

1	Enhancement of co-production of nutritional protein and carotenoids in Dunaliella
2	salina using a two-phase cultivation assisted by nitrogen level and light intensity
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### 19 Abstract

20 Microalga Dunaliella salina is known for its carotenogenesis. At the same time, it can also 21 produce high-quality protein. The optimal conditions for *D. salina* to co-produce 22 intracellular pools of both compounds, however, are yet unknown. This study investigated 23 a two-phase cultivation strategy to optimize combined high-quality protein and 24 carotenoid production of *D. salina*. In phase-one, a gradient of nitrogen concentrations 25 was tested. In phase-two, effects of nitrogen pulse and high illumination were tested. 26 Results reveal optimized protein quantity, quality (expressed as essential amino acid index 27 EAAI) and carotenoids content in a two-phase cultivation, where short nitrogen starvation 28 in phase-one was followed by high illumination during phase-two. Adopting this strategy, 29 productivities of protein, EAA and carotenoids reached 22, 7 and 3 mg/L/d, respectively, with an EAAI of 1.1. The quality of this biomass surpasses FAO/WHO standard for human 30 31 nutrition, and the observed level of  $\beta$ -carotene presents high antioxidant pro-vitamin A 32 activity.

# 33 Key words

- 34 Single-cell protein; pigment; nitrogen limitation; food; microalgae
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#### 39 **1. Introduction**

40 The global population will reach 9.3 billion by 2050, with 6.4 billion of people in urban 41 areas (Corcoran et al., 2010). The societal changes of both population and living standard 42 are leading to 50% increase of protein demand, and even 82% and 102% increase of dairy 43 and meat products by 2050, respectively (Boland et al., 2013). Along with protein 44 shortage, the deficiency of functional nutrients in food, like  $\beta$ -carotene, are causing severe 45 health problems for human. Specifically,  $\beta$ -carotene is essential for the human body due 46 to its antioxidant pro-vitamin A activity, and insufficient uptake of  $\beta$ -carotene will lead to 47 severe vitamin A deficiency, prompting human blindness and affecting immune response 48 systems (Sommer, 2001). Currently,  $\beta$ -carotene and vitamin A deficiency have become a 49 major public health concern in more than 70 countries (Sommer, 2001). To sustainably 50 fulfill the protein gap for human consumption, novel protein sources such as microalgae 51 are considered important contributions (Muys et al., 2019). Moreover, microalgae with 52 elevated carotenogenesis can further increase nutritional quality by preventing vitamin A 53 deficiency. Particularly, natural β-carotene found in microalgae, fruits and vegetables has 54 the advantage of its mixed stereoisomers of all-*trans* and 9-*cis* β-carotene, which are more 55 fat-soluble and less crystallizable than synthetic  $\beta$ -carotene (all-trans  $\beta$ -carotene) (Ben-56 Amotz, 1993). By consuming carotenoid-rich diet, potentially lower incidence of various 57 kinds of cancer can be expected (Ben-Amotz, 1993) 58 Microalgal production is conventionally aiming at one specific target, such as biomass

59 from *Chlorella*, protein from *Spirulina* and β-carotene from *Dunaliella* (Ben-Amotz, 1993).

Consequently, individual production lines are required to achieve production of multiple
target compounds. If one microalgal species possesses the ability to optimally co-produce
both high-quality protein as well as β-carotene at the same time, the efficiency in
production of multiple high-value products can be substantially increased.

64 It is well established that the microalga Dunaliella salina is one of the best sources of 65 natural  $\beta$ -carotene, which can contribute up to 14% of the cell dry weight (Aasen et al., 66 1969). Up to date, many studies have shown that stress conditions are major factors 67 enhancing the accumulation of  $\beta$ -carotene in *D. salina*, such as high light intensity, high 68 salinity, extreme temperatures and nitrogen (N) deficiency, as a protective mechanism to 69 prevent cellular damages e.g. photo-damage (Lamers et al., 2008; Marín et al., 1998). 70 While carotenogenesis by *D. salina* has been widely studied, its potential as protein source 71 has drawn limited attention. In fact, D. salina can display high protein content (up to 80% 72 of ash free dry weight), which is subjected to different cultivation conditions and growth 73 phases (Sui and Vlaeminck, 2019). Furthermore, the essential amino acid (EAA) content of 74 D. salina fulfills human requirement as indicated by FAO/WHO reference, which defines its 75 high-quality protein profile (Becker, 2007; Sui et al., 2019). When comparing with other 76 microalgae such as Chlorella and Spirulina, both protein and EAA content of D. salina have 77 either comparable or even superior values (Becker, 2007; Muys et al., 2019; Sui et al., 78 2019). Based on the characteristics of *D. salina*, it strongly appears to be a valuable 79 candidate as novel food source, containing both high-quality protein and  $\beta$ -carotene.

80 Based on both lab- and large-scale experience with cultivating *Dunaliella* for  $\beta$ -carotene 81 production, a two-phase cultivation system was proven to be successful, and has been 82 applied by commercial producers (Ben-Amotz, 1995). The concept of such two-phase 83 systems is to increase microalgal biomass level at phase-one with optimum growth 84 conditions, and to induce  $\beta$ -carotene production at phase-two with enhanced stress 85 conditions, e.g. high light intensity and N deficiency (Ben-Amotz, 1995). Moreover, it has 86 been reported that stress conditions can also contribute to the up-regulation of EAA levels 87 in plants and microalgae (Galili et al., 2016; Obata and Fernie, 2012). Specifically, high light 88 intensity and N deficiency have been shown to enhance the production of EAA, especially 89 lysine, threonine, methionine, valine and isoleucine in microalgae (Kiyota et al., 2014; Zhang et al., 2016). For D. salina, there is no report on the regulation of EAA affected by 90 91 cultivation conditions, and the combined production of two main nutritional compounds 92 from D. salina has not been studied yet (Sui et al., 2019). Instead of a process targeting 93 either  $\beta$ -carotene or protein production, a two-phase cultivation approach adopting 94 sequential stress conditions may prove to be an effective way to boost both high-quality 95 protein and  $\beta$ -carotene production from *D. salina*.

96 In this study, the impact of a gradient in N availability together with N pulses and high light

97 intensities on the dynamics of biomass, protein, EAA and  $\beta$ -carotene production in *D*.

98 salina have been explored over different growth phases in a two-phase cultivation

99 approach. This study intends to demonstrate for the first time an optimized cultivation

100 condition and harvest regime for the maximum production of both EAA and β-carotene of
101 *D. salina*.

# **2. Materials and methods**

103	2.1 Microalgal strain, two-phase cultivation approach and cultivation conditions
104	D. salina CCAP 19/18 was purchased from Culture Collection of Algae and Protozoa (CCAP,
105	Scotland, UK). Sterilized Modified Johnson's medium (Borowitzka, 1988) as standard
106	medium for <i>D. salina</i> with different N levels was used for cultivation. Six treatments
107	covering a gradient of N concentrations were tested in phase-one (N1 to N6; Table 1).
108	When the algae reached stationary phase, each treatment was divided into two further
109	conditions for phase-two: high N (NN1 to NN6) and high light intensity (NL1 to NL6; Table
110	1). All treatments were performed in triplicates for both phases. Triplicates in phase-two
111	were derived from the pooled triplicates of corresponding treatments in phase-one. The
112	initial biomass concentration in phase-one was around 40,000 cells/mL. Experiments were
113	conducted in a water bath with a controlled temperature of 25°C. Continuous light was
114	provided by fluorescent tubes at an incident irradiance of 70 $\mu$ mol photons/m <sup>2</sup> /s for
115	standard conditions, and 110 $\mu mol$ photons/m²/s for high irradiance conditions (Philips TL-
116	D 30W/33-640, the Netherlands). Mixing and aeration were given by a mixture of pre-
117	humidified air and 2% CO <sub>2</sub> . Although pH was not controlled, it was found stable (7.8-8)
118	over the experiment due to the $CO_2$ addition (Table 1).

119 2.2 Sample analyses, calculations and statistics

120	Daily samples from all treatments were analyzed directly for cell number and cell volume,
121	and preserved at -20°C for protein, carotenoids and EAA analyses at the end of the
122	experiment. A Multisizer 3 Coulter Counter was used for both cell number and volume
123	measurement. The protein content was determined following Markwell method, a
124	modified Lowry method with bovine serum albumin as standard (Markwell et al., 1978).
125	Total carotenoids content was measured according to Lichtenthaler, (1987) with 100%
126	acetone extraction:
127	Chlorophyll a $(mg/L) = 11.24 \times OD_{661.6} - 2.04 \times OD_{644.8}$
128	Chlorophyll b $(mg/L) = 20.13 \times OD_{644.8} - 4.19 \times OD_{661.6}$
129	Total carotenoids (mg/L)
130	$=\frac{1000 \times OD_{470} - 1.90 \times Chlorophyll a - 63.14 \times Chlorophyll b}{214}$
	$= \frac{1000 \times OD_{470} - 1.90 \times Chlorophyll a - 63.14 \times Chlorophyll b}{1000 \times OD_{470} - 1.90 \times Chlorophyll a}$
130	$=\frac{1000 \times OD_{470} - 1.90 \times Chlorophyll a - 63.14 \times Chlorophyll b}{214}$
130 131	$=\frac{1000 \times OD_{470} - 1.90 \times Chlorophyll a - 63.14 \times Chlorophyll b}{214}$ where OD <sub>661.6</sub> , OD <sub>644.8</sub> and OD <sub>470</sub> refer to the optical densities of the extracted supernatant
130 131 132	$= \frac{1000 \times OD_{470} - 1.90 \times Chlorophyll a - 63.14 \times Chlorophyll b}{214}$ where OD <sub>661.6</sub> , OD <sub>644.8</sub> and OD <sub>470</sub> refer to the optical densities of the extracted supernatant measured at 661.6 nm, 644.8 nm and 470 nm, respectively.
130 131 132 133	$= \frac{1000 \times OD_{470} - 1.90 \times Chlorophyll a - 63.14 \times Chlorophyll b}{214}$ where OD <sub>661.6</sub> , OD <sub>644.8</sub> and OD <sub>470</sub> refer to the optical densities of the extracted supernatant measured at 661.6 nm, 644.8 nm and 470 nm, respectively. To prepare for EAA analysis, samples were centrifuged (5,000 x g, 10 min), hydrolyzed (6M
130 131 132 133 134	$= \frac{1000 \times OD_{470} - 1.90 \times Chlorophyll a - 63.14 \times Chlorophyll b}{214}$ where OD <sub>661.6</sub> , OD <sub>644.8</sub> and OD <sub>470</sub> refer to the optical densities of the extracted supernatant measured at 661.6 nm, 644.8 nm and 470 nm, respectively. To prepare for EAA analysis, samples were centrifuged (5,000 x g, 10 min), hydrolyzed (6M HCl, 110°C, 24 hours) with vacuum and evaporated, after which samples were re-
130 131 132 133 134 135	$= \frac{1000 \times OD_{470} - 1.90 \times Chlorophyll a - 63.14 \times Chlorophyll b}{214}$ where OD <sub>661.6</sub> , OD <sub>644.8</sub> and OD <sub>470</sub> refer to the optical densities of the extracted supernatant measured at 661.6 nm, 644.8 nm and 470 nm, respectively. To prepare for EAA analysis, samples were centrifuged (5,000 x g, 10 min), hydrolyzed (6M HCl, 110°C, 24 hours) with vacuum and evaporated, after which samples were re- dissolving in 0.75mM HCl and stored at -20°C before analysis. For EAA determination, the

based on the EAA content with FAO/WHO EAA requirements for human as reference
(Oser, 1959; WHO/FAO/UNU Expert Consultation, 2007):

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$$EAAI = \sqrt[n]{\frac{aa1}{AA1} \times \frac{aa2}{AA2} \times \dots \times \frac{aan}{AAn}}$$

142 where *aan* and *AAn* are the EAA content over total protein content (mg EAA/g protein) in 143 the sample and FAO/WHO reference, respectively. An EAAI value of  $\geq$  1, 0.95-1, 0.86-0.95, 144 0.75-0.86 and  $\leq 0.75$  indicates its superior quality, high quality, good quality, useful quality 145 and inadequate quality, respectively (Kent et al., 2015). For comparison purposes, it has 146 been calculated that the EAAI value of egg and soybean are 1.65 and 1.34 separately 147 (Becker, 2007). 148 Total protein and carotenoids per liter of culture (mg/L) were defined as suspension 149 protein and carotenoids content. The protein, carotenoid and EAA productivity (mg/L/d) 150 were calculated from their suspension content (mg/L) divided by the time of cultivation 151 (days) at each sampling point. 152 Samples at the end of the two phases were analyzed for nitrate (NO<sub>3</sub>-), ammonium (NH<sub>4</sub>+) 153 and phosphate (PO<sub>4</sub><sup>3-</sup>) concentrations in the medium. As ammonium concentrations were

not detected in any treatment, they were not reported in this study. Filtered samples

- 155 were diluted with de-ionized water accordingly and a Seal QUAATRO Auto Analyzer (Seal
- 156 Analytical Inc., the Netherlands) was used for determination following standard methods
- 157 (APHA, 2012).

Multiple regression analysis in SPSS statistics 24 was used to compare data in Fig. 3. A
significance level p < 0.05 was considered as statistically different. All results were</li>
expressed as means ± standard deviations in tables and figures (apart from EAA and EAA
derived parameters). The values stated in the main text were without standard deviation
for better readability.

#### 163 **3. Results and discussion**

164 3.1 Biomass growth

165 Both the N level and light intensity greatly affected the microalgal growth. During phase-166 one, the microalgal cells reached different concentrations at stationary growth phase from 167 N1 to N4, ranging from approximately  $2 \times 10^5$  cells/mL to  $2.5 \times 10^6$  cells/mL. This was mostly contributed by the differences in initial N concentrations in the medium, which were 168 depleted at the end of phase-one (Table 2). From N4 to N6, the cell densities did not 169 170 change, despite different initial N concentrations (Fig 1A). Considering that there was still 171 N remaining in the medium after phase-one for N5 and N6, the cells in these treatments 172 were limited by another factor, while for N4 cells may have been co-limited. One likely 173 limiting factor can be phosphorus (P), as the residual P in the medium from N4 to N6 was 174 zero (Table 2). Besides, light limitation could occur as well, which will be discussed in more 175 detail later.

176 During phase-two, both higher N and higher light intensity had effect on biomass growth.

177 Specifically, from NN1 to NN3, cell numbers increased with extra N addition as a result

178 from their N starvation in phase-one. Differently, cell concentration from NN4 to NN6 did 179 not increase, and even slightly decreased after N addition, indicating that N was not the 180 limiting factor. Higher light intensity did not affect the cell concentration in NL1, most 181 likely due to the N scarcity reached during phase-one. For NL2 and NL3, higher light 182 intensity promoted cell growth, although N or P was limiting. It is possible that D. saline in 183 NL2 and NL3 might have stored both N and P inside the cell to be used for further growth under higher light intensity (Dortch et al., 1984). From NL4 to NL6, higher light intensity 184 185 did not affect much or slightly lowered the biomass, which demonstrates that light is not 186 the limiting factor. At this point, P concentration in the medium is expected to be the main 187 limiting factor which prohibited the further growth of cells (Table 2). This finding is in line 188 with previous work, where higher light intensity could not further boost cell densities of 189 Arthrospira platensis when P was depleted (Markou et al., 2012).

190 Besides cell densities, also cell sizes were subject to changes in response to the N and NN 191 treatments (Fig. 1). An oscillation pattern was visible, indicating the variations of cell sizes 192 at different growth stages. This pattern might be related to cell division, where cells grew 193 exponentially in the exponential growth phase, and reached their maximum in stationary 194 phase. These findings suggest that microalgal cells start increasing in size until they reach 195 a critical point for cell division, which can be likely affected by external energy supply e.g. 196 light, temperature and nutrients (Zachleder et al., 2016). This explains the less 197 pronounced oscillation in NN5 and NN6 even when extra N was supplied, indicating again that cells were P limited. Nonetheless in this study, higher light intensity does not seem to 198

199	result in distinct cell volume changes (Fig 1A). It has been shown that cell oscillation by
200	lighting can be species-specific, where cell volume changes are not restricted to a fixed
201	pattern (Agusti and Kalff, 1989).
202	3.2 Protein and carotenoids dynamics
203	During phase-one, all treatments from N1 to N6 showed increases of suspension protein
204	along with cell growth, and the more initial N in the medium, the more suspension protein
205	was reached at the end of phase-one, ranging from 17 mg/L in N1 to 597 mg/L in N6 (Fig

1B). The corresponding cellular protein content also reached the highest level in N6 (233

207 pg/cell) from N1 (83 pg/cell). Higher protein levels at increased N concentrations in

suspension, cell, and biomass have all been reported for various species such as Dunaliella

209 tertiolecta, Scenedesmus sp. LX1, and Chlorella sp. (Fabregas et al., 1989; Kiran et al.,

210 2016; Zhuang et al., 2018), while N starvation is well known to reduce the protein content

211 (Gao et al., 2018).

212 During the high-N treatments in phase-two, both suspension protein and cellular protein

213 were significantly boosted from NN1 to NN4. Such rises were observed very shortly after

N addition and continued until the end of phase-two (Fig 1B). The biggest increase of

suspension protein occurred in NN1 with 3460% (from 17 mg/L to 599 mg/L), which was

216 due to the extremely low protein and biomass concentration in N1. The highest

suspension protein at the end of phase-two reached 902 mg/L in NN4. For cellular protein,

similar results were found with 404% increase in NN1 reaching 419 pg/cell, which is

comparable with NN2-NN6. When microalgae are supplied with excess of substrate after a

220 period of starvation, overcompensating mechanisms can occur, in which cells are 221 triggered to uptake and store higher substrate amounts than necessary (Brown and 222 Shilton, 2014). The boost of protein production in NN1-NN4 can likely be a result of these 223 mechanisms. Another event that can occur when enough N is present in the culture is 224 called "luxury uptake", which implies the natural uptake of resources beyond necessity 225 without prior starvation. The increase in protein content of NN5-NN6 could be a result of 226 this mechanism (Brown and Shilton, 2014). Both mechanisms linking to the survival of 227 microalgae are commonly found and thoroughly described (Xie et al., 2017). Due to these 228 two phenomena, protein or N content of many microalgal species such as *Chlorella* 229 vulgaris, and also macroalgal species such as Oedogonium could be boosted with N 230 addition (Cole et al., 2015; Dortch et al., 1982; Xie et al., 2017). 231 During the higher-light treatments in phase-two, however, both suspension and cellular 232 protein level for NL1-NL4 did not substantially change, likely due to N depletion from the 233 previous phase. In NL5 and NL6, however, where N was still present, suspension and 234 cellular protein levels increased by up to 37% and 77%, respectively. Higher protein 235 content induced by higher light intensity has been observed for *Chlorella vulgaris* and 236 Chlorella pyrenoidosa (Chen et al., 2015; Ogbonna and Tanaka, 1996). However, other 237 microalgal species could also present either decreased protein content or no change at all 238 following higher light intensity, especially with low N availability (Chen et al., 2015; 239 Markou et al., 2012). Thus, the results from both high-N and higher-light treatments 240 suggest that N levels directly links to the total protein accumulation.

241 The dynamics of carotenoids generally showed the opposite pattern of protein. During 242 phase-one from N1 to N4, both suspension and cellular carotenoids accumulated, with 243 maximum 29 mg/L and 16 pg/cell reached during the stationary phases of N3 and N1, 244 respectively (Fig 1C). This trend is a consequence of N deficiency, which is one of the most 245 effective ways to induce  $\beta$ -carotene accumulation in *D. salina* (Lamers et al., 2012). The 246 cellular carotenoids showed no changes in N5 and N6, although a slight increase of 247 suspension carotenoids occurred, which was mainly due to an increase in biomass (Fig 248 1C). During the higher-N treatment in phase-two, all cellular carotenoids from NN1 to NN6 249 dropped to initial levels, around 4-8 pg/cell. Due to different biomass levels in this phase, 250 the suspension carotenoids either dropped or slightly increased, but remained in a similar 251 range from 9 to 22 pg/cell for all treatments (Fig 1C). During the higher-light treatment in 252 phase-two, both cellular and suspension carotenoids increased rapidly in NL1 to NL4. The 253 increases started shortly after the switch to higher light and continued till the end of the 254 experiment, with maximum 196% cellular carotenoids increase in NL4 and 262% 255 suspension carotenoids increase in NL2 (Fig 1C). Very little changes of carotenoids levels 256 were found in NN5, NL5, NN6 and NL6, primarily due to the presence of N, which suggest 257 N limitation to be the determining factor for effective carotenoids induction. When light 258 intensity is higher than needed for photosynthesis in *D. salina*,  $\beta$ -carotene is produced in 259 excess to overcome light stress and potential photo-oxidative damage, and it has been 260 well documented that high light intensity can significantly increase the production of  $\beta$ -261 carotene in D. salina (Lamers et al., 2010; Raja et al., 2007). This is also one important

reason why large-scale cultivation of *D. salina* is located wherever high light intensity is
expected (Ben-Amotz, 1993).

264 3.3 Protein and carotenoids productivities 265 In phase-one, protein productivities generally declined (N1 and N3), or rose and declined 266 (N2, N4-N6) towards the stationary phase (Fig 1D). These results showed that higher 267 protein productivity was obtained with high N availability. The highest protein 268 productivities reached were 7, 28, 35, 50, 50 and 49 mg/L/d, for N1 to N6, respectively, 269 and mostly occurred between exponential and linear growth phase. This is in agreement 270 with earlier work on *D. salina* tested at different salinities, pH levels and light regimes (Sui 271 et al., 2019; Sui and Vlaeminck, 2019). For treatments with N starvation at the end of 272 phase-one, namely N1-N4, higher-N treatment in phase-two (NN1-NN4) significantly 273 stimulated their protein productivity by 37-1024%. This again can be linked to N 274 overcompensating mechanism. Differently, for N5 and N6, where N was still abundant 275 after phase-one, protein productivity in phase-two (NN5-NN6) did not show notable trend 276 changes but kept declining (Fig 1D). During the higher-light treatment in phase-two, all 277 protein productivities continued to decrease, as a result from prolonged cultivation time 278 and rather stable protein content (Fig 1B, 1D). Overall, these results show that when D. 279 salina cells experience N starvation, extra N addition enhances protein productivity, likely 280 due to overcompensating mechanisms. These findings can be used to optimize the design 281 for harvesting strategy, which could increase biomass protein content. When abundant N 282 is provided, the optimal harvesting point will be around exponential to linear growth

283 phase. This recommendation, however, only applies to maximize protein quantity without284 considering its quality.

285	Differently from protein productivity, carotenoids productivities showed consistent
286	increase pattern during phase-one from N1 to N4, with a maximum of 3 mg/L/d reached
287	in N3 (Fig 1E). These high carotenoids productivities followed the pattern of their
288	suspension carotenoids contents, which depends on both biomass and cellular
289	carotenoids accumulation (Fig 1C). During higher-light treatment in phase-two, NL1 to NL4
290	all showed significant increases of carotenoids productivity, ranging from 15 to 107%. The
291	highest carotenoids productivity reached was 4 mg/L/d in NL3 (Fig 1E). In contrast, the
292	high-N treatments in phase-two contributed negatively to all carotenoids productivities
293	from NN1 to NN4, which decreased by 5 to 54% (Fig 1E). For the treatments of N5, NN5
294	and NL5, as well as N6, NN6 and NL6, low productivities without evident changes in
295	carotenoids were observed throughout the experiment, which was possibly due to the
296	presence of excess N. Generally, N starvation in <i>D. salina</i> enhances the accumulation of
297	carotenoids, which further increases with higher-light exposure. Thus, the combination of
298	N starvation and subsequent exposure to higher light can be a beneficial way to boost
299	carotenoids production.
300	3.4 EAA dynamics and productivities

A few samples have been selected for a more detailed analyses of EAA. They were the
intermediate treatment on the intersection of N and P limitation (N4) and the highest N
treatment (N6) as reference, together with the respective high nitrogen (NN4 and NN6)

304 and light (NL4 and NL6) treatments, both just after the transfer (start) and at the end of 305 the experiment (end). At the end of phase-one, N4 clearly showed higher levels of EAA 306 content, EAA productivity and EAAI relative to N6 (Fig 2A). The EAAI of N4 reached 1.3, 307 which is of superior quality for human consumption (threshold EAAI = 1), while the EAAI of 308 N6 was only 0.4, showing inadequate quality. Looking at individual EAA levels, all the EAA 309 in N4 were substantially present in higher amounts as compared to N6, exceeding 310 FAO/WHO reference except for valine (Fig 2B, 2C and Table 3). These findings indicate that 311 D. saling with EAAI of 1.3 is well suited to be incorporated into food which perfectly match 312 human requirement (EAAI = 1), and actually saving 23% of biomass, further increasing the 313 efficiency of food consumption. Protein and amino acid synthesis in microalgae naturally 314 relies on N assimilation pathways, where nitrate is transported inside the cell and 315 converted to nitrite by nitrate reductase. Ammonia is then obtained by further reduction 316 of nitrite and incorporated into glutamate/glutamine via glutamine synthase and NADPH-317 dependent glutamine: 2-oxoglutarate aminotransferase (GS/GOGAT) pathway (Alipanah et 318 al., 2015; Halsey et al., 2011; Remmers et al., 2018; Sanz-Lugue et al., 2015). 319 Glutamate/glutamine sequentially provides the critical entry point of N into cellular 320 biochemicals, which can subsequently be used for synthesis of other EAAs (Guerra et al., 321 2013). Although N availability associates closely with EAA production in microalgae, as 322 shown in this study and many others, a higher N level does not necessarily lead to higher 323 EAA production. When microalgal cells become N limited, mostly towards the stationary 324 growth phase, their major response is to preserve cellular N capacity via scavenging

mechanisms, by which EAA biosynthesis can still occur using intracellular N (Alipanah et
al., 2015; Halsey et al., 2011; Lv et al., 2017; Remmers et al., 2018; Zhang et al., 2016).
However, when N deprivation occurs in the long term, EAA synthesis is interrupted and
results in sharp decreases of protein and amino acids (Kiyota et al., 2014; Van de Waal et
al., 2010; Zhang et al., 2016).

330 Several studies have shown that N limitation or a short period of starvation can boost EAA 331 production in microalgae and macroalgae. For instance, it has been shown that all EAA 332 levels of *D. salina* SAG 184.80 were enhanced towards stationary growth phase, when N 333 became limited and then shortly starved (Sui et al., 2019). Synechocystis sp. also exhibited 334 rising levels of all EAA during short N starvation, while longer N starvation resulted in 335 dropping of EAA levels (Kiyota et al., 2014). Similarly, short N starvation contributed to the 336 production of several EAAs, especially phenylalanine, in marine microalga Isochrysis 337 zhangjiangensis, which can be the consequence of N scavenging of e.g. nucleotides and 338 rubisco protein. Nevertheless, significant decreased EAA levels were found after long-339 exposed N deficiency (Zhang et al., 2016). Besides, the macroalgae Ulva ohnoi was also 340 shown to exhibit higher proportions of alanine, serine, glycine, and the EAAs 341 phenylalanine, threonine and valine with low N concentration (Angell et al., 2014). 342 When extra N was supplied during phase-two, EAA content, EAA productivity and EAAI 343 from NN4 (start) decreased sharply, resulting in inadequate protein quality (Fig 2A). A 344 similar drop was also observed in NN6 (start). Regarding individual EAA levels, higher N 345 addition also led to their reductions, where the overall level of EAA decreased

346	dramatically from N4 to NN4 (start and end) (Fig 2B and Table 3). Overall, these results
347	clearly demonstrate that short N starvation promotes EAA accumulation in D. salina, while
348	high N levels has a negative impact. At the end of phase-two after N addition, EAA
349	content, EAA productivity, and EAAI in NN4 (end) and NN5 (end) all increased by 14-28%,
350	3-14%, and 35-55%, respectively. This again indicates that cells tend to preserve EAAs
351	towards later growth phases to maintain the N capacity and cell functions, possibly due to
352	the drop of N concentration in the environment (Fig 2A, Table 2).
353	While the complex biosynthesis of EAAs following various pathways are associated with N
354	availability, it is not always easy to link them to other environmental conditions. Here,
355	higher light intensity in phase-two seemed to have differential effects on EAA production.
356	In the higher light treatment, NL4 (start and end) did not respond positively, resulting in
357	slight decreases of EAA content, EAA productivity, and EAAI (Fig 2A). Nonetheless, protein
358	quality of the biomass at this stage still remained superior for human consumption, as
359	indicated by EAAI and relatively high amounts of individual EAAs (Fig 2A, 2B and Table 3).
360	Although <i>D. salina</i> biomass seems to maintain protein quality during the higher-light
361	treatment, thus only slight differences between NL4 (start) and NL4 (end) in EAA content,
362	the EAA productivity in NL4 (end) decreased by 46%, which was mainly attributed to
363	growth stage (Fig 2A). In NL6 (start and end), higher light intensity evidently promoted the
364	accumulation of EAA in the biomass with higher EAA content (Fig 2A). Shortly after
365	introducing higher light in NL6 (start), EAA content, EAA productivity and EAAI increased
366	with 49, 25, and 30%, respectively. At the end of phase-two in NL6 (end), such increases

367	were even 232, 152, and 189%, reaching 36%, 14 mg/L/d and 1.2, respectively. Similarly,
368	the individual EAA levels rose in NL6 from start to end (Fig 2C). The photo-acclimation
369	response of microalgae towards excess light has been well-described, and upregulation of
370	EAA during photo-acclimation was found in both microalgae and plants in response to
371	high light exposure (Davis et al., 2013; Galili et al., 2016). Davis et al., (2013) suggest that
372	the elevation of EAA could be due to an induction of <i>de novo</i> amino acid biosynthesis
373	and/or protein catabolism from photodamage. Nevertheless, the fact that EAA content in
374	NL4 (start and end) was not elevated might be related to their extreme N deficiency (Table
375	2). As stated, when N deprivation and high light were both applied to Dunaliella tertioecta,
376	most amino acids were significantly decreased (Lee et al., 2014).
377	3.5 Optimized biomass with both high-quality protein and carotenoids
378	As the parameters indicating protein quantity, protein quality and carotenoid content
379	varied substantially depending on the cultivating conditions, Fig 3 and Table 4 elucidate
380	the effects of N and light intensity on the biomass quality of <i>D. salina</i> . Nitrogen
381	concentration in the medium and light intensity, but particularly their interaction,
382	significantly affected EAAI (Table 4; Fig 3B, 3C). These findings suggest that short N
383	starvation and higher light intensity together enhance EAA production in <i>D. salina</i> , rather
384	than either factor alone. Similar results were estimated for carotenoids content, where
385	highest content was found with a combination of decreasing N and increasing light
386	intensity, while only N starvation and not higher light showed also a significant separate
387	effect (Table 4). It is noted however, that the correlation of light intensity to carotenoids

content was limited by the low sample sizes. When the effect of light intensity on
 carotenoid contents was analyzed from more samples obtained from this study, it indeed
 becomes evident that light intensity is correlated to carotenoid contents (Table 4; Fig 3G).
 Together, these results indicate that short N starvation and subsequent higher light
 intensity together are favored for carotenoids accumulation.

393 Overall, short N starvation together with higher light intensity are beneficial for the 394 production of both EAA and carotenoids. Specifically, treatment NL4 (end) presented 395 exceptional EAA enhancement and carotenoids accumulation in D. salina, indicating the 396 possibility to produce microalgal biomass with both high protein quality and anti-oxidant 397 strength. Consequently, D. saling demonstrated to be a valuable, and potentially unique 398 species as novel protein source with high anti-oxidant activity for humans. Having been 399 successfully applied in commercial production of *D. salina* using two-phase cultivation 400 system, minor modifications can be applied to optimize biomass value, where short N 401 starvation should be reached in the stationary phase in phase-one and a higher-light 402 treatment should be applied in phase-two.

Broadly looking at the market of microalgal products, microalgae are mainly supplied as
food supplement with high protein content and other nutritional values like vitamins.
Specifically for *D. salina*, the global market is mainly for high-value food and feed
supplement. To the authors' knowledge, there is no microalgal product in the food
industry which provides both high-quality protein and carotenoids at the same time. The
results from this study thus present a step further towards production of microalgae with

409 multiple high-value compounds, which largely improves the production efficiency. The

410 biomass can therefore be a superior source for food supplement and food ingredient,

411 either consumed next to food sources or be incorporated into food products.

#### 412 Conclusions

- 413 Protein quality and carotenoids content of *D. salina* could be simultaneously enhanced 414 applying a two-phase cultivation strategy. Short N starvation should be reached in the 415 stationary phase in phase-one to upregulate EAA production, and phase-two should be 416 given higher illumination to boost carotenoids production. The optimized cultivation 417 conditions resulted in production of *D. salina* biomass with an EAAI of 1.1 and cellular
- 418 carotenoids content of 24 pg/cell. This study reveals that *D. salina* is a valuable, and
- 419 potentially unique species bringing together both high-quality protein and carotenoid,
- 420 thus can be used as protein source with antioxidant pro-vitamin A effect for humans.

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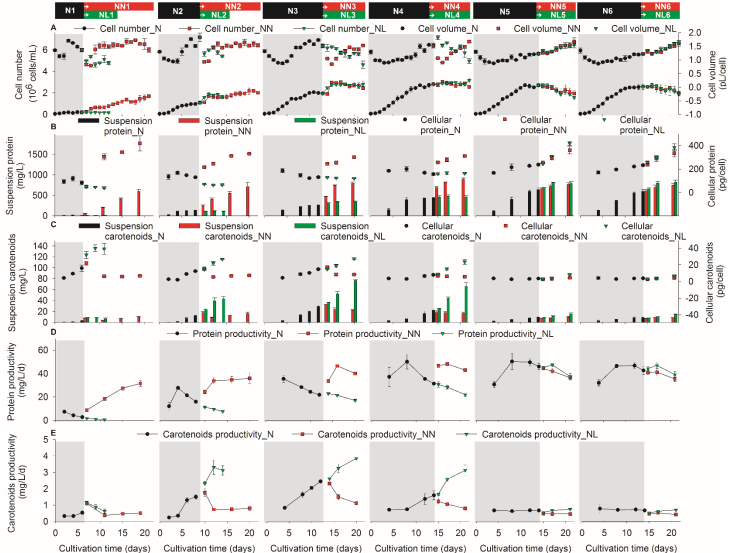
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564	Figure	captions

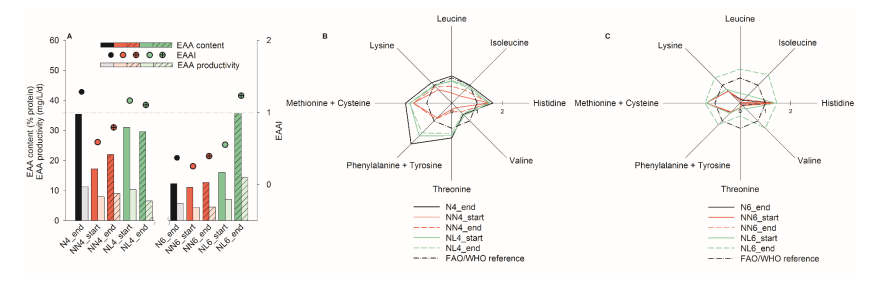
565 Fig. 1. Effect of two-phase cultivation on *D. salina*: (A) cell number and cell volume, (B) 566 suspension protein and cellular protein, (C) suspension carotenoids and cellular 567 carotenoids, (D) protein productivity and (E) carotenoids productivity. Different N levels 568 were applied in phase-one and both N addition and higher illumination were applied in 569 phase-two. Cultivation occurred at 25°C and pH 7.8-8. Data are expressed as means ± 570 standard deviation (n = 3)571 Fig. 2. Effect of two-phase cultivation on *D. salina* at the end of each phase: (A) EAAI, EAA 572 content and EAA productivity, (B) individual EAA level in treatment N4, NN4 (start and 573 end) and NL4 (start and end) and (C) individual EAA level in treatment N6, NN6 (start and 574 end) and NL6 (start and end). Different N levels were applied in phase-one and both N 575 addition and higher illumination were applied in phase-two. Cultivation occurred at 25°C 576 and pH 7.8-8. Data are expressed as means  $\pm$  standard deviation (n = 3) Fig. 3. N and light intensity on EAAI and carotenoids content in ten D. salina samples: (A) N 577 578 and light intensity vs. EAAI, (B) N vs. EAAI, (C) light intensity vs. EAAI, (D) N and light 579 intensity vs. carotenoids content, (E) N vs. carotenoids content and (F) light intensity vs. 580 carotenoids content. Highlighted blue dots represent all conditions where an EAAI above 1 581 was obtained. The corresponding carotenoids content obtained from above conditions 582 were highlighted as orange dots. Light intensity vs. carotenoids content from 24 D. salina samples was presented in (G). 583





-> N





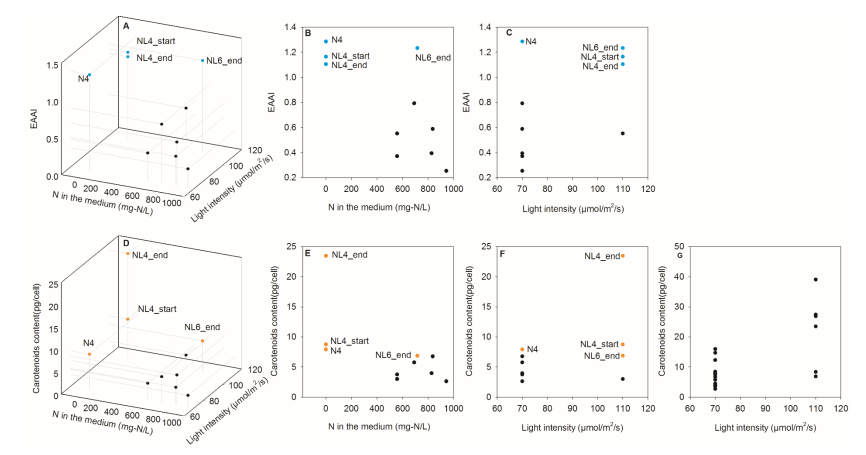


Fig. 3.

Treatment			Pha	ise-on	9		Phase-two (higher N)	Phase-two (higher light intensity)			
	N1	N2	N3	N4	N5	N6	NN1-NN6	NL1-NL6			
N concentration (mg-N/L)	1.4	11.2	22.4	44.8	179.2	716.8	932.8 n.a.				
Light intensity (µmol/m²/s)					70		110				
Erlenmeyer flask volume (mL)	.) 500 250							250			
Culture volume (mL)	400 200							200			
Temperature (°C)	25										
рН	7.8-8										
NaCl concentration (g/L)	117										
Aeration (L/h)		3.3									
n a not annlicable											

Table 1 Experimental conditions of all treatments in phase-one and phase-two

n.a. not applicable

	N1	NN1	NL1	N2	NN2	NL2	N3	NN3	NL3	N4	NN4	NL4	N5	NN5	NL5	N6	NN6	NL6
NO <sub>3</sub> -	0	1098	0	0	953	0	0	742	0	0	691	0	88	687	44	558	827	716
(mg-N/L)	(0)	(84)	(0)	(0)	(83)	(0)	(0)	(46)	(0)	(0)	(56)	(0)	(5)	(33)	(1)	(61)	(45)	(51)
PO4 <sup>3-</sup>	7.2	0	7.1	3.4	0	0	0	0	0	0	0	0	0	0	0	0	0	0
(mg-P/L)	(2.3)	(0)	(0.7)	(0.4)	(0)	(0)	(0)	(0)	(0)	(0)	(0)	(0)	(0)	(0)	(0)	(0)	(0)	(0)

Table 2 Nitrate and phosphate concentration in the medium at the end of different treatments

Standard deviation is listed in brackets

mg/g protein	Histidine	Isoleucine	Leucine	Lysine	Methionine + Cysteine	Phenylalanine + Tyrosine	Threonine	Valine	Total EAA
N4_end	24.4	<u>30.1</u>	<u>63.7</u>	<u>51.5</u>	40.4	<u>87.0</u>	<u>31.8</u>	24.8	<u>353.7</u>
NN4_start	<u>22.3</u>	12.3	23.5	33.9	<u>33.5</u>	31.1	5.7	9.9	172.3
NN4_end	<u>22.6</u>	21.8	39.2	41.4	<u>33.6</u>	34.2	7.9	19.2	220.0
NL4_start	<u>22.6</u>	28.4	52.9	43.3	<u>36.5</u>	<u>69.8</u>	<u>29.7</u>	27.3	<u>310.4</u>
NL4_end	<u>21.9</u>	26.2	50.8	<u>45.3</u>	<u>37.0</u>	<u>63.6</u>	<u>27.8</u>	22.9	<u>295.6</u>
N6_end	<u>20.7</u>	4.7	9.2	30.2	<u>29.7</u>	19.6	4.2	4.9	123.2
NN6_start	<u>20.7</u>	1.1	4.9	29.8	<u>29.7</u>	18.7	4.1	2.1	111.1
NN6_end	<u>20.7</u>	6.0	10.6	30.1	<u>29.8</u>	19.8	4.0	5.6	126.6
NL6_start	<u>20.9</u>	13.2	22.8	34.8	<u>29.7</u>	21.9	4.4	12.5	160.2
NL6_end	<u>21.8</u>	<u>47.6</u>	<u>79.6</u>	<u>64.0</u>	<u>30.6</u>	<u>46.3</u>	11.6	<u>55.0</u>	<u>356.5</u>
FAO/WHO	15	30	59	45	22	38	23	39	271

Table 3 Individual and total amino acid content of selected treatments and FAO/WHO reference

<u>Underlined</u> values indicate levels above FAO/WHO reference.

Model	R <sup>2</sup>	Significance	Corresponding figure
Dependent variable: EAAI Predictors: Nitrogen, Light <sup>1</sup>	0.582	0.047*	Fig. 3A
Dependent variable: EAAI Predictors: Nitrogen <sup>1</sup>	0.534	0.016*	Fig. 3B
Dependent variable: EAAI Predictors: Light <sup>1</sup>	0.421	0.042*	Fig. 3C
Dependent variable: Carotenoids content Predictors: Nitrogen, Light <sup>1</sup>	0.461	0.115	Fig. 3D
Dependent variable: Carotenoids content Predictors: Nitrogen <sup>1</sup>	0.426	0.041*	Fig. 3E
Dependent variable: Carotenoids content Predictors: Light <sup>1</sup>	0.211	0.182	Fig. 3F
Dependent variable: Carotenoids content Predictors: Light <sup>2</sup>	0.511	0.000*	Fig. 3G

Table 4 Correlation of N and light intensity on EAAI and carotenoids content from statistical results on data from Fig. 3.

\*: significant difference (p < 0.05)

<sup>1</sup>: statistical analyses using 10 samples with EAA contents determined in this study.

<sup>2</sup>: statistical analysis using 24 samples with carotenoids content determined in this study