Modeling Plankton Mixotrophy: A Mechanistic Model Consistent with the Shuter-Type Biochemical Approach

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Mixotrophy, i.e., the ability to combine phototrophy and phagotrophy in one organism, is now recognized to be widespread among photic-zone protists and to potentially modify the structure and functioning of planktonic ecosystems. However, few biogeochemical/ecological models explicitly include this mode of nutrition, owing to the large diversity of observed mixotrophic types, the few data allowing the parameterization of physiological processes, and the need to make the addition of mixotrophy into existing ecosystem models as simple as possible. We here propose and discuss a flexible model that depicts the main observed behaviors of mixotrophy in microplankton. A first model version describes constitutive mixotrophy (the organism photosynthesizes by use of its own chloroplasts). This model version offers two possible configurations, allowing the description of constitutive mixotrophs (CMs) that favor either phototrophy or heterotrophy. A second version describes non-constitutive mixotrophy (the organism performs phototrophy by use of chloroplasts acquired from its prey). The model variants were described so as to be consistent with a plankton conceptualization in which the biomass is divided into separate components on the basis of their biochemical function (Shuter-approach; Shuter, 1979). The two model variants of mixotrophy can easily be implemented in ecological models that adopt the Shuter-approach, such as the MIRO model (Lancelot et al., 2005), and address the challenges associated with modeling mixotrophy.

Keywords: constitutive mixotroph (CM), non-constitutive mixotroph (NCM), modeling, phytoplankton, zooplankton

INTRODUCTION

Traditionally, planktonic protists are separated into two distinct functional groups: the phototrophic phytoplankton and the phago-heterotrophic microzooplankton. However, many protists assigned to these two groups are recognized as capable of combining phototrophy and phago-heterotrophy (Flynn et al., 2013). These mixotrophic protists have been reported in all planktonic functional groups (with the notable exception of the diatoms) and they include a large diversity of mixotrophic types (Flynn et al., 2013). In some mixotrophs, growth is mainly supported by phototrophy (e.g., Cryptomonas ovata; Tranvik et al., 1989) while in others, growth
is mainly supported by phagotrophy (e.g., Poterioochromonas malhamensis; Sanders, 1991). Some mixotrophs are forced to use both trophic modes to grow and survive because some essential metabolites come specifically from one of the two metabolic pathways (e.g., the ciliate Laboea strobila; Stoecker et al., 1988); others are facultative mixotrophs (e.g., the dinoflagellate Fragilidium duplocampanaforme; Park et al., 2015). Some mixotrophs use their second trophic mode to fulfill carbon requirements (e.g., the haptophyte Chrysochromulina brevisillum; Stoecker, 1998); for others, it is a means to fulfill nutrient requirements (e.g., the dinoflagellate Procorcentrum minimum; Stoecker, 1998); and for others yet, it is a mechanism to get specific metabolites (as phospholipids; Kimura and Ishida, 1989).

Among this large diversity of mixotrophic types, a major distinction can be made between mixotrophs depending on whether they photosynthesize using their own chloroplasts or using chloroplasts acquired from their phototrophic prey (Mitra et al., 2016). Mixotrophs that actively synthesize and maintain their own chloroplasts are called constitutive mixotrophs (CMs). Mixotrophs that do not constitutively synthesize chloroplasts but photosynthesize using chloroplasts acquired from their phototrophic prey are called non-constitutive mixotrophs (NCMs). The acquired chloroplasts in NCMs remain functional for periods ranging from hours to days, depending on the type of NCM (Mitra et al., 2016). Generalist NCMs, i.e., those that acquire chloroplasts from a broad range of phototrophic prey, have a poor ability to maintain functional chloroplasts (Dolan and Pérez, 2000). By contrast, specialist NCMs, i.e., those that need to acquire chloroplasts from specific phototrophic prey, can maintain a photosynthetic activity for long periods (Stoecker et al., 2009; Hansen et al., 2013). Specialist NCMs can further be divided into those that retain the entire photosynthetic prey (endosymbiosis) and those that only retain the chloroplasts (kleptochloroplasty; Mitra et al., 2016). In a functional capacity, the CMs are closer to the strict phototrophs while the NCMs are closer to the strict phago-heterotrophs.

While mixotrophy has been reported for a long time, it is now appreciated as being much more widespread in aquatic ecosystems than initially thought. Previously, the mixotrophic status was only accorded to some dinoflagellates, forams, radiolarians, and acantharia while now, mixotrophy has been acknowledged in all eukaryote planktonic microorganism groups, except diatoms (Sanders and Porter, 1988; Burkholder et al., 2008; Flynn et al., 2013). In addition, these species are not limited to a specific habitat: mixotrophs are observed in both freshwater and marine (Sanders, 1991; Stoecker et al., 2009), oligotrophic and eutrophic systems (Burkholder et al., 2008), and from polar to equatorial regions (Zubkov and Tarran, 2008; Stoecker et al., 2009; Sanders and Gast, 2012). In coastal areas, mixotrophic flagellates can account for some 50% of the pigmented biomass (Havskum and Riemann, 1996) and chloroplast-containing ciliates can account for 40–60% of the planktonic ciliates in summer (Stoecker et al., 1987; Bernard and Rassoulzadegan, 1994). In oceanic waters, mixotrophs account for 40–95% of the bacterivory in the euphotic layer (Zubkov and Tarran, 2008).

Constitutive mixotrophy is suggested to play an important ecological role when inorganic nutrients are low (e.g., in oligotrophic systems; e.g., Arenovski et al., 1995) or unbalanced (e.g., in eutrophic systems; e.g., Nygaard and Tobiesen, 1993; Burkholder et al., 2008), and when light is limiting (e.g., during the polar night or occasionally in eutrophied systems; e.g., Bird and Kalff, 1986; Roberts and Laybourn-Parry, 1999; Jones et al., 2009). In such systems, prey ingestion provides nutrients and energy. Similarly, non-constitutive mixotrophy may be advantageous in “high light–low prey” ecosystems (e.g., in oligotrophic systems; e.g., Skovgaard, 1998; Hansen et al., 2013) because acquired phototrophy supplies carbon by photosynthesis. Finally, mixotrophy is worthy of interest in an environmental perspective because most marine harmful algae have been reported as mixotrophs (Stoecker et al., 2006; Burkholder et al., 2008; Jeong et al., 2010).

Several models have been developed with the specific aims of exploring planktonic mixotrophy from an ecophysiological or ecological (species competition) perspective. Most interest has been leveled at the potential impact of mixotrophs on the microbial food web structure and functioning, and the conditions under which mixotrophs may likely coexist with strict phototrophs and heterotrophs (Thingstad et al., 1996; Baretta-Bekker et al., 1998; Stickney et al., 2000; Jost et al., 2004; Hammer and Pitchford, 2005; Hood et al., 2006; Flynn and Mitra, 2009; Crane and Grover, 2010; Ward et al., 2011; Våge et al., 2013). By far the greater effort has been applied to CM organisms. The complexity of the model structure varies widely among such studies. The simplest models describe mixotrophy as the ability to combine both phototrophy and heterotrophy without any feedbacks or trade-offs between the two nutritional modes and organisms have a fixed stoichiometry (e.g., Hammer and Pitchford, 2005). At the other extreme, the most complex model explicitly describes the main regulative processes that occur between phototrophy and heterotrophy in the mixotroph and allows for a variation of the cellular stoichiometry (Flynn and Mitra, 2009).

Despite the potential significance of mixotrophy in aquatic systems and the existence of mathematical models describing this trophic mode, few ecological/biogeochemical models consider it explicitly (e.g., Mitra et al., 2014). Those models that do include mixotrophs demonstrate the potential for mixotrophy to significantly impact the flow of energy and nutrients in the system. In particular, primary production can potentially be increased by a factor 2 compared to a system in which mixotrophy is not considered, thanks to a shortened and more effective chain from nutrient regeneration to primary production (Mitra et al., 2014).

Considering mixotrophy in biogeochemical/ecological models is however challenging due to the large diversity of mixotrophic types, the scarcity of data allowing the parameterization of physiological processes, and the need to make the inclusion of mixotrophy into existing ecosystem models as simple as possible. Indeed, the addition of a mixotroph functional type description into ecosystem models will inevitably be tempered by the structure of the extant ecosystem model.
Regarding the diversity of mixotrophic types, this paper proposes a flexible mechanistic model featuring the two main types of mixotrophy: constitutive and non-constitutive. The two mixotrophic types have indeed different ecophysiology and different relation to their prey (the NCM being strictly dependent on the presence of prey); therefore, they affect differently the ecosystem dynamics (Mitra et al., 2016). In addition, the version describing constitutive mixotrophy can be configured to represent either CMs that favor either phototrophy or heterotrophy. Simulations have been performed to compare each mixotrophic type with the strict trophic type it most closely resembles from a functional perspective (the CM with the strict phototroph and the NCM with the strict heterotroph) in order to explore the competitive advantage of mixotrophy over strict forms.

As for the explicit inclusion of mixotrophy into existing biogeochemical/ecological models, we were mainly interested in introducing mixotrophy in a form consistent with the mechanistic structure of the AQUAPHY model (Lancelot et al., 1991); this model has been used to describe primary production via phytoplankton growth physiology within several biogeochemical models (Billen et al., 1994; Lancelot et al., 2000, 2005). A feature of AQUAPHY is that the total cellular carbon is divided into separate classes of components on the basis of their function: these comprise synthetic and structural material (i.e., photosynthetic apparatus, ribosomes, genetic material, membranes, etc.), stored carbon (i.e., carbohydrates, lipids), and carbon monomers (i.e., photosynthetic products and precursors of complex molecules). This mechanistic approach was suggested by Shuter (1979) as a means to successfully describe the observed behavior of a variety of unicellular algal species, quantitatively and qualitatively. The work is thus a first step in the implementation of mixotrophy in the biogeochemical models that adopt a “Shuter mechanistic approach” to describe the phytoplankton physiology.

**METHODS**

**Constitutive Mixotrophy**

The model of constitutive mixotrophy was constructed on the basis of the following hypotheses, summarizing the main qualitative observations related to CMs: (i) the CM is preferentially phototrophic but it can supplement its nutrient requirements (under inorganic nutrient limitation) or its carbon requirements (under light limitation) by ingesting prey (e.g., in Jones et al., 1995; Carvalho and Granéli, 2010; McKie-Krisberg et al., 2015); (ii) the maximum mixotrophic growth is not allowed to exceed the maximum phototrophic growth; (iii) the phototrophic growth has to account for at least 10% of the mixotrophic growth, such that we describe an obligate phototroph (Caron et al., 1993; Brutemark and Granéli, 2011); (iv) the nutrients regenerated by heterotrophic activity (PO$_4^{3-}$, NH$_4^+$) are re-assimilated to sustaining phototrophic growth with any surplus being excreted (Flynn and Mitra, 2009).

The model combines an adaptation of the AQUAPHY model (Lancelot et al., 1991; Ghyoot et al., 2015) for the phototrophic pathway, coupled to a simple zooplankton model (Lancelot et al., 2005) for the phago-heterotrophic path. It is important to note that, in keeping with the formulation of the AQUAPHY model, the description of mixotrophy given here describes biomass growth (i.e., mol C m$^{-3}$ time$^{-1}$) rather than per capita growth (i.e., C C$^{-1}$ time$^{-1}$) as in some other models (e.g., Flynn and Mitra, 2009). Figure 1A shows the schematic representation of the CM growth physiology, linking the phototrophic (dark gray) and heterotrophic (light gray) components. The CM model contains six state variables (Table 1) describing intracellular components on the basis of their function: the functional and structural metabolites (e.g., nucleic acids, proteins) synthesized by phototrophic and heterotrophic activities ($F_{\text{phot}}$ and $F_{\text{het}}$, respectively), soluble carbon monomers (i.e., early products of photosynthesis ($S_{\text{C}}$), carbon reserves (i.e., carbohydrates, fatty acids; $R_{\text{C}}$), intracellular soluble phosphate ($S_{\text{P}}$), and intracellular soluble inorganic nitrogen ($S_{\text{N}}$). The total C-cell biomass (mmol C m$^{-3}$) is given by the sum of $F_{\text{phot}}$, $F_{\text{het}}$, $S_{\text{C}}$, and $R_{\text{C}}$. The $F_{\text{phot}}$ and $F_{\text{het}}$ pools are assumed to have a fixed C:N:P stoichiometry, based on biochemical constraints (Geider and Laroche, 2002). Variable cellular stoichiometry is enabled by considering the additional C, P, and N accumulated as carbon monomers ($S_{\text{C}}$), carbohydrates and fatty acids ($R_{\text{C}}$), soluble inorganic phosphorus ($S_{\text{P}}$), and soluble inorganic nitrogen ($S_{\text{N}}$). The model of constitutive mixotrophy is linked to three state variables describing external inorganic nutrients—dissolved inorganic nitrogen ($\text{DIN} = \text{NO}_3^- + \text{NH}_4^+$) and phosphate ($\text{PO}_4^{3-}$)—and also the prey which in reality expresses a variable C:N:P stoichiometry (though here prey stoichiometry is fixed as the emphasis is on the description of the autecology of the mixotrophs, rather than system ecology).

Tables 2, 3 show, respectively, the nine conservation equations related to the state variables and the associated processes. Phototrophic growth ($\mu_{\text{phot}}$; Equation 17) is controlled by the concentration of the structural and functional metabolites related to phototrophic activity ($F_{\text{phot}}$), by the limitation in carbon monomers $S_{\text{C}}$—either directly produced by photosynthesis ($\psi$; Equation 20) or indirectly by $R_{\text{C}}$ catabolism (cat$_{\text{RC}}$; Equation 10)—and by the limitation in internal soluble inorganic nutrients ($S_{\text{N}}$ and $S_{\text{P}}$). The $S_{\text{C}}$ limitation is formulated by a Michaelis–Menten equation in which the substrate concentration is expressed by $X_{\text{SC}} - k_{\text{SC}}$, with $X_{\text{SC}} = \frac{S_{\text{C}}}{k_{\text{SC}}}$ and $k_{\text{SC}}$, which is the minimum value for $X_{\text{SC}}$ (assumed to be equal to the half-saturation constant for $S_{\text{C}}$ assimilation). The inorganic nutrient limitation is formulated by the Liebig's minimum law in which the limitation for each nutrient is expressed by a hyperbolic function depending on $X_{\text{SN},p}$ (the ratio between $S_{\text{N}}$ and the N,P contained in $F_{\text{phot}}$). The uptake of inorganic nutrients ($\text{upt}_{\text{DIN}}$ and $\text{upt}_{\text{PO4}}$; Equations 26 and 27) depends on the external nutrient concentration and the status of the internal nutrient reserve. The phototrophic respiration ($\text{resp}_{\text{phot}}$; Equation 22) includes costs for cellular maintenance and for synthesis of new $F_{\text{phot}}$.

Prey ingestion (graz$_{\text{p}}$; Equation 14) is controlled by the concentration of the structural and functional metabolites related to heterotrophic activity ($F_{\text{het}}$) and by prey availability; the latter
is controlled by a sigmoid (type III) function. From the ingested prey, a non-assimilated fraction is egested (egest; Equation 11) as dissolved and particulate organic matter (loss terms), a fraction is respired to meet the heterotrophic metabolic costs and is released as carbon dioxide (resp\text{het}; Equation 21), and the last fraction is assimilated and contributes directly to the heterotrophic growth (\(\mu\text{het};\) Equation 16). If the nutrient content of the prey is higher than the nutrient required by \(F_{\text{het}}\), the surplus is regenerated with no limitation (\(\mu\text{reg};\) Equation 19). When the 24 h-average \(S_C\) limitation is below a threshold value (thresh\(S_C\lim\)) set here as 0.15, we assumed that the mixotroph switches to perform increasing levels of phagotrophy to acquire carbon (in addition to nutrients if also nutrients limited) and, therefore, the maximum mixotrophic growth is equal to the phototrophic growth obtained with no limitation (\(\mu_{\text{mix}}\text{max};\) Equation 18). When the 24 h-average \(S_C\) limitation is above this threshold value, we assumed that the CM undertakes phagotrophy to acquire nutrients and, therefore, the maximum mixotrophic growth is equal to the phototrophic growth obtained with no nutrient limitation (\(\mu_{\text{mix}}\text{max};\) Equation 19). If the sum of the phototropic and the heterotrophic growth is higher than the maximum mixotrophic growth, either the grazing or the photosynthesis rate is regulated to

**TABLE 1 |** Model state variables (all expressed as a concentration in water).

| State variable | Description | Units |
|----------------|-------------|-------|
| \(\mu_{\text{phot}}\) | Functional and structural metabolites related to photothrophy | mmolC m\(^{-3}\) h\(^{-1}\) |
| \(\mu_{\text{het}}\) | Functional and structural metabolites related to heterotrophy | mmolC m\(^{-3}\) h\(^{-1}\) |
| \(S_C\) | Soluble carbon monomers (early products of photosynthesis) | mmolC m\(^{-3}\) |
| \(R_C\) | Carbon reserves (carbohydrates, fatty acids) | mmolC m\(^{-3}\) |
| \(S_N\) | Intracellular soluble inorganic nitrogen | mmolN m\(^{-3}\) |
| \(S_p\) | Intracellular soluble phosphate | mmolP m\(^{-3}\) |

**TABLE 2 |** Conservation equations for the constitutive mixotroph (CM).

| Eq. | Conservation equations | Units |
|-----|-----------------------|-------|
| 1 | \(\frac{d\mu_{\text{phot}}}{dt} = \mu_{\text{phot}} - \text{lys}_{\text{phot}}\) | mmolC m\(^{-3}\) h\(^{-1}\) |
| 2 | \(\frac{d\mu_{\text{het}}}{dt} = \mu_{\text{het}} - \text{lys}_{\text{het}}\) | mmolC m\(^{-3}\) h\(^{-1}\) |
| 3 | \(\frac{dS_C}{dt} = -\text{excr}_{\text{C}} - \text{synth}_{\text{C}} + \text{cat}_{\text{RC}} - \mu_{\text{phot}} - \text{resp}_{\text{het}} - \text{lys}_{\text{SC}}\) | mmolC m\(^{-3}\) |
| 4 | \(\frac{dS_C}{dt} = \text{cat}_{\text{RC}} - \text{lys}_{\text{RC}}\) | mmolC m\(^{-3}\) |
| 5 | \(\frac{dS_N}{dt} = \text{upt}_{\text{DIN}} + \frac{\text{upt}_{\text{Pi}}}{\text{C}_{\text{Pi}}} + \text{resp}_{\text{het}} - \text{lys}_{\text{SN}}\) | mmolN m\(^{-3}\) |
| 6 | \(\frac{dS_p}{dt} = -\text{upt}_{\text{PO}_4} + \text{excr}_{\text{SP}} + \text{lys}_{\text{SP}}\) | mmolP m\(^{-3}\) |
| 7 | \(\frac{d\mu_{\text{mix}}}{dt} = -\text{graz}\) | mmolC m\(^{-3}\) h\(^{-1}\) |

Mixotrophic growth is computed as the sum of the phototropic and the heterotrophic growth rate, but limited by the maximum mixotrophic growth. The latter is computed differently according to whether phagotrophy is used to offset a lack of carbon or a lack of nutrients. When the 24 h-average \(S_C\) limitation is below a threshold value (thresh\(S_C\lim\)) set here as 0.15, we assumed that the mixotroph switches to perform increasing levels of phagotrophy to acquire carbon (in addition to nutrients if also nutrients limited) and, therefore, the maximum mixotrophic growth is equal to the phototrophic growth obtained with no limitation (\(\mu_{\text{mix}}\text{max};\) Equation 18). When the 24 h-average \(S_C\) limitation is above this threshold value, we assumed that the CM undertakes phagotrophy to acquire nutrients and, therefore, the maximum mixotrophic growth is equal to the phototrophic growth obtained with no nutrient limitation (\(\mu_{\text{mix}}\text{max};\) Equation 19). If the sum of the phototropic and the heterotrophic growth is higher than the maximum mixotrophic growth, either the grazing or the photosynthesis rate is regulated to...
limit the mixotrophic growth to its maximum. In other terms, the mixotroph can favor either phototrophic or heterotrophic growth. The model offers the possibility to test either of these two configurations: the CM that regulates its grazing rate (i.e., tends to grow phototrophically) hereafter referred to as “Reg\textsuperscript{graz}” and the CM that regulates its photosynthesis (i.e., tends to grow heterotrophically) hereafter referred to as “Reg\textsuperscript{phot}.” In either case, we assumed that phototrophic growth has to represent at least 10% of the mixotrophic growth. This implies that phagotrophy cannot sustain growth under prolonged dark conditions because of an obligatory demand for products of photosynthesis.

TABLE 3 | Equations that describe the processes occurring in the constitutive mixotroph (CM).  

| Eq. | Process | Value | Units | Explanation |
|-----|---------|-------|-------|-------------|
| 10  | cat\textsubscript{RC} | $k\text{excr} \cdot R\text{C}$ | mmolC m\textsuperscript{-3} h\textsuperscript{-1} | Catabolism of carbon reserves |
| 11  | egest  | $fp \cdot $graz | mmolC m\textsuperscript{-3} h\textsuperscript{-1} | Egestion of non-assimilated prey |
| 12  | excr\textsubscript{G} | $k\text{excr} \cdot SC$ | mmolC m\textsuperscript{-3} h\textsuperscript{-1} | Excretion of soluble C monomers |
| 13  | excr, ($i = N, P$) | $reg_i - ret_i$ ($i = N, P$) | mmol i m\textsuperscript{-3} h\textsuperscript{-1} | Excretion of NH\textsubscript{4}\textsuperscript{+} and PO\textsubscript{4}\textsuperscript{3-} which are regenerated by the heterotrophic activity but are not retained in the reserves SN and SP (i.e., tends to grow phototrophically) hereafter referred to as “Reg\textsuperscript{graz}” and the CM that regulates its photosynthesis (i.e., tends to grow heterotrophically) hereafter referred to as “Reg\textsuperscript{phot}.” In either case, we assumed that phototrophic growth has to represent at least 10% of the mixotrophic growth. This implies that phagotrophy cannot sustain growth under prolonged dark conditions because of an obligatory demand for products of photosynthesis. |
Parameter values were selected here to describe constitutive mixotrophic nanoflagellates that graze on bacteria to fulfill nutrient or energy requirements (Table 4). Parameters related to phototrophic carbon pathways are those of phototrophic nanoflagellates as used in the MIRO model (the MIRO model being a biogeochemical model that uses AQUAPHY to represent phytoplankton growth and that describes the planktonic ecosystem of the Southern North Sea; Lancelot et al., 2005). Parameters related to phototrophic P-pathways are similar to those presented in Ghyoot et al. (2015). Parameters related to phototrophic N-pathways and parameters related to heterotrophic activity were estimated by implementing the mixotrophy model into the MIRO model and by tuning the model against observations reported in the Belgian coastal zone (Ghyoot et al., submitted). Observations used in that tuning included planktonic biomass (bacteria, nanoflagellates, diatoms, Phaeocystis colonies, microzooplankton, and copepods) and nutrient concentrations [NO\textsubscript{3}, NH\textsubscript{4}, PO\textsubscript{4}, and dissolved silica (DSi)].

Single parameter steady-state model sensitivity to parameters values was studied with the method of Haefner (1996). The model was run under “low DIN–high prey” chemostat-type conditions and a normalized sensitivity index (SI; Equation 28) based on steady-state biomass was calculated for each parameter:

\[
SI = \left( \frac{R - R_{\text{ref}}}{p - p_{\text{ref}}} \right) \frac{R_{\text{ref}}}{p_{\text{ref}}}
\]

Where \( R_{\text{ref}} \) is the value of CM biomass reached at steady-state with the reference parameter value \( p_{\text{ref}} \) (Table 4) and \( R \) is the value of CM biomass reached at steady-state with \( p \), the reference parameter increased/decreased by 25%. The SI-value is thus a measure of the relative variation of CM biomass compared to the relative variation of the parameter.

Non-constitutive Mixotrophy

We constructed the model of non-constitutive mixotrophy on the basis of available qualitative observations specific to this type of mixotrophy: (i) the photosynthetic capacity of the ingested phototrophic prey is retained for some time in the food vacuole so that it provides carbon to the mixotroph (Skovgaard, 1998); (ii) in the food vacuole, there is no replication of the functional metabolites related to phototrophy; they are only supplied by phagotrophy on phototrophic prey; (iii) the digestion rate of the ingested phototrophic prey is constant; (iv) nutrients regenerated through heterotrophic activity (PO\textsubscript{4}, NH\textsubscript{4}) can be retained for the phototrophic activity while the excess is excreted outside the cell; (v) there is no inorganic nutrient uptake (we assume these NCMs are generalist rather than specialist NCMs; the latter, such as Mesodinium and Dinophysis, are capable of using externally supplied inorganic nutrients; Hansen et al., 2013); (vi) the organism does not require some level of phototrophy to grow, meaning that we describe a facultative mixotroph (but a minimum level of phagotrophy is required for obtaining chloroplasts).

As for the model of constitutive mixotrophy, the model of non-constitutive mixotrophy combines the AQUAPHY model for the phototrophic path and a simple zooplankton model for the heterotrophic path. Figure 1B shows the schematic representation of the NCM growth physiology, showing the heterotrophic (light gray) and phototrophic (dark gray) components. While the model of non-constitutive mixotrophy deploys the same six state variables as the model of constitutive mixotrophy and operates with the same external factors (Table 1), some conservation equations (Table 5) and processes (Table 6) differ. The differences mainly rely on the fact that NCMs acquire their phototrophic capacity by ingesting phototrophic prey and they are not able to permanently maintain this capacity.

The phototrophic prey biomass (prey) is characterized by the five phototrophic compounds usually used within AQUAPHY: F\textsubscript{phot}, S\textsubscript{C}, R\textsubscript{C}, S\textsubscript{N}, and S\textsubscript{P} (we have therefore: prey\textsubscript{phot}, prey\textsubscript{SC}, prey\textsubscript{RC}, prey\textsubscript{SN}, and prey\textsubscript{SP}). The grazing on each of these five compounds (graz\(i\), with \( i = \text{F}^{\text{phot}}, \text{S}^{\text{C}}, \text{R}^{\text{C}}, \text{S}^{\text{N}}, \text{S}^{\text{P}}\); Equation 41) is computed as the grazing on the total prey biomass (prey = prey\textsubscript{phot} + prey\textsubscript{SC} + prey\textsubscript{RC}) relative to the compound concentration. Once ingested by the NCM, the five compounds are distributed to their corresponding pools. As there is no mechanism for synthesis or maintenance of F\textsubscript{phot} in the NCM configuration, F\textsubscript{phot} is exclusively supplied via grazing on phototrophic prey\textsubscript{phot} (Equation 30). S\textsubscript{C}, R\textsubscript{C}, S\textsubscript{N}, and S\textsubscript{P} are supplied by the grazing but also by other (phototrophic and heterotrophic) processes (Equations 31–34). Photosynthesis and R\textsubscript{C} catabolism supply S\textsubscript{C} in the same way as in the model of constitutive mixotrophy (\( \omega \); cf. Equation 20 and cat\textsubscript{RC}; cf. Equation 10). R\textsubscript{C} synthesis supplies R\textsubscript{C}, as in the model of constitutive mixotrophy (synth\textsubscript{RC}; cf. Equation 25), while nutrient retention supplies S\textsubscript{N} and S\textsubscript{P} (ret\(i\), with \( i = \text{N}, \text{P}\); cf. Equation 24).

The phototrophic components obtained by grazing enables phototrophic growth to be computed as for the model of constitutive mixotrophy (\( \mu^{\text{phot}} \); cf. Equation 17) but now the phototrophic growth supports the production of F\textsubscript{het} instead of F\textsubscript{phot} (Equation 29). As there is no synthesis and maintenance of F\textsubscript{phot}, phototrophic respiration includes only costs for the synthesis of F\textsubscript{het} (resp\textsubscript{phot}, Equation 44).

The phototrophic components in NCMs (F\textsubscript{phot}, S\textsubscript{C}, R\textsubscript{C}, S\textsubscript{N}, S\textsubscript{P}) are continuously degraded (de facto digested) at a constant rate (dig\(i\), with \( i = \text{F}^{\text{phot}}, \text{S}^{\text{C}}, \text{R}^{\text{C}}, \text{S}^{\text{N}}, \text{S}^{\text{P}}\); Equation 39). The digested material has three possible fates: the non-assimilated material is egested outside the cell (eges\(i\); Equation 40) as dissolved and particulate organic matter, a fraction of the assimilated material is respired (resp\textsubscript{het}; Equation 43) to meet the heterotrophic metabolic costs, and the remaining fraction is used for heterotrophic growth (\( \mu^{\text{het}} \); Equation 42). If the C:N:P stoichiometry of the digest is higher than the C:N:P stoichiometry of F\textsubscript{het}, NH\textsubscript{4} and PO\textsubscript{4} are regenerated (reg\(i\), with \( i = \text{N}, \text{P}\); Equation 45). These nutrients can be retained in S\textsubscript{N} and S\textsubscript{P} and contribute to the phototrophic growth (ret\(i\), with \( i = \text{N}, \text{P}\) calculated as in CMs; cf. Equation 24). As we assumed that inorganic nutrient uptake does not occur in NCMs, these retained inorganic nutrients are the only ones that allow phototrophic growth. If the reserves S\textsubscript{N} and S\textsubscript{P} are full, the regenerated nutrients are excreted outside the cell (exc\(e\), with \( i = \text{N}, \text{P}\) calculated as in CMs; cf. Equation 13).
TABLE 4 | Constant parameters used in the constitutive mixotroph (CM) and non-constitutive mixotroph (NCM) models.

| Parameters     | CM         | NCM        | Units                  | Explanation |
|---------------|------------|------------|------------------------|-------------|
| \( \alpha \)  | 0.002      | 0.002      | h\(^{-1}\) \(\mu\text{mol}^{-1}\) m\(^{-3}\) s | Parameter of light adaptation |
| \( \text{CChl}_{\text{phot}} \) | 2.083      | 2.083      | mmol\(\text{C} \text{mgChl}^{-1} \) | C:Chl ratio for \(F_{\text{phot}}\) |
| \( \text{DIN}_{\text{phot}} \) | 5.25       | 5.25       | \(\text{molC molN}^{-1} \) | C:N ratio for \(F_{\text{phot}}\) |
| \( \text{CP} \) | 4.1        | 4.1        | \(\text{molC molP}^{-1} \) | C:P ratio for \(F_{\text{phot}}\) |
| \( \text{S}_{\text{prey}} \) | 4.67 variable | \(\text{molC molP}^{-1} \) | C:P ratio for prey |
| \( \text{PO} \) | 84         | 66         | \(\text{molC molP}^{-1} \) | Light extinction coeff. related to chl. |
| \( \text{NP} \) | 74.67      | variable   | \(\text{molC molP}^{-1} \) | Light extinction coeff. related to chl. |
| \( \eta_{\text{self}} \) | 0.02       | 0.02       | m\(^{-1} \) | Egested fraction of ingestion |
| \( \eta_{\text{ing}} \) | 0.25       | 0.25       | – | Maximum grazing rate (at optimal T) |
| \( \eta_{\text{cap}} \) | 0.07       | 0.052      | h\(^{-1} \) | Constant of \(R_{\text{C}}\) in-Rasobiosis (at optimal T) |
| \( \eta_{\text{dig}} \) | 0.06       | 0.06       | h\(^{-1} \) | Constant digestion of ingested prey \(\text{at opt. T}\) |
| \( \eta_{\text{catabolism}} \) | 0.001      | 0.001      | – | Constant of \(S_{\text{C}}\) excretion |
| \( \eta_{\text{dig}} \) | 1.25       | 1.75       | mmol\(\text{C m}^{-3} \) | Half-saturation constant of grazing |
| \( \eta_{\text{lys}} \) | 0.0025     | 0.0025     | h\(^{-1} \) | Constant of cell autolysis \(\text{at opt. T}\) |
| \( \mu_{\text{maint}} \) | 0.00081    | –          | h\(^{-1} \) | Constant of cellular maintenance \(\text{at opt. T}\) |
| \( \eta_{\text{in}} \) | 0.1        | 0.1        | h\(^{-1} \) | Maximum photosynthesis rate \(\text{at opt. T}\) |
| \( \eta_{\text{prey}} \) | 0.2        | –          | mmol\(\text{N m}^{-3} \) | Half-saturation constant for \(\text{DIN}\) uptake |
| \( \eta_{\text{PP} \text{uptake}} \) | 0.25       | –          | mmol\(\text{P m}^{-3} \) | Half-saturation constant for PO\(_\text{uptake}\) |
| \( \eta_{\text{PP} \text{uptake}} \) | 0.06       | 0.06       | – | Half-saturation constant for \(\text{PO}_{\text{uptake}}\) |
| \( \eta_{\text{PP} \text{uptake}} \) | 0.014      | 0.014      | – | Half-saturation constant for \(\text{PP}_{\text{uptake}}\) |
| \( \eta_{\text{PP} \text{uptake}} \) | 0.02       | 0.02       | – | Half-saturation constant for \(\text{PP}_{\text{uptake}}\) |
| \( \eta_{\text{PP} \text{uptake}} \) | 0.09       | 0.09       | h\(^{-1} \) | Half-saturation constant for \(\text{DIN}_{\text{uptake}}\) regulation |
| \( \eta_{\text{PP} \text{uptake}} \) | 0.1        | 0.1        | h\(^{-1} \) | Half-saturation constant for \(\text{PO}_{\text{uptake}}\) regulation |
| \( \text{DIN}_{\text{uptake}} \) | 0.15       | –          | – | Maximum phototrophic growth rate \(\text{at opt. T}\) |
| \( \mu_{\text{PP} \text{uptake}} \) | 0.2        | –          | h\(^{-1} \) | Maximum \(\text{PP}_{\text{uptake}}\) growth rate \(\text{at opt. T}\) |
| \( \mu_{\text{PP} \text{uptake}} \) | 0.24       | –          | h\(^{-1} \) | Maximum \(\text{PP}_{\text{uptake}}\) growth rate \(\text{at opt. T}\) |
| \( \mu_{\text{PP} \text{uptake}} \) | 0.14       | 0.14       | – | Maximum \(\text{PP}_{\text{uptake}}\) growth rate \(\text{at opt. T}\) |
| \( \mu_{\text{PP} \text{uptake}} \) | 0.2        | 0.2        | – | Maximum \(\text{PP}_{\text{uptake}}\) growth rate \(\text{at opt. T}\) |
| \( \mu_{\text{PP} \text{uptake}} \) | 0.35       | 0.35       | – | Maximum \(\text{PP}_{\text{uptake}}\) growth rate \(\text{at opt. T}\) |

Values were taken from Lancelot et al. (2005), Ghyoot et al. (2015) and Ghyoot et al. (submitted).

TABLE 5 | Conservation equations for the non-constitutive mixotroph (NCM).

| Eq. | Conservation equations | Units |
|-----|------------------------|-------|
| 29  | \( \frac{d\text{PP}}{dt} = \mu_{\text{het}} + \mu_{\text{phot}} - \text{lyss}_{\text{phot}} \) | mmol\(\text{C m}^{-3} \text{h}^{-1} \) |
| 30  | \( \frac{d\text{PP}}{dt} = \text{graz}_{\text{PP}} - \text{lyss}_{\text{PP}} \) | mmol\(\text{C m}^{-3} \text{h}^{-1} \) |
| 31  | \( \frac{d\text{PP}}{dt} = \text{graz}_{\text{PP}} - \text{lyss}_{\text{PP}} \) | mmol\(\text{C m}^{-3} \text{h}^{-1} \) |
| 32  | \( \frac{d\text{PP}}{dt} = \text{graz}_{\text{PP}} - \text{lyss}_{\text{PP}} \) | mmol\(\text{C m}^{-3} \text{h}^{-1} \) |
| 33  | \( \frac{d\text{PP}}{dt} = \text{graz}_{\text{PP}} - \text{lyss}_{\text{PP}} \) | mmol\(\text{C m}^{-3} \text{h}^{-1} \) |
| 34  | \( \frac{d\text{PP}}{dt} = \text{graz}_{\text{PP}} - \text{lyss}_{\text{PP}} \) | mmol\(\text{C m}^{-3} \text{h}^{-1} \) |
| 35  | \( \frac{d\text{PP}}{dt} = \text{graz}_{\text{PP}} - \text{lyss}_{\text{PP}} \) | mmol\(\text{C m}^{-3} \text{h}^{-1} \) |
| 36  | \( \frac{d\text{PP}}{dt} = \text{graz}_{\text{PP}} - \text{lyss}_{\text{PP}} \) | mmol\(\text{C m}^{-3} \text{h}^{-1} \) |

With the selected parameters values (Table 4), the model describes a generalist non-constitutive mixotrophic microzooplankton (ciliate) that feeds on phototrophic nanoflagellates and uses their chloroplasts to photosynthesize. The value of the constant rate of degradation of acquired photosystems is 0.03 \(\text{h}^{-1} \), the same as in Flynn and Hansen (2013). The other parameters related to heterotrophic processes were estimated by implementing the mixotrophy models into the MIRO model that describes the planktonic ecosystem of the Southern North Sea and by tuning the model against observations reported in this area (Ghyoot et al., submitted). Parameters values involved in phototrophic processes are those of the phototroph prey, i.e., the nanoflagellates, as in the model of constitutive mixotrophy.
TABLE 6 | Equations that describe the processes occurring in the non-constitutive mixotroph (NCM).

| Eq. | Process | Value | Units | Explanation |
| --- | --- | --- | --- | --- |
| 38 | dig | \( \mu_{\text{dig}} \cdot \left( F_{\text{phot}} + S_{C} + R_{C} \right) \) | mmol C m\(^{-3}\) h\(^{-1}\) | Digestion of the ingested prey at a constant rate |
| 39 | \( \text{dig/\text{i}} = F_{\text{phot},i}, S_{C}, R_{C}, S_{S}, S_{p} \) | \( \mu_{\text{dig}} \cdot \left( F_{\text{phot}} + S_{C} + R_{C} \right) \) | mmol N m\(^{-3}\) h\(^{-1}\) | Digestion of the ingestion prey according to the cellular component |
| 40 | egest | \( f_{p} \cdot \text{dig} \) | mmol C m\(^{-3}\) h\(^{-1}\) | Egestion of non-assimilated prey |
| 41 | \( \text{graz/i} = F_{\text{phot},i}, S_{C}, R_{C}, S_{S}, S_{p} \) | \( g_{\text{max}} \cdot \frac{\text{pred}}{\text{pred}^2 + k_{S}^{2}} \cdot \frac{\text{pred}}{S_{C}} \cdot S_{i} \) | mmol C m\(^{-3}\) h\(^{-1}\) | Grazing |
| 42 | \( \mu_{\text{het}} \) | \( y \cdot \text{dig} \) | mmol C m\(^{-3}\) h\(^{-1}\) | Heterotrophic growth |
| 43 | \( \text{resp}^{\text{het}} \) | \( (1 - f_{p} - y) \cdot \text{dig} \) | mmol C m\(^{-3}\) h\(^{-1}\) | Heterotrophic respiration which is the carbon part of the prey that is not egested and not assimilated in \( \text{f}_{\text{het}} \) |
| 44 | \( \text{resp}^{\text{phot}} \) | \( \xi \cdot \mu_{\text{phot}} \) | mmol C m\(^{-3}\) h\(^{-1}\) | Phototrophic respiration including only costs for synthesis of new \( \text{f}_{\text{het}} \) |
| 45 | \( \text{reg, i} = N, P \) | \( (1 - f_{p}) \cdot \frac{\text{dig}}{C_{i}^{\text{phot}} + f_{\text{phot}}^{\text{het}}(i = N, P)} - \frac{\mu_{\text{het}}}{C_{i}^{\text{phot}}} \) | mmol N m\(^{-3}\) h\(^{-1}\) | Regeneration of NH\(_{3}\) and PO\(_{4}\) through heterotrophic activity (regeneration is then distributed between retention and excretion). The Ci of the ingested prey \( \left( C_{i}^{\text{phot}} + f_{\text{phot}}^{\text{het}} \right) \) is computed as: \( \text{Ci} = \frac{F_{\text{phot}} + S_{C} + R_{C}}{f_{\text{phot}} + f_{\text{phot}}^{\text{het}} + S_{i}} \) with \( i = N, P \). |

Model sensitivity to the parameter values was studied with the method of Haefner (1996) as described above for the model of constitutive mixotrophy (Equation 28).

RESULTS
Constitutive Mixotrophy
In order to explore the qualitative behavior of the CM under different environmental conditions, we ran the model of constitutive mixotrophy under steady-state conditions with various values of DIN (ranging from 0 to 20 mmol N m\(^{-3}\)) and prey (bacteria) biomass (ranging from 0 to 20 mmolC m\(^{-3}\)). These simulations were run under two contrasting photon flux densities (30 and 200 \( \mu \text{mol quanta m}^{-2} \text{s}^{-1} \)). Conditions are such that P never limits the growth and the prey is considered as “inert” (i.e., bacteria growth and metabolism is not described). The two possible regulation mechanisms (photosynthesis “\( \text{Reg}^{\text{phot}} \)” vs. grazing “\( \text{Reg}^{\text{graz}} \)” limiting the mixotrophic growth to its maximum were tested. The performance of the two CMs (“\( \text{Reg}^{\text{graz}} \)” and “\( \text{Reg}^{\text{phot}} \)” ) is compared with that of the strict phototrophic nanoflagellate when growing under the same growth conditions.

Under high photon flux density (Figure 2), the growth rate of the strict phototroph decreases to zero as DIN decreases to zero because phototrophic growth is prevented by the lack of DIN (Figure 2A). By contrast, when prey concentration is higher than 4 mmolC m\(^{-3}\), the growth rate of the two CM configurations (“\( \text{Reg}^{\text{graz}} \)” and “\( \text{Reg}^{\text{phot}} \)” ) does not vary as a function of DIN because the lack of DIN is offset by bacteria ingestion (Figures 2F,K). De facto, the model captures correctly the competitive advantage of the mixotrophs at low inorganic nutrient concentration and high prey concentration. The grazing rate of the two CMs varies in function of the external conditions: it increases when prey concentration increases and when DIN decreases (Figures 2H,M). Thanks to their grazing activity, at low DIN, the photosynthesis rate of the two CMs decreases less than the photosynthesis rate of the strict phototroph (Figures 2B,G,L). When mixotrophs ingest prey under high photon flux density, the entire part of the regenerated DIN is retained inside the cell to sustain phototrophic activity (Figures 2J,O): there is no NH\(_{3}\) excretion (Figures 2I,N). Under this photon flux density, there is no noticeable difference between the two mixotrophic configurations (CM “\( \text{Reg}^{\text{graz}} \)” vs. CM “\( \text{Reg}^{\text{phot}} \)” ).

Under low photon flux density (Figure 3), the strict phototroph cannot grow in any nutrient conditions because the carbon monomers (\( S_{C} \)) limitation (which is controlled by the light limitation) is too high to allow steady-state growth rates above zero (Figure 3A). In function of its configuration, the CM behaves differently: the CM “\( \text{Reg}^{\text{graz}} \)” is able to grow under specific DIN and prey conditions (Figure 3F) while the CM “\( \text{Reg}^{\text{phot}} \)” has a zero steady-state growth in all conditions (Figure 3K). This different behavior is explained by the initial hypothesis used to construct the model, i.e., a minimum level of phototrophic growth (involving inorganic nutrient assimilation) is needed to allow prey ingestion. When the “\( \text{Reg}^{\text{phot}} \)” configuration is used, the CM down-regulates its photosynthesis when the maximum mixotrophic growth is attained, instead of its grazing. Therefore, phototrophy is more constrained in this “\( \text{Reg}^{\text{phot}} \)” configuration and the minimum level of phototrophic growth required to allow prey ingestion is not attained. The hypothesis of a minimum level of phototrophy also explains that the CM “\( \text{Reg}^{\text{graz}} \)” cannot take advantage of its ability to ingest prey at low DIN, as phototrophy involves inorganic nutrient assimilation (Figure 3F). By comparison with the high photon flux density conditions, the regenerated DIN related to phagotrophic activity is not entirely retained inside the cell; a significant part is
FIGURE 2 | Comparison between steady-state behaviors of two different types of constitutive mixotrophs (CMs; middle and right column) and a strict phototroph (left column, A–E), under constant high photon flux density (200 μmol quanta m$^{-2}$ s$^{-1}$) and under different DIN and prey biomass. The two CMs differ by their propensity to use either phototrophy or heterotrophy to grow: CM *Reg$^{Graz}$* (middle column, F–J) tends to down-regulate its grazing rate while CM *Reg$^{Phot}$* (right column, K–O) tends to down-regulate its photosynthesis. With the physiological parameters used, the model describes a phototrophic nanoflagellate and mixotrophic nanoflagellates grazing on bacteria. Growth, photosynthesis and grazing rates are expressed in C C$^{-1}$ day$^{-1}$ while NH$_4$ excretion and N retention rates are expressed in N N$^{-1}$ day$^{-1}$.

excreted outside the cell (Figure 3I) because the demand is lower due to the low photosynthesis rate (Figure 3G). Without the hypothesis of a minimum level of phototrophic growth, the behavior of the two CMs is similar a steady-state growth rate is reached around 0.3 day$^{-1}$ when prey is available (not shown).
The sensitivity analysis conducted on the physiological parameters involved in the model of constitutive mixotrophy shows that parameter rankings constructed on the basis of the sensitivity index (SI; Equation 28) differ according to the model configuration selected, i.e., “RegGraz” or “RegPhot” (Figures 4A,B). Because grazing is the process that is regulated when the configuration “RegGraz” is selected, the parameters related to phago-heterotrophy have proportionally a higher impact on the model response (as shown on the left part of the ranking; Figure 4A). By contrast, if the configuration “RegPhot” is selected, the parameters related to phagoheterotrophy are located on the right part of the ranking, indicating that they have a lower impact than parameters related to phototrophic processes (Figure 4B).

Despite the difference of parameter appearance in the ranking, the two model configurations are generally low-sensitive to most of parameters in the range of the tested values: 17 out of 22 parameters have a SI < 0.4 for “RegGraz” and 15 out of 22 parameters have a SI < 0.4 for “RegPhot”, meaning that a 25% change of their reference value induces <10% change of the mixotroph biomass reached at steady-state.

Non-constitutive Mixotrophy
To study the behavior of the NCM, we ran the model under steady-state conditions for various prey (a phototrophic nanoflagellate) biomass (ranging from 0 to 20 mmolC m\(^{-3}\)) and for various DIN-values (ranging from 0 to 20 mmolN m\(^{-3}\)). These simulations were run under two contrasting photon flux densities (30 and 200 μmol quanta m\(^{-2}\) s\(^{-1}\)). As for the model of constitutive mixotrophy, we considered the prey as inert, meaning that the metabolism of the phototrophic prey is not active. The behavior of the NCM is compared with that of a strict heterotrophic microzooplankton under the same growth conditions.

Under high photon flux density (Figure 5), the NCM can grow at lower prey concentrations than the strict heterotroph (Figures 5A,F) because the NCM takes advantage of the photosynthetic capacity of the prey retained in the food vacuole for C supply (Figures 5B,G) and of inorganic nutrient retention from heterotrophic regeneration (Figures 5E,J). DIN concentration has no impact on the NCM growth rate because we assumed that the NCM is not able to take up inorganic nutrients; the inorganic nutrients required for phototrophic growth are only provided by nutrient recycling from prey digestion. As photosynthesis in the NCM relies exclusively on the acquired prey chloroplasts, when prey concentration is lower than 5 mmolC m\(^{-3}\), the NCM cannot grow so well because the grazing is limited by the low prey availability (Figures 5F–H). With the model configuration used, the grazing rate of the NCM is lower than the grazing rate of the strict heterotroph at prey concentration higher than 8 mmolC m\(^{-3}\) (Figures 5C,H).

The difference is explained by the phototrophic growth that contributes to the mixotrophic growth by providing photosynthetically assimilated C (Figure 5G) and inorganic nutrient from nutrient retention (Figure 5J). This illustrates the benefit of the close interactions that occur between phototrophy and heterotrophy inside the mixotrophic cell. At high prey concentrations, NH\(_4\) excretion is substantially lower for the NCM than for the strict heterotroph (Figures 5D,I). Given the low N retention observed for the NCM (Figure 5J), the difference cannot be only explained by the ability of the NCM to retain a part of the regenerated N for its phototrophic growth. The difference is actually due to a different N regeneration for the strict heterotroph and the NCM (N regeneration being defined here as the sum of NH\(_4\) excretion and N retention).

As the grazing rate of the NCM is lower than that of the strict heterotroph at high prey concentration, N regeneration is lower for the NCM. In addition, when the NCM photosynthesizes thanks to the chloroplasts acquired from its prey, it can use a part of the N inorganic pool (Sn) of the prey to grow phototrophically. This process tends to increase the C/N stoichiometry of the prey so that N regeneration issued from prey digestion is lower (Equation 45). The latter effect is particularly important at high photon flux density because the nutrient demand for phototrophic growth is higher.

Under low photon flux density (Figure 6), the growth rates of the NCM are the same as those obtained under high photon flux density (Figures 6A,F) because the decrease of photosynthesis rate in the NCM (Figure 6G) associated with the lower photon flux density is offset by an increase of grazing rate (Figure 6H). As we assumed that the NCM was a facultative mixotroph (i.e., does not require some level of photosynthesis to grow), it is not impacted by the low light. Due to its higher grazing rate, NH\(_4\) excretion and N retention by the NCM are slightly higher than under high photon flux density (Figures 6L,J).

The sensitivity analysis conducted on the physiological parameters involved in the model of non-constitutive mixotrophy (Table 4) shows that the parameters that have the highest impact on the model response (here, in terms of steady-state biomass) are those related to heterotrophic activity (Figure 4C). As the grazing directly controls the heterotrophic growth as well as the phototrophic growth, it is not surprising to observe that the half-saturation constant for grazing (k\(_{g}\)) and the maximum grazing rate (g\(_{max}\)) are the two first parameters in the ranking. Among parameters related to heterotrophic activity, the half-saturation constant for prey digestion (k\(_{dig}\)) is the most problematic to measure experimentally. Further, it is an important parameter because it directly controls the extent to which the organism is able to use photosynthesis, as a low k\(_{dig}\) means that the organism maintains the kleptochloroplasts active during a long period while a high k\(_{dig}\) means that the organism rapidly digests the kleptochloroplasts and thus, approaches the strict heterotrophic organism. However, the model is rather robust against k\(_{dig}\) changes because SI = 0.4, meaning that a 25% change of its reference value only induces a 10% change of the mixotroph biomass reached at steady-state.

Parameters related to phototrophic activity are generally of lower importance as most of them have a SI < 0.003, meaning that a 25% change of the parameter value induces a 0.075% change of the mixotroph biomass reached at steady-state. The parameter of light adaptation (α) and the maximum photosynthesis rate (k\(_{max}\)) i.e., the two parameters directly involved in the photosynthesis (Equation 20), are the only parameters related to phototrophy that have a visible impact on the model response. However, their impact is minor as the SIs are, respectively, 0.33 and 0.19.
DISCUSSION

Many field, experimental and modeling studies have highlighted the potential, if not real significance, of planktonic mixotrophy in aquatic systems (e.g., Bird and Kalff, 1986; Estep et al., 1986; Bockstahler and Coats, 1993; Hall et al., 1993; Nygaard and Tobiesen, 1993; Arenovski et al., 1995; Havskum and Riemann, 1996; Stoecker et al., 1997; Stickney et al., 2000; Carvalho and Granéli, 2010; Hartmann et al., 2012; Mitra et al., 2014). In addition, experimental studies have shown that there is a large diversity among mixotrophs, in terms of (i) planktonic groups in which mixotrophic species have been observed, (ii) prey...
ingested, (iii) obligation to feed on a specific prey (specialist or generalist mixotroph), (iv) obligation to use the two trophic modes (obligate or facultative mixotrophy), (v) proportion of phototrophy and heterotrophy involved in growth, and (vi) factors inducing the use of the additional trophic mode (carbon limitation or nutrient limitation). However, despite the large diversity, a major distinction can be made among mixotrophs according to the origin of their chloroplasts: either constitutive or acquired from ingested phototrophic prey (Mitra et al., 2016).

All mixotrophic protists can, therefore, be divided between CMs or NCMs. The aim of this work is thus to offer a model able to represent the two contrasting forms of mixotrophy and that can be easily implemented in biogeochemical/ecological models. Specifically, the offering here facilitates the implementation of these groups into the AQUAPHY model that deploys the Shuter (1979) concept. The aim was not to develop descriptions for specific organisms, but rather to provide flexible constructs in which key parameters could be safely varied (as demonstrated by sensitivity analyses, Figure 4) to enable applications as appropriate for different ecosystem scenarios.

We adopted a model structure in which the organism biomass is divided into different components (on the basis of their function in the cell, i.e., structure, synthesis, or reserve) that interact and explain the main features related to microalgae metabolism (as in Shuter, 1979). This kind of model structure offers the advantage to be particularly appropriate to represent the main interactions that occur between phototrophic and heterotrophic activities in a mixotroph. Mitra and Flynn (2010) have indeed shown that descriptions integrating physiological processes, with some degree of feedback to modulate the processes of phototrophy and heterotrophy, are needed to properly represent the qualitative behavior of mixotrophs.

Among existing mixotrophic models, only those of Stickney et al. (2000) and the “perfect beast” of Flynn and Mitra (2009) take the interactions between the two trophic modes into account; the others rely on additive descriptions of phototrophy and heterotrophy (Thingstad et al., 1996; Baretta-Bekker et al., 1998; Jost et al., 2004; Hammer and Pitchford, 2005; Crane and Grover, 2010; Ward et al., 2011; Våge et al., 2013).

In this work we presented descriptions of the two mixotroph forms: the CMs and NCMs. The two versions, describing
fundamentally different mixotroph functional types, differ in the models by the processes linking the different components (i.e., the state variables) though they do share a common structure, with similar components in most respects. In the tested conditions, the model of constitutive mixotrophy was able to reproduce the expected observed behaviors of CMs: under light limitation or nutrient limitation (here, DIN), the CM has a growth rate substantially higher than its equivalent strictly autotroph (Figures 2, 3). This is consistent with field and experimental observations showing that mixotrophs are generally dominants under these conditions (Nygaard and Tobiesen, 1993; McKie-Krisberg et al., 2015). Regarding the model of non-constitutive mixotrophy, it properly captures the competitive advantage of NCMs over the strict heterotrophs.
under light conditions, when prey are limiting (Figure 5), in accordance with the observations (Skovgaard, 1998).

In this work, we configured the CM as a nanoflagellate able to feed on bacteria while the NCM was illustrated by a microzooplankton (ciliate) that feeds on autotrophic nanoflagellates. However, parameters values can easily be adapted to describe other planktonic group or other prey. To consider another prey, the only parameter that has to be changed in the CM model is the prey C:N:P ratio (if assumed fixed). In the NCM model, all parameters values related to photoautotrophy have to be changed in order to be the same as those of the prey. The type of prey will have an impact on the amount of inorganic nutrients that is retained inside the cell or excreted outside the cell (for the CM and the NCM), and also impacts on the phototrophic capacity for the NCM. We did not consider the fact that some specialists NCMs (i.e., acquiring photosynthetic capabilities from a specific prey) are capable of replicating their acquired photosystems (Hansen et al., 2013). It has been reported, for instance, that Mesodinium rubrum had the potential to synthesize and replicate new chloroplasts at least 3–5 times when starved of prey (Hansen et al., 2013). To take this specificity into account, the constant degradation rate of ingested prey ($k_{dig}$) can be lowered to maintain the kleptochloroplasts for a longer period.

By imposing a minimum level of photoautotrophy to mixotrophic growth, we assumed that the CM is an obligate photoautotroph, but a facultative mixotroph. The fact that our CM cannot maintain a positive growth rate when both light and DIN are limiting, despite the high prey concentration, stems directly from the assigned 10% minimum level of phototrophic growth in the mixotrophic growth. However, to describe an obligate mixotroph, the only thing to do is imposing a minimum level of heterotrophy in the mixotrophic growth, in addition to the minimum level of photoautotrophy. A similar approach was enacted by Flynn and Mitra (2009). In the NCM configuration, we also assumed that mixotrophy is facultative but we assumed that photoautotrophy is facultative too (in contrast to the CM configuration). The NCM configuration could, however, be modified to match that of the CM in respect of the minimum level photoautotrophic growth required in the mixotrophic growth.

To take account that the proportion of photoautotrophy and heterotrophy involved in mixotrophic growth is not the same for all mixotrophs, we distinguished two model configurations for the model of constitutive mixotrophy: one that describes CMs using photoautotrophy as a priority (e.g., C. ovata; Tranvik et al., 1989) and another that describes CMs using heterotrophy as a priority (e.g., P. malhamensis; Sanders, 1991). The appropriate configuration can therefore be selected as appropriate for the application. When comparing the two CM model configurations (i.e., "RegGraz" vs. "RegPhot") results show that they behave similarly under high photon flux density. The different behaviors under low photon flux density are more explained by a combination of the model configuration and the hypothesis of a minimum level of phototrophy, than by the configuration controlling the dominance of heterotrophy vs. phototrophy itself.

When constructing the model of constitutive mixotrophy, we assumed that both carbon and nutrient limitations could induce phagotrophy. If only one of these factors was to be considered, the formulation of $\mu_{\text{mix}}$ (Table 3, Equations 18–19) can be changed to take only account of the factor inducing phagotrophy. For instance, if only nutrient limitation is assumed to induce phagotrophy, the formulation of $\mu_{\text{mix}}$ would be restricted to Equation (19).

In conclusion, the model structure presented in this work is able to take account of the main features and interactions between photoautotrophy and phagoheterotrophy in mixotrophs, and has enough flexibility to represent the observed diversity among mixotrophs. The mechanistic model of mixotrophy developed by Flynn and Mitra (2009)—the "perfect beast"—has these two characteristics as well. The main difference between the two models lies in the model structure; the "perfect beast" of Flynn and Mitra (2009) is based on cell quotas and results from the merging between C:N:P zooplankton and photoacclimative models, while the models presented here are based on the Shuter approach of a division of the biomass between components chosen on the basis of their function in the cell. The type of biogeochemical/ecological model in which mixotrophy will be implemented could guide the choice between either of these model structures; the models presented in this work are particularly adapted to be implemented in those biogeochemical models that use the "Shuter" mechanistic approach to describe the phytoplankton compartment but that do not take mixotrophy into account yet.

**AUTHOR CONTRIBUTIONS**

All five authors contributed to the model conception and test design. CG conducted the model analyses and prepared the manuscript while AM, CL, KF, and NG revised it.

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