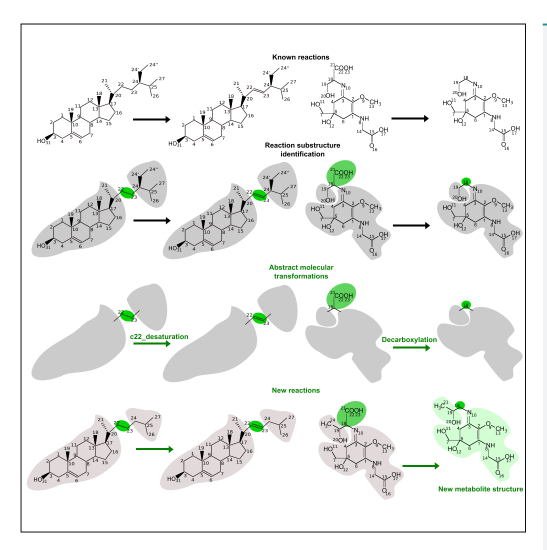
Article

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HIGHLIGHTS

Combination of metabolite profiling and genome-scale metabolic networks analysis

New method to infer biochemical reactions and metabolites in promiscuous pathways

Red algal model for sterol and mycosporine-like amino acid biosynthesis pathways

Metabolic drift if pathway variation despite conserved initial and final metabolites

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Article

Inferring Biochemical Reactions and Metabolite Structures to Understand Metabolic Pathway Drift

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SUMMARY

Inferring genome-scale metabolic networks in emerging model organisms is challenged by incomplete biochemical knowledge and partial conservation of biochemical pathways during evolution. Therefore, specific bioinformatic tools are necessary to infer biochemical reactions and metabolic structures that can be checked experimentally. Using an integrative approach combining genomic and metabolomic data in the red algal model *Chondrus crispus*, we show that, even metabolic pathways considered as conserved, like sterols or mycosporine-like amino acid synthesis pathways, undergo substantial turnover. This phenomenon, here formally defined as "metabolic pathway drift," is consistent with findings from other areas of evolutionary biology, indicating that a given phenotype can be conserved even if the underlying molecular mechanisms are changing. We present a proof of concept with a methodological approach to formalize the logical reasoning necessary to infer reactions and molecular structures, abstracting molecular transformations based on previous biochemical knowledge.

INTRODUCTION

Life is driven by a high diversity of metabolic processes, and each species or even strain may be characterized by its own metabolic particularities (e.g., Rhee et al., 2011). During evolutionary time and speciation processes, there are many ways that variations can be generated within metabolic pathways. Evolutionary models have been developed and experimentally tested to explain the arising of new pathways, but these efforts were focused on the activities of individual enzymes (Noda-Garcia et al., 2018). Changes in the metabolism may, however, also occur at a higher level of organization, notably an enzymatic replacement by non-orthologous genes encoding enzymes with identical biochemical function (Koonin et al., 1996; Figure 1, left side), or a change in enzyme order, which leads to different main biosynthetic intermediates (Figure 1, right side). This kind of variability, which we refer to as "metabolic drift" in this article, is possible due to substrate promiscuity of the enzymes and may be an important driver of evolution (Peracchi, 2018).

The concept of "drift" has previously been used in the field of animal comparative developmental biology, where it was already used to explain how morphologically similar structures can be maintained even if there are substantial variations in the molecular mechanisms underlying their formation (True and Haag, 2001). Here also, the use of "drift" is distinct from genetic drift, but, nevertheless, appropriate because chance and not selection explains how changes occur. The concept was more recently extended to plants, where such cases have been observed in leaf development (Townsley and Sinha, 2012). It was later exported to the fields of protein evolution (Hart et al., 2014) and gene expression evolution (Petit et al., 2016). Here, we hypothesize that the evolutionary concept of drift could also adequately explain the strict functional conservation of many metabolic pathways, despite variations in the underlying mechanisms (Figure 1).

To study metabolic drift by non-orthologous gene displacement (Figure 1, left part), classical comparative genomic approaches can generate hypotheses that can be experimentally checked using targeted metabolic profiling combined with inactivation of enzyme-encoding genes by reverse genetics (Markov et al., 2016; Sonawane et al., 2016). However, in case of drift by change in enzyme order (Figure 1, right part), it is necessary to combine both genomic and metabolomic data and to introduce a knowledge-based approach that implements reasoning in the manner of a biochemist. Such inter-disciplinary strategies

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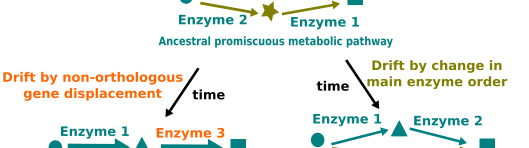
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Enzyme 2

Contemporary metabolic pathway 1

Contemporary metabolic pathway 2

Enzyme 2

Figure 1. The Hypothesis of Metabolic Pathway Drift Based on Two Possible Elementary Mechanisms

Enzyme 1

Starting from an ancestral promiscuous pathway (main pathway in teal; upper part, alternative pathway in olive green), changes can occur either by non-orthologous gene displacement (in orange, left side) or by change in main enzyme order, leading to a different intermediate metabolite (in olive green, right side). Substrate promiscuity enables the same molecular transformation to occur on different molecules, making the enzyme able to catalyze two different reactions. Promiscuity can be secondarily lost, as shown on the left side, leading to the impossibility to observe the star-shaped metabolite in contemporary metabolic pathway 1.

have already been used to design experiments for the analysis of auxotrophic mutants in yeast (King et al., 2004) or for synthetic biology, where *ab initio* pathway inference is done to find a biosynthetic route that is not necessarily present in nature (Koch et al., 2017). We hypothesize that similar tools can also be used to group biochemical reaction variants based on shared ancestry and to infer undescribed pathways that may be present in emerging model species.

To test this hypothesis, we have implemented a semi-automatic analogy reasoning approach, which we use to study two distinct metabolic pathways, the sterol pathway and the mycosporine-like amino acid (MAA) pathway, in the red alga *Chondrus crispus*. Red algae are sufficiently distant from terrestrial plants to anticipate substantial metabolic drift compared with the known pathways in organisms such as *Arabidopsis thaliana*, yet *C. crispus* has been subject to biological studies for more than two centuries (Collén et al., 2014). Notably, its genome was sequenced and annotation was performed with a focus on metabolic features (Collén et al., 2013), and there is extensive literature available describing its metabolome (Young and Smith, 1958; Saito and Idler, 1966; Laycock and Craigie, 1977; Matsuhiro and Urzua, 1992; Karsten et al., 1998; Tasende, 2000; Kräbs et al., 2004; Gaquerel et al., 2007; Banskota et al., 2014; Pina et al., 2014; Melo et al., 2015, Alcaide et al., 1968; Goldberg et al., 1982; Kremer and Kirst, 1982; Pettit et al., 1989; van Ginneken et al., 2011; Santos et al., 2015; Robertson et al., 2015; Athukorala et al., 2016; Belghit et al., 2017; Guihéneuf et al., 2018; Lalegerie et al., 2019).

The sterol metabolism was chosen as one focus point, because there is extensive knowledge about this pathway at the comparative genomics level (Desmond and Gribaldo, 2009). Furthermore, analytical standards are available for different sterol molecules, enabling their identification by mass spectrometry (MS) (Sumner et al., 2007). MAA synthesis, on the other hand, involves combination of different building blocks, and analytical standards are lacking for this class of compounds, limiting metabolite identification to putative annotated compounds based on spectral similarity with spectral libraries (Sumner et al., 2007). Therefore the reconstruction of MAA pathway raised the problem of integrating unannotated compounds that were identified uniquely based on their m/z ratio.

By implementing a semi-automatic analogy reasoning approach that integrates both metabolite and genomic data, we here propose an exhaustive model for both metabolic pathways in *C. crispus* and provide strong indications for the importance of metabolic drift in shaping these pathways. We furthermore consolidated at least part of these hypotheses with targeted metabolite profiling of metabolic intermediates predicted in the models.



Species	Reactions	Enzymes	Metabolites	Pathways	Reference
C. crispus	2,024	2,006	2,196	1,108	This study, before curation
E. siliculosus	1,977	2,281	2,132	1,101	Aite et al. (2018)
Ectocarpus subulatus	2,074	2,445	2,173	1,083	Dittami et al., 2020
A. thaliana	1,567	1,419	1,748	796	de Oliveira Dal'Molin et al. (2010)
Chlamydomonas reinhardtii	3,083	1,355	1,133	522	Imam et al., 2015

Table 1. Comparison of Global Features of Genome-Scale Metabolic Networks from Macroalgae and Other Chlorophyllian Eukaryotes

RESULTS

Genome-Scale Metabolic Network Enriches the Inferred Basic Integrated Metabolism for C. crispus

Genome-scale metabolic networks (GSMNs) are graph-based representations of enzymatic reactions assumed to occur in a given organism. In this framework, an enzymatic reaction denotes a chemical reaction that transforms one or several metabolic substrates into one or several metabolic products under the control of an enzyme that can be associated with a gene in the considered species.

We used the tool AuReMe (Automatic Reconstruction of Metabolic models) dedicated to "à la carte" reconstruction of GSMNs (Aite et al., 2018) to reconstruct a GSMN of C. crispus. This GSMN comprised in total 2,024 enzymatic or transport reactions (Table 1). Among them, 595 reactions were recovered from annotation-based searches from the C. crispus genome annotation (Collén et al., 2013) with the Pathway Tools suite (Karp et al., 2002) and the MetaCyc database (Caspi et al., 2016). In addition, 1,429 reactions were included in the network according to orthology evidences with protein sequences encoding enzyme reactions in the Arabidopsis thaliana GSMN (de Oliveira Dal'Molin et al., 2010), the Galdieria sulphuraria GSMN (based on genome data from Schönknecht et al., 2013), or the Ectocarpus siliculosus GSMN (Cormier et al., 2017; Aite et al., 2018). A biomass reaction was established based on the previous E. siliculosus data, defining a list of 33 compounds to be produced to consider the network functional (Prigent et al., 2014). According to this biomass reaction, the network was manually gap-filled to unblock the production of L-alpha-alanine with an alanine dehydrogenase reaction whose associated gene had been incorrectly annotated in the C. crispus genome. The predicted maximal growth rate was then 2.43 $g.gDW^{-1}.h^{-1}$ (gram per gram dry weight per hour). As shown in Table 1, the *C. crispus* GSMN is comparable in size with the E. siliculosus GSMN. A total of 254 pathways are complete, including those involved in central metabolism of carbohydrates, fatty acids, and amino acids, as well as those related with photosynthesis. The greatest contributor to the inferred reaction set was the phylogenetically close red microalga G. sulphuraria, which provided 1,361 reactions.

Building a Catalog of Evidenced Metabolites in C. crispus Using Metabolomic Data

To better understand the specificities of *C. crispus*, we built a catalog of metabolic compounds attested to be produced by the alga. To that goal, we reviewed the literature to collect experimental evidence of presence of all reported metabolites. To have more species-specific data on sterol and MAA synthesis we also experimentally tested for the presence of these compounds and their precursors in *C. crispus* using MS analysis. In this way, we assembled a set of 142 metabolites that are reported in Tables S1 and S2. Those metabolites broadly cover various classes of amino acids, carbohydrates, and lipids. We divided this dataset into two main categories: 85 database metabolites, which are already indexed into Metacyc (Table S1), and 57 orphan metabolites, not yet indexed (Table S2). In addition, we have acquired additional experimental data on two pathways for which molecules are in both categories: the sterol and the MAA pathways.

Our MS data confirmed previous findings and also pointed out possible additional precursors and intermediary metabolites in sterol and MAA biosynthesis pathways. More precisely, the results of these analyses of sterols are found in Table 2. In addition to confirming the presence of eight previously identified sterols (brassicasterol, campesterol, cholesterol, 7-dehydrocholesterol, desmosterol, lathosterol, β -sitosterol, and stigmasterol), we identified in *C. crispus* an immediate precursor of sterols, i.e., squalene (Figure S1). However, we did not find evidence for cycloeucalenol, ergosterol, fucosterol, and zymosterol, which are



Analyzed Compounds	Molecular Formula	Found in This Study	Previous Evidence
Brassicasterol	C ₂₈ H ₄₆ O	Yes	Saito and Idler (1966) (GC-MS), Tasende (2000) (TLC, GC-MS)
Campesterol	C ₂₈ H ₄₈ O	Yes	Tasende (2000) (TLC, GC-MS)
Cholesterol	C ₂₇ H ₄₆ O	Yes	Saito and Idler (1966) (TLC, GC-MS), Tasende (2000) (TLC, GC-MS)
Cycloartanol	C ₃₀ H ₅₂ O	No	Not reported
Cycloartenol	C ₃₀ H ₅₀ O	No	Saito and Idler (1966) (TLC), Alcaide et al. (1968) (TLC)
Cycloeucalenol	C ₃₀ H ₅₀ O	No	Not reported
7-Dehydrocholesterol	C ₂₇ H ₄₄ O	Yes	Tasende (2000) (TLC, GC-MS)
Desmosterol	C ₂₇ H ₄₄ O	Yes	Saito and Idler (1966) (TLC, GC-MS), Goldberg et al. (1982) (GC-MS)
Ergosterol	C ₂₈ H ₄₄ O	No	Not reported
Fucosterol	C ₂₉ H ₄₈ O	No	Not reported
Lanosterol	C ₃₀ H ₅₀ O	No	Saito and Idler (1966) (TLC)
Lathosterol	C ₂₇ H ₄₆ O	Yes	Goldberg et al. (1982) (GC-MS)
β-Sitosterol	C ₂₉ H ₅₀ O	Yes	Saito and Idler (1966) (GC-MS), Tasende (2000) (TLC, GC-MS)
Squalene	C ₃₀ H ₅₀	Yes	Not reported
Stigmasterol	C ₂₉ H ₄₈ O	Yes	Saito and Idler (1966) (GC-MS), Tasende (2000) (TLC, GC-MS)
Zymosterol	C ₂₇ H ₄₄ O	No	Not reported

Table 2. List of Sterols Profiled in This Study, and Comparisons with Previous Studies

TLC, thin-layer chromatography

For each compound, analytical parameters (retention time and m/z ratio) are given in Table S1.

intermediates present in other eukaryotes (Desmond and Gribaldo, 2009; Sonawane et al., 2016). We also did not find cycloartenol, contrary to a previous report in *C. crispus* using thin-layer chromatography (Alcaide et al., 1968). This negative finding is strengthened by the fact that we are able to identify the cycloartenol standard when added in algal extract (Figure S2).

Similar experiments were carried out for the MAA pathway, and corresponding data are summed up in Table 3. Using liquid chromatography-MS (LC-MS) profiling we confirmed the presence of six MAAs in *C. crispus* (see references in Table 3): asterina-330, palythene, palythine, palythinol, porphyra-334, and shinorine. In addition, we identified mycosporine-glycine in *C. crispus*. The relative abundance of MAAs varied between sampling dates (Figure 2), which is consistent with a report of MAA variation in the Galway Bay, Ireland (Guihéneuf et al., 2018). We noticed that two unknown peaks potentially corresponding to MAAs were detected in our samples. These peaks exhibit m/z ratios consistent with peaks reported in a study on 40 red algae by Lalegerie et al. (2019). Specifically, we found a peak at m/z = 270.3 that does not match with any already identified candidate MAA, which we named MAA1, and a second one at m/z = 302.3, which we named MAA2.

Secondary Metabolism Evidenced by Metabolomic Data Is Only Partially Accurately Described in the *C. crispus* GSMN

To investigate the accuracy of GSMNs to describe the synthesis pathway of secondary metabolites of importance, we studied the capability of the *C. crispus* GSMN to describe synthesis pathways of metabolic

Analyzed Compounds	Molecular Formula	Found in This Study	Previous Evidence
Asterina-330	C ₁₂ H ₂₀ N ₂ O ₆	Yes	Athukorala et al. (2016) (LC-MS-MS); Guihéneuf et al. (2018) (LC-MS)
MAA1	compatible with m/z 270.272	Yes	Lalegerie et al. (2019) (HPLC)
MAA2	compatible with $m/z = 302.3117$	Yes	Lalegerie et al. (2019) (HPLC)
Mycosporine-glycine	C ₁₀ H ₁₅ NO ₆	Yes	not reported
Palythine	$C_{10}H_{16}N_2O_5$	Yes	Karsten et al. (1998) (UV + LC-MS); Athukorala et al. (2016) (LC-MS-MS); Guihéneuf et al. (2018) (LC-MS)
Usujirene/Palythene	$C_{13}H_{20}N_2O_5$	Yes	Karsten et al. (1998) (UV + LC-MS)
Palythinol	$C_{13}H_{22}N_2O_6$ (m/z = 302.3117)	No	Karsten et al. (1998) (UV + LC-MS), Athukorala et al. (2016) (LC-MS-MS)
Porphyra-334	C ₁₄ H ₂₂ N ₂ O ₈	Yes	Athukorala et al. (2016) (LC-MS-MS)
Shinorine	C ₁₃ H ₂₀ N ₂ O ₈	Yes	Karsten et al. (1998) (UV + LC-MS), Athukorala et al. (2016) (LC-MS-MS)

Table 3. List of Mycosporine-like Amino Acids Identified in This Study and Comparisons with Previous Studies HPLC, high-performance liquid chromatography.

compounds whose presence have been evidenced in the alga. Among those 142 metabolites, only 85 (60%) were indexed in the MetaCyc database version 20.5 (Table S1). The GSMN could provide synthesis pathways for only 59 metabolites already indexed in the Metacyc database. Those metabolites were amino acids (19), fatty acids (12), aldehydes (6), aliphatic alcohols (3), carotenoids (3), ketones (2), carboxylic acids (2), a halocarbon, a galactolipid, an oxylipin, a tetrapyrrole, an MAA, a nucleotide sugar, a methylketone, and a polyol. The synthesis of the 29 remaining metabolites indexed in the MetaCyc database could not be explained by any addition of enzymatic reactions in the database (failure of the exhaustive gap-filling procedure). The metabolites belonged to the following classes: fatty acids (8), sterols (8), alkanes (2), carotenoids (2), carragheens (2), aliphatic alcohols (2), an aldehyde, and a halocarbon. For the 57 metabolites that had not yet been indexed in MetaCyc (Table S2), it was impossible to generate hypotheses on their synthesis pathways. There were galactolipids (12), oxylipins (9), fatty acids (8), MAAs (6), alcohols (6), unconventional amino acids (3), tetrapyrroles (3), alkanes (2), carboxylic acids (2), ketones (2), a heteroside, a sterol, a phospholipid, and an aldehyde.

Building a Knowledge Base of Enzymatic Reactions and Molecules in Sterol and MAA Synthesis

To enable the incorporation of the orphan metabolites that were not yet in MetaCyc and derive possible synthesis pathways for sterols and MAAs, we built two knowledge bases describing the existing knowledge about known enzymatic reactions and molecules involving sterols or MAAs (available on https://github.com/pathmodel/pathmodel). They are encoded in the Pathmodel datafiles sterol_pwy.lp and MAA_pwy.lp, both accessible in the pathmodel/data folder of the Github repository. In these knowledge bases, molecules are described by atoms (identified by a number and atom types) and bonds (identified by atom numbers and bond type) as highlighted in green on Figure 3. Atom numbers were assigned manually to ensure consistency between molecules from the same family and followed IUPAC conventions when existing (Moss, 1989). Molecules can be automatically associated with a theoretical m/z ratio, calculated using their chemical formula, as described in the program MZComputation.lp.

Enzymatic reactions, denoted by reaction, model the link between two molecules (a reactant and a product, e.g., reaction(rxn_4243, "sitosterol", "stigmasterol")). They are associated with cross-references to MetaCyc pathway IDs when possible.

The MAA knowledge base (Figure 4) contained 13 enzymatic reactions involving 12 molecules. It first contained the shinorine biosynthesis pathway (MetaCyc: PWY-7751), corresponding to the best understood

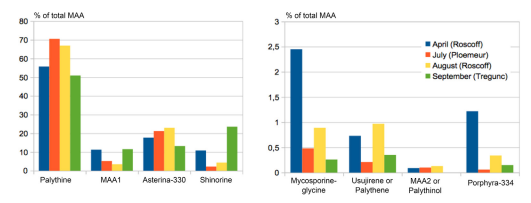


Figure 2. Composition and Seasonal Variation (MS Relative Quantification) of MAAs in C. crispus
The four most abundant compounds, including the unknown MAA1, are on the left panel. The four less abundant compounds, including MAA2, are on the right panel.

part of the pathway (Shick and Dunlap, 2002). We also encoded in the knowledge base an extended version of the amino acid C3-transfer reaction (MetaCyc: RXN-17371) to incorporate the hypothesis that the enzyme performing the C3-transfer of serine can also perform the C3-transfer of threonine, leading to porphyra-334 (Brawley et al., 2017). We also incorporated additional reactions hypothesized in the literature (Carreto and Carignan, 2011), for which either the substrate or the product was an MAA described in *C. crispus*. We finally added to the database the four orphan MAAs listed in Table S2, as well as two unknown molecules (MAA1 and MAA2) for which we had an experimental support for peaks corresponding to unassigned m/z ratios, as well as mycosporine-glycine, which was here identified in *C. crispus*.

The sterol knowledge base (Figure 5) contained 15 enzymatic reactions involving 24 molecules, including the eight unproducible sterols and the orphan molecule 22-dehydrocholesterol, which was not linked to any reaction. We encoded the «early side-chain reductase» (early SSR) pathway based on the model previously published for tomato (Sonawane et al., 2016), which was added in MetaCyc upon our request (MetaCyc: PWY18C3-1). It also included portions of the canonical plant sterol biosynthesis pathway (MetaCyc: PWY-2541; Benveniste, 2004), as well as portions of the animal sterol synthesis pathway (MetaCyc: PWY66-4, Mitsche et al., 2015).

Inferring Molecular Transformations from a Database of Enzymatic Reactions

Based on these examples, we hypothesize that these orphan molecules challenge the ability of the GSMN to produce them because of lacks in secondary metabolism synthesis pathways that enable the description of all possible molecular transformations between compounds catalyzed by a small family of enzymes. First, we define a molecular transformation to be a chemical reaction transforming a metabolite into another metabolite by operating on a specific part of the metabolite structure, called a *substructure*. We define a *substructure* to be a set of one of several atoms associated with one or several bonds in a given molecule. For instance, *substructure* ("simple bond 22-23") denotes a simple bond between atoms 22 and 23, such as in sitosterol on Figure 3A.

Importantly, an enzymatic reaction involving a single reactant and a single product can be defined as a molecular transformation by assuming that the enzyme enables the transformation of a metabolite site into another. Therefore, for each reaction involving a single reactant and a single product, we call *pair of transformed substructures* the substructure of the reactant (for instance, as shown in Figure 3A, simple bond between atoms 22 and 23 in sitosterol) and the substructure of the reaction product (for instance, as shown in Figure 3A, double bond 22-23 in stigmasterol).

Pairs of transformed substructures can be computed by removing all the atoms and bonds that are common to both reactant and product molecules, all atoms and bonds being previously ordered. For the sterol and MAA cases, we identified pairs of transformed substructures with a reasoning-based approach (see Methods) and then annotated them to describe a catalog of molecular transformations associated with known enzymatic reactions. For instance, the transformation replacing a simple bond between atoms 22

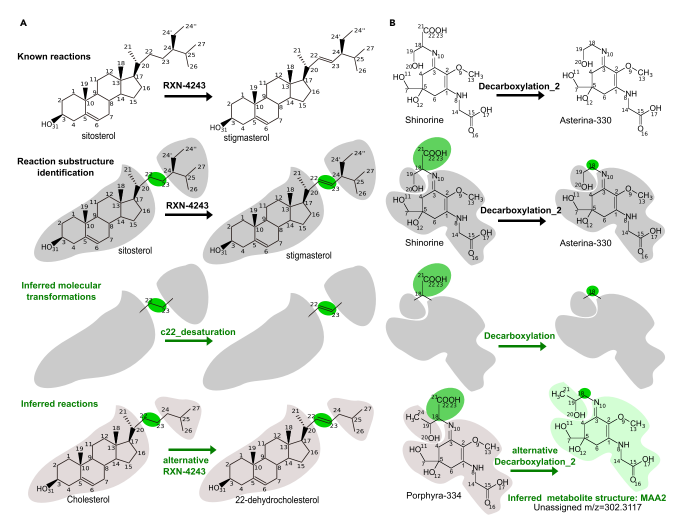


Figure 3. The Two Reasoning Methods Implemented in Pathmodel

(A) Inference of a reaction between two known molecules.

(B) Inference of reaction and metabolite structure corresponding to an unassigned m/z peak. Input data encoded in the knowledge base are in black; inferred reactions and metabolite structures are in green. In both cases, reaction substructure identification followed by inference of molecular transformations are common intermediate steps.

and 23 (as exhibited in sitosterol) into a double bond 22-23 (as exhibited in stigmasterol) is a c22 desaturation, described by the term *transformation*(c22_desaturation, simple bond 22-23, double-bond 22-23).

The complete list of molecular transformations for the sterol and MAA reactions is given in Table S5. The sterol enzymatic reaction database (15 reactions) yields 12 molecular transformations, whereas the MAA enzymatic reaction database (13 reactions) yields 9 molecular transformations. Reasoning on molecular transformations instead of enzymatic reactions, we reduce the complexity of the reaction set by abstracting a library of molecular transformations that can be applied to other known molecules belonging to the same chemical family.

Inferring Molecular Compounds and Enzymatic Reactions from a Database of Molecular Transformations

We used these databases of molecular transformations to infer putative new molecules and reactions.

To that goal, we assumed that for any molecular transformation from a substructure S1 to a substructure S2, and any pair of metabolites A and B, a *putative enzymatic reaction* can occur from the reactant A to the

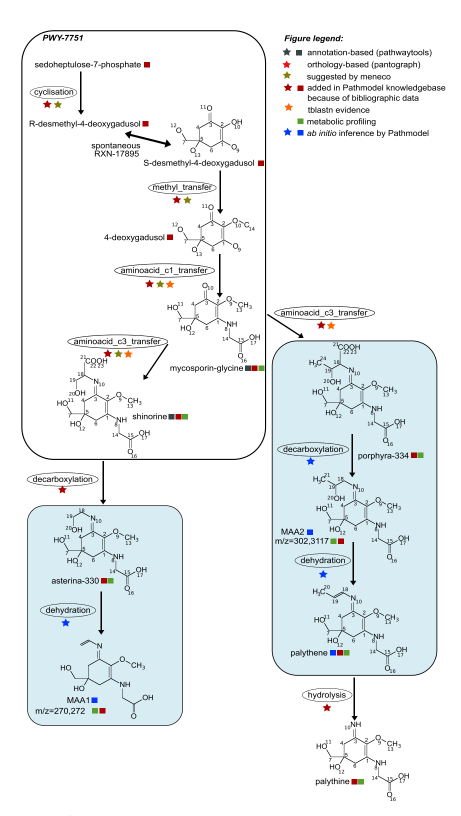


Figure 4. A Model for MAA Biosynthesis Pathway in C. crispus

PWY-7751: shinorine biosynthesis pathway. The figure legend details the various data sources integrated to infer the pathways. Stars indicate reactions, and squares indicate molecules. In light blue boxes: reactions and metabolites inferred through Pathmodel.

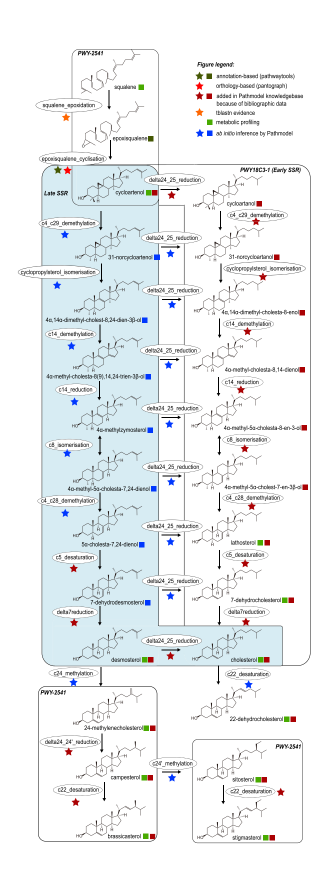




Figure 5. A Model for Sterol Biosynthesis in C. crispus

Early SSR: pathway involving an early sterol side-chain reduction (SSR), also present in solanacean plants (PWY18C3-1). In light blue box: late SSR pathway, involving a late sterol SSR, so far only described in *C. crispus*. Portions identical with the plant sterol biosynthesis pathway (PWY-2541) are also boxed. Ovals indicate molecular transformations.

product B as soon as (1) the molecule A contains the substructure S1, (2) the molecule B contains the substructure S2, and (3) the molecules A and B have identical structures (e.g., same numbered atoms and bonds) for all atoms and bonds to the exception of S1 and S2 substructures. An example is shown in Figure 3A: cholesterol and 22-dehydrocholesterol share a sterane skeleton. The only difference between these molecules is that cholesterol has a simple bond between atoms 22 and 23, whereas 22-dehydrocholesterol has a double bond between atoms 22 and 23. As the transformation from a simple bond to a double bond between atoms 22 and 23 has been evidenced in the reaction database (reaction from sitosterol to stigmasterol, Figure 3), we assume that a putative enzymatic reaction exists between cholesterol and 22-dehydrocholesterol and that it should be a c22_desaturation.

In addition, we assumed that, for a given molecular transformation, a given metabolite A and an experimental mass-to-charge ratio m/z corresponding to an unassigned peak, a *putative metabolite* B can be produced as soon as (1) the theoretical mass-to-charge ratio of B equals the experimental m/z and (2) A can be transformed into B according to a putative enzymatic reaction associated with the selected molecular transformation. An example is depicted in Figure 3B. We consider that the molecule MAA2 is a putative derived metabolite of porphyra-334 because its mass-to-charge m/z = 302.3117 corresponds to an observed peak in our measurements and that it can be obtained from porphyra-334 by removing a carboxyl group from carbon 18, a transformation named *decarboxylation*, which occurs in the enzymatic reaction from shinorine to asterina-330.

With these two assumptions, we introduce the concept of *putative synthesis pathway* computed from a source molecule A, a list of target molecules, a list of molecular transformations, a list of putative metabolites, a list of corresponding theoretical mass-to-charge m/z, and a list of forbidden molecules, for which we have analytical standards that did not match experimental peaks. From these inputs, a putative synthesis pathway is a family of putative reactions connecting the source to all targets metabolites such that (1) all reactions are consistent with the database of molecular transformations, (2) all reactants and products of the reactions are allowed either metabolites or putative metabolites matching with the allowed theoretical mass-to-charge ratio m/z, (3) no transformation associated with the pathway can produce any of the forbidden molecules, and (4) a minimal number of reactions is used to connect the source to the targets.

The Pathmodel method was developed as a prototype implementation of a semi-automatic analogy reasoning approach (Figure 3). Its aim is to infer putative synthesis pathways, to connect orphan metabolites, not yet indexed in metabolic databases, according to the known enzymatic reactions. The Pathmodel method takes as input a knowledge base including a set of known metabolites, a set of known enzymatic reactions, a set of observed mass-to-charge (m/z) ratios for unknown metabolites, an initial source metabolite, a family of targeted metabolites, and a list of forbidden molecules. Metabolites are described by their numbered atoms and bounds. Pathmodel then computes the list of molecular transformations associated with the database of enzymatic reactions and putative synthesis pathways of the targeted metabolites from the source metabolite. More precisely, for each pair of metabolites not linked by a reaction in the knowledge base, the method checks whether a molecular transformation can occur between them (deductive reasoning) and further derives from known reactions candidate metabolites corresponding to observed unassigned m/z ratios (analogical reasoning). These are the bases for iterating the selection of potential reactants or products and the inference by a reasoning component of new reaction occurrences or new metabolites. It was implemented using a logic programming approach known as *Answer Set Programming* (Lifschitz, 2008; Gebser et al., 2012, see Methods). An example of this reasoning is given in Figure 3.

Application of Pathmodel to the MAA Synthesis Pathway

We first used Pathmodel to compute the synthesis pathway for MAA1 and palythine from sedoheptulose-7-phosphate according to the MAA enzymatic reaction database described above. We also assumed that Z-palythenic acid was defined as a forbidden molecule according to the absence of a corresponding peak in *C. crispus* extracts.



The putative pathway for MAA synthesis obtained by our method is shown in Figure 4. Thanks to this approach, the knowledge base of MAA biosynthesis could be enriched with two putative molecule structures and three putative enzymatic reactions. In addition, the reactions that we compiled from the literature but are not yet indexed in MetaCyc will be submitted to the next release.

A first output of this approach is a pathway leading to a molecule structure compatible with our measured mass-to-charge ratio m/z = 270.3. It was therefore possible to infer the hypothetical structure shown in Figure 4 for this unassigned compound, named MAA1, in MS data. The predicted transformation leading from asterina-330 to MAA1 is a dehydration (in purple), a molecular transformation already observed between other MAAs such as porphyra-334 and Z-palythenic acid, a compound not identified in *C. crispus* (Figure 4).

Another output of this approach is to suggest a putative involvement of decarboxylation and dehydration transformations in MAA biosynthesis pathways. As no candidate enzymes were mentioned so far in the literature related to MAA biosynthesis pathways, we performed a semantic search on the GSMN from *C. crispus*, to identify other enzymes that may perform those molecular transformations on a serine coupled with other chemical building blocks. Serine decarboxylation indeed occurs in phospholipid metabolism and was inferred in the *C. crispus* GSMN based on orthology with *G. sulphuraria*. The candidate gene is CHC_T00008892001, annotated as phosphatidylserine decarboxylase. Interestingly, there is some evidence of catalytic promiscuity for this enzyme, enabling it to also decarboxylate a threonine residue (Heikinheimo and Somerharju, 2002). Therefore, we hypothesize that this enzyme in *C. crispus* may also perform serine/threonine decarboxylation on a serine/threonine linked to a mycosporine-glycin.

A final interesting feature of the MAA synthesis pathways is the inferred enzymatic reaction required to decarboxylate shinorine and porphyra-334 and further dehydrate their derivatives. These reactions were added to the pathway to take into account the fact that Z-palythenic acid was absent in *C. crispus* extracts. Such an absence does not support dehydration occurring before decarboxylation, as proposed in other species (Carreto and Carignan, 2011), and therefore does not allow finding any chains of reaction producing palythinol, a compound previously considered to be present in *C. crispus* based on UV + LC-MS or LC-tandem MS data (Karsten et al., 1998; Athukorala et al., 2016). We noticed, however, that there is no synthesis-based analytical standard available for verifying this prediction. To identify alternative routes for the production of porphyra-334, we ran Pathmodel by allowing the production of putative metabolites with m/z ratio of 302.3177 (the one of palythinol). Pathmodel then predicted an additional intermediate with m/z ratio of 302.3177, named MAA2 on Figure 4. From a genomic viewpoint, switching palythinol with MAA2 does not necessitate a candidate enzyme to perform hydrogenation and demethylation on an MAA-like substrate (Figure 4) and thus reduces the number of unassigned enzymatic activities to candidate genes. We thus included this alternative route in the MAA synthesis pathway that is consistent with the possible absence of Z-palythenic acid.

Application of Pathmodel to the Sterol Synthesis Pathway

The Pathmodel approach was also applied to the sterol biosynthetic pathway in *C. crispus* (Figure 5). To that goal, we used it as an enzymatic reaction database described below. The source metabolite was cycloartenol. The targeted metabolites were 22-dehydrocholesterol, brassicasterol, and stigmasterol. In addition, forbidden metabolites were ergosterol, fucosterol, and zymosterol, which are compounds with no detection result using targeted profiling with analytical standards.

We decided to use cycloartenol even if we did not find it by gas chromatography (GC)-MS for the two following reasons. First, a cycloartenol synthase from the red alga *Laurencia dendroidea* was cloned and expressed in yeast cells, where it is able to transform squalene into cycloartenol, even if the authors did not report cycloartenol identification in the whole alga by GC-MS (Calegario et al., 2016). Second, unambiguous cycloartenol derivatives are known in another florideophyte red alga, *Tricleocarpa fragilis* (Horgen et al., 2000). Therefore, we considered more parsimonious to hypothesize that cycloartenol is present and below the limit of experimental detection rather than considering that this step is performed via an unknown intermediate.

We propose two alternative synthesis pathways from cycloartenol to cholesterol, depending on when the side-chain reductase (SSR) enzyme is acting (Figure 5).



If *C. crispus* uses the « early SSR » pathway (Sonawane et al., 2016), the metabolic intermediates would be identical to tomato, but there would be an important difference concerning the enzymes. Indeed, the genes encoding SSR are duplicated in Solanaceae (tomato and potato) but not in the *C. crispus* genome or in any red algal genome and in other plants, analyzed so far (Figure S3). The unduplicated SSR from non-solanaceous plants is known to be catalytically promiscuous, and indeed Pathmodel suggested that SSR could act on all possible intermediates (Figure 5).

Another alternative synthesis pathway inferred by Pathmodel consists in producing methylated sterols through C24-methylation on desmosterol (Figure 5). This is in agreement with the identification of a methylated sterol, 24-methylenecholesterol, in *C. crispus* (Tasende, 2000) and builds on with other reports about methyltransferase catalytic promiscuity across land plants and green algae (Neelakandan et al., 2009; Haubrich et al., 2015). This option highly reduces the number of non-identified methylated intermediates. Indeed, in land plants, a first methylation, involving methyltransferases, occurs directly on cycloartenol, whereas the second one occurs later on 24-methylenelophenol, to produce methylated sterols like campesterol or brassicasterol (Benveniste, 2004) with intermediates such as cycloeucalenol or fucosterol, both compounds for which we did not find any evidence of presence. By specifying in our model not to enable the production of fucosterol, we naturally omitted this possibility. This model seems also more relevant from a quantitative viewpoint with respect to the formation of cholesterol as the main sterol, because this late methylation step would enable the production of methylated sterols using the late SSR pathway.

A Complete Set of Candidate Enzymes for the C. crispus Sterol Synthesis Pathway

To identify the enzymes associated with the sterol synthesis pathways, we carried out a comparative genomic analysis. Results are summed up in Table S6. In line with previous analyses on sterol biosynthesis gene families in eukaryotes (Desmond and Gribaldo, 2009) or more specifically in green plants (Sonawane et al., 2016), the candidate sterol synthesis enzyme set shows a mixture of conservation and divergence. Seven enzymes are encoded by genes that are conserved as 1:1 orthologs, whereas four of them either underwent lineage-specific duplications (squalene epoxidase and C-4 demethylase) or were lost or may have been replaced by distant paralogs (C24 and C24' methylases and C22 desaturases). In one case, we found no homolog of known plant or animal enzymes performing delta-7/delta-8 isomerization in the *C. crispus* genome, but we found a 1:1 ortholog of ERG2, the gene that secondarily took up this function in yeast (Desmond and Gribaldo, 2009). Another similar case in the sterol synthesis pathway occurs in diatoms, where the epoxisqualene cyclase, otherwise conserved in eukaryotes, was secondarily lost and replaced by a protein belonging to the fatty acid hydroxylase superfamily (Pollier et al., 2019). We consider the ERG2 ortholog to be the best candidate for delta-7/delta-8 isomerization in *C. crispus*, but it is also possible that this reaction is performed by an enzyme encoded by a taxonomically restricted orphan gene (Khalturin et al., 2009).

A Complete View of Algal Primary Metabolism Associated with MAA and Sterol Synthesis

The final GSMN associated with *C. crispus* is constituted of the initial GSMN, enriched with models curated in detail for the sterol and the MAA synthesis pathways. It was therefore reconstructed from both the genome data and the metabolomic data, using the extraction of molecular transformations implemented in Pathmodel (Figure 6). The final GSMN contains 2,207 metabolites, including the eight initial target sterols that are now producible based on Pathmodel inferences as well as seven formerly orphan metabolites (six MAAs and 22-dehydrocholesterol). In this way, we increased the proportion of producible targets from 69.4% (59/85) up to 80% (74/92). The final GSMN is available as a wiki website at the following address: https://gem-aureme.genouest.org/ccrgem/index.php/Main_Page. This website is open for further community curation and will serve as the central point to later integrate additional biochemical knowledge about *C. crispus*. More widely, the curation effort on those pathways will also benefit the entire GSMN community. As already done for the early SSR pathway (now MetaCyc: PWY18C1-3), we will systematically suggest the inclusion of biochemical data used in Pathmodel in the next versions of MetaCyc.

DISCUSSION

In this study we present an extensive analysis of the sterol and MAA biosynthetic pathways in the red alga *C. crispus*, integrating automatic metabolic network reconstruction, manual curation, metabolite profiling, and semi-automatic analogy reasoning approach based on molecular similarity and dissimilarity to generate hypotheses on metabolic pathways associated with secondary metabolism and metabolites,

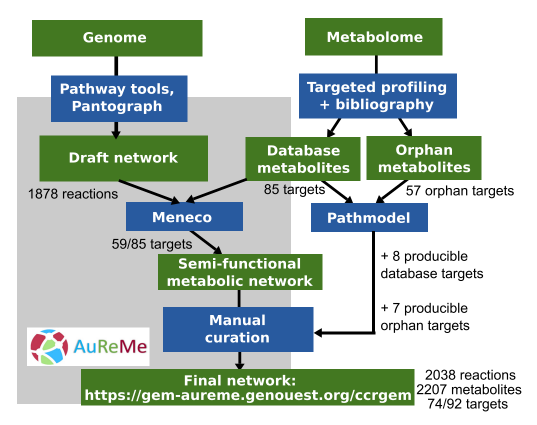


Figure 6. Reconstruction Scheme for the Genome-Scale Metabolic Network of C. crispus

Green boxes indicate starting data and resulting knowledge. Blue boxes indicate the tools that were used to analyze and integrate genomic and metabolomic data. The part overshadowed in gray indicates tools that are already integrated in the AuReMe workflow (Aite et al., 2018).

which are not described in GSMN reference databases such as Metacyc. Three main conclusions can be derived from the presented research. (1) Our findings underline the usefulness of semi-automatic analogy reasoning approaches to link orphan metabolites to existing pathways through the prediction of molecular transformations to infer reactions based on molecular similarity and dissimilarity. (2) These models support our hypothesis of drift as an evolutionary mechanism shaping the metabolism of living organisms. (3) We propose models of sterol and MAA biosynthesis in *C. crispus* and partially validate these models by metabolite profiling.

Semi-automatic Analogy Reasoning Approaches to Integrate Orphan Metabolites

Our study demonstrates that data on metabolite occurrence can be explicitly incorporated into the quality criteria for evaluating a GSMN. To deal with metabolic drift during metabolic network reconstruction, the incorporation of metabolite data is essential, because it puts further constraints on partially known pathways. Putting more emphasis on metabolites, especially the missing ones, creates methodological challenges regarding *ab initio* inferences of pathways when enzymes are not yet known, and we have shown that it is now possible to build tools to specifically address those challenges. The next issue is about the scalability of our approach. The Pathmodel version we present here is a working prototype that can already be applied to other metabolic pathways in *C. crispus* or in other organisms where genomic and metabolomic data are available.

Application of the Pathmodel Approach to Other Studies

The results presented in this study are a first step toward the further development of the underlying bioinformatic tools and their application to additional model biosynthesis pathways. Indeed, the Pathmodel tool was developed to support reasoning based on the metabolic pathway drift hypothesis to automatically infer reactions and metabolites. A key feature of the successful application of this strategy was the



precision and the quality of the biochemical and biological knowledge encoded in the reaction and metabolite databases used as entries to Pathmodel. In particular, all metabolites have to be described with a homogeneous ordering of atoms to predict molecular transformations. Generalizing this approach to any other application will similarly require interactions between chemists, biologists, and computer scientists. Further improvements should be made to minimize the burden in manually entering molecular structures. It is not yet possible to fully automate the atom numbering during metabolic reaction, due to the intrinsic complexity of metabolic pathways. For example, the split of molecules during biochemical processes can generate inconsistent numbering (Figure S5A) that can only be handled by resetting atom numbering from one step to another. This is already done in MetaCyc, but in a way that does not make possible to automatically number in the same manner atoms from different reactions that share a molecular transformation (Figure S5B). Other methods, like the CLCA approach (Kumar and Maranas, 2014) already implemented ways to compare reactions sharing molecular transformations, and thus would provide lists of candidate reactions for molecular transformations, but the atom numbering during the comparisons does not allow simultaneous atom mapping between reactant and product (Figure S5C) that we need in Pathmodel to abstract the molecular transformation. Therefore, following IUPAC atom encoding as much as possible seems to be the best way to combine atom mapping and abstraction of molecular transformations (Figure S5D).

The second key feature of Pathmodel is to be focused on a selected pathway rather than on a complete genome-scale metabolic network. The selection of the relevant pathway to be considered, for instance, from preliminary evidences extracted from metabolomic analysis, is therefore a key pre-processing step to combine and filter the predictions of Pathmodel with genomic and metabolomic data. Practically, Pathmodel can already be used on any type of incomplete biochemical pathway on a molecule class for which there is knowledge about some biochemical reactions and orphan molecules not yet connected to each other by any reaction. The main limitation will be the user time necessary to properly encode the starting knowledge base.

Metabolic Drift as a Driver for Pathway Evolution

Whatever the actual topology of the sterol and MAA pathways in *C. crispus*, each discussed hypothesis has implications regarding metabolic pathway drift. All possible sterol pathways provide further strong candidate case studies for drift by non-homologous enzyme replacement, and the pathways inferred by Pathmodel provide candidate case studies for drift by enzyme inversion. The unresolved point with the sterol pathways is that, among eukaryotes, there is no consensus yet about the ancestral order of enzymatic reactions. Experimental data are too disparate across the tree of life to enable firm conclusions on this. In that respect, the MAA pathway is interesting, because if our hypothesis about decarboxylation of porphyra-334 before dehydration is true, this would mean that an enzymatic inversion took place in other lineages where porphyra-334 is first dehydrated to Z-palythenic acid and then decarboxylated to palythene. Here the limit is that, to date, enzymes are unknown for both reactions, so the system is not yet genomically tractable. Identifying close enzymatic inversions would be important to provide a mechanism for gradual divergence of pathways. Indeed, experimental analyses on *E. coli* have shown that drastic pathway rewiring by enzyme knockout or gene overexpression can led to toxic intermediates (Kim et al., 2010).

Limitations of the Study

Our arguments for the metabolic pathway drift hypothesis rely on comparisons between pathways from *C. crispus* and from distantly related species belonging to a few other phyla, such as land plants or animals. Additional support should become available from detailed comparisons based on additional eukaryotic lineages and, when possible, from multiple species in the same phylum. Finally, it remains to be determined to what extent chance and selection contributed in generating metabolic pathway diversity.

METHODS

All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.100849.



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AUTHOR CONTRIBUTIONS

G.V.M., L.D., S.M.D., P.S., E.C., J.N., C.B., C. Leblanc, A.S., and J.C. conceived the project. J.G. conducted the sterol profiling with help from L.D., C. Leroux, C. Leblanc, J.C., and G.V.M. P.S. conduced the MAA profiling with help from C.M. G.V.M. performed the genome-scale metabolic network reconstruction with help from M.A., C.T., S.M.D, J.C., and A.S. A.B., G.V.M., and J.N. wrote the Pathmodel software. A.B., J.G., L.D., and G.V.M. curated the biosynthetic pathway models. G.V.M. wrote the manuscript with input and edits from all authors.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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Supplemental Information

Inferring Biochemical Reactions and Metabolite Structures to Understand

Metabolic Pathway Drift

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TRANSPARENT METHODS

Genome-Scale Metabolic Network reconstruction

Genome-Scale Metabolic Network (GSMN) reconstruction was performed using the AuReMe pipeline (Aite et al., 2018). A set of 85 targets coming from the literature was used as an input and is provided in Table S1. Orphan metabolites that are experimentally supported but do not have a MetaCyc ID are listed in Table S2. The process encompassed the following steps:

- 1) an annotation-based draft network was generated using the PathoLogic program from the Pathway Tools suite, using the gbk file from the *Chondrus crispus* genome annotation (Collén et al., 2013) and the metabolic reaction database MetaCyc20.5 (Caspi et al., 2016).
- 2) an orthology-based network was generated using the protein sequences and metabolic network of *Arabidopsis thaliana* (AraGEM, De Oliveira d'al Molin et al., 2010), using the Pantograph software (Loira et al., 2015) to combine the output of ortholog searches with the Inparanoid and OrthoMCL softwares.
- 3) an orthology-based network was generated using the protein sequences from the well-annotated red microalga *Galdieria sulphuraria* (Schönknecht et al., 2013) and its metabolic network reconstructed using Pathway Tools. This *G. sulphuraria* annotation-based network was then used as a template to generate a *C. crispus* network using Pantograph. We decided to build this template GSMN after realizing that the genbank file was especially annotation-rich.
- 4) an orthology-based network was also generated using the protein sequences from the version 2 of the annotated genome of *Ectocarpus siliculosus* (Cormier et al., 2017), as well as version 2 of its metabolic network (Aite et al., 2018).
- 5) the four preliminary networks were merged together in the AuReMe environment, and an additional gap-filling step was performed using Meneco (Prigent et al., 2017), constraining the network to produce the 85 metabolites from the literature that were indexed in the Metacyc database (Table S1).

Sampling of algae

For sterol analyses, samples from *C. crispus* were collected from a population on the shore at Roscoff, France, in front of the Station Biologique (48°43'38" N; 3°59'04" W). Algal cultures were maintained in 10 L flasks in a culture room at 14°C using filtered seawater and aerated with 0.22 µm-filtered compressed air to avoid CO₂ depletion. Photosynthetically active radiation (PAR) was provided by Philips daylight fluorescence tubes at a photon flux density of 40 µmol.m⁻².s⁻¹ for 10 h.d⁻¹. The algal samples were freeze dried, ground to powder using a cryogrinder and stored at

-80°C.

For MAAs analysis, more than 50 g (wet weight) of *C. crispus* were collected along the Brittany coasts (France) at Ploemeur (47°42'07" N; 3°24'31" W) in July 2013, Roscoff (48°43'38" N; 3°59'04" W) in April and August 2013, and Tregunc (47°50'25"N; 3°54'08" W) in September 2013.

Standards and reagents

Cholesterol, stigmasterol, β -sitosterol, 7-dehydrocholesterol, lathosterol (5 α -cholest-7-en-3 β -ol), squalene, campesterol, brassicasterol, desmosterol, lanosterol, fucosterol, cycloartenol, 5 α -cholestane (internal standard) were acquired from Sigma-Aldrich (Saint-Quentin-Fallavier, France), cycloartanol and cycloeucalenol from Chemfaces (Wuhan, China) and zymosterol from Avanti Polar Lipids (Alabaster, USA). The C7-C40 Saturated Alkanes Standards were acquired from Supelco (Bellefonte, USA). Reagents used for extraction, saponification, and derivation steps were n-hexane, ethyl acetate, acetonitrile, methanol (Carlo ERBA Reagents, Val de Reuil, France), (trimethylsilyl)diazomethane, toluene (Sigma-Aldrich, Saint-Quentin-Fallavier, France) and N,O-bis(trimethylsilyl)trifluoroacetamide with trimethylcholorosilane (BSTFA:TMCS (99:1)) (Supelco, Bellefonte, USA).

Standard preparation

Stock solutions of cholesterol, stigmasterol, β -sitosterol, 7-dehydrocholesterol, lathosterol (5 α -cholest-7-en-3 β -ol), squalene, campesterol, brassicasterol, desmosterol, lanosterol, fucosterol, cycloartenol and 5 α -cholestane were prepared in hexane with a concentration of 5 mg.mL⁻¹. Working solutions were made at a concentration of 1 mg.mL⁻¹, in hexane, by diluting stock solutions. The C7-C40 Saturated Alkanes Standard stock had a concentration of 1 mg.mL⁻¹ and a working solution was made at a concentration of 0.1 mg.mL⁻¹. All solutions were stored at -20°C.

Sample preparation

For sterol analyses, dried algal samples (60 mg) were extracted with 2mL ethyl acetate by continuous agitation for 1 hour at 4°C. After 10 min of centrifugation at 4000 rpm, the solvent was removed, the extracts were saponified in 3 mL of methanolic potassium hydroxide solution (1M) by 1 hour incubation at 90°C. The saponification reaction was stopped by plunging samples into an ice bath for 30 min minimum. The unsaponifiable fraction was extracted with 2 mL of hexane and 1.2 mL of water and centrifuged at 2000 rpm for 5 min. The upper phase was collected, dried under N_2 , and resuspended with 120 μ L of (trimethylsilyl)diazomethane, 50 μ L of methanol:toluene (2:1 (v/v)) and 5 μ L of 5 α -cholestane (1 mg.mL⁻¹) as internal standard. The mixture was vortexed for 30

seconds, and heated at 37°C for 30 min. After a second evaporation under N_2 , 50 μL of acetonitrile and 50 μL of BSTFA:TMCS (99:1) were added to the dry residue, vortexed for 30 seconds and heated at 60°C for 30 min. After final evaporation under N_2 , the extract was resuspended in 100 μL of hexane, transferred into a sample vial and stored at -80°C until the GC-MS analysis.

For MAAs, one gram of dried algae was extracted twice for two hours under continuous shaking with 10 mL of acetone. After 5 min of centrifugation at 3000 rpm, acetone was discarded and samples were re-extracted twice with 10 mL water/acetone (30/70, v/v) for 24 hours under continuous shaking at 120 rpm. Water/acetone supernatants were pooled, added to one gram of silica and evaporated to dryness by rotary evaporation. Extracts were then purified by silica gel chromatography column with dichloromethane/methanol mixtures and MAAs were eluted with 200 mL of dichloromethane/methanol (15/85, v/v). After rotary evaporation, samples were re-suspended in water/methanol (50/50, v/v) and filtrated using 0.45 μm syringes filter. Solution were adjusted to a final concentration of 1 mg.mL⁻¹ and stored at 3°C until LC-MS analysis.

Sterol analysis by gas chromatography-mass spectrometry

The sterols were analyzed on a 7890 Agilent Technologies gas chromatography coupled with a 5975C Agilent Technologies mass spectrometer (GC-MS). A HP-5MS capillary GC column (30 m x 0.25 mm x 0.25 µm) from J&W Scientific (CA, USA) was used for separation and UHP helium was used as carrier gas at flow rate to 1 mL.min⁻¹. The temperature of the injector was 280°C and the detector temperature was 315°C. After injection, the oven temperature was kept at 60°C for 1 min. The temperature was increased from 60°C to 100°C at a rate of 25°C.min⁻¹, then to 250°C at a rate of 15°C.min⁻¹, then to 315°C at a rate of 3°C.min⁻¹ and then held at 315°C for 2 min, resulting in a total run time of 37 min. Electronic impact mass spectra were measured at 70eV and an ionization temperature of 250°C. The mass spectra scanned from m/z 50 to m/z 500. Peaks were identified based on the comparisons with the retention times and the mass spectra (Table S3).

MAA analysis by liquid chromatography-mass spectrometry

High Resolution Mass Spectrometry was carried out on a microTOF-Q II (Bruker Daltonics, Germany) coupled to an Ultimate 3000 LC System (Dionex, Germany). Experiments were performed on a Gemini C6-Phenyl column (250 mm x 4.6 mm x 5 μm) (Phenomenex, Germany). The gradient was as follows: methanol/water (20:80, v/v) with 0.2% acid acetic for two minutes to 100 % methanol with 0.2% acid acetic in 23 minutes. The UV detector was set to 330 nm, flow rate was kept constant at 0.4 mL.min⁻¹ and column temperature set at 30°C. MS spectra were recorded in positive ESI mode with a drying gas temperature of 220°C, a nitrogen flow of 12 L.min⁻¹, a nebulizer pressure set to 60 psi, and a collision energy of 20 eV. MAAs were identified by HR-MS

on the basis of the detection of the pseudo-molecular ion $[M+H]^+$ with a m/z value varying less than \pm 0.02 Da compared to the theoretical m/z value. In the absence of commercially available standards, relative quantification of MAAs in each sample was estimated by calculating the ratio between the area under the curve of the Extracted Ion Chromatogram (EIC) corresponding to the selected MAAs and the sum of the areas under the curve of the EIC of all MAAs detected in the algal extract. The same procedure was applied to UV detection (Table S4).

Flux-balance analysis

A biomass reaction was established based on the previous *E. siliculosus* data, defining a list of 33 compounds to be produced in order to consider the network functional (Prigent et al., 2014). One compound, L-alpha-alanine, was not producible, thus blocking biomass production. This was due to the absence of the alanine dehydrogenase reaction. The corresponding enzyme (CHC_T00008930001) was present in the *C. crispus* network but annotated as an NAD(P) transhydrogenase. We completed the annotation through the manual curation form to enable it to dehydrogenate alanine and to restore producibility of the biomass (https://gem-aureme.genouest.org/ccrgem/index.php/Manual-ala_dehy).

Global metabolic networks comparisons

In order to compare the global features of the GSM from *C. crispus* with other ones, it is necessary to use the same reference database. This is the case for *E. siliculosus* and *E. subulatus* for which the reconstructions are based on MetaCyc (Caspi et al., 2016) while *A. thaliana* and *Chlamydomonas* reinhardtii are respectively from KEGG (Kanehisa et al., 2017) and BiGG (King et al., 2016). To get access to MetaCyc pathway information for *A. thaliana* and *C. reinhardtii*, their networks were mapped using the sbml_mapping function implemented in the AuReMe workflow (Aite et al., 2018). This function provides a dictionary of corresponding reactions from a database to another one using the MetaNetX cross-reference database (Moretti et al., 2016). This dictionary was then used in AuReMe to create a new genome-scale metabolic network based on the new reference database for *A. thaliana* and *C. reinhardti*. Those new networks, who are comparable in size with the published ones (+/- 10 reactions and enzymes in our counts) enabled to estimate the number of pathways as defined in MetaCyc for both species.

Ab-initio inference of metabolic reactions: implementation of a Semi-Automatic Analogy Reasoning Approach

The Pathmodel method was developed to infer new reactions based on molecular similarity and dissimilarity. This knowledge-based approach is founded on two modes of reasoning (deductive and

analogical) and was implemented using a logic programming approach known as Answer Set Programming (ASP) (Lifschitz et al., 2008; Gebser et al., 2012). It is a declarative approach oriented toward combinatorial (optimization) problem-solving and knowledge processing. ASP combines both a high-level modeling language with high performance solving engines so that the focus is on the problem specification rather than the algorithmic part. ASP expresses a problem as a set of logical rules (clauses). Problem solutions appear as particular logical models (so-called stable models or answer sets) of this set. An ASP program consists of rules $h :- b_1, \ldots, b_m$ not $b_{m+1}, \ldots, not b_n$, where each b_i and h are literals and not stands for default negation. In fact, each proposition is a predicate, encoded by a function whose arguments can be constant atoms or variables over a finite domain. The rule states that the head h is proven to be true (h is in an answer set) if the body of the rule is satisfied, i.e. b_1, \ldots, b_m are true and it cannot be proved that b_{m+1}, \ldots, b_n are true.

The main predicates used in Pathmodel to represent molecules and reactions forming a knowledge base are *bond*, *atom* and *reaction*. The theoretical m/z ratio of a molecule is determined by logical rules, which were encoded in the program MZComputation.lp.

As depicted in Figure 3, several logical rules are then applied to all possible reactions and potential reactants. These are the bases for the selection of potential reactants or products and the inference by a reasoning component of reaction occurrences or metabolites, using either deductive or analogical reasoning in the PathModel.lp program. Resulting products that do not belong to the knowledge base but that correspond to an observed m/z ratio are considered as inferred metabolites and reactions. The finally encoded reactions result from iterative interactions between analogical model construction, automated inference, and manual validation of inferred reactions with respect to experimental results.

By comparing reactants and products, the program ReactionSiteExtraction.lp characterizes two structures of the reaction site containing atoms and bonds involved in the reaction. The predicates diffAtomBeforeReaction, diffBondBeforeReaction, diffAtomAfterReaction and diffBondAfterReaction compare atoms and bonds between the reactant and the product and extract the two structures. Then these two structures are compared to the structure of all other molecules in the knowledge base (predicates siteBeforeReaction and siteAfterReaction). These predicates characterize sub-structures of the molecules that can be part of a reaction.

By deductive reasoning, the reference molecule pair of each reaction is compared to the structures of a potential reactant-product pair sharing a common chemical structure. The presence of the reaction site in the two putative molecules is checked using the predicates *siteBeforeReaction* and *siteAfterReaction*. Furthermore, if the product and the reactant have the same overall structure, except for the reaction site, the program will infer that the reaction actually occurs between the

reactant and the product.

By analogical reasoning, all possible reactions are applied to potential reactants, and resulting products are filtered using their structures and m/z ratios. The predicate *newMetaboliteName* creates all the possible products from a known molecule using all the reactions in the knowledge base. These possible metabolites are filtered using their m/z ratios, which must correspond to an unassigned m/z ratio (predicate *possibleMetabolite*) and checked if they share the same structure as a known molecule (predicate *alreadyKnownMolecule*). If they do not fit with an already known molecule, they will be added as new molecules and a new reaction variant.

Given a source molecule and a target molecule, the program will take several inference steps iteratively applying either analogical or deductive reasoning modes. To connect the source and the target molecules along a pathway, Pathmodel infers missing reactions and metabolites using a minimal number of reactions. To further constraint the number of possible pathways, a predicate *absentmolecules* was added to avoid pathways with compounds for which targeted profiling with analytical standards gives strong evidence for real absence (here ergosterol, fucosterol and zymosterol). The source code is available in the following Github repository: https://github.com/pathmodel/pathmodel

It includes a specific tutorial to replicate the analysis reported in the article:

https://github.com/pathmodel/pathmodel#tutorial-on-article-data-chondrus-crispus-sterol-and-mycosporine-like-amino-acids-pathways

De novo gene prediction and manual curation of gene sequence models

Missing genes from the sterol synthesis pathway (squalene monooxygenase and sterol C-4 methyl oxidase) were found by targeted tblastn using orthologs from other organisms as a query. The new gene predictions are provided in supplementary dataset 1 and will be included in the next version of *C. crispus* genome browser (http://mmo.sb-roscoff.fr/jbrowse/?data=data%2Fpublic%2Fchondrus). The split protein sequence of sterol delta-7 reductase was also restored as a single protein prediction, merging the two adjacent partial predictions.

Phylogenetic analyses

Collected sequences were aligned using Clustal Omega (Sievers and Higgins, 2014) and alignments were checked manually and edited with Seaview (Gouy et al., 2010). Phylogenetic trees were built using PHYML (Guindon and Gascuel, 2003) using the LG model (Le and Gascuel, 2010) with a gamma law. The reliability of nodes was assessed by likelihood-ratio test (Anisimova and Gascuel, 2006).

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Table S1 related to Fig 6. Database metabolites.

References

MetaCyC ID

Category

Usual name

dodecanoic acid	12:0 fatty acid	DODECANOATE	Santos et al., 2015 ; Robertson et al., 2015
Myristic acid	14:0 fatty acid	CPD-7836	Pettitt et al., 1989; Tasende, 2000; Van Ginneken et al., 2011; Robertson et al., 2015; Belghit et al., 2017
Pentadecanoic acid	15:0 fatty acid	CPD-8462	Santos et al., 2015. Belghit et al., 2017
Palmitic acid	16:0 fatty acid	PALMITATE	Pettitt et al., 1989; Tasende, 2000; Van Ginneken et al., 2011; Robertson et al., 2015 ; Belghit et al., 2017
Heptadecanoic acid	17:0 fatty acid	CPD-7830	Santos et al., 2015
Stearic acid	18:0 fatty acid	STEARIC_ACID	Tasende et al., 2000 ; Robertson et al., 2015
Eicosanoic acid	20:0 fatty acid	ARACHIDIC_ACID	Santos et al., 2015
Docosanoic acid	22:0 fatty acid	DOCOSANOATE*	Santos et al., 2015
Tricosanoic acid	23:0 fatty acid	CPD-7834*	Santos et al., 2015
Tetracosanoic acid	24:0 fatty acid	TETRACOSANOATE	Santos et al., 2015
Palmitoleic acid	16:1(n-7) fatty acid	CPD-9245	Pettitt et al., 1989; Tasende, 2000; Robertson et al., 2015; Belghit et al., 2017
Oleic acid	18:1(n-9) fatty acid	OLEATE-CPD	Tasende et al., 2000; Van Ginneken et al., 2011; Robertson et al., 2015 ; Belghit et al., 2017
Linoleic acid	18:2(n-6) fatty acid	LINOLEIC_ACID	Tasende et al., 2000 ; Robertson et al., 2015 ; Belghit et al., 2017
Alpha Linolenic acid	18:3(n-3) fatty acid	LINOLENIC_ACID*	Tasende et al., 2000
γ-linolenic acid	18:3(n-6) fatty acid	CPD-8117*	Robertson et al., 2015 ; Belghit et al., 2017
Octadecatetraenoic acid	18:4(n-3) fatty acid	CPD-12653*	Tasende et al., 2000 ; Robertson et al., 2015 ; Belghit et al., 2017
Arachidonic acid	20:4(n-6) fatty acid	ARACHIDONIC_ACID	Tasende et al., 2000 ; Banskota et al., 2014 ; Robertson et al., 2015 ; Belghit et al., 2017
Eicosapentaenoic acid	20:5(n-3) fatty acid	5Z8Z11Z14Z17Z-EICOSAPENTAENOATE*	Tasende et al., 2000 ; Banskota et al., 2014 ; Robertson et al., 2015 ; Belghit et al., 2017
Octanedioic acid	fatty acid	CPD0-1264*	Santos et al., 2015
Nonanedioic acid	fatty acid	CPD0-1265*	Santos et al., 2015
Cycloartenol	sterol	CYCLOARTENOL*	Saito and Idler, 1966; Alcaide et al., 1968
Cholesterol	sterol	CHOLESTEROL*	Saito and Idler, 1966; Tasende et al., 2000; Santos et al., 2015
7-Dehydrocholesterol	sterol	7-DEHYDROCHOLESTEROL*	Tasende et al., 2000
Brassicasterol	sterol	BRASSICASTEROL*	Saito and Idler, 1966; Tasende et al., 2000
Campesterol	sterol	CAMPESTEROL*	Tasende et al., 2000 ; Santos et al., 2015
24-Methylenecholesterol	sterol	24-METHYLENECHOLESTEROL*	Tasende et al., 2000
Sitosterol	sterol	SITOSTEROL*	Saito and Idler, 1966; Tasende et al., 2000; Santos et al., 2015
Stigmasterol	sterol	STIGMASTEROL*	Tasende et al., 2000
15-keto-prostaglandin E2	oxylipin	HYDROXY-915-DIOXOPROSTA-13-ENOATE*	Gaquerel et al., 2007
lutein	carotenoid	LUTEIN*	Banskota et al., 2014
Chlorophyll a	tetrapyrrole	CHLOROPHYLL-A	Melo et al., 2015; Robertson et al., 2015
all-trans-beta-carotene	carotenoid	CPD1F-129	Robertson et al., 2015
9-cis-betacarotene	carotenoid	CPD-14646	Robertson et al., 2015; Belghit et al., 2017
zeaxanthin	carotenoid	CPD1F-130	Robertson et al., 2015
2,6,6-trimethyl-1,3-cyclohexadiene-1-carboxaldehyde (safranal)	carotenoid	CPD-8669*	Pina et al., 2014
Alanine	aminoacid	L-ALPHA-ALANINE	Young et al., 1958, Belghit et al., 2017
Arginine	aminoacid	ARG	Young et al., 1958, Belghit et al., 2017
Aspartic acid	aminoacid	L-ASPARTATE	Young et al., 1958, Belghit et al., 2017
Citrulline	aminoacid	L-CITRULLINE	Young et al., 1958 ; Belghit et al., 2017

Table S1 related to Fig 6. Database metabolites.

Cystine	aminoacid	CYSTINE	Young et al., 1958
Glutamic acid	aminoacid	GLT	Young et al., 1958 ; Belghit et al., 2017
Glycine	aminoacid	GLY	Young et al., 1958 ; Belghit et al., 2017
Histidine	aminoacid	HIS	Young et al., 1958 ; Belghit et al., 2017
Isoleucine	aminoacid	ILE	Young et al., 1958 ; Belghit et al., 2017
Leucine	aminoacid	LEU	Young et al., 1958; Belghit et al., 2017
Lysine	aminoacid	LYS	Young et al., 1958; Belghit et al., 2017
Methionine	aminoacid	MET	Young et al., 1958; Belghit et al., 2017
Ornithine	aminoacid	L-ORNITHINE	Young et al., 1958; Belghit et al., 2017
Phenylalanine	aminoacid	PHE	Young et al., 1958
Proline	aminoacid	PRO	Young et al., 1958; Belghit et al., 2017
Serine	aminoacid	SER	Young et al., 1958; Belghit et al., 2017
Threonine	aminoacid	THR	Young et al., 1958; Belghit et al., 2017
Tyrosine	aminoacid	TYR	Young et al., 1958; Belghit et al., 2017
Valine	aminoacid	VAL	Young et al., 1958; Belghit et al., 2017
Shinorine	Mycosporine-like aminoacid	CPD-18778	Kräbs et al., 2004
UDP-α-D-galactose	nucleotide sugar	CPD-14553	Collén et al., 2014
D-galactosyl-1,2-diacylglycerol	galactolipid	D-Galactosyl-12-diacyl-glycerols	Banskota et al., 2014
ı-carrageenose	carrageenan	Iota-Carrageenan*	Matsuhiro et al., 1992
v-carrageenan	carrageenan	Nu-Carrageenan*	Matsuhiro et al., 1992
Glycerol	polyol	GLYCEROL	Santos et al., 2015
Heptadecane	alcane	HEPTADECANE-CPD	Santos et al., 2015
6,10,14-Trimethyl-2-pentadecanone	methylketone	CPD-7875	Santos et al., 2015
Hexadecan-1-ol	Long chain aliphatic alcohol	CPD-348	Santos et al., 2015
9-Octadecen-1-ol	Long chain aliphatic alcohol	CPD-7873	Santos et al., 2015
Docosan-1-ol	Long chain aliphatic alcohol	CPD-7845	Santos et al., 2015
Octacosan-1-ol	Long chain aliphatic alcohol	CPD-7872*	Santos et al., 2015
acetaldehyde	aldehyde	ACETALD	Pina et al., 2014
2-methypropanal	aldehyde	BUTANAL	Pina et al., 2014
Butanal	aldehyde	CPD-7031	Pina et al., 2014
3-methybutanal	aldehyde	METHYLBUT-CPD	Pina et al., 2014
Pentanal	aldehyde	CPD-9053*	Pina et al., 2014
Hexanal	aldehyde	HEXANAL	Pina et al., 2014
Benzaldehyde	aldehyde	BENZALDEHYDE	Pina et al., 2014
Ethanol	short chain aliphatic alcohol	ETOH	Pina et al., 2014
1-butanol	short chain aliphatic alcohol	BUTANOL	Pina et al., 2014
1-pentanol	short chain aliphatic alcohol	PENTANOL*	Pina et al., 2014
2-butanone	short chain ketone	ACETONE	Pina et al., 2014
3,5-octadien-2-one	short chain ketone	MEK	Pina et al., 2014

Table S1 related to Fig 6. Database metabolites.

dichloromethane	halocarbon	CPD-681	Pina et al., 2014
chloroform	halocarbon	CPD-843*	Pina et al., 2014
glycerate	carboxylic acid	GLYCERATE	Belghit et al., 2017
2-methylpropanoic acid	carboxylic acid	ACET	Pina et al., 2014
2-methylbutanoic acid	carboxylic acid	ISOBUTYRATE	Pina et al., 2014
hexane	alcane	CPD-9288*	Pina et al., 2014
2,2,4-trimethylpentane	alcane	CPD-19039*	Pina et al., 2014

^{*} non predicted in initial GSMN reconstruction

Table S2 related to Fig 6. Orphan metabolites.

Usual name	Category	References
Heneicosanoic acid	21:0 fatty acid	Santos et al., 2015
N/A	15:1 fatty acid	Robertson et al., 2015
N/A	18:1(n-7) fatty acid	Robertson et al., 2015
10-nonadecenoate	19:1(n-9) fatty acid	Belghit et al., 2017
Eicosadienoic acid	20:2(n-6) fatty acid	Robertson et al., 2015
Eicosatrienoic acid	20:3(n-6) fatty acid	Robertson et al., 2015
Docosadienoate	22:2(n-6) fatty acid	Belghit et al., 2017
Octadeca-9-enoic acid	fatty acid	Santos et al., 2015
22-Dehydrocholesterol*	sterol	Tasende et al., 2000
11-hydroxy-octadecadienoic acid (11-HODE)	oxylipin	Gaquerel et al., 2007
13-hydroxy-9Z,11E-octadecadienoic acid (13-HODE)	oxylipin	Gaquerel et al., 2007; Belghit et al., 2017
13S-hydroxy-9Z,11E,15Z-octadecatrienoic acid (13-HOTrE)	oxylipin	Belghit et al., 2017
13-oxo-9Z,11E-octadecadienoic acid (13-oxo-ODE)	oxylipin	Gaquerel et al., 2007
13-hydroxyeicosatrienoic acid (13-HETrE)	oxylipin	Gaquerel et al., 2007
13-hydroxyeicosatetraenoic acid (13-HETE)	oxylipin	Gaquerel et al., 2007
13-hydroxyeicosapentaenoic acid (13-HEPE)	oxylipin	Gaquerel et al., 2007
15-hydroxydocosahexaenoic acid (15-HDHE)	oxylipin	Gaquerel et al., 2007
11-hydroxyoctadecadienoic acid (11-HETE)	oxylipin	Gaquerel et al., 2007
Hydroxypheophytin a	tetrapyrrole	Melo et al., 2015
Pheophytin d	tetrapyrrole	Melo et al., 2015
Hydroxypheophytin d	tetrapyrrole	Melo et al., 2015
Monogalactosyldiacylglycerol 2 (MGDG2)	galactolipid	Pettitt et al., 1989
Digalactosyldiacylglycerols (DGDG)	galactolipid	Pettitt et al., 1989
Sulfoquinovosyldiacylglycerol 1 (SQDG1)	galactolipid	Pettitt et al., 1989
Sulfoquinovosyldiacylglycerol 2 (QDG2)	galactolipid	Pettitt et al., 1989
$(2S)\hbox{-}1,2\hbox{-}bis\hbox{-}O\hbox{-}eicosapentaenoyl-}3\hbox{-}O\hbox{-}\beta\hbox{-}D\hbox{-}galactopyranosylglycerol}$	galactolipid	Banskota et al., 2014
$(2S\)-1-O-eicosapentaenoyl-2-O-arachidonoyl-3-O-\beta-D-galactopyranosylglycerol$	galactolipid	Banskota et al., 2014
$(2S)\text{-}1\text{-}O\text{-}(6Z,9Z,12Z,15Z\text{-} octade catetran oyl)\text{-}2\text{-}O\text{-}palmitoyl\text{-}3\text{-}O\text{-}\beta\text{-}D\text{-} galactopyran osylgly cerol}$	galactolipid	Banskota et al., 2014
(2S)-1-O -eicosapentaenoyl-2- O -palmitoyl-3-O - β -D -galactopyranosylglycerol	galactolipid	Banskota et al., 2014
(2S)-1, 2-bis -O -arachidonoyl-3-O - β -D -galactopyranosylglycerol	galactolipid	Banskota et al., 2014
(2S)-1-O -arachidonoyl-2-O -palmitoyl-3-O - β -D - galactopyranosylglycerol	galactolipid	Banskota et al., 2014
$(2S)\text{-}1\text{-}O\text{-}eicos apentaenoyl-2-}O\text{-}palmitoyl-3-}O\text{-}(\beta\text{-}D\text{-}galactopyranosyl-6-}1\alpha\text{-}D\text{-}galactopyranosyl})\text{-}glycerol$	galactolipid	Banskota et al., 2014
$(2S) \text{-} 1\text{-}O\text{-}arachidonoyl\text{-}2\text{-}O\text{-}palmitoyl\text{-}3\text{-}O\text{-}(\beta\text{-}D\text{-}galactopyranosyl\text{-}6\text{-}1}\alpha\text{-}D\text{-}galactopyranosyl\text{)-}glycerol$	galactolipid	Banskota et al., 2014
diphosphatidylglycerol, phosphatidic acid	phospholipid	Pettitt et al., 1989
l-citrullinyl-l-arginine	aminoacid	Laycock et al., 1977
Gigartinine	aminoacid	Laycock et al., 1977
Amide N	aminoacid	Young et al., 1958
Asterina-330*	Mycosporine-like aminoacid	Athukorala et al., 2016; Guihéneuf et al., 2018

Table S2 related to Fig 6. Orphan metabolites.

MAA1* MAA2*

Palythine*

Palythene

Porphyra-334*

Isofloridoside

1-Monohexadecanoin

Tetradecan-1-ol

Octadecan-1-ol

1-penten-3-ol

2(Z)-penten-1ol

3-methylbutanoic acid

2-methylbutanal

1-octen-3-ol

2,6-dimethylpyrazine

2-propanone

acetic acid, anhydride

2,2,3-trimethylpentane

tetradecane

Mycosporine-like aminoacid This study

Mycosporine-like aminoacid This study

Mycosporine-like aminoacid Karsten et al., 1998; Athukorala et al., 2016; Guihéneuf et al., 2018

Mycosporine-like aminoacid Karsten et al., 1998

Mycosporine-like aminoacid Athukorala et al., 2016
heteroside Kremer et al., 1982

Long chain aliphatic alcohol Santos et al., 2015

Long chain aliphatic alcohol Santos et al., 2015

Long chain aliphatic alcohol Santos et al., 2015

short chain aliphatic alcohol Pina et al., 2014

short chain aliphatic alcohol Pina et al., 2014

carboxylic acid Pina et al., 2014
aldehyde Pina et al., 2014
short chain aliphatic alcohol Pina et al., 2014
short chain ketone Pina et al., 2014
short chain ketone Pina et al., 2014
carboxylic acid Pina et al., 2014
alcane Pina et al., 2014
Pina et al., 2014

^{*} incorporated in GSMN after Pathmodel

Analysed compounds	Molecular weight (g.mol-1)	RT (min)	m/z [M+H] ⁻ (TMS)
brassicasterol	398.64	25.5	470
campesterol	400.68	26.6	472
5α-cholestane	372.67	20.3	372
cholesterol	386.65	24.7	458
cycloartanol	428.75	30.5	500
cycloartenol	426.72	29.0	498
cycloeucalenol	426.73	30.4	498
7-dehydrocholesterol	384.63	25.6	456
desmosterol	384.64	25.5	456
ergosterol	396.65	26.2	468
fucosterol	412.69	28.2	484
lanosterol	426.39	27.8	498
lathosterol	386.65	25.8	458
β-sitosterol	414.39	28.0	486
squalene	410.72	19.8	482
stigmasterol	412.66	27.0	484
zymosterol	384.64	25.9	456

Table S3, related to Figure 5. Retention times and m/z ratio for analytical standards of sterols on a 7890-5975C Agilent GC-MS.

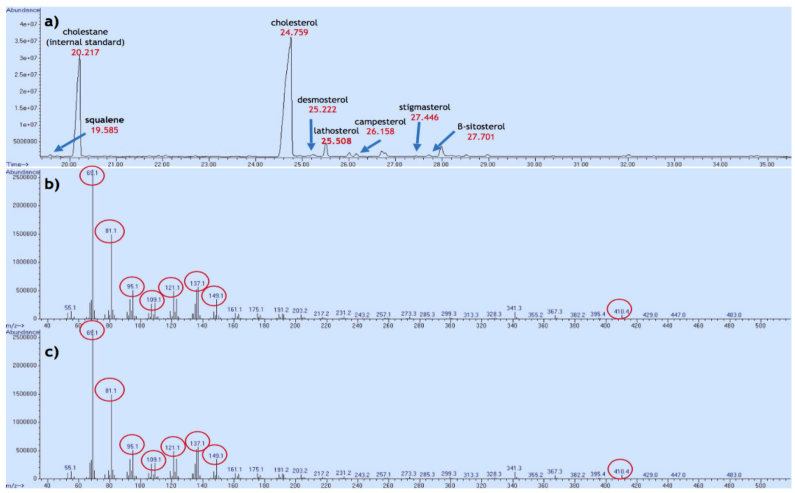


Figure S1, related to Figure 5. Identification of squalene in *C. crispus.* a) Total Ion Chromatogramm (TIC) from *C. crispus* extract. b) MS spectrum of squalene in *C. crispus* extract. c) MS spectrum of the squalene analytical standard. Main fragmentation peaks identical in both spectra are highlighted in red circles.

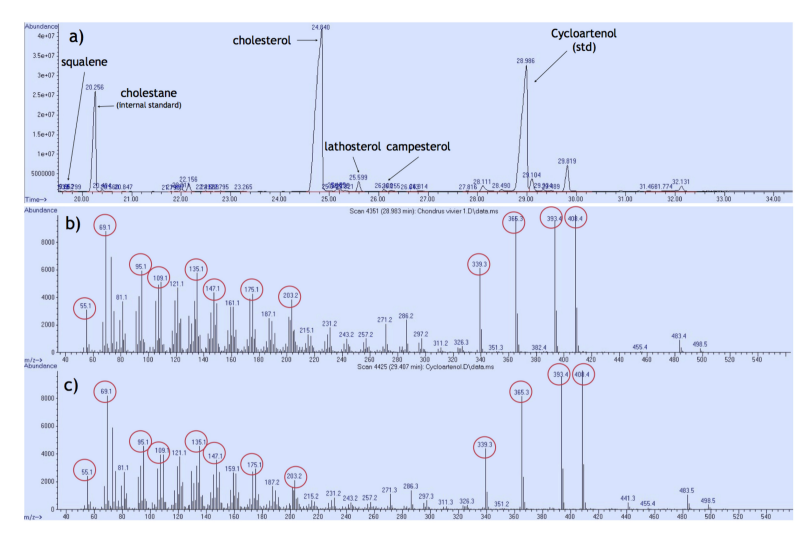


Figure S2, related to Figure 5. Control for technical detectability of cycloartenol in spiked *Chondrus crispus* extract.

a) TIC from *Chondrus crispus* extract incubated with cycloartenol. b) MS spectrum of cycloartenol standard incorporated in *C. crispus* extract. c) MS spectrum of cycloartenol standard alone. Main fragmentation peaks identical in both spectra are highlighted in red circles.

MAAs	Palythine	Mycosporine-glycine	MAA1	Isujirene/Palythene	Asterina-330	Palythinol	Shinorine	Porphyra-334
						or MAA2		
Rt (min.)	8.3	20.0	10.8	19.3	8.7	10.1	18.5	19.5
m/z [M+H]+ observed	245.1090	246.0932	271.1241	285.1401	289.1349	303.1497	333.1245	347.1399
m/z calculated	245.1132	246.0972	271.1288	285.1445	289.1394	303.1551	333.1292	347.1449
EIC (Intens. x108)							•	
C. crispus (April)	16118542	707375	3254803	209911	5116637	26945	3129533	353130
C. crispus (July)	12600749	85700	928894	36714	3788544	18560	394887	11021
C. crispus (August)	16469850	219296	857212	238033	5653618	32998	1063642	83569
C. crispus (Sept.)	11230824	56477	2546286	77636	2917730	< LOD	5199737	33580
UV (mAU)							•	
C. crispus (April)	20420	31,525	1541	< LOD	6996	< LOD	2299	117
C. crispus (July)	14578	171,83	1487	< LOD	7106	327,43	335	< LOD
C. crispus (August)	19005	242,7	2143	< LOD	9927	707,57	1245	248
C. crispus (Sept.)	12768	< LOD	989	< LOD	6367	< LOD	5128	136

Table S4, related to Figures 2 and 4. MAAs composition in *Chondrus crispus* determined by LC-UV-HRMS. Extracted Ion Chromatogramm (EIC) of selected MAAs were obtained in positive mode; UV Absorbance was recorded at 330 nm (LOD = Limit Of Detection).

Name of source reaction	Metacyc ID	Molecular transformation	Biosynthesis pathway
rxn_4282	RXN-4282	delta24_25_reduction	Sterols
c24_c29_demethylation	RXN-20433, RXN20434, RXN20435	c24_c29_demethylation	Sterols
rxn_20436	RXN-20436	cyclopropylsterol isomerisation	Sterols
rxn_20438	RXN-20438	c14_demethylation	Sterols
rxn_20439	RXN-20439	c14_reduction	Sterols
rxn_4286	RXN-4286	c8_isomerisation	Sterols
c24_c28_demethylation	RXN-20440, RXN20441, RXN20442	c24_c28_demethylation	Sterols
rxn_1_14_21_6	1.14.21.6-RXN	c5_desaturation	Sterols
rxn66_323	RXN66-323	delta7reduction	Sterols
rxn66_28	RXN66-28	delta24_25_reduction	Sterols
rxn_4021	RXN-4021	c24_methylation	Sterols
rxn_2_1_1_143	2.1.1.143-RXN	c24'_methyltransfer	Sterols
rxn_20131	RXN-20131	delta24_24'_reduction	Sterols
rxn_4243	RXN-4243	c22_desaturation	Sterols
c22_desaturation	RXN-4242 or RXN-8352	c22_desaturation	Sterols
mysa	RXN-17372	cyclisation	MAA
rxn_17366	RXN-17366	methyl_transfer	MAA
rxn_17370	RXN-17370	non-enzymatic tautomerization	MAA
rxn_17896	RXN-17896	methyl_transfer	MAA
aminoacid_C_1_transfer	RXN-17368	aminoacid_c1_transfer	MAA
aminoacid_C_3_transfer_serine	RXN-17367	aminoacid_c3_transfer	MAA
aminoacid_C_3_transfer_threonine	-	aminoacid_c3_transfer	MAA
hydrogenation	-	hydrogenation	MAA
demethylation	-	demethylation	MAA
dehydration	-	dehydration	MAA
decarboxylation_1	-	decarboxylation	MAA
decarboxylation_2	-	decarboxylation	MAA
hydrolysis	-	hydrolysis	MAA

Table S5, related to Figure 3. List of molecular transformations inferred in Pathmodel and associated source reactions.

Steps	Yeast	Human	Arabidopsis	C. crispus
squalene monoxygenation	ERG1	SQLE	SQE1-7	scaffolds 90*, 20*, 57*
oxydosqualene cyclisation	ERG7	LSS	CAS	CHC_T00008265001
C-14 demethylation	ERG11	CYP51A1	CYP51G1	CYP51G1 (CHC_T00009303001)
C-14 reduction	ERG24	TM7SF2	FK	CHC_T00003466001
C-4 demethylation	ERG25	SC4MOL	SMO1, SMO2	CHC_T00010320001, scaffold212*
delta-8, delta-7 isomerisation	ERG2	EBP	HYD1	CHC_T00001257001
C-5 desaturation	ERG3	SC5DL	STE1	CHC_T00006481001
C24 or C24' methylation	ERG6	-	SMT1, SMT2	CHC_T00009101001, CHC_T00000837001
delta-7 reduction	-	DHCR7	DWF5	CHC_T00006492-3001*
delta-24 reduction	ERG4	DHCR24	DWF1/SSR	CHC_T00002789001
C-22 desaturation	ERG5	-	CYP710	CYP805A1-C1 or CYP808A1-H1
cyclopropylsterol isomerisation	-	-	CPI1	CHC_T00002985001

Table S6, related to Figure 5. Comparative Genomic Analysis of Sterol Synthesis Enzymes. Dark blue indicates orthologous sequences, light blue indicates paralogous ones, and green indicates yeast enzymes non orthologous to animal or plant sequences but known to perform the same enzymatic reaction. Five corrected sequences and new predictions are indicated with an asterisk (*) and provided in Dataset S1.

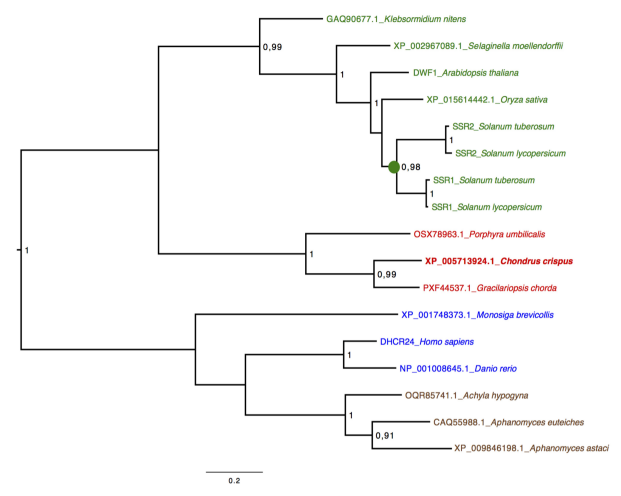


Figure S3, related to Figure 5. Maximum-likelihood tree of eukaryotic side-chain reductases. In green: protein sequences from green plants (streptophytes). The green dot indicates lineage-specific duplication in solanaceans. In red: protein sequences from red algae. In blue: protein sequences from opisthokonts (vertebrates + choanoflagellates). In brown: sequences from oomycete stramenopiles. Likelihood-ratio test values above 0.90 are indicated. Those above 0.97 are considered significant.

Dataset S1, related to Figure 5. New or edited protein sequences associated to the sterol synthesis pathway in *Chondrus crispus*.

>scaffold90:7511-6165(-) putative squalene epoxidase

RDGRRVLCVERQLYAPSGALCAPPRIVGELLQPGGYDALCRLGLADALLDIDAQVIRGYA

LFLGPRAERLPYHQPGGPDPDPDAARPQPEGRAFHNGRFLKRLREIARAHPNV

TLVEGNVLALLERDGAVVGVRYATRGNKAATAHAGLTIAC

DGCGSALRKRAAAHHHVTVYSNFHGLVLHVPALPFPNHGHVVLADPCPVLFYPISATEVR

CLVDIPSTYAGDAAEYILHTVVPQVPPPLRAPLATAVRERRSKMMPNRVMPAPA

HVVPGAVLLGDAFNMRHPLTGGGMTVALTDVELLRELLAPVPDLSDAPAVAAKLQLFYER

RKPMSTTINILANALYTLFCATDDPALRDMRAACLDYLAKGGRMTHDPIAMLGGLKPQRH

LLLAHFFAVALYGCGKALMPFPTPARLVRAWSIFRASFNIIKPLANAEGFWPLSWLPLNSL

>scaffold20:461442-460650(-) putative squalene epoxidase
LCRLGLADALLHIDAQVIRGYALFLGPRAERLPYHQPGEPDPDPAARPQPEG
RAFHNGRFLKRLREIARAHPNVTLIEGNVLALLERDGAVV
GVRYATRGNKAATAHAGLTIACDGCGSALRKRAAAHHHVTVYSNFHGLVLHVPALPFPNH
GHVVLAHPCPVLFYPISATEVRCLVDL
YILHTVVPQVPPSLRAPLATTVRERRSKMMPNRVMPAPAHVVPGAVLLGDAFNMRHPLTG
GGMTVALTDVELLRGLLAP

>scaffold57:152407-364140(+) putative squalene epoxidase RFAGPEHPSCGLKPQRHLLLAHFFAVALYGCGKALMPFPTPARPVRAWSIFRASFNFIK PLANAEGFWPLSWLPLN

LCRLGLADALLDIDAQVIRGYALFLGPRAERLPY

LCRLGLADALLHIDAQVIRGYALFLGPRAERLPYHQPGGPDPDPAARPQPEG

RAFHNCRFLKRLREIARAHPNVTLIEGNVLALLERDGAVV

GVRYATRGNKAATAHAGLTIACDGCGSALRKRAAAHHHVTVYSNFHGLVLHVPALPFPNH

GHVVLAHPCPVLFYPISATEVRCLVDL

WSTYAGDAAEYILHTVVPQVPPSLRAPLATAVRERRSKMMPNRVMPAPAHVVPGAVLLGD

AFNMRHPLTGGGMTVALTDVELLRGLLAP

>scaffold212:177405-176674(-) putative C-4 sterol methyl oxidase WDLLCRHTRAYPMFVVGCFASQLAGYFLGCAPFVLLDALRARSTPFRKIQPGKYAPRRAV FAAAAAMLRSFATVVLPLLAAGGLFIERVGISRDAPFPSPRVVLLQVAYFFLVEDFLNYW VHRALHLPWLYTRVHSVHHEYDAPFAVVAAYAHPVEVVLQALPTFAGPLMLGPHLYTLCV WQLFRNWEAIDIHSGYDHAWGLASVLPWYAGPEHHDFHHFLHSGNFASVFTWCDWAYGTD LAYE

>CHC_T00006492-3001 fusion of adjacent protein predictions CHC_T00006492001 and CHC_T00006493001; putative sterol delta-7 reductase

MLGIAAWKGFIRYGLLYDHFGEVLAFLGKFALVVTVLLYFRGIYFPTNSDSGTTSFGIVWDMWHGTELHP EIFGVSLKQLVNCRFALMGWSVAIVAFACKQREQYGYVSNSMLVSVVLQLVYIFKFFVWEAGYFNSVSLD

HSHVCLFWIYLRPLY

MVGVGAICCNYWTDKQREVFRATNGQVTIWGQKPVSIEAQYVTGDGKKRRSLLLASGWWGVSRHVNYVFE IALTFCWSVPAGGTGVIPYVYVMFLTILLTDRAYRDEVRCSEKYGKYYEEYCRLVPYKMIPGVY

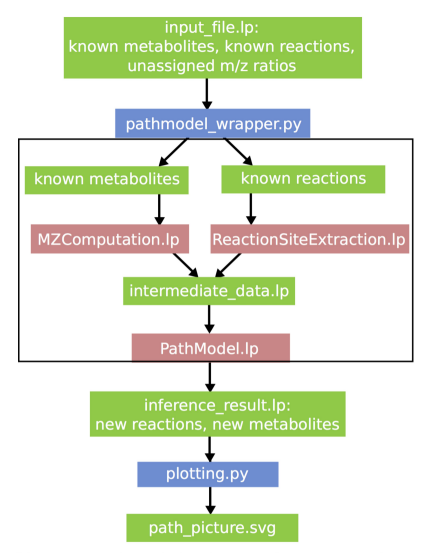


Figure S4, related to Figure 3. Architecture of Pathmodel scripts. In green: Data files, either input or result files. In red: ASP scripts. In blue: Python scripts. The black line shows the wrapping of all the scripts inside by pathmodel_wrapping.py.

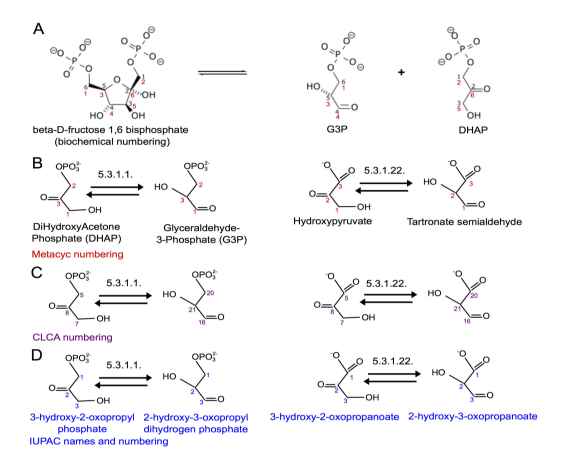


Figure S5, related to Figure 3. Comparisons of atom mapping using MetaCyc, CLCA and IUPAC. A. Biochemical origin of G3P and DHAP generates numbering inconsistencies that have to be solved by carbon atom renaming. B. Metacyc numbering does not label identically atoms from two reactions involving the same molecular transformation. C. CLCA numbering allows comparisons of reactions but not simultaneous atom mapping. D. IUPAC numbering allows both simultaneous atom mapping and comparisons of reactions.