A Two-Step Process for Improved Biomass Production and Non-Destructive Astaxanthin and Carotenoids Accumulation in Haematococcus pluvialis

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Abstract: Carotenoids extracted from microalgae have a considerable economic interest in numerous high-value markets. Natural astaxanthin has gained much interest in its powerful antioxidant properties, however, its commercial-scale production is still challenging. In this study, a simple and economical way to cultivate Haematococcus pluvialis (CCAP 34/1D) by a two-step process was investigated by exploring alternative strategies to maximise algal growth and astaxanthin yield. During the first step—improving biomass production—four nitrogen sources were tested (NH4Cl, NaNO3, Fe(III)NO3, and urea). The second step—carotenogenesis induction—was achieved by using a mix of moderate stressors that worked in synergy (i.e., mild light, nitrogen limitation, the addition of sodium acetate at 0.25% w/v), thereby minimising potential losses of the accumulated biomass caused, for example, by photobleaching or nitrogen starvation. Results showed that urea was the nitrogen source, allowing the highest cell density and growth rate. In terms of carotenogenesis induction, the use of mild stressors resulted in three out of four treatments having a relative increase in cell number (13.8–26.7%) and a concomitant increase in astaxanthin yield. Simple low-cost strategies, such as small adjustments to media recipes and synergism between mild stressors, could bring a disproportionate effect on the future successes of making algal biotechnology a widespread reality.

Keywords: nitrogen; growth medium; Haematococcus pluvialis; astaxanthin; carotenogenesis

1. Introduction

Synthetic carotenoids are ubiquitously available since they can be rapidly produced using inexpensive chemicals, therefore reducing the production costs associated with cultivating, harvesting, and extracting from a living organism [1]. Although synthetic carotenoids, including synthetically produced astaxanthin (S-AX), are more abundant on the market, and faster and cheaper to produce, their health-promoting and antioxidant properties are not as effective as their natural counterparts, hence making them a less desirable product. Astaxanthin is a powerful natural antioxidant. It can be extracted from krill, yeasts, marine bacteria, and microalgae (e.g., Chlorella zofingiensis, Chlorococccum sp.) [2,3]. The most common source of natural astaxanthin (N-AX) on the market is derived from the freshwater microalgae Haematococcus pluvialis [4]. The natural form of astaxanthin has a stronger antioxidant activity. The final product composition also differs in the two forms: natural astaxanthin (N-AX) can potentially carry other carotenoids (i.e., β-carotene, lutein, and canthaxanthin) [5], bringing additional value to the market, while the S-AX contains synthetic astaxanthin and potentially synthesises intermediates with unknown effects [6,7]. For this reason, S-AX is still not currently permitted for human consumption [8].
However, it is approved by the European Food Safety Authority (EFSA) and the United States Food and Drug Administration (US FDA) as an aquaculture feed additive.

The enhanced performance of natural astaxanthin compared to synthetic variants has made it a much more in-demand and valuable product. Currently, the market value of astaxanthin, depending on products’ purity, varies from $2500 for S-AX and >$7000 kg for N-AX derived from *Haematococcus pluvialis* [8–10]. The global market demand for both forms is estimated to exceed $880 million by 2026 and a compound annual growth rate (CAGR) of 8.2% by 2027 [11]. Furthermore, N-AX production accounts for less than 5%, of which only less than 1% is *H. pluvialis* derived [12]. The high market value is caused by the scarcity of natural astaxanthin due to its difficult production process.

*Haematococcus pluvialis* has a complex life cycle, undergoing several life stages before encysting and accumulating astaxanthin [13]. It involves different morphotypes: a green vegetative motile cell; a green vegetative palmella cell; a palmella cell with initial astaxanthin accumulation in transition to an aplanospore; and finally, an aplanospore cell with accumulated astaxanthin [14].

*H. pluvialis* industrial-scale cultivation commonly employs a two-step process: first, it aims to achieve high biomass yields; and secondly, the biomass is exposed to a stressor (e.g., high light) to induce carotenogenesis and hence the accumulation of astaxanthin [13].

In this first step, *H. pluvialis* is conventionally grown in autotrophy to reduce the risk of biological contamination, and to facilitate better operator control of the transition between green and the red phases.

There has been considerable interest in the research and development of the green phase of *H. pluvialis* during the past ten years due to the high value of N-AX and the challenges involved in *H. pluvialis* cultivation. In fact, due to its complexity, any small modification in the cultivation cycle that results in faster growth rates or higher overall biomass/astaxanthin yield can represent a significant economic improvement. The long vegetative growth contributes to the high production costs, especially in terms of nutrient supply. Among nutrients, nitrogen is needed not only for cell growth but also to produce the enzymes that catalyse astaxanthin synthesis and accumulation in *H. pluvialis*. Microalgae can uptake and assimilate different nitrogen sources (NS), i.e., nitrate, nitrite, ammonium, and urea; nitrogen assimilation utilising urea requires a different set of enzymes from nitrate, nitrite, and ammonium, which share the same metabolic pathway.

The vast majority of studies and industrial application of *H. pluvialis* cultivation have employed NaNO$_3$ as a nitrogen source. NaNO$_3$, however, is expensive and nitrogen only accounts for 16.5% of the molecular mass of the chemical, against 46% in urea containing two nitrogen atoms. For this reason, urea would be required in lower quantities for an equivalent nitrogen concentration. Attempts to cultivate *H. pluvialis* in urea commonly fail and *H. pluvialis* mostly used growth media, and does not provide urea in their formulation [15–17]. Deeper investigations and manipulations of the abiotic conditions could establish the use of urea in *H. pluvialis* growth mediums for cost-effective large-scale production.

The second phase of the two-step cultivation process is the carotenogenesis induction and can be achieved by modulating a few growth parameters: light and N depletion are commonly employed as stressors, and iron addition and osmotic stress may also enhance the accumulation of astaxanthin [18]. The astaxanthin accumulation employing autotrophic growth, however, may not have reached its maximum potential [15,16]. Currently, 4% w/v of astaxanthin accumulation in *H. pluvialis* is often considered the common threshold for comparative success [19,20]. The main difficulty in applying a stress to the *H. pluvialis* cells is to find the threshold level which minimises biomass loss and cell bleaching and maximises astaxanthin accumulation. It has been ascertained that high levels of stress, such as a sudden increase in irradiance or nitrogen starvation, may damage cells, decreasing the final biomass yield and the amount of recoverable astaxanthin [21]. For example, high light intensity can lead to photoinhibition [22], while nitrogen depletion affects photosynthesis and cellular metabolism [23–25].
In this study, *H. pluvialis* (CCAP 34/1D) was selected for a two-step cultivation based upon a previously published strain screening by Butler [26]. In our study, an initial optimisation of the growth conditions was carried out for both the green and red phases of the strain CCAP 34/1D: nutrients concentration and pH were manipulated as shown in Supplementary Materials 1. Afterward, different NS were tested to enhance biomass productivity during the green phase of growth.

To induce the red phase, the drawbacks of conventionally applied stringent individual stressors that were mentioned above were considered. We hypothesised that multiple moderate stressors could be simultaneously applied for a possible synergistic effect whilst preserving the biomass accumulated in the green phase.

Based on previous works, a mix of stressors (e.g., combining higher irradiance, salinity, organic carbon sources) were tested (Supplementary Materials 2); the mix of stressors that allowed the highest biomass yield preservation coupled with the greatest accumulation of carotenoids was selected to induce the red phase, i.e., buffered JM medium (without the addition of NaNO$_3$, but retaining the NS from Ca(NO$_3$)$_2$·4H$_2$O and (NH$_4$)$_6$Mo$_7$O$_{24}$·4H$_2$O modified with the addition of 0.25% (w/v) sodium acetate and exposed to 250 µmol photons m$^{-2}$ s$^{-1}$ for 7 days.

The dual purpose of this study is to find a simple low-cost strategy to increase the rate of vegetative growth and biomass accumulation, followed by an enhancement of astaxanthin accumulation without any deleterious effects upon biomass yield. The efficiency of the employed induction method and the pigments’ characterisation was assessed by HPLC analysis in the green and the red phases. The overarching goal was to find an economic and easily implementable manner to cultivate *H. pluvialis* CCAP 34/1D for astaxanthin production.

2. Materials and Methods
2.1. Cultivation and Growth Conditions

The algal strain *Haematococcus pluvialis* CCAP 34/1D was obtained from the Culture Collection of Algae and Protozoa (CCAP), at The Scottish Association for Marine Science (SAMS, UK). Cultures were grown in Jaworski’s media (JM) with the pH adjusted to 7.5 ± 0.2 and buffered with 10 mM HEPES. Nitrogen type was varied and supplied as either sodium nitrate, iron (III) nitrate, ammonium chloride, or urea at a final N concentration of 941 µM, equivalent to that of the nitrogen concentration only derived from the sodium nitrate contained original JM recipe.

Starter cultures were maintained in a control temperature (CT) room at 20 °C, under a 16/8 h L/D (light/dark) regime at 75 µmol photon m$^{-2}$ s$^{-1}$ photosynthetically active radiation (PAR), without shaking or aeration. Prior to experimentation, cultures underwent a period of acclimation, where they were maintained under experimental conditions, outlined above, for 7 days. After this time, a calculated volume of culture was removed from the pre-acclimated cultures and centrifuged at 3000× g for 15 min, the supernatant discarded and the cell pellet resuspended in 500 mL of media. All experiments were performed in triplicate, with an initial cell density of 4 × 10$^5$ cells/mL. Cultures were maintained in conditions outlined above for 15 days, for vegetative growth.

2.2. Carotenogenesis

Astaxanthin production was induced at day 15 (after vegetative growth) by centrifuging the cells, as described above, and resuspending in a modified JM medium: containing 0.25% (w/v) sodium acetate and NaNO$_3$ depleted but with the original concentration of Ca(NO$_3$)$_2$·4H$_2$O (84 µM) and (NH$_4$)$_6$Mo$_7$O$_{24}$·4H$_2$O (0.8 µM), as described in the JM recipe. Astaxanthin induction was performed for 7 days at the same growth conditions as above, except for the light irradiance which was increased to 250 µmol photon m$^{-2}$ s$^{-1}$ PAR.
2.3. Techniques of Enumeration of H. pluvialis and Growth Cell Monitoring

Three different enumeration techniques were employed: the first one by the hemocytometer taking advantage of microscopic observation to count the cells and assess their morphological state at the same time. The other two techniques were cell counting methods which can be useful for repeated cell counts (CASY cell counter) or to follow cells’ growth by measuring the absorbance at the spectrophotometer, which is required while running an experiment with small volumes which cannot be sampled daily or contaminated. For the first two methods, cultures were sampled every day in a Laminar-flow cabinet (MSC advantage, Thermo Scientific). Cellular integrity/health and the degree of encystment were monitored using a hemocytometer; cell enumeration was performed using a CASY cell counter (Casy TT, Roche Innovatis) using a volume of 1 mL or 0.5 mL, depending on the density of the culture, and afterward, diluted in 9 mL of a solution 0.9% NaCl. An analytic volume of 100 µL was measured three times through a 50 µm aperture. When direct counting was not possible, cell enumeration was assessed spectrophotometrically (Heλios Gamma spectrophotometer UV-vis, Thermo Scientific) at 750 nm (OD<sub>750</sub>), after a linear calibration curve was created between OD<sub>750</sub> and cells/mL (Figure S4 in Supplementary Materials 2).

2.4. HPLC Analysis

HPLC analyses were performed on cultures during both their green and red stages of growth. Cultures were harvested on cultivation day 14 (prior to astaxanthin induction) and at day 21 (7 days following induction). In both instances, cells were centrifuged at 3000 × g for 15 min; pellets of sub-samples were then flash-frozen in liquid nitrogen, stored at −80 °C, and then freeze-dried (ALPHA 1-2 LD plus freeze dryer, Christ).

The lyophilised biomass was weighed, and the pigments extracted by employing a modified protocol by Sarada (2006) [27]. Briefly, 3 mg of dry weight (DW) biomass was resuspended in 1 mL of 2N HCl and incubated at 70 °C for 2 min. The pigment extraction was facilitated by using glass beads while vortexing (diameter of 0.25–0.5 mm, Roth, Karlsruhe, Germany). Samples were then cooled at −80 °C for five minutes, to exploit the thermic stress to favour the cell breakage. After thawing, samples were then centrifuged at 5000 rpm for 3 min (Thermo Fisher Sorvall Legend X1R Refrigerated Benchtop Centrifuge), the supernatant was removed, and the HCl-treated samples were washed twice in 2 mL deionized water to neutralise the pH, pelletized, and finally resuspended in 1 mL of extraction solvent (95% methanol, buffered with 2% ammonium acetate with the addition of the internal standard) and stored at −20 °C for 15 min. The extraction procedure was repeated until the algal cell debris was almost colourless and several short (30 s) vortexing steps were alternated between each resuspension in the extraction solvent.

Supernatants were collected after centrifugation (5000 rpm, 3 min, at 4 °C) and syringe filtered (porosity, 0.47 um). The extracts were injected into a high-performance liquid chromatography (HPLC) system (Waters Alliance 2695) equipped with a cooled autosampler and a photodiode array detector (Waters PDA 2998). Chromatographic separation was carried out using a 4.6 mm × 100 mm column (Sunfire, Waters) equipped with a 3.9 mm × 5 mm pre-column (Sunfire, VanGuard, Waters) for reverse-phase chromatography during a 30 min elution program. The elution gradient was run as follows with solvent A (80/20 v/v methanol and ammonium acetate 1 M) and solvent B (60/40 v/v methanol and acetone): the time gradient was 0 min 80% A, 20% B; 10 min 0% A, 100% B; 20 min 0% A, 100% B; 30 min 80% A, 20% B with a flow rate of 1.0 mL/min and a sample volume of 100 µL [28]. The synthetic carotenoid trans-β-Apo-8′-carotenal (Sigma) was used as an internal standard in each sample to improve the precision of the quantification and to detect eventual instrumental errors.

A chromatographic and spectral library was created from standard pigments (DHI lab, Hørsholm, Denmark). Pigments in the extracts were identified by comparison of chromatographic and spectral data recorded with standard pigments, using Empowered software (Waters). In the absence of the pigment standard, canthaxanthin and carotenoid-
like pigments were quantified by the ISTD method (internal standard analysis), relating the ratio of the standard peak area and the pigment peak area to their respective amounts, as the standard concentration did not change. The absorbance spectra (400–800 nm) and retention times were also checked to validate the output given by the software, and the pigments’ concentrations were obtained from the signals in the photodiode array detector at 440 nm [29,30]. For some pigments, concentrations were summed: chlorophyll a and its allomeric and epimer forms. Values are expressed as mean values ± standard error of the triplicates.

2.5. Data Analysis

Algal growth expressed as the specific growth rate (µ) was determined using Equation (1):

$$\mu \left( \text{d}^{-1} \right) = \frac{\ln(x_1) - \ln(x_2)}{t_1 - t_2}$$

(1)

where x1 and x2 represent the number of cells per mL at the beginning and the end of the log phase, at the days t1 and t2, respectively.

All data were expressed as the mean ± standard deviation (SD) of measurements obtained from three biological replicates (n = 3). The significance of differences among the means of different treatments was determined by one-way or two-way ANOVA (if one or two independent variables were analysed in the treatments) with Tukey’s post hoc analysis (when comparing every mean with every other mean) or Šidák (when comparing sets of means) for multiple comparisons. Levels of significance were set at p < 0.05; all statistical analyses were performed using GraphPad Prism, Version 6.

3. Results

The first part of the study aimed at finding the optimal NS in the JM medium supplemented with HEPES buffer 10 mM for H. pluvialis CCAP 34/1D. In the second part, the same cultures were exposed to moderate stressors (i.e., mild light, nitrogen limitation, and sodium acetate at 0.25% w/v in the modified JM) to investigate if astaxanthin accumulation was induced without a biomass loss.

3.1. Growth with Different Nitrogen Sources

The highest growth rate (0.4 ± 0.011) and cell density (4.98 × 10^5 ± 0.66 cells/mL) were achieved when urea was the sole NS, whereas the lowest growth rate (0.28 ± 0.02 d^{-1}) was found when NaNO_3 was supplied, as in standard JM, which may be considered as a positive control (Figure 1). Cultures grown in FeNO_3 or NH_4Cl showed similar values of growth rate and maximal cell density to those values obtained when cultivated in NaNO_3 as the only NS (Figure 1).

3.2. Assessment of Cell Density Following Application of Stress Conditions

Cultures grown in different NS were harvested and resuspended in the modified JM medium (without the NaNO_3 addition, but with the original Ca(NO_3)_2·4H_2O and (NH_4)_6Mo_7O_24·4H_2O) with the addition of 0.25% (w/v) sodium acetate and exposed to 250 µmol photons m^{-2} s^{-1} for 7 days (Supplementary Materials 2).

The cell density increased during the induction process in all the treatments except for the one in NH_4Cl (−12.6%), which was statistically different compared to the biomass previously obtained in NaNO_3 (p = 0.0084). Although the cell number increase was highest when cells grew in FeNO_3 (+26.7%), followed by those in NaNO_3 (+16.6%) and urea (+13.8%), the relative change in FeNO_3 was significantly greater only when compared to the changes in urea and NH_4Cl grown cultures (Figure 2).
Figure 1. *H. pluvialis* CCAP 34/1D maximum growth rate and cell density in different NS. Error bars are standard deviations of the means, *n* = 3. The letters indicate a significant difference in growth media treatments for each measured response (i.e., growth rate or maximum cell density) (*p* < 0.05).

Figure 2. The relative change of *H. pluvialis* CCAP 34/1D cell number is based on the number of cells/mL at the beginning and end of the 7-day induction process, and thereby representative of the cell number change during the red phase alone. Error bars are standard deviations of the means, *n* = 3. Different letters indicate significant differences among the different nitrogen source treatments (*p* < 0.05).
3.3. Pigment Profiles in Green and Red Growth Phases

Table 1 shows the pigment composition of the biomass, measured using HPLC analysis at the end of the green and red phases. The representative chromatograms per each treatment are included in Supplementary Materials 3. The major pigments identified were in agreement with those recorded elsewhere in the literature [28,30]. The number of peaks identified in the red phase were less than those observed in the green phase. In most instances, the chlorophyll $a$ was the main peak in both phases, with the exception of urea, where the main pigment at the end of the green phase was unidentified but had carotene-like properties and astaxanthin in the red phase. The major pigments showing the greatest peak areas were carotenoid-like pigments, lutein, astaxanthin, chlorophyll $a$, $\beta$-carotene, pheophytin, canthaxanthin, and antheraxanthin. Minor pigments, present in traces, were not quantified: chlorophyll $b$ and echinenone. Carotenoid-like pigments and canthaxanthin, whose standards were not available, were recognised by comparing the samples peaks with those found in literature and spectral library databases [31].

The addition of sodium acetate 0.25% ($w/v$) and the increase of irradiance led to the accumulation of astaxanthin in all the treatments, with final absolute values ranging from $0.313 \pm 0.008$ to $0.711 \pm 0.143$ mg/g, with the exception of cells previously cultivated with $\text{FeNO}_3$, where its content reduced from $2714 \pm 0.585$ to $1353 \pm 0.366$, coincident with the degradation of chlorophylls and lutein.

The shift of pigment composition was more evident in the urea and $\text{NaNO}_3$ grown samples, while cells whose vegetative growth was supplied with $\text{FeNO}_3$ and $\text{NH}_4\text{Cl}$ were less affected by the induction method applied, except for the astaxanthin content drop in the iron treatment. The chlorophylls content dropped in all the treatments except for $\text{NH}_4\text{Cl}$.

A high variety of unknown pigments was found in the green phase of urea cultivated cells: traces of echinenone were found in two out of three replicates; lutein and chlorophyll decreased, while the astaxanthin increased significantly from 0 to the $0.313$ mg/g DW in the red phase. Neoxanthin was found in two out of three $\text{NaNO}_3$ samples with a concentration of $0.025 \pm 0.006$ mg/g DW.

Other pigments, such as canthaxanthin, were detected in the green phase of $\text{FeNO}_3$ and $\text{NaNO}_3$.

When considering the carotenoids/chlorophyll ratio (Figure 3), the ratio changed remarkably between the two phases in urea-grown cells and in the $\text{NaNO}_3$ control ($p < 0.001$). The change in ratio was greater in urea (1.55 ± 0.26) followed by $\text{NaNO}_3$ (0.84 ± 0.33), $\text{NH}_4\text{Cl}$ (0.092 ± 0.012), and $\text{FeNO}_3$ (−0.15 ± 0.06). This ratio change may, in part, be due to a difference in chlorophyll concentration following induction: $\text{NaNO}_3$ −70%, $\text{FeNO}_3$ −38%, Urea −91%, $\text{NH}_4\text{Cl}$ +23%. 
Table 1. Quantitative pigment composition (in mg/g DW), obtained by HPLC analysis, of *H. pluvialis* cells at the end of the green and red phases stationary phase, in cultures previously supplied with equimolar concentrations of nitrogen from different sources. Values are reported as the mean value (*n* = 3) ± 1 S.D. The letters indicate significant differences in the concentration of individual pigments between biomass cultivated using different NS at either green or red phase (*p* < 0.05), while the differences between the green and the red phase in each treatment are denoted by * (p < 0.05). Pigments denoted with † have been quantified based upon the relative peak area of an internal standard (trans-β-Apo-8′-carotenal).

|        | Lutein † | Astaxanthin † | Chlorophyll a † | β-Carotene † | Phaeophytin a † | Antheraxanthin † | Canthaxanthin | Carotenoid-like | Lutein † | Astaxanthin † | Chlorophyll a † | β-Carotene † | Phaeophytin a † | Antheraxanthin † |
|--------|----------|---------------|-----------------|--------------|----------------|-----------------|--------------|----------------|----------|---------------|----------------|--------------|----------------|----------------|
| **NaNO₃** | 0.078 ± 0.022 a | 0.227 ± 0.077 a | 2.682 ± 0.417 b | 0.037 ± 0.008 a | 0.256 ± 0.077 a | nd              |              |               | 1.4 ± 0.05 a | 1.09 ± 1.16 b | 0.048 ± 0.022 a | 0.711 ± 0.143 a | 0.012 ± 0.015 a | 0.007 ± 0.008 a |
| **FeNO₃** | 2.714 ± 0.148 b | 1081 ± 0.041 b | 2.26 ± 0.12 b  | 0.076 ± 0.04 b | 1.07 ± 0.041 b | nd              |              |               | 1533 ± 0.347 b | 0.039 ± 0.013 b | 0.222 ± 0.084 b | nd               |              |              |
| **Urea**  | 0.278 ± 0.029 b | 0.256 ± 0.077 a | 0.224 ± 0.102 a | 0.037 ± 0.008 a | 0.256 ± 0.077 a | nd              |              |               | 0.09 ± 0.032 a | 0.053 ± 0.002 b | 0.012 ± 0.018 b | 0.181 ± 0.035 a | 0.005 ± 0.010 a | nd               |
| **NH₄Cl** | 0.072 ± 0.036 a | 11.32 ± 0.058 b | 0.033 ± 0.008 a | 0.105 ± 0.023 a | 0.037 ± 0.008 a | nd              |              |               | 1386 ± 0.996 b | 0.023 ± 0.011 A | 0.040 ± 0.013 a | 0.052 ± 0.031 A |
Other biochemical analyses were performed, including total protein and carbohydrates (Table S1 in Supplementary Materials 3).

4. Discussion

This study demonstrates the possibility of successfully growing the strain *H. pluvialis* CCAP 34/1D in urea as the principal NS, reaching a high biomass yield and using a low-cost renewable source of nitrogen (Figure 1). The stressors used in our study had a significant effect on carotenoid accumulation, with a minimal negative impact upon the cell number obtained during vegetative growth (Figure 2). Namely, all the treatments, barring FeNO₃, increased the astaxanthin accumulated and all, except NH₄Cl, showed a further increase in the cell number following exposure to stressors (Table 1).

Previous reports showed that the best performance of *H. pluvialis* in terms of growth rate, cell density, and long vegetative stage, without premature encysting, is usually achieved supplying NaNO₃ and/or KNO₃ among possible NS [32,33], with results differing among the strains and the nitrogen concentrations.

In this study, cells grown in urea had a significantly greater growth rate (µ = 0.4) \( (p < 0.0001) \) and maximum cell density \( (4.98 \times 10^5 \text{ cells/mL}, p < 0.0046) \) when compared to cultures grown in any other source of nitrogen tested. Studies investigating growth in different NS are few and have reported conflicting results. For instance, in Cifuentes et al. [34] when comparing different NS including urea, a rate of 0.47 d⁻¹ was achieved in NaNO₃ (a maximum cell density of \( 1.4 \times 10^5 \text{ cells/mL} \)) while 0.44 d⁻¹ was in urea (a maximum cell density of \( 4.5 \times 10^4 \text{ cells/mL} \)), data that was considerably lower than the highest cell density reached in our study. Additionally, Goksan et al. [35] reached the maximum growth rate \( (0.26 \text{ d}^{-1}) \) in NaNO₃ and found out that their strain did not grow under urea as sole NS. Many studies comparing these different NS, found *H. pluvialis* growing optimally in sodium nitrate, which is three and eight times more expensive than...
ammonium and urea, respectively [36]. In another study, Colusse et al. [37] calculated that NaNO₃ accounts for 83.3% of the total chemical cost of all those used to make BG11 medium.

Urea (CO(NH₂)₂) is an organic compound, thus it provides additional C, which is important for the green-red transition and the accumulation of astaxanthin. As a low-cost N-source, urea could enhance the economics of the whole process. The replacement of sodium nitrate with urea has been attempted for other species cultivation, e.g., *Spirulina platensis* with the same purpose of making its cultivation more cost-effective by reducing the input cost without sacrificing the production efficiency [38,39].

Upon improving the green phase of cultivation in terms of biomass productivity, the astaxanthin induction was achieved by applying a mix of stressors at moderate intensity and assessed through a previous screening (Supplementary Materials 2), where the aim was to find the compromise between the highest cell concentration and the highest carotenoids accumulation. This was attained using modified JM (without the addition of extra NS in the form of NaNO₃ but keeping the original recipe containing Ca(NO₃)₂ and (NH₄)₆Mo₇O₂₄·4H₂O) with the addition of 0.25% (w/v) sodium acetate and exposed to 250 μmol photons m⁻² s⁻¹ for 7 days.

Previous studies on the nutrition of *H. pluvialis* have shown that acetate appears to be an important carbon source, enhancing both growth and carotenogenesis [40,41]. This was confirmed in our study and was used in the astaxanthin induction medium. Orosa et al. [42] tested different concentrations of sodium acetate, observing that 0.25% (w/v) acetate led to carotenoids accumulation and biomass increase, while higher concentrations (>0.5% w/v) caused growth inhibition, including a reduction in the chlorophyll a content and cell number decrease, whereas, Cifuentes et al. [34] recorded that the addition of sodium acetate (0.15% w/v) did not produce any increase in cell numbers but induced cyst formation. Zhang et al. [43] exposed the cultures to 10 mM (0.08% w/v) sodium acetate under solar irradiation, obtaining astaxanthin yields more than two-fold compared to the control, after 6 days. Moreo, in other studies, the induction process in mixotrophy led to a decrease in cell density, and in the present study, induction conditions did not stop algal growth. This result can be explained by two factors: (i) the resuspension of the 14 days old cultures in the fresh NaNO₃ depleted buffered medium with the application of a moderate intensity stressor, which could have contributed to further growth; and (ii) the moderate increase of irradiance and the addition of an organic carbon source at 0.25% w/v, did not reduce the cell density, elongating the growth curve and concomitantly enhancing the degree of cell encystment.

Finding moderately stressful conditions is, indeed, essential to guarantee cell survival, e.g., the limitation of nitrogen instead of depletion, or exposure to a moderate increase in light [13]. The presence of nitrogen is essential for the cell division rate and for the accumulation of secondary carotenoids, and in addition, the synthesis of astaxanthin requires nitrogen for continuous protein synthesis in order to support its hyper-accumulation [44]. In the JM original recipe, the nitrogen from NaNO₃ accounts for 941 μM, and in this study, it was eliminated from the induction medium. Meanwhile, the additional sources of nitrogen contained within the original JM recipe, including Ca(NO₃)₂ × 4H₂O (84 μM N) and (NH₄)₆Mo₇O₂₄·4H₂O (0.8 μM N) were retained and their presence was considered enough to maintain background levels of cell metabolism and growth.

When an NS (NaNO₃ in our case) remained available for the cultures, chlorophylls, and total carotenoid concentrations remained at constant levels. However, under nitrogen deficiency, a rapid chlorophyll decline and carotenoid increase per cell occurred—this was avoided in this study due to the medium refresh without the total lack of nitrogen source.

As shown in the Supplementary Materials 3, the protein amount remained nearly constant, thanks to the presence of nitrogen in the medium (Table S1 in Supplementary Materials 3). Other authors found a decrease of proteins and a consequent decrease in photosynthetic activity [45,46]. The increase in the carbohydrate content is an indicator
of the carotenogenesis induction, being the initial precursors in the carotenoids pathway (Table S1 in Supplementary Materials 3) [47].

The carotenoids detected by the HPLC analysis varied from the primary carotenoids commonly found in *H. pluvialis*, such as lutein and β-carotene, to the less frequent antheraxanthin and canthaxanthin [48]. The significant increase of the carotenoid/chlorophyll ratio between green and red phases of cells grown with urea and NaNO₃ was mainly caused by a concomitant reduction of chlorophylls after the induction and a concomitant increase in the carotenoids value (Figure 3). On the other hand, it was observed that cultures grown in FeNO₃ and NH₄Cl, started to encyst before the induction process, from day 10 and 12, respectively, which was not observed in the cultures grown with either urea or NaNO₃. The presence of astaxanthin in the green phase of the FeNO₃ and NH₄Cl treatments confirmed the presence of red cysts already formed before the induction process, as observed under the microscope. Although iron is a necessary element in photosynthesis and it is used to enhance the algal growth during the green phase [49], free ferrous (Fe²⁺) represents a hazard to the Fenton reaction process and generates reactive oxygen species (ROS), thus augmenting *H. pluvialis* astaxanthin synthesis [49,50]. Iron availability, in our case, may have caused a premature accumulation of astaxanthin to help prevent cellular damage caused by ROS production. The FeNO₃, contrarily to the control in NaNO₃, caused premature cell encystment and astaxanthin accumulation, accounting for the highest amount of astaxanthin reached in our treatments (2.71 ± 0.58 mg/g), showing that even if light is essential for its accumulation, other stressors can be equally or even more efficient. Additionally, the main issue regarding NH₄-N nutrition is the conversion of the reduced ion to NH₃ according to chemical equilibrium, and ammonia, which is a toxic nitrogen form and has a direct negative impact on the photosynthetic apparatus of microalgae [40], especially at high N concentrations. Algae exhibit varying tolerances to the NH₄-N nutrition [51].

The HPLC analysis confirmed this different behaviour of *H. pluvialis* cells in the different treatments. In general, the green phase tends to accumulate chlorophylls and primary carotenoids, such as lutein and neoxanthin, in the chloroplasts. The accumulation of astaxanthin was concomitant to the presence of other carotenoids, which are part of the astaxanthin metabolic pattern (involving a step-wise movement from β—carotene—echinenone—canthaxanthin—astaxanthin) [52], and detected only at the end of the green phase. While at the end of carotenogenesis, the presence of astaxanthin was dominant among the carotenoids. Contrarily, at the end of the green phase, other carotenoids were detected: canthaxanthin was present in iron and nitrate treatments; neoxanthin in two replicates of NaNO₃; and urea-treatment presented many unidentified peaks, carotene-like, and echinenone. Canthaxanthin is a precursor of astaxanthin, especially present in the orange cells when they are transitioning from between the green and the red phases of growth; implying the onset of secondary carotenoid accumulation, a part of the astaxanthin metabolic pathway [24]. In fact, in green algal cells, chlorophyll *a*, chlorophyll *b*, and lutein are the main pigments, while small amounts of other carotenoids, such as neoxanthin, violaxanthin, astaxanthin, lutein-like, and canthaxanthin, also exist in green algal cells [53].

5. Conclusions

This study highlights the advantages of a modified JM medium where urea is the primary NS, not only for the enhanced growth of *H. pluvialis* but also for the low-cost sustainable source of nitrogen which could be economically employed at the industrial level.

It also scans a broader image, including an innovative method to induce astaxanthin by a combination of abiotic factors acting as stressors of moderate intensity without the loss of biomass. The combination of the following factors, including nitrogen limitation, the increase of light, and sodium acetate addition, was found to be the best. Different nitrogen sources showed a different mosaic of pigments that converged in the astaxanthin accumulation after exposure to the induction conditions.
Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/app12031261/s1, Figure S1: Growth rate of cultures in each different medium 3N-BBM+V+V, BG11, MWC, JM at 6, 6.5, 7, 7.5, 8 pHs; Figure S2: H. pluvialis 34/1D grown in different N/P ratios: Control with NaNO₃ 2.6/1; 1/1; 5/1; 10/1; Figure S3: Effect of sodium acetate 0%, 0.25%; 0.5%; 1%; 1.5%; 1.75%; 2% on carotenoids accumulation (OD₄₈₀) and cell density (OD₇₅₀) under ML and HL; Figure S4: H. pluvialis 34/1D calibration curve used for cell quantification; Figure S5: Representative HPLC chromatograms of H. pluvialis green (A) and red (B) phases of growth, when cultivated under FeNO₃, NaNO₃, NH₄Cl, or Urea as the principal nitrogen source in Jaworski’s Media; Table S1: Biochemical table reporting the mean value (% DW) ± S.D. within samples (n = 3); Table S2: Ordinary one-way ANOVA and Tukey’s multiple comparison test, with a single pooled variance were performed for H. pluvialis; Table S3: The relative change of H. pluvialis CCAP 34/1D cell number based on the number of cells/mL at the beginning and end of the 7 day induction process; Table S4: Quantitative pigment composition (in mg/g DW), obtained by HPLC analysis, of H. pluvialis cells at the end of the green phase, in cultures previously supplied with equimolar concentrations of nitrogen from different sources; Table S5: For canthaxanthin, a t-test was used to determine if there is a significant difference between the means of two groups, NaNO₃ and FeNO₃; Table S6: Quantitative pigment composition (in mg/g DW), obtained by HPLC analysis, of H. pluvialis cells at the end of the red phases stationary phase, in cultures previously supplied with equimolar concentrations of nitrogen from different sources; Table S7: Pigments variation between the green and the red phase; Table S8: Ordinary two-way ANOVA and Sidak’s multiple comparison test, with individual variances computed for each comparison, were performed; Table S9: Carotenoids/Chlorophylls ratio analysis between the green and the red phase.

Author Contributions: A.R., A.N. and M.E.R. conceived and planned the project, and designed the experiments. A.R. conducted the experiments, and generated, collated, and processed the data. A.R., A.N., B.J. and M.E.R. interpreted the results. A.R. wrote the manuscript. A.R., A.N., B.J. and M.E.R. reviewed and edited the manuscript. A.N., B.J. and M.E.R. supervised the project and acquired the funding. All authors have read and agreed to the published version of the manuscript.

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3. Li, F. Accumulation of Astaxanthin Was Improved by the Nonmotile Cells of Haematococcus pluvialis CCAP 34/1D under ML and HL; Figure S4: H. pluvialis 34/1D calibration curve used for cell quantification; Figure S5: Representative HPLC chromatograms of H. pluvialis green (A) and red (B) phases of growth, when cultivated under FeNO₃, NaNO₃, NH₄Cl, or Urea as the principal nitrogen source in Jaworski’s Media; Table S1: Biochemical table reporting the mean value (% DW) ± S.D. within samples (n = 3); Table S2: Ordinary one-way ANOVA and Tukey’s multiple comparison test, with a single pooled variance were performed for H. pluvialis; Table S3: The relative change of H. pluvialis CCAP 34/1D cell number based on the number of cells/mL at the beginning and end of the 7 day induction process; Table S4: Quantitative pigment composition (in mg/g DW), obtained by HPLC analysis, of H. pluvialis cells at the end of the green phase, in cultures previously supplied with equimolar concentrations of nitrogen from different sources; Table S5: For canthaxanthin, a t-test was used to determine if there is a significant difference between the means of two groups, NaNO₃ and FeNO₃; Table S6: Quantitative pigment composition (in mg/g DW), obtained by HPLC analysis, of H. pluvialis cells at the end of the red phases stationary phase, in cultures previously supplied with equimolar concentrations of nitrogen from different sources; Table S7: Pigments variation between the green and the red phase; Table S8: Ordinary two-way ANOVA and Sidak’s multiple comparison test, with individual variances computed for each comparison, were performed; Table S9: Carotenoids/Chlorophylls ratio analysis between the green and the red phase.

Author Contributions: A.R., A.N. and M.E.R. conceived and planned the project, and designed the experiments. A.R. conducted the experiments, and generated, collated, and processed the data. A.R., A.N., B.J. and M.E.R. interpreted the results. A.R. wrote the manuscript. A.R., A.N., B.J. and M.E.R. reviewed and edited the manuscript. A.N., B.J. and M.E.R. supervised the project and acquired the funding. All authors have read and agreed to the published version of the manuscript.

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