facilitating the downstream processing to recover and valorize the PNSB biomass rich in nutrients for biorefinery purposes.

# 4.2 A high, simultaneous removal of C-N-P nutrients was achieved by the PNSB biomass

High performances of organic matter (96% COD removal at a volumetric rate of 1.1 kg COD  $d^{-1}$  m<sup>-3</sup>), ammonium (77% N-removal at 113 g N  $d^{-1}$  m<sup>-3</sup>), and orthophosphate (73% P-removal at 15 g P d<sup>-1</sup> m<sup>-3</sup>) removal were obtained under operation with a single anaerobic reaction phase using the PNSB process. Conventionally, a sequence of anaerobic, anoxic, and aerobic conditions is needed for full BNR in activated sludge or granular sludge (Barnard & Abraham, 2006; Merle K. de Kreuk et al., 2005). The main difference relies that with PNSB single organisms can remove all nutrients by assimilation into the biomass by making use of photonic energy. BNR activated sludges make use of different microbial guilds of nitrifiers, denitrifiers, polyphosphate- and glycogen-accumulating organisms among others to remove all nutrients biologically. In activated sludge or granular sludge SBRs, the different redox conditions should be alternated to this end. Hence, this PNSB SBR process is a very interesting compact alternative to conventional BNR systems, that enables an enhanced removal of all nutrients in a single reaction phase by managing one single predominant microbial guild, thus simplifying considerably the microbial resource management.

With the PNSB biomass, the SBR process becomes simpler in terms of sequencing operation by feeding, anaerobic reaction, settling, and withdrawal. In practice, a fill/draw phase can be envisioned in function of the settling properties of the PNSB biomass. This can result in a SBR system operated by alternation of fil/draw and reaction phases only. Energy-wise, aeration is not

needed in a PNSB process, resulting in possible electricity savings. In the case of sunlight use, electricity savings will be substantial. The tank will have to be equipped with light filters to supply IR light and select for PNSB as predominant phototrophs in the mixed culture. The irradiance of 375 W  $m^{-2}$ applied in this bench-scale photoSBR is high versus of practical operation window. Sunny regions of Europe are typically characterized by an annual average sunlight irradiance of 150 W m<sup>-2</sup> (Posten, 2009). Nonetheless, light can be provided synthetically in photobioreactors using, e.g. immersed LED devices. In the case of 'artificial' supply of IR light, e.g., with LEDs, the process economics will have to be balanced with the electrical power needed to provide the irradiance needed to run the process. Biofilm formation on lightemitting tubes or light-emitting floating carriers will necessitate periodical cleaning to remediate shading, such as conventionally done for the maintenance of sensors used for process monitoring and control. The aim of this study was not to optimize the reactor design. Further thermoeconomical analysis will have to be conducted to determine the optimum irradiance to supply. This is analogical to the comparison of stirring performances in bench-scale reactors versus full-scale systems. There is room to study PNSB processes at different illumination intensities and their impact on the system responses such as enrichment grades, biomass concentrations, aggregation levels, and nutrient removal performances. Recent studies published on purple phototrophic bacteria have involved irradiances of ca. 50 W m<sup>-2</sup> (Hülsen et al., 2016; D. Puyol et al., 2017) which is about 8-times lower than the one used here at bench. However, no study has yet come with clear information on irradiance cutoffs and light patterns related to the microbial performance of PNSB in the mixed-culture and the economics of pilot and full-scale PNSB processes.

The volumetric removal rate of up to 1.1 kg COD d<sup>-1</sup> m<sup>-3</sup> achieved under the operation of SBR3 is comparable to the ranges of 0.8-2.5 kg COD d<sup>-1</sup> m<sup>-3</sup> reported for the PAnMBR (Hülsen et al., 2016), 0.2-1.4 kg COD d<sup>-1</sup> m<sup>-3</sup> for a continuous-flow stirred-tank reactor without separation of PNSB biomass (Alloul et al., 2019), and 1.2-3.2 kg COD d<sup>-1</sup> m<sup>-3</sup> for conventional BNR activated sludge processes (Tchobanoglous *et al.*, 2003). It was nonetheless higher compared to aeration reactors, anaerobic ponds, and oxidation ditches (Tchobanoglous *et al.*, 2003). Further enhancement of the COD loading rate and removal rate will be achieved by decreasing the SBR cycle time.

The difference in COD:N:P assimilation ratio between SBR1-2 (100:9.2:1.2 m/m/m) and SBR3 (100:6.7:0.9) resulted from the doubling of the acetate load in the influent. These COD:N:P assimilation ratios were in the range of ratios of 100:5.1-7.1:0.9-1.8 that have been characterized during growth of purple phototrophic bacteria (Hülsen *et al.*, 2016a; Puyol *et al.*, 2017b). As comparison basis, a COD-N-P assimilation ratio of 100:5:1 is theoretically used for activated sludge (Henze et al., 2000).

The oscillating periods of higher ISS content (>40%) in the PNSB biomass during SBR3 may underlie an interesting accumulation of inorganics in the biomass, *e.g.*, as intracellular polyphosphate. Similar ISS fractions of 30-40% have been widely detected in biomasses engineered for an enhanced biological phosphorus removal (EBPR) (Weissbrodt et al., 2014). Ecophysiological elucidation of PNSB populations for phosphorus removal is of scientific and technological interests.

# 4.3 Acetate and wavelength gradients can trigger microbial selection in the PNSB guild

Phototrophic organisms are widespread in natural and anthropogenic environments. In the 16S rRNA gene amplicon sequencing analysis of the flocculating activated sludge here used as inoculum, around 4% of the total reads was reported to belong to the genus Rhodobacter, and sequences corresponding to *Rhodopseudomonas* genus were also detected, highlighting the presence of PNSB in conventional sludges. Potentially, due to their versatile metabolism, different genera of PNSB can be constitutively present in activated Phototrophs derive the energy for their metabolism from the sludges. conversion of photons energy to chemical energy. The lack of requirements of substrate utilization to generate energy and external electron acceptors to catabolize acetate give to phototropic organisms a selective advantage over general chemotrophic organisms. PNSB absorb light in the IR spectrum to generate energy. The extensive IR irradiation along with a continuous stirring of the system led to the penetration of the light in the reactor, providing a selective pressure for the enrichment of PNSB. Spectrophotometric measurements of the biomass by wavelength scans from 300 to 900 nm revealed absorbance peaks characteristics for carotenoids and bacteriochlorophylls in PNSB. Peaks at 805 and 850 nm are typical for bacteriochlorophylls detected in vivo from cells of Rhodopseudomonas capsulata (Michael T Madigan & Gest, 1979), whereas a peak around 865 nm is typical for Rhodobacter (Zubova et al., 2005). These peaks were detected across the whole experimental period, indicating the presence and selection of PNSB organisms in the process. Pigments are excellent biomarkers of phototrophic populations, and provide specificity to distinguish between them (Stomp et al., 2007). Wavelength scan analyses are therefore very efficient for a rapid measurement (at min level) of the selection of PNSB.

The 16S rRNA gene amplicon sequencing analysis provided insights at higher resolution on the composition of the PNSB guild and underlying selection phenomena. Amplicon sequencing revealed a consistent enrichment of PNSB after already the first 40 h of batch. The initial enrichment of *Rhodobacter* followed by selection of *Rhodopseudomonas* and then *Blastochloris* can be

explained by competition phenomena across substrate and wavelength gradients between these genera inside the guild of PNSB.

Okubo & Hiraishi (2007) have reported a preferential selection of *Rhodobacter* under high acetate concentration (5-20 mmol L<sup>-1</sup>, *i.e.*, 320-1280 mg CODs L<sup>-1</sup>) due to its low affinity for acetate, while *Rhodopseudomonas* was enriched at lower concentrations  $(0.5 - 1 \text{ mmol L}^{-1}, i.e., 32-64 \text{ mg CODs L}^{-1})$ . The initial 40-h batch was fully loaded with acetate across the whole reaction period, making this condition favourable to select for *Rhodobacter*. Instead, *Rhodopseudomonas* harbours a higher affinity (*i.e.*, lower affinity constant K<sub>s</sub> of 0.11 mM for *Rhodopseudomonas* vs 0.23 mM for *Rhodobacter*) (Okubo & Hiraishi, 2007) for acetate, enabling this population to grow more efficiently than *Rhodobacter* under acetate-limited conditions. During SBR1 and SBR2, the carbon source became progressively depleted after 1.5 h of reaction phase, leaving other 2.5 h of starvation period at low residual acetate concentration. This provided *Rhodopseudomonas* with a competitive advantage for growth.

The competition between Rhodobacter and Rhodopseudomonas may also be governed by their growth rate and thus the SRT in the system. Populations of *Rhodobacter* have displayed a higher maximum growth rate (1.8-2.2 d<sup>-1</sup> in an enrichment and 2.3-3.8 d<sup>-1</sup> with isolates) about 2.6 times faster than Rhodopseudomonas on VFA (Alloul et al., 2019). Batch regimes primarily select on growth rate: organisms deploy their maximum growth rate across most of a batch period during which substrate concentrations are mostly not limiting (Rombouts et al., 2019a). The organism with the highest growth rate that can be activated under the actual operation conditions is therefore preferentially selected. This underlay the selection for Rhodobacter first prior to the progressive establishment of *Rhodopseudomonas* along the progressive increase in SRT. Light availability can also be accounted as responsible for the selective enrichment of Rhodobacter or Rhodopseudomonas. As mentioned, bacteriochlorophylls from *Rhodobacter* and *Rhodopseudomonas* absorb at different wavelengths (800-850 vs. 865 nm respectively). At the beginning of the operation, the biomass concentration in the reactor was lower compared to following phases (*i.e.*, SBR3). The initial higher availability of light at shorter wavelengths can have led to the selection of Rhodobacter, and similarly, the shadowing effect of the biomass can have acted as a natural filter, resulting in the selection of *Rhodopseudomonas*.

The genus *Blastochloris*, which appeared during SBR3, harbours bacteriochloropyll b (BChl b) instead of BChl a in *Rhodobacter* and *Rhodopseudomonas*. BChl b absorbs lower photonic energy at higher wavelengths (1020-1030 nm) (Hoogewerf et al., 2003) and can, therefore, interestingly survive at higher cell densities (here, around 1.5 g VSS L<sup>-1</sup>) with lower light penetration in the bulk liquid. Absorbance of the incident IR light increased across reactor operation with the development of a biofilm dominated by *Rhodopseudomonas* on the reactor wall and with a high concentration of

PNSB biomass of up to 3.8 g VSS  $L^{-1}$  that accumulated in the reactor. According to the Beer-Lambert law, the accumulation of *Rhodobacter and Rhodopseudomonas* in the reactor and wall biofilm resulted in the absorbance of the higher-energy wavelengths in the 800-850 nm range of the IR light supplied, thus acting as wavelength filter. The lower-energy wavelengths not absorbed by *Rhodobacter* or *Rhodopseudomonas* were still available in the bulk-liquid, where *Blastochloris* could have absorbed them. The conjunction of the relatively high irradiance of 375 W m<sup>-2</sup> and high SRT of 11 d were likely favorable for *Blastochloris* selection. Physiological characterization of this genus is needed in order to better predict its competition with other members of the diverse PNSB guild like *Rhodobacter* and *Rhodopseudomonas* among others.

Overall, substrate gradients, light gradients, biofilm formation, and bioaggregation were identified as factors that triggered population selection and dynamics in the PNSB enrichment. Different lineages therefore act *de concert* inside the guild of PNSB, providing metabolic redundancy and process resilience in the case of regime shifts in the process.

### 4.4 PNSB mixed cultures from bench toward process development

The development of a lab-scale SBR system enriched for PNSB opens the doors for a possible upscaling of the process. A high nutrient capture was coupled with the production of a PNSB-rich biomass. Such biomass can be valorized for, e.g., proteins or PHAs productions (Alloul et al., 2018; Honda et al., 2006; Hülsen et al., 2018b). The high settling ability of the biomass allows an easier solid-liquid separation of the latter from the treated water either in a compact external settler or directly in the SBR tank. This provides a definite downstream processing advantage over suspended biomass. It also overcomes the use of membrane filtration modules. The SBR regime resulted in the efficient aggregation of PNSB, underlying an enhanced settling ability and accumulation of biomass in the system. The SRT is an important process variable to control toward a stable bioprocess (Morgenroth & Wilderer, 1999). This becomes even more important in the perspective of harnessing the phosphorus removal capability of the PNSB biomass: cells saturated with phosphorus have to be effectively removed from the system such as conventionally performed by purge of excess sludge to maintain robust activated sludge or granular sludge processes operated for EBPR (Barnard & Abraham, 2006; Weissbrodt et al., 2013). The growth rate and affinity for the substrate are further important parameter to manage the selection of PNSB populations in either batch or continuous-flow reactor regimes, respectively. Similarly, light irradiance is a key operational variable since it constitutes the primary energy source for PNSB. Light penetration and distribution are directly linked to the reactor geometry. In surface water ecosystems, IR light photons are typically consumed over the first 30 cm depth. Following the Beer-Lambert law, the absorbance of light will substantially increase with the biomass concentration. Shallow reactor systems can be opportune. SBR regimes can easily be transferred from stirred-tank to any reactor design, like raceway systems (or also known as carrousel plants) currently under investigation for green and purple phototrophic mixed-culture processes (Alloul et al., submitted). The application of substrate gradients via SBR or plug-flow reactor configurations can foster biomass aggregation to sustain efficient S/L separation for biomass recovery on top of nutrient capture.

## 5. Conclusions

We investigated at bench the possibility to establish a mixed-culture PNSB process for nutrient capture from wastewater in a photobioreactor operated as a traditional simple and flexible stirred-tank SBR. This work led to the following three main conclusions:

SBR process conditions stimulated aggregation and accumulation (as high as 3.8 g VSS  $L^{-1}$ ) of a PNSB-enriched mixed culture in a fast-settling biomass that removed all nutrients biologically in a single reaction stage. The formation of compact aggregates facilitated S/L separation.

Nutrient removal was substantial by assimilation in the biomass, reaching simultaneously 96% of organic matter at 1.1 kg COD d<sup>-1</sup> m<sup>-3</sup>, 77% of ammonium at 113 g N d<sup>-1</sup> m<sup>-3</sup>, and 73% of orthophosphate at 15 g P d<sup>-1</sup> m<sup>-3</sup>, *i.e.*, comparable to BNR activated sludge processes. Under non-COD-limiting conditions, the process reached the nutrient discharge limits set by the European Union.

The PNSB guild accounted for as high as 90% of the bacterial community of the sludge (*i.e.*, amplicon sequencing dataset), enabling a simple management of the microbial resource. A sequential selection between the genera *Rhodobacter*, *Rhodopseudomonas*, and *Blastochloris* was detected inside PNSB, allowing for functional redundancy in the microbiome and highlighting the microbial ecology of PNSB populations across light wavelengths.. Next investigations should elucidate competition phenomena along growth rates, substrate affinities, and wavelength gradients across the mixed liquor.

For engineering practice, process analysis should cover the technological and economical aspects related to light supply in the bioreactor. Besides wastewater treatment, the value of the PNSB-based mixed-culture SBR process will reside in opportunities for water and resource recovery by valorization of the retained, concentrated, and nutrient-rich PNSB biomass.

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## Supplementary material Supplementary material 1: Symbols and Equations

# Table SM1.1. Symbols used in calculation and model computation formula. Substrate concentrations were expressed in COD (organic matter), N-NH<sub>4</sub><sup>+</sup> (ammonium), and P-PO<sub>4</sub><sup>3-</sup> (orthophosphate) based units. Biomass concentrations were expressed in COD based units.

Symbols	Units	Definition
Cs	kg S m <sup>-3</sup> or mg S L <sup>-1</sup>	Concentration of substrate in the reactor (state variable)
$C_{S,inf}$	kg S m $^{\!\!-\!\!3}$ or mg S L $^{\!\!-\!\!1}$	Concentration of substrate in the influent
C <sub>S,0</sub>	kg S m $^{\!\!-\!\!3}$ or mg S L $^{\!\!-\!\!1}$	Concentration of substrate at beginning of reaction phase
C <sub>S,end</sub>	kg S m $^{\text{-3}}$ or mg S L $^{\text{-1}}$	Concentration of substrate at end of reaction phase
$C_{S,\text{eff}}$	kg S m $^{\text{-3}}$ or mg S L $^{\text{-1}}$	Concentration of substrate in the effluent
Cx	kg X m $^{\!\!-3}$ or mg X L $^{\!\!-1}$	Concentration of biomass in the reactor (state variable)
$C_{X,\mathrm{eff}}$	kg X m $^{\!\!-3}$ or mg X L $^{\!\!-1}$	Concentration of biomass in the effluent
HRT	d or h	Hydraulic retention time
Ks	kg S m $^{\text{-3}}$ or mg S L $^{\text{-1}}$	Half-saturation affinity constant for substrate
ms	$kg \ S \ d^{\text{-1}} \ kg^{\text{-1}} \ X$	Biomass maintenance rate
$M_{S}$	kg S	Mass of substrate
$M_{\rm X}$	kg X	Mass of biomass
$\mu$ or $q_X$	$kg X d^{-1} kg^{-1} X = d^{-1}$	Biomass specific growth rate
N <sub>cycles</sub>	cycles d <sup>-1</sup>	Number of SBR cycles per day
$\eta_{\rm S}$	%	Percentage of nutrient removal
qs	$kg \ S \ d^{\text{-1}} \ kg^{\text{-1}} \ X$	Biomass specific rate of substrate consumption
$Q_{inf}\!=\!Q_{eff}\!=\!Q$	m <sup>3</sup> cycle <sup>-1</sup> or L cycle <sup>-1</sup>	Volume of influent fed and effluent withdrawn per SBR cycle
Qsample	m <sup>3</sup> cycle <sup>-1</sup> or mL cycle <sup>-1</sup>	Volume of mixed liquor samples collected during reaction phase
Q <sub>purge</sub>	m <sup>3</sup> cycle <sup>-1</sup> or mL cycle <sup>-1</sup>	Volume of mixed liquor purged at the end of reaction phase $(Q_{purge} = 0 \text{ if SRT} \text{ let freely evolve})$
r <sub>s</sub>	kg S d <sup>-1</sup> m <sup>-3</sup>	Apparent volumetric rate of nutrient removal
Rs	kg S d <sup>-1</sup>	Total rate of nutrient removal
SRT	d	Sludge retention time
t	d or h	Time
t <sub>cycle</sub>	h or d	SBR cycle time length
V	m <sup>3</sup> or L	Volume
$\mathbf{V}_{\mathrm{inf}}$	m <sup>3</sup> or L	Volume of influent
$V_r$	m <sup>3</sup> or L	Working volume of the reactor
$V_{\text{eff}}$	m <sup>3</sup> or L	Volume of effluent
VER	%	Volume exchange ratio

 Table SM1.2. Equations used in calculations and model computations. Definitions and units of symbols are available in Table SM2.1 above.

$$HRT = \frac{V_r}{Q_{inf} N_{cycles}}$$
(eq. 1)

Sludge retention time in a SBR, SRT (d)

$$SRT = \frac{C_x V_r}{(Q_{eff} C_{x,eff} + Q_{purge} C_x + Q_{sample} C_x) N_{cycles}}$$
(eq. 2)

Apparent volumetric rate of nutrient removal, r<sub>s</sub> (kg S d<sup>-1</sup> m<sup>-3</sup>)

$$rs = \frac{(Cs, \inf Qinf - Cs, eff Qeff) Ncycles}{Vr} = \frac{(Cs, \inf - Cs, eff) Q Ncycles}{Vr}$$
(eq. 3)  
$$\equiv \frac{Cs, 0 - Cs, end}{tcycle}$$

with:

$$Cs, 0 = Cs, \inf \frac{Vinf}{Vr} + Cs, end \frac{Vr - Vinf}{Vr} = Cs, \inf VER - Cs, end (1 - VER)$$

$$Cs, inf = Cs, 0 \frac{1}{VER} - Cs, end \frac{1 - VER}{VER}$$

Cs, eff = Cs, end

$$VER = \frac{Vinf}{Vr} = \frac{Veff}{Vr}$$

Percentage of nutrient removal,  $\eta_{s}$  (%)

$$\eta s = \frac{Cs, inf - Cs, eff}{Cs, inf} 100 = \left(1 - \frac{Cs, eff}{Cs, inf}\right) 100$$

$$= \frac{Cs, 0 - Cs, end}{Cs, 0 - Cs, end (1 - VER)} 100$$
(eq. 4)

Total rate of nutrient removal, R<sub>s</sub> (kg S d<sup>-1</sup>)

$$|Rs| = \frac{dMs}{dt} = Vr \frac{dCs}{dt} = Vr rs$$
(eq. 5)

Substrate consumption balance during a batch phase (at constant V<sub>r</sub>)

$$\frac{dC_s}{d_t} = q_s C_x = q_{s,max} \frac{C_s}{C_s + K_s} C_x$$
(eq. 6)

Biomass production balance during a batch phase (at constant V<sub>r</sub>)

$$\frac{dC_x}{d_t} = \mu C_x = (q_s - m_s) Y_{sx,max} C_x =$$

$$(eq. 7)$$

$$(q_{s,max} \frac{C_s}{C_s + K_s} - m_s) Y_{sx,max} C_x$$

Herbert-Pitt equation for substrate allocation for growth and maintenance

Biomass specific rates of substrate consumption,  $q_s$  (kg S d<sup>-1</sup> kg<sup>-1</sup> X) Biomass specific growth rate,  $\mu$  (kg X d<sup>-1</sup> kg<sup>-1</sup> X = d<sup>-1</sup>) Biomass maintenance rate,  $m_s$  (kg S d<sup>-1</sup> kg<sup>-1</sup> X)

$$q_s = \frac{1}{Y_{x/s}}\mu + m_s$$
 or  $\mu = (q_s - m_s) Y_{x/s}$  (eq. 8)

$$\mu_{max} = (q_{s,max} - m_s) Y_{x/s} \approx q_{s,max} Y_{x/s} \quad \text{with } q_{s,max} \gg m_s \qquad (\text{eq. 9})$$

## Supplementary material 2

Correlation between VSS and absorbance measurements

The correlation between the absorbance measurement and the VSS concentration of the PNSB-enriched biomass is described in Figure SM1.1 hereafter. It displays the experimental data of three measurement series with the linear regression line. Such a correlation analysis should checked on regular basis across a long-term experimental period in function of the composition of the biomass. It should also be done for every new experiment, since the content of pigments and of intracellular storage compounds of the PNSB biomass can vary from case to case, based on environmental conditions tested. Absorbance measurements of biomass are mainly applicable with biomass in suspension and at relatively low concentrations. It was mainly valid here for the first 40-h batch period and SBR1. As soon as biomass aggregates, traditional gravimetry measurements via TSS, ISS and VSS as described in standard methods are more effective. This was applied on SBR2 and SBR3. One should nonetheless keep in mind that gravimetry measurements do not differentiate cells and organic intracellular polymers such as PHAs.



Figure SM2.1. Correlation between the absorbance at 660 nm and the VSS concentration of the PNSB-enriched biomass, based on dilution series of mixed liquor taken from the reactor  $(2 \times, 4 \times, 8 \times, 16 \times$  diluted). The linear regression of serie 1 is y = 0.3995x, serie 2 is y = 0.3856 x and serie 3 is y = 0.4124 x. For absorbance measurements, mixed-liquor samples were diluted to fit in the range from 0.2-1.2 absorbance units. On this abosrbance window the correlation to VSS was considered as linear. Absorbance data were mainly accurate measurements for the 40-h batch and SBR1 where the biomass was low concentrated and in suspension. In SBR2 and SBR3, the biomass aggregated and VSS measurements were much more accurate.

## Supplementary material 3

Dynamics of nutrient and biomass concentration and composition



Figure SM3.1. Evolutions of COD (A), ammonium (B) and orthophosphate (C) removal across SBR1 (1st month), SBR2 (2nd-3rd months), SBR3 (4th-5th months). Concentrations at begin (black dots) and end (white dots) of the reaction phases of the SBRs are displayed. While acetate was fully removed, the remaining COD primarily related to EDTA present in the medium (ca. 50 mg COD  $L^{-1}$ ).

## Supplementary material 4 Parameter fit in Aquasim along the 40-h batch and SBRs 1-3



x-axes = Time (h)

Figure SM4.1. Conversion dynamics during the 40-h batch and during the batch reaction phase of selected cycles in SBR1, SBR2, and SBR3 used for the computation of basic kinetic and stoichiometric parameters. Experimental data of biomass growth and COD depletion are displayed with the fitted Aquasim model along the batch reaction phase.

Chapter 4

Syntrophy between fermentative and purple phototrophic bacteria to treat and valorize carbohydrate-based wastewater

This chapter is deposited in a modified version as preprint with contributions of: Marta Cerruti, Guillaume Crosset-Perrotin, Mythili Ananth, Julius Laurens Rombouts and David G. Weissbrodt 2021. Syntrophy between fermentative and purple phototrophic bacteria for carbohydrate-based wastewater treatment. *bioRxiv*, preprint, 2021.05.13.444055. https://doi.org/10.1101/2021.05.13.444055



## Abstract

chemoorganoheterotrophic bacteria (FCB) Fermentative and purple photoorganoheterotrophic bacteria (PPB) are two interesting microbial guilds to process carbohydrate-rich wastewaters. Their interaction has been studied in axenic pure cultures or co-cultures. Little is known about their metabolic interactions in mixed cultures. We aimed to harness the competitive and syntrophic interactions between FCB and PPB in mixed cultures. We studied the effect of reactor regimes (batch or chemostat) and illumination modes (continuous irradiation with infrared light, dark, or light/dark diel cycles) on glucose conversions and the ecology of the process. In batch, FCB (>80% of sequencing read counts) outcompeted PPB, under both dark and continuous infrared light conditions. In chemostat under continuous and alternating irradiance, three FCB populations of Enterobacteriaceae, Lachnospiraceae and Clostridiaceae were enriched (>70%), while Rhodobacteraceae relatives of PPB made 30% of the community. Fermentation products generated from glucose were linked to the dominant FCB. Continuous culturing at a dilution rate of 0.04 h<sup>-1</sup> helped maintain FCB and PPB in syntrophy: FCB first fermented glucose into volatile fatty acids and alcohols, and PPB grew on fermentation products. Direct supply of carboxylates like acetate under infrared light selectively enriched for PPB (60%) independent of reactor regimes. Engineering the association between FCB- and PPB-based can help treat and valorize carbohydrate-based aqueous waste feedstocks.

## 1. Introduction

In nature, microorganisms have evolved to occupy diverse functional niches (Deines et al., 2020). They form metabolic associations to grow under nutrientlimiting conditions (Pearman et al., 2008). Wastewater treatment plants (WWTPs) use microbial communities to remove relatively high concentrations of organics (from 200-500 mg COD L<sup>-1</sup> in municipal wastewater treatment to 30-50 g COD L<sup>-1</sup> in anaerobic digestion) and other nitrogenous and phosphorous nutrients. WWTPs deal with complex compositions of organic matter like cellulose and undigested fibers from food. In agri-food industrial wastewaters, most of organics are derived from simple carbohydrates (*e.g.*, glucose, fructose or xylose) and polymeric sugars such as starch, cellulose and lignocellulosic derivates (Ghosh et al., 2017).

The wastewater treatment sector transitions to develop bioprocesses that combine nutrient removal with bioproduct valorization. Environmental biotechnologies involving mixed cultures are studied to treat and valorize nutrient-rich wastewaters by unravelling the biochemical mechanisms that can drive microbial conversions and products formations (Kleerebezem & Van Loosdrecht, 2007). The efficiency of mixed-culture processes is based on proper selection of microbial guilds needed to metabolize the wastes.

Two guilds across chemo- and photoorganoheterotrophic groups are particularly attractive to process carbohydrate- and carboxylate-rich feedstocks, namely fermentative chemoorganoheterotrophic bacteria (FCB) and purple phototrophic bacteria (PPB).

FCB like *Clostridium*, *Enterobacter*, *Lactobacillus* degrade carbohydrates through fermentation pathways, producing a spectrum of valuable compounds such as carboxylates (*e.g.*, volatile fatty acids – VFAs, lactic acid), ethanol and  $H_2$ . Their microbial niches are harnessed in non-axenic mixed-culture fermentation processes to valorize, *e.g.*, lignocellulosic sugars into targeted fermentation products like lactate or ethanol (Rombouts et al., 2020).

PPB like *Rhodobacter*, *Rhodopseudomonas*, *Rhodospirillum* are versatile microorganisms, involving a diversity of phototrophic and chemotrophic metabolisms depending on environmental and physiological conditions (Table 1). They grow preferentially as photoorganoheterotrophs, harvesting their energy from infrared (IR) light and using a variety of carbon sources from carboxylates to carbohydrates to alcohols (Puyol et al., 2017). PPB can also grow as chemotrophs, they can achieve high biomass yields up to 1 g  $COD_X g^{-1} COD_S$  (Puyol & Batstone, 2017). PPB can thus be exploited to valorize anabolic products like biomass, single-cell proteins used in animal/human feed, or industrially relevant compounds like carotenoids. PPB can also photoferment, coupling glucose degradation to H<sub>2</sub> production.

	Fnorm		Flactron	Ovidized	Carhon	Flactron	Daducad
Metabolism	THEFT			OMINIZO	Cal DUL	FICCH OIL	nonnou
	source		donors	e-donors	source	acceptors	e-acceptors
	(photo-/chemo-)	_	(organo-/litho-)		(hetero-/auto-)		
Under light conditions							
Anoxygenic	Light photons		Reduced	Oxidized	Organic	Endogenous	Biomass
photoorganoheterotrophy			organic	organic, CO <sub>2</sub>		compound, CO <sub>2</sub> fixation	
Anoxygenic photolithoautotrophy	Light photons		Reduced inorganic	Oxidized inorganic	Inorganic (CO <sub>2</sub> )	Endogenous compound,	Biomass
			${ m H}_2^{}$ ${ m H}_2^{}{ m S}$ ${ m Fe}^{2+}$	${ m H_2O}_{ m SO_4^{2-}}$		CO <sub>2</sub> fixation	
	T isht aboto		NO <sup>2</sup>	NO <sup>3</sup>		Fudoromore	Diaman.
Uxygenic nhotolithoautotronhv	Light photons		$H_2O$	$O_2$	Inorganic (CO <sub>2</sub> )	Endogenous	Biomass
						$CO_2$ fixation	
Photofermentation	Light photons		Excess electrons		ı	$\mathrm{H}^{+}$	$\mathrm{H}_2$
(not growth associated)			energized from substrate				
Under dark conditions							
Aerobic-respiring	Chemical r	.edox	Reduced	$CO_2$	Organic	$O_2$	$H_2O$
chemoorganoheterotrophy	reaction by aerobic respirati	on	organic				Biomass
Anaerobic-respiring	Chemical r	edox.	Reduced	$CO_2$	Organic	NO <sup>3-</sup> SO <sup>2-</sup>	N <sub>2</sub> s <sup>0</sup> HS <sup>-</sup> H s
	anaerobic respir	ation	or 5mirs			400	Biomass
Fermentative	Chemical r	edox	Reduced	Pyruvate	Organic	Endogenous	Fermentation
chemoorganoheterotrophy	reaction	by	organic (e.g.,	or $CO_2$		compound	products
	fermentation		fermentable sugars)				( <i>e.g.</i> , acetate, H <sub>2</sub> ).
			ò				Biomass

Table 1. Metabolic versatility of purple phototrophic bacteria (PPB). PPB can grow by combining a diversity of substrates and redox conditions. PPB grow primarily as photoorganoheterotrophs using VFAs as preferential carbon sources. but they are able to grow also on glucose with low growth rates. The metabolisms targeted in this study for the conversion of glucose by combination of fermentation and photoorganoheterotrophy are highlighted in grey.

Combinations of fermentative chemoorganoheterotrophy, and photofermentation photoorganoheterotrophy processes have been investigated to degrade, convert and valorize carbohydrates in different configurations using pure cultures and synthetic co-cultures of FCB and PPB. First, a two-stage process of dark fermentation and photoorganoheterotrophy has been carried in two separate reactors. In the upstream unit, a pure culture of FCB degrade carbohydrates into VFAs that are fed in the downstream unit to produce a PPB biomass in pure culture (Ghimire et al., 2015). Second, in a single-stage dark fermentation process, a pure culture of PPB has been used to convert glucose and other organics into  $H_2$ . This has necessitated longer contact times compared to the two-stage fermentation process and the requirement of a pre-sterilized feed stream and axenic environment (Abo-Hashesh & Hallenbeck, 2012). Third, in single-stage dark- and photo-fermentation process, attempts have been made to co-cultivate defined FCB and PPB to simultaneously degrade carbohydrates and convert the produced VFAs to biomass (Rai & Singh, 2016). Despite the advantages of this synthetic co-culture approach for lower investment and operation costs compared to two-stage fermentation process, it still requires an axenic environment that impacts process economics. Open mixed-culture biotechnologies associating FCB and PPB can become beneficial to treat and valorize carbohydrate-based wastewaters. Compared to traditional industrial biotechnologies, open mixed cultures do not require sterilization of the inflow and tank, therefore substantially reducing the costs (Kleerebezem & Van Loosdrecht, 2007). Based on ecological principles, microorganisms can be selected to specifically target substrate degradation, bioproduct valorization or both.

Little is known about the selection phenomena and community dynamics in open mixed cultures of FCB and PPB, and the impact of substrate composition, environmental conditions, and reactor regimes thereof. Competition and metabolic interactions between these microbial guilds have not been unraveled yet. One can also wonder whether PPB, due to their wide metabolic versatility, could be selectively enriched in a carbohydrate-fed mixed culture to complete the network of metabolisms from glucose fermentation full to photoorganoheterotrophy, or whether FCB would more efficiently ferment glucose, prior to supplying fermentation products for the PPB to grow.

Here, we evaluated the ecological association of PPB and FCB in carbohydratefed mixed cultures. We addressed the selection mechanisms, competitive and syntrophic interactions of the two microbial guilds across reactor regimes (batch, chemostat) and IR light irradiance patterns (continuous light, continuous dark, dark/light alternation) using either glucose or acetate as model carbohydrate and carboxylate substrates, respectively, that harbor the same degree of reduction (4 mol e<sup>-</sup> C-mol<sup>-1</sup>). With these microbial ecology insights, we tested the treatment of a carbohydrate-rich synthetic wastewater in a continuous single-stage mixed-culture process assembling FCB and PPB in syntrophy, by ecological engineering.

## 2. Material and methods

To evaluate microbial competition between PPB and fermenters, two mixedculture bioreactor regimes (batch and continuous culturing), two substrates with the same degree of reduction per carbon mole (acetate and glucose), and three light patterns (continuous illumination, continuous dark, and light / dark alternation in a 16 h / 8 h sequence) were applied. The reactor performances were measured by kinetic and stoichiometry endpoints (rates, yields). Microbial selection and community dynamics were tracked by 16S amplicon sequencing. The experimental design is shown in Table 2.

Reactor regime	Organic substrate	Illumination	Input concentration of organic matter (gCOD L <sup>-1</sup> )
Batch	Acetate	Continuous	10
Batch	Acetate	No (dark)	10
Batch	Glucose	Continuous	10
Batch	Glucose	No (dark)	10
CSTR	Acetate	Continuous	5 / 10
CSTR	Glucose	Continuous	5
CSTR	Acetate	16h light / 8h dark	10
CSTR	Glucose	16h light / 8h dark	10
CSTR	No carbon sources	16h light / 8h dark	-

 Table 2. Combinations of operational conditions tested to address selection, competition, and interaction mechanisms between FCB and PPB.

### 2.1 Composition of cultivation media

Under all conditions, the cultures were provided with a mineral medium composed as in Cerruti et al. (2020). The cultivation medium was buffered at pH 7.0 with 4-(2-hydroxyethyl)-1piperazineethanesulfonic acid (HEPES) at 4 g  $L^{-1}$ . Sodium acetate and glucose were used as carbon sources, in a concentration of 5 or 10 g COD  $L^{-1}$  depending on experimental conditions (Table 2), mimicking concentrations of moderately loaded agri-food wastewaters. Acetate was provided as  $C_2H_3O_2Na\cdot 3H_2O$  (18.28 g  $L^{-1}$ ) and glucose as  $C_6H_{12}O_6\cdot H_2O$  (9.68 g  $L^{-1}$ ).

#### 2.2 Batch operational conditions

Batch tests were first operated to study the competition of PPB and FCB in discontinuous mode. PPB biomass from an in-house enrichment culture grown and maintained in a sequencing batch reactor (SBR) (Cerruti et al., 2020) was used to inoculate 100-ml anaerobic batch bottles. The cultures were flushed with 99% argon gas to maintain anaerobic conditions (Linde, NL, >99% purity). Every batch bottle was duplicated. Half of the batch bottles were exposed to dark. The other half were exposed to IR light (> 700 nm) supplied by a halogen lamp (white light) with a power of 120 W whose wavelengths below 700 nm were filtered out by two Black 962 Infrared Transmitting Perspex Acrylic Sheets (Black Perspex 962, Plasticstocktist, UK). Every batch was incubated in a closed shaker (Certomat® BS1, Sartorius Stedim Biotech, Germany) at  $30\pm1$  °C in the dark and at  $37\pm1$  °C under IR light: these temperature differences originated from the heat emitted from the lamps. The flasks were agitated at 170 rpm. The cultures were fed with 10 g COD L<sup>-1</sup> of either glucose or acetate.

#### 2.3 Continuous culturing conditions

Competition and syntrophy of FCB and PPB under continuous culturing were tested and tuned in a 2.5-L single-wall, continuous-flow stirred-tank reactor (CSTR) with 2 L of working volume connected to a controller system (In-Control and Power units, Applikon, Netherlands). The pH was controlled at 7.0 with NaOH at 0.25 mol L<sup>-1</sup> and HCl 1 mol L<sup>-1</sup>. The temperature was maintained at 30°C with a finger-type heat exchanger connected to a thermostat (WK 500, Lauda, Germany).

Similar to the batches, the reactor was irradiated from two opposite sides with two incandescent halogen lights of 120 W filtered to select for IR wavelengths  $\lambda$  > 700 nm using Black Perspex 962 sheets (Plasticstocktist, UK) to promote PPB growth. An effective incident light intensity of 270 W m<sup>-2</sup> was measured on the reactor wall with a pyranometer (CMP3, Kipp & Zonen, The Netherlands).

The reactor was inoculated with an in-house PPB enrichment, and started under batch mode. After 48 h it was switched to continuous mode. To enrich for PPB, the dilution rate was set to  $0.04 \text{ h}^{-1}$ , *i.e.*, less than half of the biomass-specific

maximum growth rate  $(\mu_{max})$  of the representative PPB genus *Rhodopseudomonas* (0.15 h<sup>-1</sup>) (Cerruti et al., 2020).

Four combinations of energy patterns and carbon sources were then tested in the mixed-culture CSTR: (*i*) continuous light with acetate, (*ii*) continuous light with glucose, (*iii*) 16/8 h light/dark alternation cycles with acetate, and (*iv*) 16/8 h light/dark alternation cycles with glucose. Light/dark alternation was controlled with an automatic switch time controller (GAMMA, The Netherlands).

After 3 months of continuous cultivation, the carbon source feed was stopped, while phosphate and nitrogen sources were maintained for a period of 4 days (*ca.* 4.5 hydraulic retention times – HRTs), in order to test/confirm the hypothesis that PPB can grow on fermentation products produced by FCB.

### 2.4 On-line mass spectrometry analysis of off-gas composition

The off-gas of all CSTR experiments was connected to a mass spectrometer (Thermofisher, Prima BT Benchtop MS) which was used to measure on-line the production rates of  $CO_2$  and  $H_2$  (used in the simulations).

### 2.5 Sampling of mixed liquors for biomass and liquid phase analyses

All mixed liquor samples were collected in 2-mL Eppendorf tubes, centrifuged at 10000 x g for 3 min. The biomass pellet was separated from the supernatant and stored at -20°C until further analysis. The supernatant was filtered with 0.2  $\mu$ m filters (Whatman, US), and stored at -20°C until further analysis. From the batch cultures, the samples were collected every 24 h. From the CSTR, 5 to 9 samples were collected for each condition along the experiments.

### 2.6 Measurement of biomass concentrations

Biomass concentrations were measured over the time course of the batch and continuous reactors by spectrophotometry (Biochrom, Libra S11, US) through absorbance at 660 nm (A<sub>660</sub>). A calibration curve was established to correlate A<sub>660</sub> to the concentration of volatile suspended solids (VSS):  $c(g VSS L^{-1}) = 0.61 A_{660}$  (-). The VSS concentrations were measured by taking samples from the mixed liquor, filtering them using 0.2 µm filters (Whatman, 435 USA), drying the wet filters at 105 °C in a stove for 24 h, prior to incinerating the dried filters at 550 °C in an oven for 2 h, and weighted.

#### 2.7 Wavelength scans of biomass samples to rapidly track PPB selection

Wavelength scan of biomass samples were done using a spectrophotometer (DR3900, Hach, Germany) to evaluate the presence of the typical PPB photopigments (carotenoids at 400-500 nm and bacteriochlorophyll at ca 850 nm). Absorbance profiles were measured at each wavelength between 320 nm and 999 nm.

#### 2.8 Amplicon sequencing analysis of bacterial community compositions

Genomic DNA was extracted using DNeasy Powersoil microbial extraction kits (Qiagen, Hilden, Germany) accordingly to manufacturer's instructions. The DNA extracts were quantified by fluorometry (Qubit; Thermofisher, US), and stored at -20°C (pending analysis) and transferred at -80°C for long-term storage. The compositions of the bacterial communities of the mixed liquors were characterized from the DNA extracts by V3-V4 16S rRNA gene-based amplicon sequencing as detailed in Cerruti et al. (2020). Aliquots of the DNA extracts were submitted to Novogene (UK) for library preparation and amplicon sequencing. The fastq files of the sequencing datasets provided from Novogene were analysed with the QIIME2 pipeline (Bolyen et al., 2019), using the Greengenes database. The sequences were deposited in the online database NCBI under the Bioproject code ID PRJNA759396.

## 2.9 Chromatography analysis of substrate conversions and fermentation products

The depletion of substrates (acetate and glucose) and the formation of fermentation products (succinate, propionate, formate, lactate, acetate, butyrate) were monitored using high-performance liquid chromatography (HPLC) (Waters, 2707, NL) equipped with an Aminex HPX-87H separation column (BioRad, USA) maintained at  $60^{\circ}$ C. The H<sub>3</sub>PO<sub>4</sub> (1.5 mmol L<sup>-1</sup>) eluent was supplied at a flowrate of 0.6 mL min<sup>-1</sup>. The compounds were detected by refraction index (Waters 2414, US) and UV (210 nm, Waters 2489, US). Ethanol was quantified through gas-chromatography as in Rombouts et al. (2019).

#### 2.10 Solver for pathway utilization prediction

Based on the observed yields, a mathematical simulation was used to evaluate the contribution of the fermentative and phototrophic pathways to the carbon metabolism. The model was constructed in Microsoft Excel 16.48 minimizing the residual sum of squares (SSQ) error between modelled and measured yields:

$$\min(SSQ) = \min\left(\sqrt{\sum_{i=1}^{n} \frac{(\hat{y_i} - y_i)^2}{n}}\right)$$
(1)

where  $\hat{y}_i$  is the modelled yield value and  $y_i$  is the measured yield value, and *n* is the amount of metabolic compounds that are considered.

The fermentation products yields (observed yields) were calculated as

$$y_i = Y_{p/s} = \frac{q_p}{q_s} \tag{2}$$

With  $q_p$  and  $q_s$  the biomass specific rates.

The solver estimated the contribution of the singular pathways (Table 3) to the overall metabolism as fractions. The modelled yields were estimated by summing the products between the pathway fractions and their stoichiometric coefficients. The process was automatically reiterated to minimize the SSQ between the observed and the estimated yields through the Generalized Reduced Gradient (GRG) Nonlinear solver available in Excel. The pathways that were used in the model varied per enrichment, depending on the microbial community structure, assuming only the activity of relevant catabolic and anabolic pathways. To simulate the carbon metabolism in batch enrichments the equations 1.1-1.9 were used. To simulate the pathways used in the CSTRs, reactions 1.10-1.11 were also added (Table 3). The model output indicated, as percentage, the contribution of the individual pathways to the total carbon, oxygen, hydrogen and electrons metabolism. The flux distribution of the different pathways predicted from the model is represented a semi-quantitative way in Figure 4.

		Glucos	Ethanol	Acetate	Propionate	Lactate	Butyrate	Succinate	Formate	Hydrogen	Carbon dioxide	Biomass	Biomass	Protons	Water
Reactions	Pathways	e C <sub>6</sub> H <sub>12</sub>	C <sub>2</sub> H <sub>6</sub> O	$C_2H_3O_2$	$C_3H_5O_2$	C <sub>3</sub> H <sub>5</sub> O	$C_4H_7O_2$	$C_4H_4O_4$	$CH_2O_2$	$\mathrm{H}_2$	$CO_2$	FCB CH <sub>1.8</sub> O <sub>0.</sub>	PPB	$\mathrm{H}^{\dagger}$	$\rm H_2O$
		06				3						5			
1.1)	Glucose $\rightarrow$ Biomass + CO <sub>2</sub>	-1.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	-0.10	2.00	4.00	0.00	5.00	0.00
1.2)	Glucose $\rightarrow$ Ethanol + CO <sub>2</sub>	-1.00	2.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	2.00	0.00	0.00	0.00	0.00
1.3)	Glucose + $\rightarrow$ Acetate + Ethanol + CO <sub>2</sub>	-1.00	1.00	1.00	0.00	00.00	0.00	0.00	2.00	0.00	0.00	0.00	0.00	1.00	-1.00
1.4)	Glucose → Butyrate + CO <sub>2</sub>	-1.00	0.00	0.00	0.00	0.00	1.20	00.00	0.00	0.00	1.20	0.00	0.00	1.20	1.20
1.5)	Glucose → Succinate	-1.00	0.00	00.0	0.00	0.00	0.00	1.71	0.00	0.00	-0.86	0.00	0.00	3.43	0.86
1.6)	Glucose → Lactate	-1.00	0.00	0.00	0.00	2.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	2.00	0.00
1.7)	Lactate → Acetate + Propionate + CO <sub>2</sub>	0.00	0.00	1.00	2.00	-3.00	0.00	0.00	00.0	0.00	1.00	0.00	00.0	0.00	1.00
1.8)	Formate $\rightarrow$ CO <sub>2</sub> + H <sub>2</sub>	0.00	0.00	0.00	0.00	0.00	0.00	0.00	-1.00	1.00	1.00	0.00	0.00	0.00	0.00
1.9)	Acetate → Biomass + CO <sub>2</sub>	0.00	0.00	-1.00	0.00	0.00	0.00	0.00	0.00	-0.02	1.00	0.00	1.00	2.00	-0.38
1.1)	$CO_2 + H_2O \rightarrow$ Biomass + $H_2$	0.00	0.00	00.0	0.00	0.00	0.00	0.00	00.0	0.00	-1.00	1.00	1,00	-5.00	1.60

Table 3. Stoichiometric matrix of the putative reactions included in the numerical model of metabolic reactions along FCB and PPB pathways. The reactions 1.1-1.9 were used to evaluate the contribution of each pathway under batch conditions. The reactions 1.10 and 1.11 were included in the simulation of the CSTRs.

## 3. Results & Discussion

The batch and continuous reactor experiments revealed key insights into the microbial community assembly principles that govern the selection, competition and interaction of FCB and PPB in open mixed cultures in function of organic substrates, light patterns, and reactor regimes.

## 3.1 Selection and competition of PPB and FCB in batch cultures

## 3.1.1 In mixed-culture batches, PPB were selectively enriched on acetate and IR light

In batches, the acetate-fed cultures grew only when subjected to light, reaching a biomass concentration of 1 g VSS L<sup>-1</sup> at a biomass specific growth rate ( $\mu$ ) of 0.0411 ± 0.0007 h<sup>-1</sup> and with a yield of biomass on substrate (Y<sub>X/S</sub>) of 0.60 ± 0.09 C-mmol<sub>x</sub> C-mmol<sub>s</sub><sup>-1</sup>. Little growth was reported in the six days of cultivation for the acetate cultures in the dark ( $\mu = 0.006$  h<sup>-1</sup>).

The inocula of the batches originated from a PPB in-house enrichment culture operated and maintained as SBR in parallel to the experiments. The inoculum seeded in the acetate-fed batches presented a relative abundance above 60% (*vs.* total amplicon sequencing read counts) of the purple non-sulfur genus *Rhodopseudomonas*. In the acetate-fed batches, *Rhodopseudomonas* sp. reached as high as 80% of the total community after 6 days of cultivation under IR light. Despite the low growth, chemoorganoheterotrophs of the family of the *Moraxellaceae* showed the highest relative abundance (80%) in the little biomass obtained on acetate in the dark (Figure 1A).





Figure 1. A: Microbial community composition in batches. PPB are dominant at the beginning of the cultivation (ACE INIT and GLU INIT). After 6 days of cultivation under acetate feed, PPB were dominant under light conditions (ACE LIGHT). Under acetate-dark conditions there was little grow from the genus Moraxellaceae. Under glucose feed, independently of the light conditions, the family Enterobacteriaceae was dominant. B: pH evolution in the glucose-fed batches. Both under light and under dark conditions the pH dropped to 4 in the first 2 days of incubation.

Acetate is one of the preferred substrates for PPB (Blankenship et al., 1995). It is assimilated into biomass by many PPB species, through the tricarboxylic acids cycle (Pechter et al., 2016). Acetate and IR light selected for *Rhodopseudomonas*, belonging to the *Xanthobacteraceae* family, under acetate and light conditions. Among the PPB, *Rhodopseudomonas* has the highest biomass-specific maximum growth rate ( $\mu_{max}$ ) on acetate (0.15 h<sup>-1</sup>) (Cerruti et al., 2020) and therefore was predominant at the end of the batch cultivation. The growth rate in the acetate-fed cultures in the dark was 7 times lower than under light; the biomass concentration after 6 days was 270 times lower in the dark than under light. In the dark, acetate can only be assimilated by PPB in presence of suitable electron acceptors (*e.g.*, S<sup>0</sup> or SO<sub>4</sub><sup>2-</sup>) in order to produce ATP for growth (Madigan et al., 1982). In the batches, little external electron acceptor was present in the medium (3 mmol SO<sub>4</sub><sup>2-</sup> L<sup>-1</sup>).

## 3.1.2 In mixed-culture batches, FCB outcompeted PPB on glucose, independent of light supply

In the glucose-fed batches, higher biomass concentration (1 g VSS L<sup>-1</sup>) and growth rate ( $\mu$  of 0.070 ± 0.001 h<sup>-1</sup>) were reached in the dark than in the light (0.4 g VSS L<sup>-1</sup>; 0.020 ± 0.002 h<sup>-1</sup>).

The inoculum of the glucose-fed batches was dominated by the purple sulfur genus *Thiobaca*, due to an unexpected variation of the microbial community composition in the parent reactor. In the glucose-based batches, the genus of *Enterobacter* was predominant (>90%) both under light and dark (Figure 1A).

The *Enterobacteriaceae* family is widely predominant in anaerobic environments in the presence of non-limiting concentration of monomeric carbohydrates, such as glucose (Wijffels et al., 2003). Rombouts et al. (2019) reported an enrichment of 75% of the genus *Enterobacter* in a glucose-fed SBR, in nutrient, temperature and pH conditions similar to the one here presented.

pH was not controlled in batch cultivations. During glucose fermentations, pH dropped to 4 within 48 h, despite the presence of the HEPES buffer (Figure 1B).

### 3.1.3 Ethanol was the main fermentation product in glucose-fed batches

In irradiated acetate-fed batches, acetate was fully depleted after 6 days of incubation.

In glucose-fed batches, the biomass yields on glucose were  $0.052 \pm 0.002$  C-mmol<sub>x</sub> C-mmol<sub>s</sub><sup>-1</sup> under light and  $0.134 \pm 0.011$  C-mmol<sub>x</sub> C-mmol<sub>s</sub><sup>-1</sup> in the dark (Figure 2).

After 6 days of dark batch fermentations, ethanol was the main fermentation product with a yield of  $0.20 \pm 0.10 \text{ C-mmol}_{et} \text{ C-mmol}_{s}^{-1}$ , followed by succinate  $(0.07 \pm 0.03 \text{ C-mmol}_{succ} \text{ C-mmol}_{s}^{-1})$  and acetate  $(0.03 \pm 0.01 \text{ C-mmol}_{ace} \text{ C-mmol}_{s}^{-1})$ .

In the irradiated glucose-fed batches, 30% of the initial glucose was still present in the reaction broth after 6 days, and carbon balances could not be properly closed (>90% recovery). Ethanol was the main fermentation product  $(0.10 \pm 0.07 \text{ C-mmol}_{s}^{-1})$ , followed by lactate  $(0.05 \text{ C-mmol}_{lac} \text{ C-mmol}_{s}^{-1})$  and succinate  $(0.04 \pm 0.02 \text{ C-mmol}_{succ} \text{ C-mmol}_{s}^{-1})$ .



Figure 2. Yields of the glucose-fed cultures. Under batch conditions, the main fermentation products were ethanol and succinate. Under light conditions, glucose was not fully depleted. Under CSTR conditions, the main fermentation products were ethanol, formate and acetate.

A solver enabled to evaluate the theoretical contribution of 8 putative glucose fermentation pathways to the overall carbon and electron balances. In batches, the phototrophic pathways were not included in the simulation, since the relative abundance of PPB was very low (~1%), being outcompeted by FCB with glucose (Figure 1A). The predicted contribution of the individual pathways to the overall yields of the batches under light and dark were comparable. In batches, the dominant metabolic pathway was identified as homo-ethanol production from glucose (Table 3, Eq. 1.2). This catabolism is associated with the fermentative *Enterobacteriaceae* (Jang et al., 2017).

Wavelength scans are commonly used as an indirect measure to rapidly check for the presence/selection of PPB in the cultures, since photopigments absorb photons with specific wavelengths. Absorbance peaks around 400-500 nm represent the carotenoids series, and the peaks around 850 nm the bacteriochlorophyll a (bchl a) (Stomp et al., 2007). In the acetate-based irradiated batches, the aforementioned peaks were clearly detected at 450 nm

for carotenoids and 800-850 nm for bchl a. In the acetate-fed dark batches, almost no growth was reported within the 6 days of incubation but peak for photopigments was detected at 800 and 850 nm. No photopigment characteristic peaks were detected in the spectra of the biomasses cultivated in both the dark and irradiated glucose-fed batches, (Figure 3A/B).

Overall, in anaerobic batches, PPB are predominantly selected with acetate and IR light. FCB outcompete PPB on glucose independent of light or dark conditions.



Figure 3. Wavelength scan from 300 to 1000 nm. Absorbance units normalized for the biomass concentration (as A.U. at 660nm). A: acetate-fed batches. The typical PPB peaks (at ca 850 nm) were clearly visible under light conditions. B: glucose-fed batches. No peak was present. C: continuous light glucose-fed CSTR. The presence of PPB was confirmed by the presence of their typical peaks. D: dark / light glucose-fed CSTR. The presence of PPB was confirmed by the presence of their typical peaks.

### 3.1.4 Fermentative Enterobacteriaceae were enriched in batches because of higher growth rates

In batch,  $\mu_{max}$  is a key factor driving selection mechanisms in substrate-rich environments (Winkler et al., 2017), if no significant intermediate storage of substrate is displayed. FCB populations of the genus *Enterobacter* present a  $\mu_{max}$  of 0.45-0.80 h<sup>-1</sup> in mineral medium conditions with moderate sugar concentrations (<10 g<sub>COD</sub> L<sup>-1</sup>) (Rombouts et al., 2019). When fermenting

carbohydrates like glucose in pure culture, *Rhodopseudomonas capsulata* grows at a  $\mu_{max}$  of 0.014 h<sup>-1</sup> (Conrad & Schlegel, 1978). In batches, where the substrate is not limiting across most of the conversion time, the organisms with higher substrate uptakes rates get selected above organisms with high biomass yields (Rombouts et al 2019). According to the Herbert-Pirt relation, when the substrate uptake is directly coupled with growth, the organisms with the highest growth rate dominate. Using these studies, the genus *Enterobacter* is estimated to exhibit a  $\mu_{max}$  about 32 times higher than *Rhodopseudomonas* on glucose, making FCB successful to outcompete PPB for glucose in batch.

In both irradiated and dark glucose-fed batches, populations of the family of Enterobacteriaceae were enriched (>90%). Enterobacteriaceae relatives ferment sugars through either the Embden-Meyerhof pathway or the pentose phosphate pathway, with a net production of ethanol (Jang et al., 2017). Ethanol can be used as substrate for growth by PPB, but with a strict pH regulation at 7.0 (Inui et al., 1995). Because of the production of lactate (pKa = 3.8) – a major byproduct of the glucose fermentation pathway to ethanol (Converti & Perego, 2002) – and insufficient pH buffering, the pH dropped to 4.0 within the first 2 days of glucose fermentation (Figure 1B). FCB can still grow at low pH such as below 5.5 (Tsuii et al., 1982). PPB grow instead in a pH range between 6.5 and 9.0, with an optimum at 7.0 (Pfennig, 1977). The low pH (pH = 4.0) resulting in the glucose-fed batches may have prevented PPB to grow on alcohols and VFAs produced by *Enterobacter*, explaining their low relative abundance ( $\sim 1\%$ ). We postulate that with a strict pH control of the batch cultures to 7.0, PPB should be able to grow on fermentation products of FCB. The genus Enterobacter can grow in a range of temperatures between 20 and 45°C (Gill & Suisted, 1978), with an optimum growth at 40°C (Tanisho, 1998). The temperature differences reported here for the batches incubated under dark (30°C) and light (37°C) (*i.e.*, ca, 7°C difference, resulting from the heat emitted by the lamps) should not have substantially affected its specific growth rates. On the other hand, light irradiance has been used as a method to prevent the proliferation of fermentative organisms (Gwynne & Gallagher, 2018). Since the illuminated cultures were supplied with IR light, although less energetic than

white light, we postulate an inhibitory action on FCB, possibly explaining the 3.5-times lower growth rates measured under light than in the dark. The effective inhibition mode of action of photons on cellular and molecular level remains to be elucidated by future works.

## *3.2 Unravelling the selection and syntrophy of FCB and PPB in continuous cultures*

Under continuous-flow regime, four different light and substrate conditions were imposed, namely: (i) acetate under light; (ii) acetate under dark/light cycles; (iii) glucose under light; and (iv) glucose under dark/light cycles. The pH was controlled at 7.0 to favor the enrichment of PPB.

*3.2.1 Ethanol was the main fermentation product in the glucose-fed CSTR as well* 

In glucose-fed CSTRs, the biomass yield over glucose was  $0.22 \pm 0.02$  C-mmol<sub>x</sub> C-mmol<sub>s</sub><sup>-1</sup> under continuous illumination and  $0.45 \pm 0.12$  C-mmol<sub>x</sub> C-mmol<sub>s</sub><sup>-1</sup> under dark/light cycles. Ethanol was the main fermentation product (0.18 ± 0.06 under continuous light and 0.11 ± 0.10 C-mmol<sub>et</sub> C-mmol<sub>s</sub><sup>-1</sup> under dark/light cycles). In the continuously irradiated CSTR, acetate (0.21 ± 0.15 C-mmol<sub>ace</sub> C-mmol<sub>s</sub><sup>-1</sup>) and butyrate (0.06 ± 0.05 C-mmol<sub>but</sub> C-mmol<sub>s</sub><sup>-1</sup>) were also produced. In the dark/light CSTR, formate (0.11 ± 0.01 C-mmol<sub>form</sub> C-mmol<sub>s</sub><sup>-1</sup>) and butyrate (0.10 ± 0.01 C-mmol<sub>but</sub> C-mmol<sub>s</sub><sup>-1</sup>) were present in the fermentation product spectrum (Figure 2).

In CSTR, assimilation of acetate by photoorganohetrotrophy and  $CO_2$  fixation (photosynthesis) by photolithoautotrophy were added to the solver matrix to evaluate the contribution of PPB to the total carbon and electron balances. Under all CSTR conditions, the major fermentation pathway involved fermentation of glucose through acetyl-CoA (Figure 4), up to 83% under continuous illumination and 55% under dark/light cycles. In the continuously irradiated CSTR, the solver estimated a high contribution of the photolithoautotrophic pathway enabling PPB to recycle  $CO_2$  as extra carbon source, and of the photoheterotrophic pathway under dark/light cycles.



Figure 4. Pathways involved under the different reactor regimes. The thickness of the arrows represents the predicted yield toward a specific metabolic route, based on the calculations of the solver. The reactions with the blue background are performed by FCB.

The reaction with the purple background is performed by both FCB and PPB. The reactions with the pink background are performed by PPB. A: batch conditions with continuous illumination. B: batch conditions in the dark. C: CSTR under continuous illumination. D: CSTR conditions under light / dark cycles. Under batch conditions the pathways used were mainly directed toward biomass and ethanol production. Under CSTR ethanol was the main fermentation product, but the production of propionate, succinate and lactate was predicted. The phototrophic pathways were not included for the batch conditions but were included in the carbon and electron balance of the CSTRs. The conversion of lactate and acetate to butyrate (1.9 of Table 3) was not included in the diagram as the solver predicted 0% flux through it under all conditions.

#### 3.2.2 PPB were enriched in the acetate-fed CSTR

In the continuously illuminated CSTR fed with acetate, PPB formed the dominant guild ( $68 \pm 21 \%$ ) (Figure 5). The genus *Rhodopseudomonas* was predominant at the beginning of the CSTR (65%). Its relative abundance decreased to 40% after one week of cultivation (8.5 HRTs), while the genus *Rhodobacter* got concomitantly enriched (from 13 to 23%). *Rhodobacter* was further enriched once the dark/light cycles were applied, reaching 65% after 15.6 HRTs, while *Rhodopseudomonas* remained stable below 10%.

### 3.2.3 FCB were enriched in the glucose-fed CSTR

In the CSTR, FCB accounted above 50% under glucose feed, regardless of the light condition. In CSTR, dark/light cycles impacted the selection of FCB. Under continuous illumination, the family of *Lachnospiraceae* was predominant (around 30 %, with a peak at 75%), followed by the family of *Enterobacteriaceae* (15%) (Figure 5). Under dark/light cycles, *Enterobacteriaceae* (32  $\pm$  17%) and *Clostridiaceae* (18  $\pm$  15%) relatives were predominant.

*Rhodobacter* was the only PPB to significantly persist in the glucose-fed CSTR (19.5  $\pm$  14 %) (Figure 5) independently of the illumination regime. The persistence of PPB in the glucose-fed CSTRs was further proven by the wavelength scan. Peaks for carotenoids (400-450 nm) and bacteriochlorphyll a (800 and 850 nm) photopigments were identified for both the continuous illumination and dark/light cycles (Figure 3C/D).



Figure 5. Relative abundance of five representative families under CSTR conditions based on 16s rRNA gene amplicon sequencing analysis.. Under acetate conditions, PPB were dominant community, with a shift from Rhodopseudomonas under continuous illumination to Rhodobacter under light / dark conditions. Under glucose feed, the FCB were dominant, especially the genera Enterobacteriaceae and Lachnospiraceae.

## 3.2.4 The microbial composition of the fermentative guild defined the spectrum of fermentation products

Clostridiaceae, Enterobacteriaceae and Lachnospiraceae relatives are all able to ferment glucose. They compete for glucose through different fermentation pathways (Grimmler et al., 2011; Horiuchi et al., 2002). Lachnospiraceae and *Clostridiaceae* are phylogenetically and morphologically heterogeneous families belonging to the phylum of Firmicutes (De Vos et al., 2005). Glucose fermentation through acetyl-CoA is typical of Clostridiaceae (Aristilde et al., 2015). In human gut microbiota, they contribute to sugar fermentation to lactate and short-chain fatty acids production (Venegas et al., 2019). Butyrate was produced only in the CSTR under continuous illumination  $(0.06 \pm 0.05 \text{ C-mmol})$ C-mmol<sub>s</sub><sup>-1</sup>). The *Lachnospiraceae* and *Clostridiaceae* families ferment glucose into primarily butyrate, acetate and ethanol (Rombouts et al., 2019b; Temudo et al., 2008). Butyrate is produced through the butyril-CoA dehydrogenase gene, responsible for the conversion of crotonyl-CoA to butyryl-CoA and the butyryl-CoA:acetyl-CoA transferase (Venegas et al., 2019). According to the NCBI database (NCBI, 2021), the gene encoding for this enzyme is present in the phylum of Firmicutes that comprises Clostridiaceae and Lachnospiraceae relatives, but not in the proteobacterial family of the *Enterobacteriaceae*, explaining the absence of butyrate in the glucose-fed batches.

In the dark/light CSTR, the relatively high production of propionate (0.10  $\pm$  0.01 C-mmol C-mmol<sub>S</sub><sup>-1</sup>, *i.e.*, 17 times higher than under continuous

illumination) can be linked to the abundance of *Clostridiaceae*, which are propionate producers (Johns, 1952). *Clostridium* species can present a  $\mu_{max}$  of 0.25 h<sup>-1</sup> (Gomez-Flores et al., 2015) with primary fermentation products including ethanol, butyrate, acetate and propionate (Lamed et al., 1988). *Clostridium beijerincki* decreases its H<sub>2</sub> production when exposed to light intensities above 200 W m<sup>-2</sup>, with a shift from a preferential production of butyric acid to acetic acid (Zagrodnik & Laniecki, 2016). Continuous illumination in the CSTR might have inhibited *Clostridiaceae* and selected for other *Firmicutes* like *Lachnospiraceae*.

Formate and propionate production was 10 times higher under dark/light conditions compared to continuous illumination. Acetate production was instead 12 times lower under dark/light cycles compared to continuous illumination (Figure 2). This may link to a higher relative abundance of *Enterobacteriaceae* relatives under dark/light cycles (31%) compared to the continuous illumination (18%) (Figure 5). *Enterobacteriaceae* present a  $\mu_{max}$  between 0.5-1 h<sup>-1</sup> (Khanna et al., 2012; Martens et al., 1999) and are known to ferment sugars to primarily ethanol, with lactate and acetate as major byproducts (Converti & Perego, 2002).

### 3.2.5 PPB metabolized the fermentation products released by FCB

After 3 months of CSTR operation, we tested the hypothesis that PPB grow on the fermentation products of FCB. Glucose supply in the feed was stopped while nitrogen and phosphate were still provided over 4 days (*i.e.*, 4.5 HRTs). Along the 4 days of operation, the relative abundance of the PPB shifted from 10 to 50% (Figure 6A). The biomass concentration sharply decreased already 2 h after stopping the glucose feed, from 80 C-mmol<sub>X</sub> L<sup>-1</sup> and stabilizing at a concentration of  $15.7 \pm 2$  C-mmol<sub>X</sub> L<sup>-1</sup>. Butyrate, ethanol, formate, lactate, propionate and succinate were washed out of the system at the imposed dilution rate (0.04 h<sup>-1</sup>). Acetate was depleted in 50 h, *i.e.*, 4 times faster than the theoretical washout rate (Figure 6B). In absence of glucose as carbon source, the highly active fermentative organisms were not growing. The increase in the PPB relative abundance proved their interaction with FCB, showing that PPB utilize VFAs as preferred substrate and can selectively grow on FCB fermentation products.



Figure 6. A: Microbial composition after the stop to the glucose feed. Over time, the FCB relative abundance decreased over time, while the Rhodobacteraceae increased. B: Fermentation products concentration after the glucose feed-stop. After 55 h all the fermentation products were depleted.

## 3.3 Metabolic strategies for tuning the microbial assembly of FCB and PPB

Microorganisms use a combination of adaptive reactions to coexist with other organisms. Based on the adaptation techniques, two major types of organisms can be identified (Andrews & Harris, 1986), namely r- and K-strategists. The r-strategists show high growth and conversion rates, in low-populated and resources-rich environments. K-strategists thrive in highly populated environments, where the resources are limited. They exhibit lower growth rates but high substrate conversion yields.

Under the different reactor regimes, the microbial communities followed this postulate. *Enterobacteriaceae* were predominant under non-limiting conditions (batches) and can be classified as r-organisms. *Clostridiaceae* and *Lachnospiraceae* showed high growth rates in nutrient-limiting environments (chemostat) and can be classified as K-strategists. Among the PPB guild, *Xanthobacteraceae* like *Rhodopseuodmonas* showed a high-rate behavior, dominating in substrate-rich environments. *Rhodobacteraceae* showed high efficiency typical of K-organisms, being able to survive to nutrient-limited environments, as the CSTRs (Table 4).

Substrates	Maximum growth r	ate $\mu_{max}$ (h <sup>-1</sup> )			
	Fermentative cheme (FCB)	oorganoheterotroph	uic bacteria	Purple photoorgan bacteria (PPB)	oheterotrophic
	Enterobacteriaceae	Lachnospiraceae	Clostridiaceae	Rhodopseudomonas	Rhodobacter
Glucose	0.3 - 1	0.3	0.25	0.014	0.1
	(Hacona of al	(Soto-Martin et	(Gomez-	(Conrad & Schlegel,	(Imam et al.,
	(11100111 et al.	al., 2020)	Flores et al.,	1978)	2011)
	2004; Khanna		2015)		
	et al., 2012)				
Acetate				0.15	0.05
				(Cerruti et al., 2020)	(Kopf &
					Newman,
					2012)
Selection	r-strategist	K-strategist	K-strategist	r-strategist	K-strategist
strategy	(on glucose)	(on glucose)	(on glucose)	(on acetate)	(on acetate)

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Table 4. Reportedmaximum growth ratesof the main FCB andPPB in the systemsunder glucose andacetate feed andadaptation strategies.

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Kinetic parameters define the microbial selection mechanisms in CSTRs (Kuenen, 2019). In the CSTRs, the dilution rate set to retain PPB in the system was relatively low (0.04 h<sup>-1</sup>) compared to the maximum growth rates of the fermentative organisms (0.5-1 h<sup>-1</sup>). In a continuous culture, the substrate (here glucose) is the limiting compound, and the organisms with higher affinity for it get predominant (Andrews & Harris, 1986). Within the guild of FCB, *Clostridiaceae* and *Lachnospiaraceae* are expected to harbor a lower affinity constant (K<sub>s</sub>), outcompeting *Enterobacteriaceae*. The controlled

pH (7.0) and appropriate dilution rate (0.04  $h^{-1}$ ) sustained the persistence of PPB. Notably, *Rhodobacteraceae* were enriched in the glucose-fed CSTRs regardless the illumination conditions. We propose here a syntrophic association of FCB and PPB for glucose conversion toward CO<sub>2</sub> and biomass (**Figure 4**). Glucose was first converted into acetate, ethanol, formate and lactate, and lactate was further transformed into acetate and propionate. All the fermentation products can be potentially used by PPB for growth.

### 3.3.1 Syntrophic associations between FCB and PPB

The syntrophy between FCB and PPB is not only important at an ecological level but can also be exploited for wastewater treatment. FCB are used to treat food and agricultural wastes, to degrade sugars and produce biofuels as bioethanol and hydrogen gas, or carboxylates like VFAs (L. Li et al., 2015; Thapa et al., 2019). PPB, thanks to high biomass yields (Alloul et al., 2018), harbor a major industrial potential for the production of antioxidants and single-cell proteins.

## 3.3.2 Advantages of single-sludge and two-sludge processes for the treatment and valorization of carbohydrate-based aqueous feedstocks by FCB and PPB

The single-step bioprocess implemented here by combining the metabolisms of FCB and PPB in one single sludge can help to simultaneously treat carbohydrate-rich wastewaters, factually removing the organic pollutants, and enriching organisms with high industrial potential (*i.e.* for single cell proteins and photopigments production). An integrated single-stage process combining fermentation and photoorganoheterotrophy allows to treat carbohydrates in only one tank, therefore minimizing capital and operating expenditures. The illumination conditions did not affect the syntrophy between FCB and PPB. A discontinuous illumination can further reduce the operational costs compared to a continuously irradiated system (Qin et al., 2018). Understanding interactions between the FCB and PPB will help design efficient processes for carbohydrates-based wastewater treatment and valorization, in example toward single cell proteins production. If the aim is to maximize and valorize the PPB biomass, a two-sludge process can be efficient to first ferment carbohydrates in a first tank selecting for FCB and supply the fermentation products (whose spectrum can be engineered by mastering the selection of fermentative microbial niches) to a subsequent photoorganoheterotrophic compartment to select for, e.g., an edible PPB biomass, both tanks being operated as selective open mixed cultures.

## 4. Conclusions

We elucidated the competitive and syntrophic phenomena between FCB and PPB in mixed culture. FCB degrade glucose into VFAs, lactate and ethanol. Acetate was preferentially assimilated into biomass by PPB. Combination of acetate and IR light supply selectively enriches for PPB. The operational conditions are crucial to establish an interaction between FCB and PPB guilds. FCB are more efficient, fast-growing fermenters than the metabolically versatile PPB. In glucose-fed batches, FCB outcompeted PPB under the action of both faster growth rate and pH drop. In glucose-fed CSTRs, PPB got enriched up to 30% in syntrophic association with FCB. An appropriate dilution rate  $(0.04 \text{ h}^{-1})$  and pH regulation to 7.0 enabled to enrich PPB in glucose-rich environments by enabling their metabolic coupling with FCB.

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# Supplementary material 1

#### Off-gas analysis of CO<sub>2</sub> production rate

The  $CO_2$  partial pressure in the off-gas of the CSTR was monitored to evaluate the growth state of the biomass. For PPB,  $CO_2$  production is a side product of acetate assimilation (Sadaie et al., 1997).

In acetate-fed CSTRs,  $CO_2$  production was constant under continuous illumination (0.0111 ± 0.0004 C-mmol<sub>CO2</sub> C-mmol<sub>s</sub><sup>-1</sup>) (Figure S1/A). When dark / light cycles were applied, the CO<sub>2</sub> production decreased to 0 C-mmol h<sup>-1</sup> during the 8 h dark periods and increased again to 0.3 C-mmol h<sup>-1</sup> during the 16 h light phases (Figure S1/B). The CO<sub>2</sub> production during the light phases indicated that acetate was continuously assimilated, whereas the sudden decrease during the dark periods (1/3 less within the first hour of dark) indicated that the metabolic activity was switched off. An oscillation in the CO<sub>2</sub> production rate was already observed in *Rhodopseudomonas* pure cultures under CSTR (Cerruti et al., 2020). In both cases, when the substrate was provided in excess, PPB grew with a growth rate close to  $\mu_{max}$  (0.1 h<sup>-1</sup>) during the light periods. In the dark, PPB halted their metabolic activity.

Both under continuous illumination and dark/light cycles, a change in  $CO_2$  production was observed 24-48 h after the switch from acetate feed to glucose feed. In the CSTR under continuous illumination, a peak of  $CO_2$  production reached 1.2 C-mmol h<sup>-1</sup> 24 h (ca 1 HRT) after the switch to the glucose feed and stabilized at 0.6 C-mmol h<sup>-1</sup> 2 days later (Figure S1/A). Under light / dark cycles a peak of  $CO_2$  production (up to 5.6 C-mmol h<sup>-1</sup>) was observed after two full cycles (3.8 HRTs from the glucose switch). After 8 HRTs from the switch to the glucose feed, the  $CO_2$  production rate presented oscillations between light and dark periods. The  $CO_2$  production rate increased of 0.5 C-mmol h<sup>-1</sup> at the beginning of the light phase and it decreased of 0.4 C-mmol h<sup>-1</sup> in every dark period (Figure S1/B). This behavior was observed for 8 HRTs, until the pH dropped to 2 due to a technical failure.



Figure S-1. A: CO<sub>2</sub> production rates under acetate-fed and glucose-fed CSTR continuous light conditions. An increase of ca 1 mmol h<sup>-1</sup> was observed 3 days after the switch to glucose feed, and then stabilized. B: CO<sub>2</sub> production rates under acetate continuous light, acetate dark / light and glucose dark / light conditions. Under acetate continuous illumination CO<sub>2</sub> was produced at constant rate. Under acetate dark / light CO<sub>2</sub> was produced only during the light periods. Under glucose dark / light conditions, a peak in production rate was observed after 3 days from the switch to the glucose-feed. The production rates increased during dark periods and decreased during light periods.

## Supplementary material 2



Detailed amplicon sequencing analysis of the four CSTR conditions

Figure SI-2: Relative abundance of the microbial genera analysed. During the acetate-fed CSTR under continuous illumination, Rhodopseudomonas was the dominant genus. Once the light/dark cycles were applied, the community shifted toward toward the genus Rhodobacter. Under glucose-fed CSTR, the FCB of the genera Lachnoclostridium, Clostridium and Enterobacter were dominant.

# Supplementary material 3



### Product yields in glucose-fed batches and CSTRS

Figure SI-3: Observed and calculated yields of product formations on glucose (C-mol / C-mol) for the mixed-culture batches and CSTRs. The calculated yields were obtained by fitting the metabolic model built based on the reactions in Table 3 by minimizing the sum of squares according to Equation 1 provided in the article.

Chapter 5

# Effect of light intensities on growth and photopigment content of a mixed culture of purple phototrophic bacteria

This chapter will be deposited in a modified version as preprint with contributions of: Marta Cerruti, Jeong-Hoon Kim, Martin Pabst, Mark C. M. van Loosdrecht and David G. Weissbrodt

# Abstract

Purple bacteria (PPB) are a guild of phototrophic organisms that thrive on aquatic environments in presence of infrared (IR) light. PPB are metabolically hyper-versatile, but they use the photoheterotrophic metabolism as preferential growth mode. Photons captured from infrared light (IR) are essential in the PPB metabolism for the providence of cellular bioenergy. Thanks to their hyper versatile metabolism and high biomass yields over substrate, PPB are promising candidate for recovery of resources from wastewater. Photopigments like bacteriochlorophylls and carotenoids are high-value compounds that can be recovered from PPB biomass. Light is a pivotal parameter to control and design PPB-based recovery facilities. So far, impact of light intensity has only been studied in pure cultures of PPB, showing that light is a limiting factor for cell growth. However, the effects of light intensities on the PPB physiology and selection in mixed cultures remain unclear. Furthermore, the enzymatic mechanisms governing the adaptation strategies to different levels of photonic energy have not been elucidated.

Here, we studied the effect of IR light irradiance ranging from 0 to 350 W m<sup>-</sup>

<sup>2</sup> on a PPB enrichment under anaerobic sequencing batch reactor regime fed with acetate as electron donor and carbon source. Higher irradiance resulted in higher growth and carbon removal. An empirical mathematical model allowed relating the growth rate of PPB to irradiance. The cellular concentration in bacteriochlorophyll a, the pigment responsible for light harvesting, and the carotenoids, the accessory pigments, was higher (6 mg<sub>bchl</sub>  $g_{DW}^{-1}$  and 3 mg<sub>lycopenel</sub>  $g_{DW}^{-1}$ ) at an incident light of 30 W m<sup>-2</sup>.

Amplicon sequencing revealed that the PPB remain dominant under all conditions (>80%), even in the absence of light. The metaproteomic analysis revealed that PPB adapt to light intensity by modulating the enzymes responsible for energy capture through the phototrophic complexes, in particular regulating the cytochrome formation and the oxidative phosphorylation pathways. The results provide in-depth phylogenetic and metabolic information on the light irradiance impacts on PPB to sustain the design of light-driven, anaerobic mixed-culture PPB processes for wastewater treatment of wastewater and the valorization of bioproducts like pigments.

**Keywords:** Anaerobic wastewater treatment, water resource recovery, PPB, irradiance effects

## 1. Introduction

Biological wastewater treatment is an established technique to treat polluted waters before discharge in the environment or re-use. Several microbial processes have been harnessed to achieve biological nutrient removal (BNR) and recovery of resources embedded in the wastewater *e.g.*, conventional activated sludge, biofilm systems, granular sludge, anaerobic processes, and phototrophic systems (Pell & Wörman, 2006). Within latest incentives to reach circular economy goals, technologies are developed to capture nutrients and valorize the microbial biomass (Guest et al., 2009; Kehrein et al., 2020)

Purple phototrophic bacteria (PPB) obtain increasing interest for wastewater treatment combined with capture of nutrients into a valuable biomass (Alloul et al., 2018; Hülsen et al., 2018b; D. Puvol et al., 2019). These anoxygenic photoorganoheterotrophs can use numerous organic and inorganic substrates as carbon and/or electron sources, growing with biomass yields over substrate close to unity by thriving on light energy (Capson-Tojo et al., 2020). PPB biomasses can be valorized for biogas and bioplastic productions, as fertilizer, and feed additive. Central to their phototrophic metabolism, PPB synthetize photopigments to capture the light and produce cellular energy (Yurkov & Beatty, 1998). Bacteriochlorophylls and carotenoids, placed in the intracellular membranes, harvest light. Bacteriochlorophylls present high quantum yields: almost all the energy from the photons is transferred and used for chemical energy production (Wraight & Clayton, 1974). Carotenoids are accessory pigments, that transfer light to the reaction centers and protect the cells from photooxidation damage (Cerullo et al., 2003). Both types of photopigments can find industrial applications. Bacteriochlorophylls can for instance be used as biomarkers in drug development (Pantiushenko et al., 2015). Carotenoids of natural and artificial synthesis, are used as food additives and food dyes (Cardoso et al., 2017).

Artificial light supply accounts for one of the major costs in photobiosystems (Blanken et al., 2013), and therefore its effects on the microbial bioprocess need to be carefully estimated. In their growth physiology, phototrophic organisms can face three main regimes driven by light intensity, namely: light limitation (growth linked to light intensity), and light saturation (Carvalho et al., 2011). PPB responses to light variations have been extensively studied in pure cultures and membrane extracts (Imam et al., 2015; Kuo et al., 2012). They adapt the number and disposition of the light harvesting complexes to the environmental conditions (Jackson et al., 2012; Muzziotti et al., 2017). The growth rates of PPB are as well dependent on light intensity (Katsuda et al., 2004), increasing with irradiance up to a saturation threshold (325  $\mu$ mol photons s<sup>-1</sup> m<sup>-2</sup> for purple

sulfur bacteria and at around 60  $\mu$ mol photons s<sup>-1</sup> m<sup>-2</sup> for purple non-sulfur bacteria) (Ritchie, 2013; Ritchie & Mekjinda, 2015).

Light is attenuated by the biomass and by the specific photopigment content (Lee et al., 2015). Shading effects have a substantial impact on the light available for the cells to grow, factually decreasing it. These effects are often neglected in PPB studies, where only the incident light is measured. An efficient growth of PPB in mixed culture and nutrient removal was achieved at low light intensities (1.4 W m<sup>-2</sup>, or 11 µmol photons s<sup>-1</sup> m<sup>-2</sup> of IR light) (Dalaei et al., 2020), but the minimum quantum requirement (the minimal irradiance necessary for growth) and the 'affinity constant for light', are not reported for PPB. Despite the pivotal importance of light for PPB growth, little is known about the PPB microbial community structure and robustness under light variations.

Here, we evaluated the effect of 9 different light intensities from 350 to 0 W m<sup>-2</sup> on the PPB community structure and growth kinetics, along with photopigment content. Using this wide range of light intensities, we evaluated the growth limits of a PPB enrichment, linked to photopigment content and enzymatic functionality. The knowledge developed on PPB community adaptation strategies to variations in energy supply will help develop PPB-based technologies for water resource recovery.

# 2. Material and methods

#### 2.1 Reactor setup and operational conditions

A 2.5-L sequencing batch photobioreactor with 2.0-L working volume connected to a controller system (In-Control and Power units, Applikon, Netherlands) was used to cultivate the PPB enrichment on an acetate-based synthetic wastewater. The photobioreactor consisted of a cylindrically shaped vessel with a curved bottom made of borosilicate glass (transmittance  $\geq$  90% in the wavelength range of the filtered light (Xia et al., 2017)). Temperature was controlled at 30° C through a finger-type heat exchanger connected to a thermostat (WK 500, Lauda, Germany). Argon gas (99% purity, Linde, NL) was continuously sparged in the reactor (excluding the settling phase) to ensure anaerobic conditions. Uniform mixing was provided with a custom-made anchor stirrer. Silicon blades were attached to the anchor stirrer to wipe the inner surface of the reactor and minimize biofilm growth on the reactor. Residual biofilm growth on probes and stirrer was removed once a week.

The system was operated in a continuously-illuminated sequencing batch reactor (SBR) mode with an 8-h cycle composed of 5 min of feed, 281 min of reaction, 4 min of purging of mixed liquor, 210 min of settling, 5 min of discharge of the supernatant, 5 minutes of idle. The purge volume was adjusted based on the

conditions to maintain an SRT of 31 h. The volume exchange ratio was set to 50% of the working volume.

The cultures were provided with a synthetic wastewater medium composed of (per L): 0.914 g C<sub>2</sub>H<sub>3</sub>O<sub>2</sub>·3H<sub>2</sub>O, 0.14 g KH<sub>2</sub>PO<sub>4</sub>, 0.21 g K<sub>2</sub>HPO<sub>4</sub>, 1 g NH<sub>4</sub>Cl, 2 g MgSO<sub>4</sub>·7H<sub>2</sub>O, 1 g CaCl<sub>2</sub>·2H<sub>2</sub>O, 1 g NaCl, as well as 2 mL of trace elements and 2 mL of vitamins solutions. The stock solution of vitamins was composed of 200 mg thiamine–HCl, 500 mg niacin, 300 mg  $\rho$ -amino-benzoic acid, 100 mg pyridoxine–HCl, 50 mg biotin and 50 mg vitamin B12 per liter. The trace elements solution was made of (per L): 1100 mg Na<sub>2</sub>EDTA·2 H<sub>2</sub>O, 2000 mg FeCl<sub>3</sub>·6 H<sub>2</sub>O, 100 mg ZnCl<sub>2</sub>, 64 mg MnSO<sub>4</sub>·H<sub>2</sub>O, 100 mg H<sub>3</sub>BO<sub>3</sub>, 100 mg CoCl<sub>2</sub>·6 H<sub>2</sub>O, 24 mg Na<sub>2</sub>MoO<sub>4</sub>·2 H<sub>2</sub>O, 16 mg CuSO<sub>4</sub>·5 H<sub>2</sub>O, 10 mg NiCl<sub>2</sub>·6 H<sub>2</sub>O and 5 mg NaSeO<sub>3</sub>. The cultivation medium was buffered at pH 7.0 with 4 g L<sup>-1</sup> 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES).

The SBR was started with an in-house mother mixed culture of PPB maintained under non-limiting IR light at 350 W m<sup>-2</sup> (Cerruti, Stevens, et al., 2020a). The experiments at high irradiances (350, 264, 175 and 87 W m<sup>-2</sup>) and the experiments at low irradiances (30, 15, 7, 3 and 0 W m<sup>-2</sup>) were subsequently performed in two different periods over the year. Between these two sets of experiments the SBR was maintained at 350 W m<sup>-2</sup>, for a period of 4 months.

#### 2.2 Infrared light irradiance and measurements

The photobioreactor was placed in a dark hood to prevent external light penetration. Light was provided from two opposite sides of the reactor with halogen lamps (Philips Plusline ES 120W R7S 230V, Philips, NL), filtered through two Black 962 Infrared Transmitting Perspex Acrylic Sheets (Black Perspex 962, Plasticstocktist, UK) for supplying infrared (IR) wavelengths ( $\lambda$ ) > 700 nm to promote PPB growth. A dimmer was used to tune the light intensity to nominal setpoints (GAMMA, NL). Incident IR light intensity at the surface of the reactor was measured with a pyranometer (CMP3; Kipp & Zonen, NL). To evaluate the effects of different light intensities, the biomass was subjected to 9 different incident light conditions: 350, 264, 175, 87, 30, 15, 7, 3 and 0 W m<sup>-2</sup>. To convert the measured light intensity (W m<sup>-2</sup>) to photon flux (µmol photons s<sup>-1</sup> m<sup>-2</sup>), the average of the photon flux per wavelength was used. To calculate it, the equation (1) was used:

$$\Phi = \frac{I \cdot \lambda \cdot 10^6}{N \cdot h \cdot c} = \frac{\mu mol \, photons}{m^2 \cdot s} \tag{1}$$

with  $\Phi$  photon flux (µmol photons s<sup>-1</sup> m<sup>-2</sup>), *I* irradiance (W m<sup>-2</sup>, *i.e.*, J s<sup>-1</sup> m<sup>-2</sup>),  $\lambda$  wavelength (m), *N* Avogadro number (6.02 · 10<sup>23</sup> photons mole<sup>-1</sup>), *h* Planck's constant (6.63 · 10<sup>-34</sup> J s), *c* speed of light (2.88 · 10<sup>8</sup> m s<sup>-1</sup>).

The emission spectrum of the filtered light is presented in Supplementary material 1.

#### 2.3 Specific light supply rate

The specific light supply rate (rEX) defines the amount of light available per gram of biomass. It was calculated as in (Sforza et al., 2015)

$$rEX = \frac{PFD \cdot A}{X_{av} \cdot V}$$
(2)

with *PFD* the photon flux (µmol photons m<sup>-2</sup> s<sup>-1</sup>), *A* the irradiated surface (calculated from the reactor geometry) (m<sup>2</sup>),  $X_{av}$  the average biomass concentration in the cycles for each condition (g L<sup>-1</sup>), *V* the reactor working volume (L). The correlation between incident light and rEX is presented in Supplementary material 2.

#### 2.4 Sample collection

Each light condition was maintained for 14 days (10.8 STRs). Samples for community composition analysis, photopigment concentration and acetate removal were collected from day 10-11 to day 14 after the beginning of a new light intensity condition, corresponding to 7.7 to 10.8 SRTs from the beginning of the new phase.

#### 2.5 Measurement of biomass concentration

The biomass concentrations in the mixed liquor obtained at the different IR-light irradiances were measured by spectrophotometry (Biochrom, Libra S11, US) through absorbance at 660 nm (Abs<sub>660</sub>). A calibration curve was established to correlate Abs<sub>660</sub> to the biomass concentration (g VSS):  $c = 1.4 \cdot Abs_{660}$ . Volatile suspended solids were measured as in standard methods (Water Environment Federation, 1999).

# 2.6 Analysis of bacterial community compositions by amplicon sequencing

The compositions of the bacterial communities present in the mixed liquors under the different IR light irradiances were characterized from the biomass samples by V3-V4 16S rRNA gene-based amplicon sequencing as detailed in Cerruti et al., 2020. The DNA extracts were sent to Novogene (UK) for library preparation and amplicon sequencing using an Illumina 600N sequencer. The fastq files provided by Novogene were analysed with the QIIME2 pipeline (Bolyen et al., 2019). The sequencing results are at NCBI database under the BioProject number ID PRJNA799236.

#### 2.7 Extraction and analysis of photopigments

Bacteriochlorophyll and carotenoid photopigments were extracted and their mass fractions quantified and compared from the PPB biomasses obtained at the different IR light irradiances.

#### 2.7.1 Photopigments extraction

Prior to pigments extraction, biomass was freeze-dried for 12 h at -50  $^{\circ}$ C and at a pressure of 0.05 mbar, to remove water from the pellets and facilitate polar solvents penetration.

Hexane / methanol / water in proportions of 2:1:1 (v:v:v) were used to extract the photopigments. Hexane and methanol were mixed and pre-chilled at 4°C overnight. 1200  $\mu$ l of this solution were added to the biomass in a cold and dark environment. The cells were vortexed with intermittent cooling on ice to promote the extraction of non-polar compounds. Once completely resuspended, the cells were incubated in ice for 5 min. Water was added to promote the separation of the photopigments (non-polar) from cells debris, and the samples were centrifuged for 5 min at 17000 x g, resulting in two separate phases. The upper phase, non-polar, consisting of methanol and hexane, was collected for spectrophotometric quantification of the photopigments.

#### 2.7.2 Quantification of photopigments

Concentration of the photopigments in the samples was calculated according to the Lambert-Beer equation

$$A = \varepsilon_{\lambda} \cdot C_p \cdot D \tag{3}$$

Where *A* is the measured absorbance at the peak wavelength (lycopene = 473 nm, bchl = 776 nm),  $\varepsilon_{\lambda}$  the molar absorption coefficient (lycopene = 1.72\*10^5  $M^{-1} g_{photopigment}^{-1}$ , bchl = 6\*10^4  $M^{-1} g_{photopigment}^{-1}$  (Bóna-Lovász et al., 2013)) of the compound at specific wavelength  $\lambda$ , *Cp* is the molar concentration of the photopigment and *D* is the pathlength of light (0.5 cm).

To calculate the photopigment mass fraction, the photopigments molar concentration was divided by the bacterial mass before the extraction and multiplied for the final volume of solvent (hexane) used for the elution

# 2.8 Calculation of growth kinetic and stoichiometric parameters *Growth rate*

Biomass specific growth rates  $\mu$  (h<sup>-1</sup>) were calculated from the mixed culture as:

$$\mu = ln(\frac{X_1}{X_0}) \cdot \frac{1}{T_1 - T_0} \tag{4}$$

with  $X_0$  and  $X_1$  the biomass concentration at the beginning and at the end of the reaction phase, respectively, and  $T_0$  and  $T_1$  the beginning of the reaction phase and end of the growth period, respectively.

The growth rates were fitted to the light intensity with a generic logistic function (Eq. 5) through the lsqcurvefit function of MATLAB (R2018b, Mathworks, Natick, MA, <u>www.mathworks.com</u>). It is a nonlinear least-square solver that finds the coefficients that best fit a given nonlinear function.

$$y = \frac{a}{1 + e^{(-b(x-c))}}$$
(5)

#### 2.9 Metaproteomic analysis

High-resolution analysis of the metaproteome of the mixed culture was performed on selected biomass samples collected at the end of the experimental periods at incident irradiances of 350, 87 and 15 W m<sup>-2</sup> to evaluate the differences in pathway expression. The protein extraction followed by shotgun metaproteomic analysis, pre-filtering, sequence randomisation, Unipept submission and compositional analysis is described in detail in (Kleikamp et al., 2021).

The area of the peak for each protein was normalized for the total area of each sample, and expressed as its percentage. A statistical analysis was carried on the triplicates using Rstudio (RStudio samples Team (2020).http://www.rstudio.com/). The 87 W m<sup>-2</sup> condition was used as reference point. The protein average and the foldchange was calculated. A t-test was used to evaluate the significance in protein expression difference between the three conditions. Proteins with a p-value below 0.05 were marked as significantly differently expressed, comparing 350 W m<sup>-2</sup> vs 87 W m<sup>-2</sup> conditions and 15 W  $m^{-2} vs 87$  W  $m^{-2}$  conditions. Proteins with a p-value < 0.05 and a log<sub>2</sub>-foldchange > 0.5 or < -0.5 were considered over- or underexpressed compared to the reference set (87 W m<sup>-2</sup>), respectively.

The UniProt accession identifiers were translated to KEGG IDs through BlastKOALA (BlastKOALA, https://www.kegg.jp/blastkoala/) server. Protein classification was done according to the KEGG database. A manual classification of the entries not represented in the KEGG database was further performed.

## 3. Results

#### 3.1 Biomass growth and acetate consumption

Incident IR light intensities supplied at the surface of the anaerobic sequencing batch photobioreactor impacted biomass growth and nutrient capture by the PPB mixed culture. Under high light intensities of 350 and 264 W m<sup>-2</sup>, the biomass reached a concentration plateau of  $2.8 \pm 0.2$  g<sub>DW</sub> L<sup>-1</sup> (Table 1) within the first 70 min of cycle operation (Figure 1A). At medium light intensities of 175 and 87 W m<sup>-2</sup>, the biomass grew to lower concentrations and continuously during the 4 h of cycle operation. At lower light intensities of 30 - 0 W m<sup>-2</sup>, the biomass growth was almost negligible (Figure 1A).

Conversion-wise, at high light intensities (350 W m<sup>-2</sup>), acetate was fully depleted within the first hour of reaction (Figure 1B). At 264 and 175 W m<sup>-2</sup>, acetate reached a residual concentration of  $0.05 \pm 0.06$  mmol L<sup>-1</sup> and  $0.04 \pm 0.10$  mmol L<sup>-1</sup> at the end of the reaction phase (Table 1). At 87 W m<sup>-2</sup>, acetate was linearly decreasing over the 4 h of incubation, reaching a final concentration  $0.27 \pm 0.05$  mmol L<sup>-1</sup>. At 30 W m<sup>-2</sup>, at the end of the incubation time  $2.34 \pm 2.02$  mmol L<sup>-1</sup> of acetate was present at the end of the reaction phase. At the end of the reaction phase of the low light (15-0 W m<sup>-2</sup>) conditions acetate was still present ( $3.10 \pm 0.6$  mmol L<sup>-1</sup>).

				Con	dition (W	m <sup>-2</sup> )			
	0	n	۲	15	30	87	175	264	350
Biomass* $(g_{DW} L^{-1})$	0 09+0 02	0 15±0 01	0 75±0 05	0.61±0.06	1 28+0 02	2 04+0 11	90.0+25.5	2 65±0 04	2 94+0 48
Acetate* (mmol $L^{-1}$ )	20-00-0 89 0+62 £	95.0+85.5	3 00+1 33	00.0-10.0	1 37+0 49	50.0+20.0	0.04+0.10	90.0+50.0	
Growth rate (h <sup>-1</sup> )	0.03±0.03	0.01±0.01	0.01±0.01	0.02±0.01	0.01±0.01	0.02±0.00	0.10±0.01	0.18±0.03	0.22±0.05
х •									
*Concentration rr <sup>+</sup> BDL: below dett Values are prese deviations.	easured at the ection limit inted as avery	end of the re age of the fo	action phase. our cycles ree	corded and c	orresponding	standard		for each condition	Table 1: Biomass and acetate concentration at the end of the reaction phase and specific growth rates calculated



Figure 1: Biomass (A) and acetate (B) concentration along the SBR cycles. Values are expressed as the average value for each time point. Samples were collected for three cycles. At high light intensities (350-87 W m<sup>-2</sup>) acetate was fully consumed at the end of the reaction cycles. At low light intensity (30-0 W m<sup>-2</sup>) acetate was still present at the end of the reaction phase and biomass growth was negligible.

#### 3.2 Biomass growth rates followed a logistic distribution

The growth rates decreased with the light intensity, from  $0.22 \pm 0.05$  h<sup>-1</sup> at  $350 \text{ W} \text{ m}^{-2}$  to  $0.03 \pm 0.01 \text{ h}^{-1}$  at 87 W m<sup>-2</sup> to  $0.008 \pm 0.006 \text{ h}^{-1}$  at 3 W m<sup>-2</sup>. The growth rates followed a logistic distribution, both considering the specific light supply rate (rEX) (Figure 2A) and the incident light (Figure 2B). It was possible to identify two conditions: acetate- and light- limited conditions. Considering the specific light supply rate, the cultures which had no acetate at the end of the reaction phase were acetate-limited and had a rEX above 10  $\mu$ mol<sub>photons</sub> s<sup>-1</sup> g<sub>DW</sub><sup>-1</sup>, corresponding to an incident light of 87 W m<sup>-2</sup>. Below 10  $\mu$ mol<sub>photons</sub> s<sup>-1</sup> g<sub>DW</sub><sup>-1</sup> the cultures were light limited, and growth was reduced compared to the higher irradiances. The logistic curve (Eq. 5) fitted through the data allowed to identify three parameters: a, corresponding to the maximum growth rate  $(\mu_{max}, h^{-1})$ , b, a correction factor, and c, corresponding to a 'half saturation constant' for light. The factor a was identified at 0.24 and 0.22  $h^{-1}$ for the rEX and the incident light curvess respectively. The factor c, at 34  $\mu$ mol<sub>photons</sub> s<sup>-1</sup> g<sub>DW</sub><sup>-1</sup> in the rEX fit and 189 W m<sup>-2</sup> in the incident light fit. In Eq. 5, x was defined as the light intensity (both as rEX and as incident irradiance). and *y* the specific growth rate.



Figure 2: Biomass growth rates (black circles) and acetate concentration (red squares) in relation to rEX (A) and incident light intensity (B). The growth rates fitted a logistic regression (dashed line), that allowed to identify the  $\mu_{max}$  for the PPB enrichments (dotted line). Based on the acetate concentration at the end of the reaction phase, an acetate- and a light-limited condition were identified.

## 3.3 Community composition

The taxonomic compositions of the bacterial communities obtained under the different irradiance treatments were measured by 16S rRNA gene amplicon sequencing. PPB were highly enriched, accounting for  $81 \pm 9$  % of the total reads under all irradiance conditions, except under dark conditions (*i.e.*, 0 W m<sup>-2</sup>) (Figure 3A). Under dark conditions, the PPB enrichment decreased to 40%, while the genus *Dechloromonas* increased to 22%. Uncultured genera of the family of the *Rikenellaceae* accounted for 8%. The metaproteomic analysis confirmed the amplicon sequencing data. In the three samples analysed by metaproteomics (350, 87 and 15 W m<sup>-2</sup>), more than 75 % of the proteins were assigned to genera belonging to the PPB guild, namely *Rhodopseudomonas*, *Rhodobacter*, *Blastochloris*, *Thiobaca* (Figure 3B).



Figure 3: A: amplicon sequencing results for community composition. PPB genera (Blastochloris, Rhodobacter, Rhodopseudomonas and Thiobaca, red bars) were the dominant guild under all condition, accounting for ca 70% of the total community, excluding under dark condition. B: metaproteomic analysis of the community composition for selected samples. These results confirmed the amplicon sequencing results. C: PPB concentration in the bulk liquid increased with the increase of incident light intensity. D: Distribution of the main PPB genera based on amplicon sequencing results under high (350-87 W m<sup>-2</sup>) and low (30-0 W m<sup>-2</sup>) conditions. Notably, the genus Thiobaca was present only under high light conditions.

The concentration of PPB in the bulk liquid decreased with the decrease of the incident light, from  $2.38 \pm 0.07 \text{ g}_{PPB} \text{ L}^{-1}$  at 350 W m<sup>-2</sup> to  $0.03 \pm 0.00 \text{ g}_{PPB} \text{ L}^{-1}$  under dark conditions (Figure 3C).

Among the PPB, *Thiobaca* was the dominant organism at high light intensities (350-87 W m<sup>-2</sup>), with a relative abundance of  $56\pm 6\%$ . *Rhodopseudomonas* and *Blastochloris* reached a relative abundance of  $17 \pm 4\%$  of  $7 \pm 4\%$ . Under low light conditions (30-3 W m<sup>-2</sup>) *Rhodopseudomonas* and *Blastochloris* were the dominant populations, with  $54 \pm 8\%$  12 ± 6 % respectively. The genus *Thiobaca* was not detectable under these conditions (Figure 3D.) These differences might be attributed to the different inocula in the reactor under high and light irradiation, as explained later.

#### 3.4 Photopigment mass fraction

Wavelength scans between 320 and 1100 nm were recorded to determine the photopigment content in the whole cells of the biomass and in the pigments extract. Peaks in the wavelengths of the bacteriochlorophylls (800-900 nm) and carotenoids (between 400-500 nm) were detected under all conditions, including under dark, both in the extract and in the whole cells (Figure 4A/B). In the whole cells, under high light conditions (350-87 W m<sup>-2</sup>) two peaks were detected at 800 and 890 nm, corresponding to the bacteriochlorophylls (bchl). At low light intensities (30-0 W m<sup>-2</sup>), the bchl peaks were detectable at 805 and 866 nm. In the carotenoid area, peaks were detected at 322 and 370 nm under all conditions (Figure 4A).

In the extracts, the peaks for bchl were detected at 776 nm under all conditions. In the carotenoids area, peaks were present at 511, 475, 446 and 389 nm (Figure 4B). The bchl mass fraction at low light intensities  $(3.8 \pm 1.7 \text{ mg } \text{g}_{DW}^{-1} \text{ at } 0.30 \text{ W m}^{-2})$  was about 4.6 times higher than under high irradiance  $(0.8 \pm 0.2 \text{ mg } \text{g}_{DW}^{-1} \text{ at } 87.350 \text{ W m}^{-2})$ . Similarly, the representative carotenoid (lycopene) showed a higher concentration at lower light intensities, with a maximum of about 3 mg  $\text{g}_{DW}^{-1}$  at 30 W m<sup>-2</sup> (Figure 4C).



Figure 4: A: Wavelength scan of PPB photopigments extract resuspended in hexane. The highest absorption peak was visible at 30 W m<sup>-2</sup>. B: Wavelength scan of PPB whole cells. The Absorption peaks in the bchl area shifted when changing high and low light intensities. C: The photopigments (bchl and lycopene) concentration in the cells was higher at lower light intensities compared to high light intensities.

#### 3.5 Metaproteomics analysis

The metaproteomic analysis identified 1471 proteins. The complete metaprote-omic profile of the analysed samples is presented in Supplementary material 4. A number of 1099 of the proteins (75% of the total) belonged to the four main PPB genera identified in the amplicon sequencing analysis, namely Thiobaca, Rhodopseudomonas, Rhodobacter and Blastochloris. Under all conditions, the most abundant proteins belonged to the phototropic machinery (reaction center proteins and cytochromes) and carbon metabolism. Following KEGG classification, the proteins for the phototrophic machinery ac-counted for  $22.5 \pm 0.7$  % of the total PPB proteins. The proteins of the reaction center (log<sub>2</sub>-foldchange =  $0.97 \pm 4.61$ ) and the light harvesting complexes ( $\log_2$ -foldchange = 3.58 ± 2.52) were significantly more expressed in the 15 W  $m^{-2}$  condition compared to the 87 W  $m^{-2}$  condition. In contrast with the results of the photopigments extraction, proteins involved in the photopigment formation were under-expressed at 15 W m<sup>-2</sup> compared to the 87 W m<sup>-2</sup> condition. At high light intensities (350 W m<sup>-2</sup>), proteins involved in phototrophic carbon fixation (log<sub>2</sub>-foldchange =  $0.09 \pm 1.88$ ) and oxidative phosphorylation were more expressed compared to the 87 W m<sup>-2</sup> condition. In the carbon metabolism, at 15 W  $m^{-2}$  proteins involved in PHA metabolism were overexpressed compared to the 87 W m<sup>-2</sup> (log<sub>2</sub>-foldchange =  $0.84 \pm$ 4). Similarly, also the enzyme involved in the glyoxylate cycle, with a  $log_2$ -foldchange of 0.61± 2.75 were overexpressed. At 350 W m<sup>-2</sup>, no significant difference was reported in the expression of the carbon metabolism enzymes compared to the 87 W m<sup>-2</sup> condition.



Figure 5: A: Heatmap representing the average percentage of the protein expressed in each metabolic category based on KEGG classification. The heatmaps represents only carbon metabolism and phototrophic pathways. B: Differential regulation of the different pathways under 15 and 350 W m<sup>-2</sup> compared to the 87 W m<sup>-2</sup> condition. Oxidative phosphorylation and photosynthesis pathways were upregulated in the 350 W m<sup>-2</sup> condition and downregulated in the 15 W m<sup>-2</sup> conditions compared to the 87 W m<sup>-2</sup> condition.

### 4. Discussion

#### 4.1 Photopigments content was inversely proportional to light intensity

For phototrophic organisms, energy is derived by light. Based on the irradiation conditions, PPB adapt the number and disposition of photopigments in their membranes to maximize the light harvest (Brotosudarmo et al., 2011; Gardiner et al., 1993). In line with previous observations (S. Liu et al., 2019), we report an increase in photopigment content (both bacteriochlorophylls and carotenoids) with a decrease of light intensity. In fact, bacteriochlorophyll and carotenoids mass fraction was respectively 4.6 and 10 times higher under low light conditions compared to high light conditions. The variation in the photopigment content of the biomass can be attributed to different factors. The decrease of photopigments is an acclimatization strategy to high light intensities. Muzziotti et al., 2016 report that Rhodopseudomonas increases the bacteriochlorophyll mass fraction by a factor 6 and the carotenoids mass fraction by a factor 2.5 when light intensity changes from 250 to 1500  $\mu$ mol<sub>phtotons</sub> m<sup>-2</sup> s<sup>-1</sup> (corresponding to ca 33 and 190 W m<sup>-2</sup>) under anaerobic conditions. By reducing the antenna size and consequently the amount of light absorbed, PPB protect the cells from photodamage induced by high light intensities (Bayon-Vicente et al., 2020; Muzziotti et al., 2017). Furthermore, the phototrophic growth is linked to the photopigment content and activation state of the reaction centers, that harvest

light and convert it into chemical energy (Saikin et al., 2014). To compensate for the lower energy supply, PPB increased the mass fraction of photopigments.

The variation in photopigment mass fraction might also be linked to the taxonomic composition of the enrichment. The microbial community was different in the two experimental set-ups (high and low light intensities, Figure 3D). At low light intensities, the community was enriched for the genus *Rhodopseudomonas*. At high light intensities, instead, the most abundant genus was *Thiobaca*. Each organism has a different distribution of the photopigment, both spatial and of type (Bullough et al., 2009; Timpmann et al., 2014). To maximize the efficiency of energy capture, PPB combine protein and pigments in the light-harvesting complexes and reaction center (Fixen et al., 2016). The differences in protein binding of the pigments influence the stability of the complexes (Mizoguchi et al., 2012), and potentially have affected the extraction yields in our enrichments. However, the taxonomic difference between *Thiobaca* and *Rhodopseudomonas* cannot be interpreted as a selection mechanism caused by light irradiance, such as explained in the next section.

#### 4.2 PPB were enriched under all conditions

PPB formed the dominant guild under all conditions, confirming the possibility to efficiently enrich them on acetate as electron donor and carbon source and with IR light as energy source (Alloul et al., 2019; Cerruti et al., 2020). The supply of IR light selectively enriches for PPB and prevents the growth of other phototrophs, such as green / blue phototrophic organisms that extract their energy on other wavelengths. This prevents formation of oxygen and ensures all organic matter is available for the PPB. The PPB enrichment was confirmed both at genetic (amplicon sequencing) and functional level (metaproteomics).

The taxonomic variation from a *Thiobaca*- to a *Rhodopseudomonas*-dominant community was due to the inoculum rather than to a light effect. The low light conditions were applied to the same parent reactor 4 months after the high light conditions. The predominance of the genus *Rhodopseudomonas* at high light intensities (> 300 W m<sup>-2</sup>) was repeatedly reported in other SBR systems (Cerruti et al., 2020, Cerruti et al, in preparation). In a batch system, the selective pressure is determined by the growth rates (Winkler et al., 2017). No information is currently available for the growth kinetics of Thiobaca. Pure cultures of *Rhodopseudomonas* present higher growth rates ( $\mu_{max} = 0.15 \text{ h}^{-1}$ (Cerruti, Ouboter, et al., 2020)) compared to other PPB pure cultures like *Rhodobacter* or *Blastochloris* under acetate feed (0.10 and ca 0.03  $h^{-1}$ respectively ) and was therefore predominant under SBR conditions (LaClair, 2006; Zavala et al., 2019). We were not able to detect what did cause the presence of *Thiobaca* in the initial mother culture. Potentially, the competition between purple sulfur bacteria (Thiobaca) and purple non-sulfur bacteria (Rhodopseudomonas, Rhodobacter, Blastochloris) might be linked to the

accumulation of sulfur compounds (not measured in this study). Competition phenomena between PNSB and PSB should be further researched in the future.

The PPB enrichment was stable under all IR light intensities, with a relative abundance above 75%, except for the dark condition (0 W m<sup>-2</sup>). In presence of IR-light, PPB show relatively high growth rates, ranging between 0.03 and 0.3 h  $^{1}$  (Alloul et al., 2019). Few organisms can grow under anaerobic conditions with acetate as sole carbon and electron source, since acetate is already a fermentation product. Acetoclastic organisms, like the slow-growing methanogenic archaea, present a  $\mu_{max}$  of 0.004-0.013 h<sup>-1</sup> (Kotsyurbenko et al., 2019). In the reactor, the SRT was controlled at 31 h, implying that the minimum growth rate to be retained in the bulk liquid was set to  $0.03 \text{ h}^{-1}$ . The taxonomic analyses showed that no methanogens were detected both at 16S rRNA gene level and at protein level. Under dark anaerobic conditions, PPB cannot grow on acetate without external electron acceptors (Yen & Marrs, 1977), which were not present in the cultivation medium. Possibly, the high relative abundance of PPB under dark conditions can be attributed to the biomass retention during the settling phase. The chemotrophic growth of Dechloromonas (present at 22 % under dark conditions) explains the apparent higher growth rates measured for the biomass under dark condition compared to other low light intensities. Under low-light conditions, the growth rates were almost equal to the SRT (0.03 h<sup>-1</sup>). Iasimone et al., 2018 report an increase in the length of the lag phase with the decrease of light intensity. Possibly, at low light intensities the PPB enrichment was still in the lag phase. An increase of the reaction length might lead to a biomass increase, and consequently higher carbon uptake and photopigment recovery.

#### 4.3 Enzymatic expression changed with light intensity

In the proteomic analysis, it emerged that at low light conditions the proteins responsible for the light harvest were overexpressed compared to higher light intensities. These results match the hypothesis that, with lower irradiances, cells require to produce more light harvesting complexes to be able to sustain the energy requirements. Interestingly, the proteins for the formation of accessory pigments were expressed under all conditions at the same level.

Enzymes involved in the electron transport chain and ATP generation were overexpressed at high light intensities, similarly to what reported for microalgae (Toyoshima et al., 2019). The higher ATP production rate can have enhanced the carbon fixation (photosynthesis) pathway, whose enzymes were overexpressed at high light conditions (350 W m<sup>-2</sup>). In fact, the Calvin-Benson-Bassham cycle is a known mechanism of electron reallocation (McKinlay & Harwood, 2010). The energy-expensive  $CO_2$  fixation regulates the redox balance by increasing the fixation rates when an excess of ATP is produced through the electron transport chain (Alsiyabi et al., 2019). Poly-hydroxyalkanoates

(PHA) formation in PPB is usually linked to redox stress conditions (Bayon-Vicente et al., 2020; Cerruti et al., 2020). The enzymes linked to the PHA granules were upregulated in the low light condition (15 W  $m^{-2}$ ), namely proteins belonging to the phasin family. However, enzymes for PHA formation were not detected. The function of the phasin proteins under this condition is unclear.

The high standard deviations reported for the log2-foldchange can be attributed to the microbial community composition. For example, at low light intensities, where *Rhodopseudomonas* was the most abundant genus, 284 proteins belonging to *Rhodopseudomonas* were over-expressed compared to the 87 W m<sup>-2</sup> condition, 58 were not significantly different and only 17 were downregulated. The opposite was reported for *Thiobaca* (the most abundant genus at high light intensities): only 2 of its proteins were upregulated at low light conditions, whereas 268 were downregulated and 32 not significantly different.

The over-expression of the glyoxylate cycle enzymes in the 15 W m<sup>-2</sup> condition can also be linked to the community composition. Carbon metabolism in PPB is specie-specific, meaning that the different organisms use different pathways to catabolize acetate. For instance, the genus *Rhodopseudomonas* utilizes the glyoxylate shunt to metabolize the acetate (Albers & Gottschalk, 1976), whereas *Rhodobacter* uses the EMC pathway (Alber et al., 2006). An enrichment of *Rhodopseudomonas* in the reactor can explain an enrichment of the glyoxylate cycle proteins at low light intensities.

#### 4.4 Growth rates are dependent to light availability

Numerous kinetics models have been constructed to correlate the growth rates of phototrophic organisms to light intensity and substrate availability (Lee et al., 2015). Light and nutrients are complementary factors contributing to growth kinetics of phototrophs. Under light-saturating conditions, concentrations of the limiting substrate (within electron donor and C-P-N nutrients), defines the growth rates, as described by the Monod model (Doran, 2013). Under nutrient-saturating conditions, growth rates are instead controlled by light intensity. At light saturation level, the maximum growth rate is achieved (Undurraga et al., 2016). Above this point, growth is inhibited (photoinhibition). Below the saturation level, growth is light-limited. At present, the definition of a univocal kinetic model for phototrophic growth is not possible, as numerous factors influence the growth rates (Lee et al., 2015). Furthermore, the kinetic parameters to define saturation and inhibition constants are species-specific (Huesemann et al., 2013), further complicating the task.

For the PPB biomass growth two conditions were found, light limited and acetate limited growth. At high light intensities (350-175 W m<sup>-2</sup>), the complete depletion of the carbon source corresponded to the stop of the biomass growth, indicating that the acetate was the limiting compound. Under these conditions, the biomass grew with a growth rate close to the calculated  $\mu_{max}$  (0.10-0.22 h<sup>-1</sup>). At low light intensities, the biomass displayed growth rates almost 8 times lower

than under high irradiance. The presence of acetate at the end of the reaction phase suggested that the culture was light-limited. In our study, the biomass growth rates could be approximated by a logistic expression. The logistic function has been used for more than one century in ecology to describe the growth of populations (Kendrick & Kesava Pai, 1911; Verhulst, 1838). More recently, it has been used to describe the growth of some PPB (Eroğlu et al., 2004; Eroğlu et al., 2008; Koku et al., 2003). In the logistic model, the growth of a population is described in terms of intrinsic rate of increase, or  $\mu_{max}$ , and the carrying capacity. In our study, the fitted  $\mu_{max}$  was 0.22-0.25 h<sup>-1</sup>, in the range of the reported  $\mu_{max}$  for PPB (Hunter et al., 2009). The carrying capacity was defined at a rEX of 10  $\mu$ mol<sub>photons</sub> s<sup>-1</sup> g<sub>DW</sub><sup>-1</sup>, or an incident light of 87 W m<sup>-2</sup>. These values can be considered as the half saturation constant for light ( $K_1$ ). The  $(K_1)$  of the rEX is particularly important for process design, as it allows to determine the limit for biomass growth. Our result suggests that light is governing all the other metabolic processes, by providing energy to the cells, and an ATP balance would be required to further sustain this hypothesis. ATP production and consumption in PPB have been described in metabolic models (Golomysova et al., 2010), but an accurate measurement of the ATP produced by PPB per photon absorbed has not vet been reported.

#### 4.5 Light intensity impacted PPB systems

In phototrophic systems, light provides energy to the cells, that use it for growth. To design a phototrophs-based process, it is necessary to accurately manage the light attenuation, as it impacts the reactor performances. The Lambert-Beer equation describes the light attenuation due to biomass concentration and photopigment content, but it is lacking the refraction and scattering component. On the other hand, the specific light supply rate describes how much light each gram of cell can experience but does not consider the optical effects. Numerous models have been developed to link nutrients and light availability to the biomass kinetic parameter (reviewed in Lee et al., 2015). These models are based on pure culture studies, and do not consider the interactions, as competition, mutualism or synthrophysm, between organisms in a community.

In the PPB enrichment, we found that the specific growth rates were correlated to the light intensity available for the cells through a logistic regression. The data fitting provided insights on the kinetic parameters of the PPB culture, as the limit for photoinhibition phenomena ( $\mu_{max} = 0.25 \text{ h}^{-1}$  at 80  $\mu$ mol<sub>photons</sub> s<sup>-1</sup> g<sub>DW</sub><sup>-1</sup>) and the half saturation constant for light (K<sub>L</sub>= 10  $\mu$ mol<sub>photons</sub> s<sup>-1</sup> g<sub>DW</sub><sup>-1</sup>). Growth kinetic models are needed to understand PPB growth and therefore to optimize the cultivation condition. For scale-up purposes, mathematical modelling proves to be a useful tool to predict the microbial behavior. Future research should address a multivariate predictive analysis of the kinetic parameters of PPB enrichments.

## 5. Conclusions

PPB form a complex microbial guild, whose metabolism is primarily photoheterotrophic. By changing the incident light, we elucidated the mechanisms of adaptation of a PPB community to variation in energy supply. Our findings contribute to paving the way for PPB-based bioprocesses.

We can conclude that:

The PPB guild was enriched above 70% of the total community under all conditions in presence of IR light. In the dark, it was still dominant, but only at 40%.

PPB growth can be either nutrient-limited or light-limited. At a flux rate (rEX) above 10  $\mu$ mol<sub>photons</sub> s<sup>-1</sup> g<sub>DW</sub><sup>-1</sup>, the enrichment was acetate-limited. Below this point it was light limited. Notably, this value is important for process design.

PPB growth rates fitted a logistic function. This allowed to identify the maximum growth rate ( $\mu_{max} = 0.25 \text{ h}^{-1}$ ) and the half saturation constant for light ( $K_L = \mu mol_{photons} \text{ s}^{-1} \text{ g}_{DW}^{-1}$ )

Photopigment mass fraction was ca 5 times higher at low light intensities compared to high light intensities. This can be due both to physiological responses and to community composition.

PPB adapted their enzymatic expression to respond to the light stress. At low light intensities the proteins involved in light-harvesting processes were more expressed, whereas at high light intensities PPB activated the mechanisms for stress response.

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Supplementary materials

# Supplementary material 1 Spectrum of the lamps with and without filter



SI 1: The light provided to the reactor was filtered with filter sheets. Only the wavelengths above 750nm were reaching the surface of the reactor.

## Supplementary material 2 Correlation between incident light and specific light supply rate



SI 2: the incident light and the specific light supply rate (rEX) were linearly correlated. The rEX was calculated as in Material and Methods section of this chapter, and the average biomass concentration of all measured cycles under each condition was used.

# Supplementary material 3 Correlation between absorbance units and gVSS L<sup>-1</sup>



SI 3: Correlation between the absorbance at 660 nm and the VSS concentration of the PPB-enriched biomass, based on dilution series of mixed liquor taken from the reactor  $(2 \times, 4 \times, 8 \times, 16 \times$  diluted). It displays the experimental data of three measurement series with the linear regression line. The linear regression of the triplicate measurement was y = 1.4x. For absorbance measurements, mixed-liquor samples were diluted to fit in the range from 0.2-1.0 absorbance units. On this absorbance window the correlation to VSS was considered as linear.

# Supplementary material 4 Amplicon sequencing analysis



The microbial community analysis for each condition was analysed in duplicate. The results of the amplicon 16S rRNA gene sequencing are shown as relative abundance of the genera for each sample. An online-only version of the amplified reads is available

# Supplementary material 5

### Metaproteomic analysis, online version only

The results of the metaproteomic analysis are shown as the peak area for each identified protein. Only proteins with 2 or more peptides are here presented, to assure accuracy.

Chapter 6

General conclusions
## Harnessing the metabolic versatility of purple bacteria

This thesis aimed to explore the metabolic versatility of purple phototrophic bacteria (PPB), with respect to the application of anaerobic, mixed-culture, photobioprocesses for wastewater treatment and valorization. PPB-based wastewater technologies are at their early stage of development, currently mostly relating to bench-scale ideas and technology readiness levels of 2-4. The metabolic insights on PPB gathered with pure-culture studies over the last 80 vears are far to be translated into practice, for both axenic industrial biotechnologies or open environmental bioengineering processes. After early characterizations of purple bacteria and their metabolism(s) in Delft by van Niel, primary environmental applications of PPB take origin in sunlight-driven ponds used to passively treat sewage residues such as reported from Japan as early as the 1970's. By being able to combine multiple phototrophic and chemotrophic modes of growth, PPB are one of the most versatile guilds of microorganisms in nature. While growing preferentially by photoorganoheterotrophy - e.g., using volatile fatty acids (VFAs) as electron donors and carbon sources, and infrared light (IR) wavelengths as energy source - their metabolism is impacted and tuned by dynamics in environmental conditions such as light / dark cycles, light intensity, substrates, and redox conditions. Understanding the mechanisms governing the microbial metabolisms and their interactions with other populations is crucial to develop and maintain mixed-culture processes for biobased environmental and refinery services.

#### 6.1 Excess electrons are first allocated toward PHA, and secondly toward H<sub>2</sub>

Poly- $\beta$ -hydroxyalkanoates (PHAs) and biohydrogen (H<sub>2</sub>) are two of the major products that have been considered to be valorized from PPB process. Both compounds are an electron sink in the metabolism, implying that the production of one compound prevents the formation of the other one. Using the model PPB Rhodopseudomonas palustris that was isolated from an enrichment culture performed in this thesis, and applying quantitative biotechnology methods, we elucidated in Chapter 2 the mechanisms of electron allocation between PHA and H<sub>2</sub>. Light and dark cycles can indirectly influence the direction of the electron flux from their supply with acetate toward one or the other product, besides biomass. Under light-saturating conditions and continuous regime, the duration of the light or dark phases determines the achievement of a steady state in the photoorganoheterotrophic metabolism of *Rhodopsdeudomonas*. Long (16 h) light phases lead to complete substrate depletion and steady conditions. Short (8 h) light phases prevent substrate limitation. The excess of carbon accumulated during the dark periods is redirected toward PHA. In growing cells, H<sub>2</sub> results only as a minor electron sink, secondary to PHA production. The experimental results were backed up by a mathematical model that captured the metabolic dynamics of PPB under light/dark diel cycles.

#### 6.2 PPB are enriched and aggregated in simple SBR systems

In Chapter 3, we investigated the possibility to enrich a PPB mixed culture in a simple and efficient way, and potentially stimulating their aggregation in a dense biomass. Continuous IR irradiation combined with acetate supply selectively enriched for PPB up to 90% of the bacterial community, regardless of the imposed sludge retention time (SRT). The PPB biomass was able to simultaneously remove nutrients (C, N, P) to levels complying to the current European regulation, by assimilation in the sludge in a single-stage reactor. The use of a sequencing batch reactor (SBR) operation strategy allowed to accumulate high biomass concentrations (as high as 3.8 g VSS L<sup>-1</sup>, *i.e.*, about 10 times higher than generally reported) forming dense bioaggregates with relatively fast settling ability. The formation of aggregates was stimulated by substrate gradients generated under SBR operation, and facilitated the solidliquid separation required for downstream processing (*i.e.*, suppressing the need for energy-expensive membranes). We showed the feasibility of a PPB-based SBR process for full biological nutrient removal, when fed with VFAs (i.e., prefermented wastewater).

#### 6.3 PPB and FCB form a syntrophic interaction

About 19% of the total aqueous wastes derive from food and agricultural sources. The chemical oxygen demand (COD) of these streams is substantially higher (up to 22  $g_{COD} L^{1}$ ) compared to municipal wastewater (*ca.* 500 mg<sub>COD</sub>  $L^{2}$ (Asgharnejad et al., 2021; Roeleveld & van Loosdrecht, 2002). Food-industry wastewater, in particular derived from fruit and vegetable processing, is usually rich in sugars h-1. In this scenario, and without pH control, fermentative chemotrophic bacteria (FCB) outcompete PPB, thanks to higher growth and carbon assimilation rates. The choice of an appropriate reactor regime is crucial to establish an efficient syntrophic enrichment for the application of PPB for nutrient removal and valorization of carbohydrate-based wastewater feedstocks. In Chapter 4 we evaluated the competitive and syntrophic mechanisms between FCB and PPB in mixed culture. PPB present slower growth rates (ca  $\mu$ =0.2 h<sup>-1</sup> (Conrad & Schlegel, 1977; T. Liu et al., 2014)) under glucose-fed conditions, even in pure cultures, and were therefore outcompeted by FCB ( $\mu$ =0.5 h<sup>-1</sup> (Rombouts et al., 2019a) in batch cultures. In continuous cultures an appropriate dilution rate of 0.04 h<sup>-1</sup> and pH regulation to 7.0 allowed the enrichment of PPB (up to 30%) in glucose-rich environments by enabling their metabolic coupling with FCB. The combination of the fermentative metabolism of FCB and photoorganoheterotrophic metabolism of PPB can become attractive to treat and valorize fermentable agri-food wastewater. Different configurations of singlesludge or two-sludge processes can be designed depending on treatment and valorization targets. If the aim is to valorize PPB biomass and/or intracellular products like single-cell proteins or photopigments, a two-sludge process can be efficient to first pre-ferment the carbohydrates into VFAs and feed them to a second SBR to produce a highly enriched PPB biomass.

#### 6.4 Light intensity governs the physiology of PPB enrichments

PPB utilize light as energy source, and VFAs as electron donors and as carbon source to build biomass. The quantum requirements for PPB have been evaluated in pure cultures, but in mixed cultures the limits of irradiation in PPB ecophysiology had not been clarified. In Chapter 5 we evaluated the robustness of a PPB enrichment in an acetate-fed (*ca* 1 g COD  $L^{-1}$ ) SBR fed exposed to decreasing IR light intensities from  $350 \text{ W} \text{ m}^2$  to darkness. We concluded that PPB can still be highly enriched at about 80% of the community also at very low light intensities, close to 1 W m<sup>-2</sup> (*i.e.*, highlighting the robustness of selective enrichments of PPB with VFAs and IR light), while the effective biomass growth is substantially lower (0.22  $h^{-1}$  at high light intensities and 0.01  $h^{-1}$  at low light intensities). Two different windows of operation were identified in function of IR light and substrate availabilities. Under incident irradiances higher than 87 W m<sup>-2</sup> or 10  $\mu$ mol<sub>photons</sub> s<sup>-1</sup> g<sub>DW</sub><sup>-1</sup>, the performance of the biosystem is governed by acetate limitation. Below this value, light limitation prevails. Under anaerobic conditions, PPB compete for acetate mainly with acetoclastic methanogenic organisms, but which present lower growth rates. PPB take advantage of the ability to derive energy from light, and compensate the lack of irradiation by increasing the production of photosynthetic complexes under low light intensities.

#### 6.5 PPB are promising organisms for water resource recovery

From the extensive investigations performed on PPB ecology and ecophysiology from pure to mixed cultures in this thesis, the following six key conclusions can be drawn:

1) The selective enrichment of PPB is enhanced by anaerobic conditions, IR light, VFA supply: pre-fermenting wastewater feeds can be efficient to selectively produce a PPB biomass in mixed-culture process.

2) When enriched and stabilized under these conditions, PPB form an extremely robust and resilient community.

3) SBR operation helps densify PPB bioaggregates and produce a concentrated, dense fast-settling biomass that enhances volumetric conversion rates, drives full nutrient removal in a single-stage process, and facilitates solid/liquid separation and downstream processing.

4) Light management is crucial for PPB growth and simultaneous nutrient removal: below 10  $\mu$ mol photons  $g_{DW}^{-1}$  PPB become light-limited, and growth is substantially limited.

5) PPB can establish syntrophic interactions with fermentative organisms for the conversion and valorization of carbohydrate-based wastewater feedstocks under careful operation of reactor regimes (*e.g.*, by tuning the dilution rate in a single-stage continuous-flow system or by separating the fermentation and photoorganoheterotrophic processes in two SBRs).

Overall, PPB are promising organisms for combined wastewater treatment and resource recovery.

Compared to other recovery schemes, PPB present some advantages: *i*) PPB get their energy from IR radiation, even at very low radiations. They assimilate almost all the carbon and electron sources for biomass growth. This is particularly important for bioproduct formation and recovery, where the biomass production needs to be maximized; *ii*) thanks to their hyper-versatile metabolism they can grow on numerous substrates. PPB grow preferentially as photoheterotrophs, and compared to microalgal processes, they do not require symbiosis with chemotrophic bacteria to produce CO<sub>2</sub>. *Iii*) PPB are a robust community, that can easily be enriched and aggregated when fed with VFAs and under IR light irradiation. Under these conditions, contaminations and culture crashes are prevented, and a continuity of the enrichment is assured. Iv) The high nutrient removal abilities of PPB can be combined with the production of high-value compounds, like PHAs, H<sub>2</sub>, single-cell proteins or photopigments.

Factually, PPB-based processes can be implemented for water resource recovery. In particular, they can be considered as a new-generation biophotoprocesses, that follows the principles of circular economy. They can be implemented for closed- loop biotechnology, allowing an efficient photo-based nutrient capture in wastewaters combined with a resource upcycle.

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# Outlook

The metabolic versatility of PPB confers them a substantial ecological advantage that has yet to be fully harnessed for environmental biotechnology and biorefinery services. While PPB-based wastewater treatment processes are promising, their development is still at their early stages. Some recent pilot-scale industrial applications exemplify a growing interest for PPB-based WWT and recovery. However, some considerations must be taken for up-scaling mixed-cultures processes, by making a strong link to the ecology and ecophysiology of PPB.

### Considerations for scaling up PPB-based processes

#### 1.1 Selection of the carbon and electron sources

Some considerations must be taken to efficiently exploit the extreme metabolic versatility of PPB and to enrich them. The supply of different carbon sources and electron donors is known to impact PPB biomass growth and yields. For example, a mixture of VFAs resulted in higher biomass yields and growth rates compared to the supply of only one compound (Alloul et al., 2019). On the other hand, the use of carbohydrates-rich waters resulted in a low PPB enrichment (30% vs. 90%) compared to VFAs feed, and a residual carbon concentration (*ca* 50 C-mmol L<sup>-1</sup>) (Chapter 4). In sugar-rich streams, the use of PPB communities is probably not the most efficient process for total carbon and nutrient removal. To enrich a PPB biomass, is advisable to use two separate treatment units: one first fermentation tank enriched in FCB, and a second unit enriched in PPB, to achieve full nutrient removal and a high enrichment of high-value PPB biomass. The use of PPB in a secondary unit, following a pre-fermentation process can be very efficient to convert carbohydrates into VFAs and supply the latter to selectively foster PPB biomass growth and nutrient capture.

#### 1.2 Selection of light

Light is arguably the foremost important parameter to grow phototrophic cultures, and a careful analysis of the light type and intensity must be considered in regard to an upscale process.

#### 1.2.1 Homogenization of the units

Light can be described through photometric (illuminance), radiometric (irradiance) and quantum units (photons flux density). In literature, the selection of the unit depends on the discipline, tradition, instrument availability and publication. Some of the units can be directly converted into other units (*i.e.*, lux and lumen). In other cases, the conversion between units is dependent on the light wavelength and energy input of the photons (*i.e.*, W m<sup>-2</sup> and mol phot m<sup>-2</sup> s<sup>-1</sup>), but the information about the light spectrum is often not provided. The problem of unit uniformity has been identified since the 1980's, and attempt

have been made to provide conversion tables (Thimijan & Heins, 1983). However, up to the present day, the scientific community has not yet agreed on a common unit to describe light supply in photobioprocesses. This complicates the comparison and the reproduction of the results.

To have a fast and efficient development of PPB-based and other phototrophic biotechnologies, it is necessary to harmonize the way to measure and report light inputs. One key parameter and unit for light should be set for scale up. Furthermore, light intensity is sometimes calculated to the source, without considering the biomass concentration in the biosystem. This way, attenuation phenomena are not evaluated, and the light available to the cultures is overestimated. For engineering practice, it is advisable to measure light as specific light supply rate ( $\mu$ mol<sub>photons</sub> s<sup>-1</sup> g<sub>DW</sub><sup>-1</sup>), as light attenuation due to biomass concentration and light energy are considered. With this unit, it is not necessary to specify the type of light emission, and allows to directly link irradiance with microbial kinetic parameters.

#### 1.2.2 Reactor geometry

Light attenuation in the photobioreactors is a crucial parameter to estimate. Several studies have evaluated light attenuation in phototrophic systems, and numerous models have been developed to describe phototrophic growth in function of light attenuation. Light penetration in a liquid medium is related to the path length, the biomass concentration and the photopigments content: at high biomass concentrations, such as achieved in bioreactors of this thesis, light penetrates only a few centimeters.

Under bench-scale conditions, an accurate control of light is possible, thanks to reactor geometry and operation (*i.e.* by using continuous cultures with high dilution rates, Chapter 4). In pilot and full-scale systems, reactor geometry might be a limiting factor for the development of PPB-based and phototrophic technologies in general.

Some bench-scale systems utilize membrane reactors (Hülsen et al., 2014), where PPB biofilm are grown attached to a surface. This allows an efficient solid/liquid separation, ideal for wastewater treatment. However, this type of system might not be efficient for biomass recovery, increasing the energy costs of the separation (Fasaei et al., 2018). In this thesis, we proposed the use of an anchor stirrer equipped with silicon blades to remove the biofilm on the reactor walls. It was efficient to clean-up wall biofilm formation simultaneous to stirred-tank reactor operation at bench scale. At pilot / full scale, such system might be too expensive.

Microalgae are often cultivated in tubular photobioreactors, to maximize the light penetration. A similar approach has been taken to cultivate PPB at bench-scale (Carlozzi et al., 2006). However, PPB aggregate and form thick biofilms on the reactor walls (Chapter 2). This can prevent light penetration, and consequently the development. In the present pilot plants, PPB are enriched in

tanks, ponds or raceway reactors (Deep Purple project; SEMiLLA). This operational condition maximizes the surface exposed to light compared to the reactor volume, therefore allowing a higher irradiation of the PPB biomass. Further studies on the ideal reactor geometry for PPB cultivation are needed, to combine light penetration with biomass growth and aggregation, while keeping an anaerobic or microaerophilic environment.

#### 1.2.3 Light type

The type of wavelengths provided to the cultures is as important as the light intensity. PPB take advantage of low-energy photons irradiations, in the near and far infrared spectrum depending on the species.

Artificial LED irradiation in microalgal cultures can cost up to 11 \$  $kg_{DW}^{-1}$ , factually decreasing the applicability of the process to larger scales (Blanken et al., 2013). For PPB, it has been calculated that the LED-lamps irradiation cost of \$ 1.9  $g_{DW}^{-1}$  would still be higher than the potential revenues of PPB products (Capson-Tojo et al., 2020).

To selectively provide IR light to enrich for PPB, several options can be considered: i) Use of IR emitting lamps: expensive and with a very short half-life; *ii*) Use of LEDs: cheaper compared to IR lamps, LEDs emit only at specific wavelengths. In pure cultures, the choice of a specific wavelength will promote either growth either photopigments formation, depending on the species (Kuo et al., 2012; Zhou et al., 2015); in mixed cultures it will influence the community selection processes. With a combined use of different LEDs, it is possible to achieve almost a full light spectrum, potentially allowing the simultaneous enrichment of several PPB species. *iii*) Filter natural light: IR light account for almost 50% of the solar radiation. Selecting for long wavelengths would prevent the growth of green-blue photorophs in PPB reactors. Using the filters to seal the reactor would also allow the creation of an anaerobic environment, promoting PPB growth. The use of light filters to seal ponds or raceway reactors should be further evaluated, especially in geographical areas naturally subjected to high solar radiation.

### **Bioproducts recovery**

#### 2.1 PPB bioproducts currently studied: H<sub>2</sub>, PHA, proteins

In recent years, the quest for 'green' products has increased. Current research on the valorization of PPB biomass focusses mainly on the production of bioplastics (PHA), biogas ( $H_2$ ), and single cell proteins.

PHAs are a promising substitute to oil-based plastics. They are recyclable and biodegradable. PHA-producing bacteria can store up to 90% of the cell weight as biopolymer, therefore being efficient biorefineries (Johnson et al., 2009). PHAs have found commercial use and are currently used also to produce fully

biodegradable plastic bottles and packaging systems (Bugnicourt et al., 2014). Nonetheless, the stability of the bioplastic depends on the form and purity of the compounds. Microbial metabolism can be engineered to tune the production of either poly- $\beta$ -hydroxybutyrate or poly- $\beta$ -hydroxyvalerate in axenic cultures, increasing the production costs due to sterility requirements (Chen & Jiang, 2017). Alternatives that do not require the separation of the two compounds are currently explored, e.g., for self-healing concrete (Vermeer et al., 2021). PHA production in PPB pure cultures has been maximized to about 90% of the

biomass dry weight (Montiel-Corona & Buitrón, 2021). In PPB mixed cultures, the highest PHA production reported was equivalent to only the 20% to the biomass dry weight (Luongo et al., 2017). The downstream process of extraction and purification is the limiting factor for a broad PHAs use, being laborious and expensive (Rodriguez-Perez et al., 2018). The relatively low PHA production from PPB mixed cultures reduce the interest in PPB for application.

 $H_2$  is an energy carrier, that rose interest for its potential as biofuel. Biohydrogen production utilizes renewable resources, in a carbon-neutral process with low environmental impact (Wu et al., 2012). PPB produce hydrogen gas to dissipate electron excess via a non-growth-related processes. The  $H_2$  production by PPB means achieved relatively high yield over substrate (up 78% of the theoretical yield, extensively reviewed in Akkerman et al., 2002). However, they have low volumetric production rates and require large volumes of substrates and large surface area photobioreactors (Hallenbeck, 2014). Despite the substantial research on photofermentation, the TLR for this process is still at level 1-3.

Single-cell protein designates microorganisms such as bacteria, fungi or microalgae with high-protein content that can be further used as feed for aquaculture, animal feed or even human consumption (Anupama & Ravindra, 2000; Daniel Puyol et al., 2017). PPB have a high protein content (up to 70% of the dry weight) compared to other bacteria (30-60%) (Honda et al., 2006; Nasseri et al., 2011). Combined with high biomass yields over substrate, a constitutive production of antioxidants, and lack of toxicity, PPB are an efficient source of protein for animals and fish feedstocks. Single-cell protein production is currently probably the most promising application of PPB mixed cultures, as it allows a direct use of the biomass.

#### 2.2 Perspective applications: Carotenoids

Carotenoids are a class of photoactive molecules, present in almost all phototrophic organisms, from higher plants to bacteria. They are accessory pigments, harvesting light at 350-450 nm and transferring the energy to the reaction centres (Cerullo et al., 2003; Michael T. Madigan et al., 2015). Additionally, carotenoids protect the cells from photooxidation (Sipka & Maróti, 2018).

Carotenoids are currently used as food additives, in pharmaceutical preparations and the cosmetic industry (Barreiro & Barredo, 2018a). They are harnessed both for their antioxidant properties as well as for their intense yellow-red coloration, making them suitable as dyes (Cardoso et al., 2017). Carotenoids are remarkable not only for their role in light harvesting, but also for their structural importance, antioxidant properties, and action in preventing photo-damage. In humans, carotenoids reduce the risk of cardiovascular disfunctions and age-related diseases (Rao & Rao, 2007). It is estimated that the world carotenoid market will reach 6222.6 metric tons per year and a worth of \$ 2.0 billions by 2022, with a yearly increase in the demand equal to 5.7 % from 2017 (Bogacz-Radomska et al., 2020).

Despite the growing demand, production and extraction of carotenoids remains a challenge. In 2019, 80-90% of the world carotenoid production was fulfilled by chemical synthesis (Saini & Keum, 2019). However, the demand for naturally produced compounds is increasing among the consumers. Natural carotenoids are primarily extracted from plants, flowers, fruits, and seeds. Due to the high costs, seasonality, and the geographic determinants of the plant production, microbial alternatives are explored (Bogacz-Radomska et al., 2020). Microbial platforms for carotenoids production are gaining interest due to the lower costs, linked to lower land requirements, year-round harvest and higher growth rate compared to plants (Barreiro & Barredo, 2018b). Only three microalgae genera are currently used for industrial carotenoid production, Dunaliella, Haematococcus, and Scenedesmus, but several other microorganisms, including fungi, are currently under examination as potential biofactories (Bonet et al., 2016).

Downstream processing is a bottleneck in the carotenoids production. Traditional extraction methods include the use of organic solvents, such as dichloromethane, hexane, methanol, ethanol, dimethyl ether, diethyl ether, toluene, 2-propanol, n-butanol, heptane, acetonitrile, tetradecane, dodecane, hexadecane, or tetrahydrofuran, some of them being problematic / forbidden for implementation at industrial scale (The European Parliament and the Council, 2018). Independently from the extraction yields, these compounds are often expensive, and considered unsafe for human health and the environment. Emerging technologies, such as ionic liquid extraction, enzyme-assisted extraction, and supercritical CO<sub>2</sub> (SC-CO<sub>2</sub>) extraction gain interest as 'green' alternatives for carotenoids recovery (Richmond, 2003). SC-CO<sub>2</sub> extraction utilizes CO<sub>2</sub> at high pressure and constant temperature to extract lipophilic molecules. Its usage is advisable over organic solvents, as it is not environmentally toxic, and its disposal is safe. The density of the SC-CO<sub>2</sub>, and therefore its polarity, can be controlled by tuning temperature and pressure conditions. This allows the extraction of different type of molecules (Baskar et al., 2019). The extraction at low temperatures (31 °C) avoids degradation and maintains the functional structure of thermosensitive compounds, like carotenoids (Shi et al., 2007). In algal systems, SC-CO<sub>2</sub> has been repeatedly applied for the extraction of carotenoids and photopigments (Goto et al., 2015).  $CO_2$  has been used pure, or in combination with co-solvents, like ethanol, to increase the extraction yields (Da Porto et al., 2014).

In the context of a circular economy, the use of PPB as wastewater biofactories for carotenoids production has not been extensively explored yet. Under anaerobic conditions, PPB naturally produce carotenoids as part of their photosynthetic machinery. In PPB, carotenoids extraction has been performed only through organic solvents. In collaboration with the Hanze University of Applied Sciences in Groningen, The Netherlands, we evaluated the feasibility of a SC-CO<sub>2</sub> extraction for PPB carotenoids. We cultivated Rhodopseudomonas palustris in an axenic, SBR system. We provided 0.9 g L<sup>-1</sup> of sodium acetate as C-source, and high light intensities (300 W m<sup>-2</sup>) as energy sources. High irradiation was chosen to promote biomass growth rather than photopigment production. PPB biomass was separated from the liquid phase by centrifugation, freeze-dried overnight, and grinded to powder. Two different extractions were attempted, with and without addition of ethanol as co-solvent (Table 1). The extract was then collected by vacuum evaporation. A 1% decrease in mass was reported after extraction with pure  $CO_2$ . The extraction yield of carotenoids from the PPB biomass (i.e., mass extracted / mass PPB (dry weight) before extraction \* 100) was 1.57% m/m. An overall mass recovery of 100.67% was achieved. When ethanol was used as co-solvent, the extraction yield decreased to 0.5%. The mass of the PPB sample after extraction was higher than before the extraction, potentially indicating the presence of residual ethanol in the sample.



Wet biomass Dry biomass



Extract after SC-CO<sub>2</sub>

Figure 1: Wet and dry biomass before extraction. Photopigment extracted with SC-CO<sub>2</sub> method

	SC-CO <sub>2</sub> extraction	SC-CO <sub>2</sub>
	1	extraction 2
Pressure (bar)	300	300
Temperature (°C)	40	40
CO <sub>2</sub> flow rate (g min <sup>-1</sup> )	20	20
Extraction duration (h) Ethanol	3	3
(cosolvent) flow rate		
(ml min <sup>-1</sup> )	-	0.2
Post-flushing under		
pressurization*	No	Yes
Post-flushing at ambient**	Yes	Yes

Table 1: SC-CO<sub>2</sub> extraction conditions for carotenoids recovery from PPB biomass

\* flushing the extraction unit (in the absence of extractor) at 300 bar; 40 °C; 20 g/min  $CO_2$ ; 0.2 ml/min EtOH; 30 min

\*\* flushing the extraction unit (in the absence of extractor) at ambient condition; 2 ml/min EtOH; 30 min

Spectrophotometric analysis was used to qualitatively determine the extracted carotenoids. Volumes of 200  $\mu$ L of extracted pigments dissolved in ethanol were used to perform a wavelengths scan, from 240 to 999 nm. Ethanol was used as measuring blank. One clear peak was visible at 470 nm (Figure 2). The absence of peaks in the 600-700 nm area indicated the absence of bacteriochlorophyll pigments, and therefore a selective extraction of carotenoids.



Figure 8: Spectrophotometric analysis of the SC-CO2 extraction products. One clear peak is visible at 470 nm, but no peak is present in the 600-700 nm area, indicating the selective extraction of carotenoids.

A precise determination of the main carotenoids extracted was not possible without HPLC or HPLC-MS analyses. In fact, the carotenoids absorption peaks are very close one to the other, and spectrophotometric analysis doesn't provide sufficient resolution (Clementson & Wojtasiewicz, 2019). As ethanol is a slightly polar compound, and the extract solution was homogeneous, we can hypnotize the presence of carotenoids with polar components. Further and more in-depth analysis of the exact composition of the SC-CO<sub>2</sub> extract should be done, to assess the quality and quantity of the singular components.

The market for naturally produced carotenoids is continuously increasing. As an example, lycopene is one of the most commonly used food-grade pigments (Nigam & Luke, 2016).

Lycopene had a market worth of 107 million \$ in 2020, and it is expected to grow to 187 million \$ by 2030 (Bogacz-Radomska et al., 2020). The increasing demand for naturally produced compounds and additives can be filled by the microbial production of carotenoids without using plant-based technologies. In this regard, our results showed that PPB can produce almost 8 mg of carotenoids per gram of dry weight biomass (Chapter 6). As a comparison, tomato produce about 24.1 mg of carotenoids per 100 g of fresh product. Considering that the water percentage in tomatoes is 95% of the fresh product weight, the carotenoids content corresponds to 4.8 mg per g of dry product weight. With a carotenoids production almost 2 times higher than tomato, PPB are promising candidates as carotenoids biofactories.

Coupling carotenoids production with residual streams treatment has the potential to recover high-value products while recovering natural resources. European authorities have defined a framework to reclassify waste streams as secondary raw materials. As stated, waste can be re-converted into resources if: *i) the substance or object is commonly used for specific purposes*;

there is an existing market or demand for the substance; ii) the use is lawful (the substance or object fulfils the technical requirements for the specific purposes and meets the existing legislation and standards applicable to products); iii) its use will not lead to overall adverse environmental or human health impacts ".

In this framework and as emerged from this thesis, PPB can be used to treat VFA-rich aqueous waste streams. The high biomass concentration, and their natural propension to aggregate can be advantageous for intracellular high-value product formation and separation. Light sources and intensity of irradiation should be carefully considered for process design, specifically for carotenoids production, when biomass formation and photopigment content in the cells have a tradeoff.

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## Acknowledgments

It is a common metaphor to describe a Ph.D. as a rollercoaster.

Looking back, I would rather describe it as a journey through a jungle: challenging, and impossible to undertake alone.

For this reason, I would like to warmly thank all the people that accompanied me during these four years of scientific and personal growth.

My first and biggest 'thank you' goes to my promotors, David and Mark, for giving me the opportunity to work in your group.

**David**, thank you for believing in me when I was nothing but a scared shy student. For all you thaught me. For your guidance, but most of all, thank you for all the freedom you gave me to develop my project.

**Mark**, thank you for all the scientific discussions we had. I learnt a lot from you, and there is nothing more important in life than keep learning. Aside from science, thank you for opening my eyes on the importance of being curious and passionate while doing research. I took it for granted, but it is not.

Secondly, thanks to all the students that worked with me in this project. To the students that I had the pleasure to directly supervise, **Heleen**, **Jeong Hoon**, **Guillaume**, **Mythili**, **Viktor**, **Daniela**, **Casper**, and to the students I have interacted with, **Berber**, **Charlotte**, **Reetu**, **Luuk**, **Wesley**. You challenged me to do the best I could to be a good mentor. You thaught me a lot, and this thesis would have been very different without your help. Thank you.

Thank you to all the incredible scientists that I had the pleasure to interact with and learn from during these years: Abbas, Bing, Cristian, Daniel, Dimitry, Gijs, Meng, Martin, Merle, Pascale, Robbert, Siegfried, Sirous, Yuemei. Your expertise and your advice have been precious.

Thanks to the whole EBT team.

In particular, to our technicians' team: **Rob**, **Dirk** and **Marcel** for the incredibly useful assistance in reactor setup, and for your endless patience when I was asking for the impossible. **Apilena**, **Astrid**, **Ben**, **Cor**, **Jannie**, **Marcel**, **Max**, **Miranda**, **Patricia**, **Zita**: thank you so much for the technical support.

To all the EBT fellows, Aina, Ali, Ana-Maria, Chris, Dana, Danny, Diana, Felipe, Francesc, Gerben (and thanks a lot for all your help!), Hugo, Jure, Laura, Lemin, Leonor, Marissa, Maxim, Monica, Morez, Nina, Rodoula, Sam, Simon, Stefan, Timmy, Viktor, Vojtech, for making EBT an eclectic, dynamic, and fun place to work in. It has been great sharing the labs with you. To the BT-PhD committee members: Albert, Anna, Britte, Chema, Florence, Stefan: what a team we were! It has been incredibly fun organizing with you serious and less serious events.

My office mates. **Michele** and **Jules**: from the moment I arrived, you guys took me under your wings, and you have been my mentors ever since. Thank you for all the discussions we had, you have been an inspiration for me. **Andres, Rhodi**, **Maria-Paula, Ruben**, **Ahmed**, thanks for making our office a fun place to be.

Being an expat made me realize the importance of having a solid network of friends you can count on at any time. The memories we created together will forever remain with me. I am not good with words, and I know that this short paragraph is not enough to express all the affection, gratitude and love I have for you. You are a piece of my hearth.

A special acknowledgment to my paranymphs, David and Alexandra.

**David**, you have been a happiness pillar for me during these years, and I can't imagine going through the PhD without you. You have been the most amazing friend, housemate, and colleague. We shared the best and the worst moment, and I couldn't be more grateful for all the time together. Gracias, primo.

Alexandra, from the first time we talked, I've always found funny to see how different yet similar we are. I am just lucky to have met you, because I found a wonderful human, a great friend, and an incredible force of nature. You rock girl.

To my Delft family, because with you I always felt at home. **Daniel, Fabio**, **Florence, Georg, Ingrid, Maria-Paula, Mariana, Marina, Michele, Sergio**: I couldn't be more grateful for meeting you. We have been 'partners in crime', and I cherish every moment together. Words can't properly describe how much I love you all, so I will just say 'thank you people, lots of love'.

**Mamma** e **Papà**, grazie per avermi sempre supportata, a distanza, in qualsiasi parte del mondo io fossi e qualsiasi cosa decidessi di fare. Spero di avervi resi fieri di me, e spero di continuare a farlo sempre.

Andrea, you have always been by my side, even while houndreds of km apart. You supported me during the darkest moments, and you were there in the happiest times. You challenged me. You made me grow. I am extremely grateful for having you in my life. Grazie Amore, ti amo.

# Biography



Marta Cerruti was born in Venice, Italy, on June 19th 1992.

In 2013 she completed her Bachelor's degree in Biotechnology from University of Torino, Italy. Passionate about environmental preservation, she started her MSc degree program in Plant Biotechnology, from University of Turin, Italy.

During her studies, she discovered a particular curiosity for microbial processes, that she explored first during an internship in plant-microbe interactions at INRA Institute, France, and then during her MSc end-project on a model organism found in wastewater treatment plants, at EPFL, Switzerland.

In 2017 she started her PhD program at TU Delft, the Netherlands, on purple bacteria for wastewater treatment, combining her interest in phototrophic systems, microbial communities and environmental preservation. The four years of work are summarized in this book.

After the PhD, she started working at ETH Zurich, Switzerland, as a postdoctoral researcher on gut microbial ecology.

# List of publications

2021 Synthrophy between fermentative and purple phototrophic bacteria for carbohydrate-based wastewater treatment; bioRxiv https://doi.org/10.1101/2021.05.13.444055 M.Cerruti, G. Crosset-Perrotin, M. Ananth, J. Rombouts, D. Weissbrodt

Plant-Wide Systems Microbiology for the Wastewater Industry;
Environmental Science: Water Research & Technology
https://doi.org/10.1039/D1EW00231G
M. Cerruti, B. Guo, R. Delatolla, N. de Jonge, A. Hommes - de Vos van Steenwijk,
P. Kadota, C. Lawson, T. Mao, M. Oosterkamp, F. Sabba, M. Stokholm-Bjerregaard,
I. Watson, D. Frigon, D. Weissbrodt

Control tools to selectively produce purple bacteria for microbial protein in raceway reactors; Environmental Science and Technology https://doi.org/10.1021/acs.est.0c08204 A. Alloul, M.Cerruti, D. Adamczyk, D. Weissbrodt, S. Vlaeminck

2020 Enrichment and Aggregation of Purple Non-sulfur Bacteria in a Mixed-Culture Sequencing-Batch Photobioreactor for Biological Nutrient Removal From Wastewater; Frontiers in Bioengineering and Biotechnology https://doi.org/10.3389/fbioe.2020.557234 M. Cerruti, B. Stevens, S. Ebrahimi, A. Alloul, S.E. Vlaeminck, D. Weissbrodt

> Effects of light/dark diel cycles on the photoorganoheterotrophic metabolism of *Rhodopseudomonas palustris* for differential electron allocation to PHAs and H<sub>2</sub>; bioRxiv https://doi.org/10.1101/2020.08.19.258533 M. Cerruti, H. Ouboter, V. Chasna, M. van Loosdrecht, C. Picioreanu, D.Weissbrodt

2019 Drivers of Bioaggregation from flocs to biofilms and granular sludge; Environmental Science: Water Research & Technology https://doi.org/10.1039/C9EW00450E H.Aqeel, D.Weissbrodt, M. Cerruti, G. Wolfaardt, B.M. Wilén, S.N. Liss

## Conference contributions

2020: **IWA Biofilms,** Indiana, USA (online conference) – oral presentation Tracking metabolic versatility in biofilms: modelling photoorganoheterotrophic and photolithoauttrophic niches of a single purple phototrophic bacterium

#### 2019 **KNWM Microbial Ecology Fall Meeting,** Wageningen, the Netherlands – oral presentation Purple non-sulfur phototrophic bacteria: role of reactor regime and light intensities in the association with fermenters for sugar-based wastewater treatment

**IWA MEWE** Hiroshima, Japan – poster presentation Substrate and light impacts on novel anaerobic associations of fermenters and purple non-sulfur bacteria for waste-based carbohydrate valorization

**IWAlgae** Valladolid, Spain – oral presentation Purple non-sulfur bacteria: can we drive association with fermenters to treat sugar-based aqueous waste?

**IWA Anaerobic digestion,** Delft, the Netherlands – poster presentation How does light intensity effect the physiology of purple bacteria and nutrient removal prom wastewater under anaerobic sequencing batch reactor regime?

**Swiss Microbial Ecology Meeting,** Lausanne, Switzerland – oral presentation Effects of light / dark diel cycles on the versatile metabolism of purple non-sulfur phototrophic bacteria and their translation for water resource recovery

2018 **IWA EcoSTP,** London, Canada – oral presentations Sequencing batch enrichment and metabolic drivers for water resource recovery using purple non-sulfur bacteria'

Introduction to ecogenomics and ecophysiology data and their integration with engineering

2017 Swiss Society for Microbiology annual joint meeting, Basel, Switzerland – poster presentation *Tetrasphaera japonica*, a model to study the metabolism of polyphosphate-accumulating organisms in real wastewater treatment plants?