The Metabolic Building Blocks of a Minimal Cell

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Simple Summary: Manufacturing artificial living cells would open endless research possibilities in basic and applied sciences. With this motivation, many research groups are developing methodologies to construct a stable minimal cell that is capable of achieving metabolic homeostasis, reproducing, and evolving in a controlled environment. Using as a template the gene set for a minimal cell proposed previously by Gil and coworkers, we have put together a network depicting its inferred minimal metabolism needed for life. This network has been further compressed as a metabolic Directed Acyclic Graph (m-DAG) in order to better visualize its topology and to find its essential reactions (i.e., critical reactions to maintain the network’s connectivity). We have also compared this minimal m-DAG to those of the smallest natural genome known until now and a synthetic minimal cell created in the laboratory. The modeling of m-DAGs based on minimal metabolisms can be a first approach for the synthesis and manipulation of minimal cells.

Abstract: Defining the essential gene components for a system to be considered alive is a crucial step toward the synthesis of artificial life. Fifteen years ago, Gil and coworkers proposed the core of a putative minimal bacterial genome, which would provide the capability to achieve metabolic homeostasis, reproduce, and evolve to a bacterium in an ideally controlled environment. They also proposed a simplified metabolic chart capable of providing energy and basic components for a minimal living cell. For this work, we have identified the components of the minimal metabolic network based on the aforementioned studies, associated them to the KEGG database and, by applying the MetaDAG methodology, determined its Metabolic Building Blocks (MBB) and reconstructed its metabolic Directed Acyclic Graph (m-DAG). The reaction graph of this metabolic network consists of 80 compounds and 98 reactions, while its m-DAG has 36 MBBs. Additionally, we identified 12 essential reactions in the m-DAG that are critical for maintaining the connectivity of this network. In a similar manner, we reconstructed the m-DAG of JCVI-syn3.0, which is an artificially designed and manufactured viable cell whose genome arose by minimizing the one from Mycoplasma mycoides JCVI-syn1.0, and of “Candidatus Nasua deltocephalinicola”, the bacteria with the smallest natural genome known to date. The comparison of the m-DAGs derived from a theoretical, an artificial, and a natural genome denote slightly different lifestyles, with a consistent core metabolism. The MetaDAG methodology we employ uses homogeneous descriptors and identifiers from the KEGG database, so that comparisons between bacterial strains are not only easy but also suitable for many research fields. The modeling of m-DAGs based on minimal metabolisms can be the first step for the synthesis and manipulation of minimal cells.

Keywords: metabolic networks; minimal gene set machinery; directed acyclic graphs; minimal cells
1. Introduction

One of the most ambitious aspirations of modern biology is to synthesize artificial living cells. Manufacturing a cell opens endless research possibilities, both in basic and advanced sciences, and it would be critical and a turning point in fields from medicine to evolutionary biology. To reduce the levels of difficulty on this task, most efforts are focused on the synthesis of minimal cells. On the one hand, they will help by increasing our understanding of living systems; on the other hand, they can be used as capsules for the introduction of genetic material to customize cells for applied purposes [1]. Several complementary paths have been followed in search of the proper technology and methods to design this fabricated cell. The most commonly used are the bottom–up and top–down approaches [2–4].

The bottom–up approach consists of the assembly, piece by piece, of each non-living biological component (i.e., a self-replicating nucleic acid, a metabolic machinery, and an encapsulating structure; [5]) in order to get a system that could be considered alive. The resulting products of this approach are called “protocells” [6,7]. No comparable system has been successfully constructed yet, but there have been developments on this front, with the designing of more refined cell-like compartments [8].

The top–down approach consists of deconstructing living cells [4,9]. Taking modern cells with reduced genomes as a starting point, it aims at further simplifying them by removing dispensable genetic material. Experimental (genome-wide analyses by massive transposon mutagenesis, antisense RNA, and systematic gene knockout) and computational approaches (including comparative genomics, comparative proteomics and in silico cell modeling) have been used to characterize a set of essential and sufficient genes to compose a living cell, that is, the core of a minimal bacterial genome [10]. Experimentally, genes are considered to be essential based on indirect evidence from systematic and genome-wide inactivation or the inhibition of each individual gene present in a genome (compiled in http://www.essentialgene.org/ [11]). Comparative genomics has also been broadly used, assuming that genes that are common between distant organisms are prone to be essential [12]. In addition, naturally reduced genomes from bacteria with a host-associated lifestyle have been used for comparisons regarding gene content, because they must be approaching a minimal genome [13,14]. The merging of these studies demonstrated the relevance of considering that essential functions can be performed by alternative and unrelated (non-orthologous) gene products. Comparative studies only retrieve genes involved in functions for which there is no alternative in nature (e.g., the complex translational machinery), while a minimal genome must also include all genes essential to maintain metabolic homeostasis [15].

There is a third approach for the construction of a minimal genome that searches for the biochemical and modular description of well-defined pathways needed to perform all essential functions [16]. Despite some major challenges needing to be addressed, this approach allows a function-by-function debugging to reach self-replication, and it suggests a good starting point for the ultimate synthesizing of a minimal genome able to sustain an artificial minimal cell. The potentiality of chemically synthesizing genomic segments or complete genomes and confining them into pre-existing cells has revolutionized the study of minimal cells [17]. The design of a truly minimal genome and its metabolic network can also benefit from computational whole-genome sequence rewriting and a design-build-test in silico approach, preceding the chemical synthesis of a customized genome [18].

A cohesive metabolic network proposal can lead the path to the synthesis of minimal cells. A minimal cell would depend on a minimal set of anabolic pathways to convert and assemble its biomolecule building blocks with the use of the energy and nutrients available in the environment, to reach metabolic homeostasis, and to achieve cellular growth and reproduction. Nevertheless, there is scientific consensus regarding the existence of a variety of minimal metabolic schemes that are ecologically dependent and able to sustain a universal genetic machinery [19]. The simplest cell should be chemoorganoheterotrophic (i.e., an organism using organic compounds as carbon and energy sources), living in
a nutrient-rich medium, in which the major metabolites (glucose, fatty acids, nitrogenous bases, amino acids, and vitamins) must be available without limitation, since this cell would not be able to synthesize them. Nevertheless, considering the adaptability of bacterial heterotrophic metabolisms, different metabolic schemes can be envisaged. The metabolic chart proposed by Gil and coworkers in 2004 [15] using a top–down approach, by performing a comprehensive analysis of all previous computational and experimental attempts to define a minimal genome, was based on the metabolic functions that were preserved in highly reduced genomes completely sequenced at that time, from endosymbiotic mutualistic or parasitic bacteria. The proposed core of the minimal genome encoded the costless pathways that would allow the cell to perform the selected metabolic functions. In order to maintain a coherent metabolic functionality, some pathways that were not present in some of the reduced genomes used in the aforementioned study were also incorporated, because their lack reflected a high dependence of their hosts. Likewise, the group of Craig Venter also explored this area and presented their list of essential genes for a minimal bacterium in 2006 [20]. Both sets of genes and the coherence of this metabolic network were further explored by Gabaldón et al. (2007) [19].

Metabolic networks determine the physiology and biochemistry of a cell. They are made of three components: the metabolic pathways, the chemical reactions involved in the metabolism, and the regulatory interactions of these reactions. Metabolic networks tend to be highly complex, even for simple organisms. For example, if we consider the metabolism of porphyrin and chlorophyll which is present in some animals, plants, fungi, bacteria, and archaea, we get a metabolic pathway map of 135 nodes and 181 edges in the reference pathway in the KEGG database (pathway: map00860). A pathway map with so many components is very difficult to visualize, especially when we are interested in the pathway topology. To this extent, it is highly advantageous to suitably reduce the number of nodes in order to visualize the network more precisely. Alberich and coworkers (2017) designed a methodology called MetaDAG [21], which consists of the contraction into a single node of those reactions that are strongly connected in the genome-wide reaction graphs. In this way, the resulting graph is a Directed Acyclic Graph (DAG), called a metabolic DAG (m-DAG), that preserves the network topology (i.e., the original relations between reactions) while it allows easy human exploration and visualization. One advantage of directed acyclic graphs is that they do not have cycles repeatedly producing and consuming the same metabolite. This methodology also creates reaction graphs and m-DAGs from multiple genomes, which can be used to calculate the core- and pan-metabolisms of a group of bacteria of interest as well as compare genomes by their m-DAGs in a novel manner. The MetaDAG methodology can also be of importance for large in silico analyses. By compressing metabolic networks and making them “simpler”, algorithms and computer analyses could also be less time consuming. Just as important, less computational resources would be needed, making it easier for researchers to work with a large number of genome wide m-DAGs, bacterial consortia m-DAGs, multiple symbiosis analyses, or even environmental metabolomics.

For the current work, we constructed the minimal metabolic network from the theoretical minimal gene set machinery revised in Gabaldón and coworkers (2007) [19], and compared it to the smallest genome of a live organism known to date [22], and to the genome of a semisynthetic bacteria produced by Craig Venter’s group in 2016 [17]. Despite the great efforts being done to homogenize gene and enzyme names in databases, due to how they have been discovered and described throughout history, some of their names are still associated with taxonomically related organisms. For this reason, to avoid any remaining biases toward any group of organisms and any need for synonym lists, we propose a minimal metabolic network defined by reactions and compounds instead of genes. Moreover, another of the advantages of our methodology is that it is essentially universal, since it uses homogenous identifiers and descriptors, so that researchers can easily associate the involved reactions and compounds to genes of bacterial genomes with different phylogenetic backgrounds, even to synthetic genomes as proven in this study.
Finally, it can also be applied to bacterial consortia in order to detect the metabolic interactions between partners and communities.

2. Materials and Methods

2.1. Inference of Minimal Metabolic Networks

The metabolic networks for this study were inferred from the reviewed version of the theoretical minimal genome described by Gabaldón et al. (2007) [19], the genome of “Ca. Nasuia deltocephalinicola” str. NAS-ALF [22] (which is also publicly available in the new version of the SymGenDB [23]), and the genome of JCVI-syn3.0, which is an artificial viable cell created by Hutchison and coworkers [17]. We first searched for all protein-coding genes in each genome for which an enzymatic activity has been assigned and then searched for the corresponding reactions in KEGG.

2.2. Reconstruction of the Directed Acyclic Graph of Metabolic Networks

Using the above obtained information, which is a set of reactions for each metabolic network, we generated the corresponding reaction graph that models the relationship between reactions in terms of shared metabolites. A reaction graph, denoted by $RG = (R, E_R)$, is a directed graph with a set of nodes $R$ that are reactions and whose edges are defined as follows: there is an edge pointing from reaction $R_i$ to reaction $R_j$ if, and only if, a metabolite produced by reaction $R_i$ is a substrate in reaction $R_j$. The fact that it is a directed graph establishes a natural production/consumption order between two reactions—that is, what is produced by $R_i$ is then consumed by $R_j$. Before generating the directed graph, we manually curated it to remove redundancies (enzymes encoded by orthologous genes).

In order to analyze the reaction graph in a visually friendly manner, we used the MetaDAG methodology [21]. In a reaction graph, two reactions $R_i$, $R_j$ are said to be biconnected if there is a path in each direction between them. A strongly connected component of a reaction graph is a subgraph such that every pair of reactions in it are biconnected. These strongly connected components are contracted in a single node. The reactions that are not biconnected to any other reaction become a node by themselves. Each node is called a Metabolic Building Block (MBB for short), and the MetaDAG software automatically assigns an ID to each MBB. When each MBB is contracted to a single vertex, the resulting quotient graph is a metabolic Directed Acyclic Graph (m-DAG for short). Thus, the m-DAG is defined as follows: its nodes are the MBBs obtained from the reaction graph, and there is an edge between two MBBs, $MBB_1$ and $MBB_2$, if there is an edge in the reaction graph from a reaction in $MBB_1$ to a reaction in $MBB_2$. We denote by $Gm$ the m-DAG, thus $Gm = (N, E)$ where $N$ is the set of MBBs and $E$ is the edges between them such that

$$(MBB_1, MBB_2) \in E \iff \exists R_i \in MBB_1 \land \exists R_j \in MBB_2 \land (R_i, R_j) \in E_R$$

MBBs contracting only one reaction and whose removal disconnects the reaction graph are considered essential reactions because they are crucial to maintain the network’s connectivity.

3. Results and Discussion

3.1. Theoretical Minimal Metabolic Network

The first step toward the creation of the minimal metabolic network was to extrapolate the list of genes and enzymes belonging to the set presented by Gabaldón and coworkers (2007) [19] (Figure 1 and Table S1) to obtain KEGG reaction identifiers (IDs). We used the complete reaction, compound, and enzyme database from KEGG and created the reaction graph by joining the reactions where metabolites were shared (see Section 2.2 for the complete explanation). The idea behind using the complete KEGG catalog is to avoid biases toward a specific phylogenetic group of bacteria.
Figure 1. Interaction graph of the proposed theoretical minimal metabolic network adapted from Gabaldón et al. (2007) [19]. Line colors denote metabolic categories: yellow, glycolysis; orange, pentose phosphate pathway; pink, phospholipid metabolism; green, nucleotide metabolism; blue, coenzyme metabolism. The two glycolytic steps in which ATP is produced by substrate-level phosphorylation are depicted with thicker red arrows, and correspond to reactions R01512 and R00200 in Table 1. The reaction graph of this same network is presented in Figure 2 for comparison.
Table 1. Reactions, enzymes, and compounds of the minimal metabolic network presented in Figure 2. Reversible reactions are denoted by the superscript r. MBB IDs are the identification numbers of the metabolic building blocks to which each reaction is contracted into, according to the MetaDAG analysis (Figure 3).

| Substrate KEGG ID | Reaction ID | Enzyme Name (E.C. Number) | Definition | Product KEGG ID | MBB ID |
|-------------------|-------------|---------------------------|------------|----------------|--------|
| C00020            | R00127      | adenylate kinase (2.7.4.3) | ATP + AMP ↔ 2 ADP | C00008        | 0.15   |
| C00882            | R00130      | dephospho-CoA kinase (2.7.1.24) | ATP + Dephospho-CoA → ADP + CoA | C00010        | 0.2    |
| C00455            | R00137      | nicotinamide-nucleotide adenylyltransferase (2.7.7.1) | Diphosphate + NAD+ ↔ ATP + Nicotinamide d-ribonucleotide | C00003        | 0.80.1.0 |
| C00015            | R00156      | nucleoside-diphosphate kinase (2.7.4.6) | ATP + UDP ↔ ADP + UTP | C00075        | 0.77.4.0 |
| C00105            | R00158      | UMP/CMP kinase (2.7.4.14) | ATP + UMP ↔ ADP + UDP | C00015        | 0.77.4.0 |
| C00061            | R00161      | FAD synthase (2.7.7.2) | ATP + FMN → Diphosphate + FAD | C00016        | 0.10   |
| C00018            | R00173      | pyridoxal phosphatase (3.1.3.74) | Pyridoxal phosphate + H₂O → Pyridoxal + Orthophosphate | C00250        | 0.11   |
| C00073            | R00177      | methionine adenosyltransferase (2.5.1.6) | ATP + L-Methionine + H₂O → Orthophosphate + Diphosphate + S-Adenosyl-L-methionine | C00019        | 0.12   |
| C00020 + C0013    | R00190      | adenine phosphoribosyltransferase (2.4.2.7) | AMP + Diphosphate ↔ Adenine + 5-Phospho-alpha-D-ribose 1-diphosphate | C00147 + C00119 | 0.78.1.0 |
| C00074 + C00008   | R00200      | pyruvate kinase (2.7.1.40) | ADP + Phosphoenolpyruvate → ATP + Pyruvate | C00022        | 0.9    |
| C00144            | R00332      | guanylate kinase (2.7.4.8) | ATP + GMP ↔ ADP + GDP | C00035        | 0.77.4.0 |
| C00044            | R00430      | pyruvate kinase (2.7.1.40) | GTP + Pyruvate ↔ GDP + Phosphoenolpyruvate | C00035        | 0.77.4.0 |
| C00055            | R00512      | (d)CMP kinase (2.7.4.25) | ATP + CDP ↔ ADP + GDP | C00112        | 0.77.4.2 |
| C00255            | R00549      | riboflavin kinase (2.7.1.26) | ATP + Riboflavin → ADP + FMN | C00061        | 0.13   |
| C00112            | R00570      | nucleoside diphosphate kinase (2.7.4.6) | ATP + CDP ↔ ADP + CTP | C00063        | 0.77.4.2 |
| C00075            | R00571, R00573 | CTP synthase (6.3.4.2) | ATP + UTP + Ammonia → ADP + Orthophosphate + CTP | C00063        | 0.77.4.6 |
| C00378            | R00619      | thiamine diphosphokinase (2.7.6.2) | ATP + Thiamine → AMP + Thiamin diphosphate | C00068        | 0.14   |
| C00631            | R00658      | enolase (4.2.1.11) | 2-Phospho-D-glycerate ↔ Phosphoenolpyruvate + H₂O | C00074        | 0.77.4.0 |
Table 1. Cont.

| Substrate KEGG ID | Reaction ID | Enzyme Name (E.C. Number) | Definition | Product KEGG ID | MBB ID |
|-------------------|-------------|---------------------------|------------|----------------|--------|
| C00186            | R00703      | lactate dehydrogenase (1.1.1.27) | (S)-Lactate + NAD+ ↔ Pyruvate + NADH + H+ | C0022 | 0.0 |
| C00093            | R00842 | sn-glycerol-3-phosphate dehydrogenase (1.1.1.94) | sn-Glycerol 3-phosphate + NAD+ ↔ Glycerone phosphate + NADH + H+ | C00111 | 0.77.4.0 |
| C00093 + C00040   | R00851 | arylamine N-acetyltransferase (2.3.1.15) | sn-Glycerol 3-phosphate + Acyl-CoA → 1-Acyl-sn-glycerol 3-phosphate + CoA | C00681 | 0.8 |
| C00415            | R00936 | dihydrofolate reductase (1.5.1.3) | Dihydrofolate + NADH + H+ ↔ Tetrahydrofolate + NAD+ | C00101 | 0.79.0 |
| C00037 + C00143   | R00945 | glycine hydroxymethyltransferase (2.1.2.1) | 5,10-Methylenetetrahydrofolate + Glycine + H2O ↔ Tetrahydrofolate + L-Serine | C00065 + C00101 | 0.79.0 |
| C00105            | R00966 | uracil phosphoribosyltransferase (2.4.2.9) | UMP + Diphosphate ↔ Uracil + 5-Phospho-alpha-D-ribose 1-diphosphate | C00106 + C00119 | 0.77.4.0 |
| C00117            | R01049 | phosphoribosylpyrophosphate synthetase (2.7.6.1) | ATP + d-Ribose 5-phosphate ↔ AMP + 5-Phospho-alpha-D-ribose 1-diphosphate | C00119 | 0.77.4.0 |
| C00117            | R01056 | ribose-5-phosphate isomerase (5.3.1.6) | d-Ribose 5-phosphate ↔ d-Ribulose 5-phosphate | C00199 | 0.77.4.0 |
| C00118            | R01061 | glyceraldehyde-3-phosphate dehydrogenase (1.2.1.12) | D-Glyceraldehyde 3-phosphate + Orthophosphate + NAD+ ↔ 3-Phospho-D-glyceroyl phosphate + NADH + H+ | C00236 | 0.77.4.0 |
| C05378            | R01070 | fructose-1,6-bisphosphate aldolase (4.1.2.13) | beta-D-Fructose 1,6-bisphosphate ↔ Glycerone phosphate + d-Glyceraldehyde 3-phosphate | C00111 + C00118 | 0.77.4.0 |
| C00131            | R01138 | pyruvate kinase (2.7.1.40) | dATP + Pyruvate ↔ dADP + Phosphoenolpyruvate | C00206 | 0.78.1.1 |
| C00119 + C00242   | R01229 | hypoxanthine phosphoribosyltransferase (2.4.2.8) | Guanine + 5-Phospho-alpha-D-ribose 1-diphosphate ↔ GMP + Diphosphate | C00144 | 0.77.4.0 |
| C00361            | R01858 | pyruvate kinase (2.7.1.40) | dGDP + Phosphoenolpyruvate → dGTP + Pyruvate | C00286 | 0.6 |
| C00008            | R02017 | ribonucleoside diphosphate reductase (1.17.4.1) | Thioredoxin + ADP → dADP + Thioredoxin disulfide + H2O | C00206 | 0.78.1.2 |
| C00035            | R02019 | ribonucleoside diphosphate reductase (1.17.4.1) | GDP + Thioredoxin → dGDP + Thioredoxin disulfide + H2O | C00361 | 0.77.4.7.0 |
| Substrate KEGG ID | Reaction ID | Enzyme Name (E.C. Number) | Definition | Product KEGG ID | MBB ID |
|-------------------|-------------|---------------------------|------------|----------------|--------|
| C00112 | R02024 | ribonucleoside diphosphate reductase (1.17.4.1) | Thioredoxin + CDP → dCDP + Thioredoxin disulfide + \( \text{H}_2\text{O} \) | C00705 | 0.77.4.5 |
| C00197 | R01512 | phosphoglycerate kinase (2.7.2.3) | ATP + 3-Phospho-D-glycerate ↔ ADP + 3-Phospho-D-glyceroyl phosphate | C00236 | 0.77.4.0 |
| C00631 | R01518 | phosphoglycerate mutase (2.3-diphosphoglycerate-independent) (5.4.2.12) | 2-Phospho-D-glycerate ↔ 3-Phospho-D-glycerate | C00197 | 0.77.4.0 |
| C00199 | R01529 | ribulose-phosphate 3-epimerase (5.1.3.1) | d-Ribulose 5-phosphate ↔ d-Xyulose 5-phosphate | C00231 | 0.77.4.0 |
| C00118 + C05382 | R01641 | transketolase (2.2.1.1) | Sedoheptulose 7-phosphate + D-Glyceraldehyde 3-phosphate ↔ d-Ribose 5-phosphate + d-Xyulose 5-phosphate | C00117 + C00231 | 0.77.4.0 |
| C00630 + C00416 | R01799 | phosphatidate cytidyltransferase (2.7.7.41) | CTP + Phosphatidate → Diphasphate + CDP-diacylglycerol | C00269 | 0.5 |
| C00065 + C00269 | R01800 | phosphatidylserine synthase (2.7.8.8) | Diacylglycerol + L-Serine → CMP + Phosphatidylserine | C00055 + C02737 | 0.4 |
| C00279 + C00111 | R01829 | fructose-1,6-bisphosphate aldolase (4.1.2.13) | Glycerone phosphate + D-Erythrose 4-phosphate → Sedoheptulose 1,7-bisphosphate | C00447 | 0.77.4.0 |
| C00118 + C05345 | R01830 | transketolase (2.2.1.1) | beta-D-Fructose 6-phosphate + D-Glyceraldehyde 3-phosphate ↔ D-Erythrose 4-phosphate + D-Xyulose 5-phosphate | C00231 + C00279 | 0.77.4.0 |
| C00363 | R02093 | nucleoside diphosphate kinase (2.7.4.6) | ATP + dTDP ↔ ADP + dTTP | C00459 | 0.81.0 |
| C00364 | R02094 | thymidine monophosphate kinase (2.7.4.9) | ATP + dTMP ↔ ADP + dTDP | C00363 | 0.81.0 |
| C00365 | R02098 | thymidine monophosphate kinase (2.7.4.9) | ATP + dUMP ↔ ADP + dUDP | C01346 | 0.77.4.1 |
| C00143 + C00365 | R02101 | thymidylate synthase (2.1.1.45) | dUMP + 5,10-Methylenetetrahydrofolate → Dihydrofolate + dTMP | C00364 + C00415 | 0.79.0 |
| C00040 + C00681 | R02241 | 1-acyl-sn-glycerol-3-phosphate acyltransferase (2.3.1.51) | 1-Acyl-sn-glycerol 3-phosphate + Acyl-CoA → Phosphatidate + CoA | C00416 | 0.7 |
| C00458 | R02325 | dCTP deaminase (3.5.4.13) | dCTP + H_2O → dUTP + Ammonia | C00460 | 0.77.4.4 |
Table 1. Cont.

| Substrate KEGG ID | Reaction ID | Enzyme Name (E.C. Number) | Definition | Product KEGG ID | MBB ID |
|-------------------|-------------|---------------------------|------------|----------------|--------|
| C00705            | R02326      | nucleoside diphosphate kinase (2.7.4.6) | ATP + dCDP ↔ ADP + dCTP | C00458 | 0.77.4.3 |
| C01346            | R02331      | nucleoside diphosphate kinase (2.7.4.6) | ATP + dUDP ↔ ADP + dUTP | C00460 | 0.77.4.1 |
| C02737            | R02055      | phosphatidylserine decarboxylase (4.1.1.65) | Phosphatidylserine → Phosphatidylethanolamine + CO₂ | C00350 | 0.3 |
| C00504            | R02235 | dihydrofolate reductase (1.5.1.3) | Folate + NADH + H⁺ ↔ Dihydrofolate + NAD⁺ | C00415 | 0.79.0 |
| C03150            | R02324      | ribosynicotinamide kinase (2.7.1.22) | ATP + Nicotinamide-beta-riboside → ADP + Nicotinamide D-ribonucleotide | C00455 | 0.80.0 |
| C00031            | R02738      | protein-Npi-phosphohistidine—D-glucose phosphotransferase (2.7.1.199) | Protein N(pi)-phospho-L-histidine + D-Glucose → Protein histidine + alpha-D-Glucose 6-phosphate | C00668 | 0.15 |
| C00668            | R02740      | glucose-6-phosphate isomerase (5.3.1.9) | alpha-D-Glucose 6-phosphate ↔ beta-D-Fructose 6-phosphate | C05345 | 0.77.4.0 |
| C00831            | R02971      | pantetheine kinase (2.7.1.34) | ATP + Pantetheine → ADP + Pantetheine 4’-phosphate | C01134 | 0.16 |
| C00864            | R03018      | pantothenate kinase (2.7.1.33) | ATP + Pantothenate → ADP + D-4’-Phosphopantothenate | C03492 | 0.19 |
| C01134            | R03035      | pantetheine-phosphate adenylyltransferase (2.7.7.3) | ATP + Pantetheine 4’-phosphate ↔ Diphosphate + Dephospho-CoA | C00882 | 0.1 |
| C03492            | R04231      | phosphopantothenoylcysteine synthetase (6.3.2.5) | CTP + D-4’-Phosphopantetheine + L-Cysteine ↔ CMP + Diphosphate + (R)-4’-Phosphopantothenoyl-L-cysteine | C04352 | 0.18 |
| C04079            | R04391      | pantothenate kinase (2.7.1.33) | ATP + N-((R)-Pantothenoyl)-L-cysteine ↔ ADP + (R)-4’-Phosphopantothenoyl-L-cysteine | C04352 | 3415 |
| C05345            | R04779      | 6-phosphofructokinase (2.7.1.11) | ATP + beta-D-Fructose 6-phosphate ↔ ADP + beta-D-Fructose 1,6-bisphosphate | C05378 | 0.77.4.0 |
| C04352            | R03269      | phosphopantothenoylcysteine decarboxylase (4.1.1.36) | (R)-4’-Phosphopantothenoyl-L-cysteine → Pantetheine 4’-phosphate | C01134 | 0.17 |
| C05382            | R01843      | 6-phosphofructokinase (2.7.1.11) | ATP + Sedoheptulose 7-phosphate ↔ ADP + Sedoheptulose 1,7-bisphosphate | C00447 | 0.77.4.0 |
Figure 2. The reaction graph of the proposed theoretical minimal metabolic network represented in Figure 1, obtained using data from the KEGG database. The yellow filled circles are the reactions with their KEGG ID and E.C. numbers, and the purple filled circles are the reverse reaction of the yellow filled circles, when appropriate. Line colors denote metabolic categories. A full-size representation can be seen as Figure S1.
This methodology gave us a resulting reaction graph with some redundancies (i.e., different enzymes encoded by orthologous genes participating in the same metabolic pathways), so we manually curated this graph to include only one copy of each reaction and their corresponding metabolites needed for a functional cell. The reaction graph obtained is composed of 98 reactions and 80 metabolites (Figure 2). The fact that our model replicates almost entirely the figure of Gabaldón et al. (2007) [19] (Figure 1), validates our methodology. Table 1 presents the complete list of reactions, substrate, and product compounds as well as their KEGG identifiers used to reconstruct the minimal metabolic network.
3.2. The MetaDAG Methodology: Analysis of the Composition and Connectivity of a Network at a Glance

Despite the fact that the reaction graph of the theoretical minimal organism constructed in this work has only 98 reactions and 80 metabolites, it is difficult to visualize the detailed relationships between the reactions that make up the network’s connectivity (Figure 2). To solve this problem, we used the MetaDAG methodology [21] to generate an m-DAG of the manually curated reaction graph. An m-DAG is a suitable reduction of a metabolic network. Namely, the reactions that are connected by multiple paths, which are the strongly connected components of the metabolic network, are contracted into one single MBB, which can be considered a robust subgraph in the reaction graph. Moreover, those MBBs that only represent a reaction that is not biconnected to any other reaction are essential to maintain the network connectivity. In this sense, the m-DAG provides a modularity of the reaction graph that keeps the information of robustness and connectivity of the metabolic network.

The m-DAG we obtained from the minimal metabolic reaction graph (Table 1, Figure 3) has a total of 36 nodes, 25 of them corresponding to single reactions (yellow nodes) and 11 to contracted MBBs (gray nodes). Clearly, there are seven connected components in this network, the biggest one covering the central metabolism of the hypothetical minimal organism, while the rest are the reactions that synthesize the essential cofactors needed for the proper functionality of the complete cell.

In addition, essential reactions (i.e., those whose removal reduces the network’s connectivity increasing the number of connected components) can be easily identified using this approach (hexagons with double lines in Figure 3). Table 2 is a list of the 12 essential reactions we found in the minimal metabolic network under study and the metabolic pathways where they participate. They are involved in purine and pyrimidine metabolism, glycerophospholipid metabolism, glycolysis and pantothenate, and CoA biosynthesis. Purines and pyrimidines are the most abundant metabolic substrates for all living organisms. They are essential components for the synthesis of DNA and RNA, and they also participate in the biosynthesis of energy nucleotides and are vital cofactors for cell survival and reproduction. Hence, purines and their by-products widely participate in biological processes. Glycerophospholipids are pivotal structural components of the cell membranes, but they are also precursors of many essential biological molecules and participate in cell signaling and other cellular processes [24]. Glycolysis is the first step in the breakdown of glucose to extract energy for cellular metabolism by creating high-energy molecules. It is considered an ancient metabolic pathway [25], and its prevalence in organisms is nearly ubiquitous.

### Table 2. Essential reactions of the m-DAG constructed from the theoretical minimal gene set machinery needed for life.

| Reaction ID       | Metabolic Pathway                                      |
|-------------------|--------------------------------------------------------|
| R02019            | Purine metabolism                                     |
| R02017            |                                                        |
| R00571/R00573     | Pyrimidine metabolism                                 |
| R02024            |                                                        |
| R02325            |                                                        |
| R00851/R02241     | Glycerolipid metabolism, Glycerophospholipid metabolism|
| R01799/R01800     | Glycerophospholipid metabolism                         |
| R00200            | Glycolysis, part of the pyruvate metabolism             |
| R04231/R03269     | Pantothenate and CoA biosynthesis                      |


We consider that what we call “essential reactions”, easily highlighted by the MetaDAG methodology, can be of crucial importance in many fields of research. Probably, the most logical and of vital importance is the idea that these reactions can help choose enzymes as potential drug targets, since the removal of these reactions breaks metabolic pathways, which can lead to the unviability of a cell. Considering that m-DAGs take into account complete genomes, and even complementary genomes (they can be calculated for two or more genomes together, to simulate complementary metabolic pathways within consortia), the resulting essential reactions are trustworthy in a sense that researchers might overlook an enzyme doing the same job as the one highlighted and, if they find it, it would be a new discovery not previously described for a specific metabolic pathway.

3.3. The m-DAG of “Candidatus Nasuia Deltocephalinicola”

In the case of a minimal metabolic network, each item included in the list of reactions and compounds is hypothetically essential for survival. When we extrapolate these results to living organisms possessing natural minimized genomes, such as pathogens or mutualist endosymbiotic bacteria, we should consider that their metabolism is a patchwork dependent on the host and, in many cases, also dependent on other bacteria with which they live in consortia. Therefore, the study of their networks’ connectivity has the potential of pointing out genes encoding critical steps that connect the different partners in a given pathway. Subsequently, the genes that encode those reactions can become targets for genetic engineering, and/or for mechanisms intended to regulate the cell metabolism; additionally, they might also have the potential to destroy the stability of the relationship, even killing the undesired organism in a parasitic relationship.

In order to compare the in silico minimal m-DAG with the m-DAG from a living organism with a naturally reduced genome, we constructed the m-DAG of “Ca. Nasuia deltocephalinicola” str. NAS-ALF (from now on referred to as Nasuia for simplicity; Supplementary Figure S2), one of the obligate endosymbiotic bacteria of the aster leafhopper Macrosteles quadrilineatus [22]. This endosymbiont possesses the smallest natural genome known so far, comprising 112,091 bp and only 138 protein-coding genes identified. The metabolic data needed to generate this m-DAG, including the complete list of its enzymes, reactions, and compounds were also obtained from the KEGG database (Table S2). Nasuia’s m-DAG comprises 29 nodes included in 12 connected components, with 7 MBBs and 22 single reactions. Regarding the single reactions, five are essential (summarized in Table 3).

Table 3. Essential reactions of the m-DAG of “Ca. Nasuia deltocephalinicola” str. NAS-ALF.

| Reaction ID | Metabolic Pathway                                                  |
|------------|--------------------------------------------------------------------|
| R09372     | Selenocompound metabolism                                         |
| R00443     | Purine metabolism, Glycerophospholipid metabolism                  |
| R03012     | Histidine metabolism                                              |
| R01163     | Histidine metabolism                                              |
| R01288     | Cysteine and methionine metabolism, Sulfur metabolism             |

It has been estimated that more than 60% of insects possess symbiotic bacteria inside their body tissues, and/or very often in a specialized cell type called bacteriocyte [26]. When these bacteria become endosymbionts, they lose their ability to interact with other organisms. Additionally, they become dependent on their respective hosts, and their genome is significantly reduced by the deletion of genes that become redundant or that are not needed in a rich environment such as the one they encounter within their hosts [15,27]. In addition, even though the niche is significantly rich for them, the insect host generally has a very incomplete diet by feeding on plant sap or seeds, or blood from mammals, so the bacteria become their helpers for the production of essential amino acids, fatty acids, or vitamins [28,29]. The essential reactions of Nasuia’s m-DAG reveal exactly that. This organism works as a factory of the vitamins and amino acids that M. quadrilineatus
needs to survive. Moreover, this bacterium is part of a consortium with “Candidatus Sulcia muelleri” str. ALF [22]. It is widely accepted that the endosymbiotic relationship between insects and bacteria, dating from 10 to several hundred millions of years, allowed the proliferation of insects and their diversification in almost any ecological niche [30,31]. Obviously, if the reactions that link the metabolic routes disappear (either naturally or due to targeted modification of those genes), this association would be affected to the point of the possible death of the host.

A direct comparison between the reactions and compounds that make up the in silico m-DAGs of the theoretical minimal cell and Nasuia would not be significant due to their dissimilar lifestyles. What we can easily assess is the topology of the networks. At first glance, it is striking that the smallest genome found in nature has fewer nodes than the in silico m-DAG. The dependence of this endosymbiotic bacteria to its host and to its second co-obligate endosymbiont explains this phenomenon.

3.4. The First Semisynthetic Viable Cell and Its m-DAG’s Reconstruction

To complete our comparative analysis, we constructed the m-DAG of JCVI-syn3.0, which is an artificially designed and manufactured viable cell whose genome arose by minimizing the one from Mycoplasma mycoides JCVI-syn1.0 created by Hutchison et al. in 2016 [17]. To do so, we used the list of enzymes presented in their article and converted it into a list of reactions and compounds, compared them to our minimal metabolic network (Table S3), and created the reaction graph of JCVI-syn3.0 and its eventual m-DAG (Figure 4).

JCVI-syn3.0 m-DAG is formed by 34 connected components, with a total of 70 nodes, 54 of them corresponding to single reactions, and 16 contracted MBBs. Ten reactions are essential (summarized in Table 4), that is, indispensable to maintain the connectivity of the network.

Table 4. Essential reactions of the m-DAG of JCVI-syn3.0.

| Reaction ID | Metabolic Pathway                        |
|-------------|----------------------------------------|
| R02024      | Pyrimidine metabolism                  |
| R01663      |                                        |
| R02059      | Amino sugar and nucleotide sugar metabolism |
| R00765      |                                        |
| R00200      | Glycolysis, part of the pyruvate metabolism |
| R00189      | Nicotinate and nicotinamide metabolism  |
| R03346      |                                        |
| R01799      | Glycerophospholipid metabolism          |
| R01801      |                                        |
| R02239      |                                        |

Once again, the essential reactions are involved in the metabolism of nucleotides, phospholipids, and coenzymes, even though there are significant differences between the list of reactions included in the reconstruction of JCVI-syn3.0 and the metabolic minimal network (Supplementary Table S3). JCVI-syn3.0 has 155 reactions included in its reaction graph, while our minimal network reaction graph has only 63 (98 when taking reverse reactions into account). The explanation for these differences is that the minimal network defined by Gil and coworkers (2004) [15] considers the minimal bacterium to live in a controlled and nutrient-rich environment, while JCVI-syn3.0 includes some metabolic pathways that are essential for the specific necessities of M. mycoides, its reproduction, and its survival. Interestingly enough, two reactions are essential for both networks (R02024 and R00200), while others participate closely in the same pathways (e.g., R01800 and R01801), which may be useful information for genetic engineering purposes.
Figure 4. m-DAG of the metabolism of JCVI-syn3.0. Single reactions appear in yellow, contracted MBBs in grey, and the essential reactions as hexagons with double lines.

3.5. Resemblance of the MBBs of the Minimal m-DAGs

In order to contrast the MBBs of the three m-DAGs constructed in this study, Table 5 shows the correspondence among them. The list of enzymes and the definition of each reaction is presented in Supplementary Table S4.
Table 5. Comparison of the MBBs of the three networks under study. Every row lists the reactions belonging to the corresponding MBB and the enzymes involved in those reactions. The list includes only MBBs composed of at least three reactions (reverse included) or with fewer reactions but that are shared by at least two of the networks under study.

| Model Cell | MBB ID | # Reactions | Reaction ID |
|------------|--------|-------------|-------------|
| Minimal cell | 0.77.4.0 | 21 | R00156 (2.7.4.6), R00158 (2.7.4.22), R00332 (2.7.4.22), R00430 (2.7.1.40), R00658 (4.2.1.11), R00842 | R00844 (1.1.1.94), R00966 (2.4.2.9), R01049 (2.7.6.1), R01056 (5.3.1.6), R01061 (1.2.1.12), R01070 (4.1.2.13), R01229 (2.4.2.8), R01512 (2.7.2.3), R01518 (5.4.2.12), R01529 (5.1.3.1), R01641 (2.2.1.1), R01829 (4.1.2.13), R01830 (2.2.1.1), R01843 (2.7.1.11), R02740 (5.3.1.9), R04779 (2.7.1.11) |
| | 0.77.4.1 | 2 | R02098 (2.7.4.9), R02331 (2.7.4.6) |
| | 0.77.4.2 | 2 | R00512 (2.7.4.25), R00570 (2.7.4.6) |
| | 0.77.4.7.0 | 1 | R02019 (1.17.4.1) |
| | 0.78.1.0 | 2 | R01127 (2.7.4.3), R0190 (2.4.2.7) |
| | 0.78.1.1 | 1 | R01138 (2.7.1.40) |
| | 0.78.1.2 | 1 | R01207 (1.17.4.1) |
| | 0.79.0 | 4 | R00936, R0099 (1.5.1.3), R00945 (2.1.2.1), R01641 (2.7.1.11), R02101 (2.1.1.45), R02235, R02236 (1.5.1.3) |
| | 0.80.1.0 | 1 | R01537 (2.7.7.18) |
| | 0.81.0 | 2 | R02093 (2.7.4.6), R02094 (2.7.4.9) |
| JCVI-syn 3.0 | 0.39 | 6 | R00014 (2.1.4.1), R00230 (2.3.1.8), R00315 (2.7.2.1), R02569 (2.3.1.12), R03270 (1.2.4.1), R07618 (1.8.1.4) |
| | 0.40 | 3 | R01126 (3.1.3.5), R01132 (2.4.2.8), R01833 (2.4.2.1) |
| | 0.41 | 3 | R02142 (2.4.2.8), R02297 (2.4.2.1), R02719 (3.1.3.5) |
| | 0.42 | 2 | R00921 (2.3.1.8), R01353 (2.7.2.1) |
| | 0.77.4.2.0 | 1 | R00512 (2.7.4.25) |
| | 0.77.4.7 | 49 | R00158 (2.7.4.22), R00289 (2.7.7.9), R00291 (5.1.3.2), R00332 (2.7.4.8), R00430 (2.7.1.40), R00505 (5.4.99.9), R00658 (4.2.1.11), R00959 (5.4.2.5), R00966 (2.4.2.9), R01015 (5.3.1.1), R01049 (2.7.6.1), R01056 (5.3.1.6), R01057 (5.4.2.7), R01058 (1.2.1.19), R01061 (1.2.1.12), R01066 (4.1.2.4), R01067 (2.2.1.1), R01068 (4.1.2.13), R01070 (4.1.2.13), R01227 (3.1.3.5), R01229 (2.4.2.7), R01229 (2.4.2.8), R01512 (2.7.2.3), R01518 (5.4.2.12), R01529 (5.1.3.1), R01641 (2.2.1.1), R01819 (5.3.1.8), R01827 (2.2.1.2), R01829 (4.1.2.13), R01830 (2.2.1.1), R01843 (2.7.1.11), R01967 (2.7.1.113), R01968 (3.1.3.5), R01969 (2.4.2.1), R02018 (1.17.4.1), R02019 (1.17.4.1), R02090 (2.7.4.8), R02098 (2.7.4.9), R02099 (2.7.1.21), R02102 (3.1.3.5), R02102 (3.1.3.89), R02147 (2.4.2.1), R02568 (4.1.2.13), R02739 (5.3.1.9), R02740 (5.3.1.9), R02749 (5.4.2.7), R03321 (5.3.1.9), R04779 (2.7.1.11) |
| | 0.77.4.8 | 5 | R01664 (3.1.3.5), R01664 (3.1.3.89), R01665 (2.7.4.25), R01666 (2.7.1.74), R01667 (3.6.1.12) |
| | 0.78.1 | 12 | R00127 (2.7.4.3), R00183 (3.1.3.5), R00185 (2.7.1.74), R00190 (2.4.2.7), R01138 (2.7.1.40), R01547 (2.4.2.1), R01547 (2.4.2.11), R02017 (1.17.4.1), R02088 (3.1.3.5), R02089 (2.7.1.76), R02557 (2.4.2.1) |
| | 0.79.1 | 6 | R00942 (6.3.2.17), R00945 (2.1.2.1), R01220 (1.5.1.5), R01655 (5.3.4.9), R03940 (2.1.2.9), R04241 (6.3.2.17) |
| | 0.80.1 | 4 | R0137 (2.7.7.18), R01271 (2.4.2.12), R02294 (2.4.2.1), R02323 (3.1.3.5) |
| | 0.81.1 | 4 | R01567 (2.7.1.21), R01569 (3.1.3.5), R01569 (3.1.3.89), R02094 (2.7.4.9) |
| Nasuia | 0.77.0 | 2 | R00435 (2.7.7.6), R00441 (2.7.7.6) |
| | 0.78.0 | 2 | R00375 (2.7.7.7), R00376 (2.7.7.7) |

Reversible reactions are denoted by the superscript r. Reactions depicted in blue are shared by the minimal and the synthetic (JCVI-syn3.0) metabolic networks. #: number of.
4. Conclusions

The construction of the minimal metabolic reaction graph and its consequent m-DAG presented in this work can be of great use in the field of synthetic biology. The composition of compounds and reactions that we present can easily be extrapolated to any phylogenetically diverse bacteria of interest considering that we did not focus specifically on genes. Chemistry and molecular biology technologies are also thriving. Thus, the in silico design of bacteria with the small number of metabolic genes described in this paper may be more feasible than previously thought.

Supplementary Materials: The following are available online at https://www.mdpi.com/2079-7737/10/1/5/s1, Figure S1: Full size representation of the reaction graph of the proposed theoretical minimal metabolic network represented in Figure 2; Figure S2: The m-DAG of “Ca. Nasua deltoccephalinicola” str. NAS-ALF; Table S1: List of enzymes and reactions modified from Gabaldón et al. (2007) [19]; Table S2: Reactions and compounds that make up the m-DAG of “Ca. Nasua deltoccephalinicola” str. NAS-ALF; Table S3: Reactions included in the reconstruction of the JCVI-syn3.0 reaction graph and the minimal organism constructed for this work and the pathways in which each reaction (can) participates. Table S4: Names of the enzymes and definition of each reaction involved in the comparison of the MBBs of the three networks under study.

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References
1. Moya, A.; Gil, R.; Latorre, A.; Peretó, J.; Garcillán-Barcia, M.P.; De La Cruz, F. Toward minimal bacterial cells: Evolution vs. design. FEMS Microbiol. Rev. 2009, 33, 225–235. [CrossRef] [PubMed]
2. Luisi, P.L. Toward the engineering of minimal living cells. Anat. Rec. Adv. Integr. Anat. Evol. Biol. 2002, 268, 208–214. [CrossRef] [PubMed]
3. Luisi, P.L.; Ferri, F.; Stano, P. Approaches to semi-synthetic minimal cells: A review. Naturwissenschaften 2005, 93, 1–13. [CrossRef] [PubMed]
4. Xavier, J.C.; Patil, K.R.; Rocha, I. Systems Biology Perspectives on Minimal and Simpler Cells. Microbiol. Mol. Biol. Rev. 2014, 78, 487–509. [CrossRef]
5. Stano, P.; Luisi, P.L. On the Construction of Minimal Cell Models in Synthetic Biology and Origins of Life Studies. In Design and Analysis of Biomolecular Circuits; Springer Science and Business Media LLC: Berlin/Heidelberg, Germany, 2011; pp. 337–368.
6. Bedau, M.A.; Parke, E.C.; Tangen, U.; Hantsche-Tangen, B. Social and ethical checkpoints for bottom-up synthetic biology, or protocells. Syst. Synth. Biol. 2009, 3, 65–75. [CrossRef]
7. Mantri, S.; Sapra, K.T. Evolving protocells to prototissues: Rational design of a missing link. Biochem. Soc. Trans. 2013, 41, 1159–1165. [CrossRef]
8. Stano, P. Is Research on Synthetic Cells. Moving to the Next Level? Life 2018, 9, 3. [CrossRef]
9. Glass, J.I.; Merryman, C.; Wise, K.S.; Hutchison, C.A.; Smith, H.O. Minimal Cells—Real and Imagined. Cold Spring Harb. Perspect. Biol. 2017, 9, a023861. [CrossRef]
10. Gil, R. The Minimal Gene-Set Machinery. In Encyclopedia of Molecular Cell Biology and Molecular Medicine; Wiley: Hoboken, NJ, USA, 2014; pp. 1–36.
11. Luo, H.; Lin, Y.; Gao, F.; Zhang, Z.; Zhang, R. DEG 10, an update of the database of essential genes that includes both protein-coding genes and noncoding genomic elements: Table 1. *Nucleic Acids Res.* 2014, 42, D574–D580. [CrossRef]

12. Acevedo-Rocha, C.G.; Fang, G.; Schmidt, M.; Ussery, D.W.; Danchin, A. From essential to persistent genes: A functional approach to constructing synthetic life. *Trends Genet.* 2013, 29, 273–279. [CrossRef]

13. Gil, R.; Silva, F.J.; Zientz, E.; Delmotte, F.; González-Candelas, F.; Latorre, A.; Rausell, C.; Kamerbeek, J.; Gadou, J.; Hölldobler, B.; et al. The genome sequence of Blochmannia floridanus: Comparative analysis of reduced genomes. *Proc. Natl. Acad. Sci. USA* 2003, 100, 9388–9393. [CrossRef] [PubMed]

14. Mushegian, A.; Koonin, E.V. A minimal gene set for cellular life derived by comparison of complete bacterial genomes. *Proc. Natl. Acad. Sci. USA* 1996, 93, 10268–10273. [CrossRef] [PubMed]

15. Gil, R.; Silva, F.J.; Peretó, J.; Moya, A. Determination of the Core of a Minimal Bacterial Gene Set. *Microbiol. Mol. Biol. Rev.* 2004, 68, 518–537. [CrossRef] [PubMed]

16. Jewett, M.C.; Forster, A.C. Update on designing and building minimal cells. *Curr. Opin. Biotechnol.* 2010, 21, 697–703. [CrossRef] [PubMed]

17. A Hutchison, C.; Chuang, R.-Y.; Noskov, V.N.; Assad-Garcia, N.; Deerinck, T.J.; Ellisman, M.H.; Gill, J.; Kannan, K.; Karas, B.J.; Ma, L.; et al. Design and synthesis of a minimal bacterial genome. *Science* 2016, 351, aad6253. [CrossRef] [PubMed]

18. Venetz, J.E.; Del Medico, L.; Wolfe, A.; Schächle, P.; Bucher, Y.; Appert, D.; Tschan, F.; Flores-Tinoco, C.E.; Van Kooten, M.; Guennoun, R.; et al. Chemical synthesis rewriting of a bacterial genome to achieve design flexibility and biological functionality. *Proc. Natl. Acad. Sci. USA* 2019, 116, 8070–8079. [CrossRef]

19. Gil, R.; Silva, F.J.; Peretó, J.; Moya, A. Structural analyses of a hypothetical minimal metabolism. *Philos. Trans. R. Soc. B Biol. Sci.* 2007, 362, 1751–1762. [CrossRef]

20. Glass, J.I.; Assad-Garcia, N.; Alperovich, N.; Yooseph, S.; Lewis, M.R.; Maruf, M.; Iii, C.A.H.; Smith, H.O.; Venter, J.C. Essential genes of a minimal bacterium. *Proc. Natl. Acad. Sci. USA* 2006, 103, 425–430. [CrossRef]

21. Alberich, R.; Castro, J.A.; Llabrés, M.; Rodriguez, P.P. Metabolomics analysis: Finding out metabolic building blocks. *PLoS ONE* 2017, 12, e0177031. [CrossRef]

22. Bennett, G.M.; Moran, N.A. Small, Smaller, Smallest: The Origins and Evolution of Ancient Dual Symbioses in a Phloem-Feeding Insect. *Genome Biol. Evol.* 2013, 5, 1675–1688. [CrossRef]

23. Reyes-Prieto, M.; Vargas-Chávez, C.; Llabrés, M.; Palmer, P.; Latorre, A.; Moya, A. An update on the Symbiotic Genomes Database (SymGenDB): A collection of metadata, genomic, genetic and protein sequences, orthologs and metabolic networks of symbiotic organisms. *Database* 2020, 2020. [CrossRef] [PubMed]

24. Alvarez, A.F.; Georgellis, D. Bacterial Lipid Domains and Their Role in Cell Processes. In *Biogenesis of Fatty Acids, Lipids and Membranes*; Springer Science and Business Media LLC: Berlin/Heidelberg, Germany, 2019; pp. 575–592.

25. Romano, A.; Conway, T. Evolution of carbohydrate metabolic pathways. *Res. Microbiol.* 1996, 147, 448–455. [CrossRef]

26. Weinert, L.A.; Araujo-Jnr, E.V.; Ahmed, M.Z.; Welch, J.J. The incidence of bacterial endosymbionts in terrestrial arthropods. *Proc. R. Soc. B Biol. Sci.* 2015, 282, 20150249. [CrossRef] [PubMed]

27. Toft, C.; Andersson, S.G.E. Evolutionary microbial genomics: Insights into bacterial host adaptation. *Nat. Rev. Genet.* 2010, 11, 465–475. [CrossRef] [PubMed]

28. Douglas, A.E. The microbial dimension in insect nutritional ecology. *Funct. Ecol.* 2009, 23, 38–47. [CrossRef]

29. Vigneron, A.; Masson, F.; Vallier, A.; Balmard, S.; Rey, M.; Vincent-Monéget, C.; Aksoy, E.; Aubailly-Giraud, E.; Zaidman-Rémy, A.; Heddi, A. Insects Recycle Endosymbionts when the Benefit is Over. *Curr. Biol.* 2014, 24, 2267–2273. [CrossRef]

30. Tamas, I. 50 Million Years of Genomic Stasis in Endosymbiotic Bacteria. *Science* 2002, 296, 2376–2379. [CrossRef]

31. McCutcheon, J.P.; McDonald, B.R.; Moran, N.A. Convergent evolution of metabolic roles in bacterial co-symbionts of insects. *Proc. Natl. Acad. Sci. USA* 2009, 106, 15394–15399. [CrossRef]