1	Convergence in amino acid outsourcing between animals and predatory bacteria
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11 Abstract

All animals have outsourced about half of the 20 proteinogenic amino acids (AAs). We recently 12 13 demonstrated that the loss of biosynthetic pathways for these outsourced AAs is driven by energy-saving selection. Paradoxically, these metabolic simplifications enabled animals to use 14 costly AAs more frequently in their proteomes, allowing them to explore sequence space more 15 freely. Based on these findings, we proposed that environmental AA availability and cellular 16 respiration mode are the two primary factors determining the evolution of AA auxotrophies in 17 18 animals. Remarkably, our recent analysis showed that bacterial AA auxotrophies are also governed by energy-related selection, thereby roughly converging with animals. However, 19 20 bacterial AA auxotrophies are highly heterogeneous and scattered across the bacterial phylogeny, making direct ecological and physiological comparisons with the animal AA 21 22 outsourcing model challenging. To better test the universality of our model, we focused on Bdellovibrionota and Myxococcota-two closely related bacterial phyla that, through aerobic 23 respiration and a predatory lifestyle, best parallel animals. Here, we show that Bdellovibrionota, 24 driven by energy-related selection, outsourced a highly similar set of AAs to those in animals. 25 This sharply contrasts with Myxococcota, which exhibit far fewer AA auxotrophies and rarely 26 show signatures of energy-driven selection. These differences are also reflected in 27 Bdellovibrionota proteomes, which are substantially more expensive than those of 28 Myxococcota. Finally, we found evidence that the expression of costly proteins plays a crucial 29 role in the predatory phase of the Bdellovibrio life cycle. Together, our findings suggest that 30

Bdellovibrionota, through their obligate predatory lifestyle, exhibit the closest analogy to the
AA auxotrophy phenotype observed in animals. In contrast, facultative predation, as seen in
Myxococcota, appears to substantially limit the evolution of AA auxotrophies. These crossdomain convergences strongly support the general validity of our AA outsourcing model.

Keywords: bacteria; predation; amino acids; animals; auxotrophy; energetics; selection;
evolution

37

38 Introduction

The biosynthesis of amino acids (AAs) is a fundamental biochemical process essential for 39 sustaining all life on Earth. However, despite the universal metabolic demand for all 20 40 proteinogenic AAs, some organisms have lost the ability to synthesize certain AAs 41 42 endogenously. The most well-known example is animals [1-3], which have lost the capacity to produce approximately half of the complete AA set. A similar pattern of auxotrophies has 43 independently emerged in other eukaryotic groups, such as certain amoebae and euglenozoans 44 [1,2]. While most bacteria remain fully prototrophic [4], many bacterial lineages display 45 varying degrees of AA auxotrophy [4-6]. 46

47 It has long been speculated that the ability to acquire AAs from the environment influences the evolution of AA auxotrophies [2]. For instance, experiments have shown that bacteria 48 49 auxotrophic for an externally supplemented AA can gain a selective advantage over fully prototrophic counterparts [7]. However, a robust theoretical framework for this phenomenon 50 51 remained elusive. In our previous work, we addressed this gap by proposing a model that explains the evolution of AA outsourcing through several key factors: an AA is more likely to 52 53 be lost if (i) its biosynthesis is highly energy-demanding, (ii) it has a low pleiotropic effect, (iii) it is abundantly available in the environment, and (iv) the organism relies on efficient aerobic 54 respiration for energy production [1]. 55

Most importantly, we demonstrated that there is a constant selective pressure to outsource the synthesis of energetically costly AAs to the environment. Surprisingly, this leads to an increased usage of these expensive AAs in proteomes, allowing animal proteins to explore sequence space more freely [1,8]. To determine whether these global patterns are also valid in bacteria, we investigated AA auxotrophies across bacterial phylogeny and found that energyrelated selection also plays an important role in shaping AA outsourcing in bacteria [5]. However, bacteria exhibit far greater metabolic and ecological diversity than animals [4,9,10],
which necessitates testing our AA outsourcing model in specific bacterial groups whose
lifestyles more closely parallel those of animals.

One such group is the phylum Bdellovibrionota, which includes several aerobic or 65 66 microaerophilic species with an obligate predatory lifestyle [11–13]. Within this phylum, predation occurs via two distinct strategies: epibiotic predation, where the bacterium attaches 67 to the prey cell and leeches nutrients from its surface, and endobiotic predation, where the 68 bacterium enters the prey cell and forms a bdelloplast, within which it feeds, grows, and divides 69 [11–13]. The life cycle of Bdellovibrionota consists of two transcriptionally distinct phases: the 70 71 attack phase, during which the bacterium seeks and attaches to prey, and the feeding phase, characterized by growth and replication [12,14]. 72

Another bacterial group exhibiting animal-like behaviors is the phylum Myxococcota, a close relative of Bdellovibrionota [15,16]. Myxococcota are aerobes known for their complex multicellular behaviors, including social movement in coordinated "wolf packs," fruiting body formation, and predation [17]. However, unlike Bdellovibrionota, Myxococcota are not obligate predators—they can scavenge nutrients from dead organic matter and survive periods of starvation through sporulation [18–20].

The predatory lifestyle evolved independently in Bdellovibrionota and Myxococcota [16], leading to vastly different phenotypic outcomes [11–13,18–20]. Thus, these two bacterial groups provide an ideal system to test the predictive power of our AA outsourcing model, as they allow us to directly link the evolution of AA auxotrophies in bacteria to distinct ecological and physiological traits [1,5,8].

Here, we demonstrate that Bdellovibrionota evolved a remarkably animal-like set of AA auxotrophies, accompanied by an increase in relative proteome costs. Our findings indicate that energy-related selection played a key role in shaping these auxotrophies and that the attack and feeding phases of their life cycle exhibit distinct energy dynamics at the transcriptomic level. Surprisingly, this pattern does not hold for Myxococcota, which show fewer auxotrophies and lower proteome costs, suggesting a fundamental difference in how different reliance on a predatory lifestyle shapes metabolic evolution across these two groups.

91

93 **Results**

We suspected that the animal-like ecophysiology of Bdellovibrionota and Myxococcota 94 95 imposes energy-related selection on their AA metabolism, leading to reductions in AA biosynthetic pathways similar to those observed in animals. To test this hypothesis, we first 96 97 estimated the completeness of AA biosynthesis pathways in 89 Bdellovibrionota and 203 Myxococcota high-quality proteomes, which we retrieved from NCBI (Table S1). To assess 98 AA biosynthesis pathway completeness, which represents the likelihood of a given pathway 99 being present, we used the MMseqs2 clustering approach [5,21]. Heatmap representations 100 clearly revealed significantly lower AA completeness scores in Bdellovibrionota compared to 101 102 Myxococcota (Figure 1, File S1).

Most Bdellovibrionota showed reductions in pathway completeness scores for nine amino acids 103 (AAs) that fall on the expensive end of the biosynthesis cost distribution (Figure 1, File S1). 104 This set of AAs with low completeness scores largely overlaps with those that are auxotrophic 105 in animals [1], with one notable exception-lysine, which appears to be prototrophic in 106 Bdellovibrionota (Figure 1, File S1). In contrast, the pattern of AA biosynthesis pathway 107 108 reduction in Myxococcota is markedly different. Most Myxococcota remain prototrophic for the majority of AAs, with only valine, leucine, and isoleucine biosynthesis pathways 109 110 consistently exhibiting reductions (Figure 1, File S1). An exception to this trend is observed in two Myxococcota species-Vulgatibacter incomptus and Pajaroellobacter abortibovis-111 which display a substantial number of AA auxotrophies (Figure S1). Taken together, these 112 findings suggest that while some reductions of expensive AA occur in Myxococcota, some 113 ecological factors likely prevent them from outsourcing most of their costly AA biosynthesis 114 pathways. 115



Increasing biosynthesis cost

Figure 1. Completeness score of AA biosynthesis pathways in Bdellovibrionota and Myxococcota. We created a database of 89 Bdellovibrionota and 203 Myxococcota proteomes to get a comprehensive overview of AA dispensability in these groups. Full figure is shown in File S1. We retrieved all enzymes involved in AA biosynthesis from the KEGG and MetaCyc databases (reference collection) and searched for their homologs within our

Bdellovibrionota/Myxococcota database using MMseqs2 (see Methods). For each AA, we showed a completeness score, which represents the percentage of enzymes within a pathway that returned significant sequence similarity matches to our reference collection of AA biosynthesis enzymes. In the case of AAs with multiple alternative pathways, we showed the results only for the most complete one.

To globally test whether energy-related selection influences the observed reductions in AA 127 biosynthesis pathways of Bdellovibrionota and Myxococcota, we correlated the average AA 128 auxotrophy index with opportunity costs calculated under high respiration mode [5], a metric 129 that estimates the impact of AA biosynthesis on the cell's energy budget. To obtain the average 130 AA auxotrophy index for a given AA, we first subtracted the completeness score from 1 131 (Material and Methods, Equation 1) and then averaged the resulting AI values across all 132 considered proteomes. We detected a significant correlation between higher biosynthesis costs 133 and the loss of AA biosynthetic ability in Bdellovibrionota (Figure 2A). This suggests that 134 selection driven by energy management shaped the global pattern of auxotrophies in 135 136 Bdellovibrionota. As might be expected considering heatmap pattern (Figure 1), this broadscale analysis did not detect a positive correlation in Myxococcota, which are prototrophic for 137 most AAs (Figure 2B). 138



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Figure 2. Correlation between AA biosynthesis cost and the average AA auxotrophy index. We estimated the AA auxotrophy index (AI), a measure which equals one minus completeness score (see Materials and Methods, Equation 1), for 89 Bdellovibrionota (a) and 203 Myxococcota (b) proteomes and calculated the average value for every AA. We correlated this value with the opportunity cost of each AA, calculated for high respiration mode (see Materials and Methods). Pearson correlation coefficient and p-value are shown on the graph.

To further evaluate whether the observed reductions in AA biosynthesis pathways in Bdellovibrionota and Myxococcota result from energy-optimizing selection, we analyzed AA auxotrophy patterns at the species level. To achieve this, we first explicitly determined the AA auxotrophy status of each species by transforming the completeness score to binary values (auxotrophic/prototrophic) (see Materials and Methods, Table S2). Although this procedure

reduces the information contained in the completeness scores, it allowed us to test the impact 151 of energy-related selection more directly by explicitly defining auxotrophic and prototrophic 152 AAs. In principle, the loss of even a single enzyme within a biochemical pathway could render 153 that pathway nonfunctional. Thus, we assigned auxotrophy status to any AA whose biosynthesis 154 155 pathway was incomplete. Using this binary-transformed dataset, we statistically compared opportunity costs between auxotrophic and prototrophic AAs within each species [1]. In 156 157 addition, we applied a permutation-based selection test to assess the probability that the 158 observed constellation of auxotrophic AAs evolved under energy-related selection [1].

On average, Bdellovibrionota species exhibited 7.89 auxotrophic AAs per species (Table S2). 159 160 The comparison between auxotrophic and prototrophic AA sets using the Mann-Whitney nonparametric test shows that auxotrophic AAs have significantly higher opportunity costs in 92% 161 of the 89 tested Bdellovibrionota species (Table S2; File S2). For illustration, we singled out 162 the results of this comparison for the type strains of Bdellovibrio bacteriovorus and 163 Pseudobdellovibrio exovorus, representing endobiotic and epibiotic lifestyles, respectively 164 165 (Figure 3). It is evident that, regardless of feeding ecology, auxotrophic AAs are significantly more expensive than prototrophic ones (Figure 3A,B). The permutation-based selection test 166 revealed similar global trends, detecting that energy-related selection impacted the observed 167 distribution of auxotrophic AAs in 61% of Bdellovibrionota species (Table S2; File S2). For 168 instance, the average opportunity cost of the auxotrophic AA sets observed in Bdellovibrio 169 170 bacteriovorus and Pseudobdellovibrio exovorus falls at the right end of the distribution of all possible permutations (Figure 3C,D). This indicates, just like in animals, that energy-optimizing 171 172 selection governed the outsourcing of auxotrophic AAs in these bacteria.

The trends are quite different in Myxococcota, where an average of 2.79 AAs are auxotrophic 173 (Table S2). The Mann-Whitney non-parametric test shows that auxotrophic AAs have 174 significantly higher opportunity costs in only 9% of the 203 tested species (Table S2; File S2). 175 In comparison, the permutation-based selection test revealed similar results, indicating that 176 energy-related selection impacted AA auxotrophies in only 7% of Myxococcota species (Table 177 S2). A prominent representative of Myxococcota that shows the impact of energy-related 178 selection on AA auxotrophies is Pajaroellobacter abortibovis, a species whose pathogenic 179 ecology differs from the facultative predatory lifestyle of most other Myxococcota (File S1, 180 Table S2, File S2). Taken together, these findings suggest that energy-optimizing selection 181 related to AA auxotrophies is rather rare among Myxococcota compared to Bdellovibrionota. 182



Figure 3. Comparisons of auxotrophic and prototrophic AA sets in *Bdellovibrio bacteriovorus* 184 (A, C) and *Pseudobdellovibrio exovorus* (B, D). (A, B) The comparison of opportunity costs in 185 high respiration mode between auxotrophic and prototrophic AA groups was tested by the 186 Mann-Whitney U test with continuity correction. The corresponding W-value, p-value, and 187 effect size (r) are depicted in each panel. The X symbol represents the mean. Individual AAs 188 are shown by one-letter symbols. (C, D) Permutation analyses of opportunity costs (selection 189 tests) were performed by calculating the average opportunity cost of every possible permutation 190 for the number of auxotrophic AAs identified in a given species (n = 8 B. bacteriovorus, n = 9191 P. exovorus). The proportions of these averages are shown in histograms. The obtained 192 distribution represents the empirical probability mass function (PMF). The value in red denotes 193 the average opportunity cost value of the EAA sets observed in nature. The p-value was 194 calculated by summing the proportions that correspond to average opportunity cost values equal 195

to, or more extreme, than the observed value. Low p-values indicate a high probability thatenergy-related selection drove the loss of auxotrophic AA biosynthesis capability.

198 The fact that Bdellovibrionota converge to an animal-like set of amino acid (AA) auxotrophies, while closely related Myxococcota remain mainly prototrophic, allows us to further test the 199 200 predictions of our model [1]. In our previous work, we showed that animals code for significantly more expensive proteomes, compared to Choanoflagellates, their sister group 201 which is mainly prototrophic [1]. We proposed that animals have more expensive proteomes 202 than Choanoflagellates, likely because they consume less energy on AA biosynthesis and are 203 thus able to maintain a larger number of expensive auxotrophic AAs in their proteomes [1]. 204 205 Using a non-parametric test to compare the opportunity costs (high respiration mode) of an average AA in bacterial proteomes (Material and Methods, Equation 2), we recovered an 206 analogous result: the proteomes of the more auxotrophic Bdellovibrionota are significantly 207 more expensive than those of the more prototrophic Myxococcota (Figure 4), underscoring the 208 universality of energy-related selection on proteome composition. 209



Figure 4. The comparison of the opportunity cost (OC) of an average AA in proteomes of Bdellovibrionota and Myxococcota. The opportunity cost under high respiration mode (OC) of an average AA in a proteome represents a weighted mean of AA biosynthesis energy costs where the frequencies of twenty AAs in the proteome act as weights. The differences in energy costs between the two groups were shown by boxplots and the significance of these differences

was tested by the Mann-Whitney U test with continuity correction. We depicted the corresponding W-value, p-value, and effect size (r) in each panel. The X symbol represents the mean. The list of Bdellovibrionota (89) and Myxococcota (203) whose proteomes are included in calculations is available in Supplementary Data X.

220 Finally, we used the published transcriptome data to examine the energetics of the two distinct phases in the life cycle of *Bdellovibrio bacteriovorus*: the attack phase and the growth 221 phase [14]. For each transcript, we calculated its frequency in the transcriptome at a given life 222 cycle phase [22]. We then multiplied this transcript frequency by the opportunity cost of an 223 average amino acid (AA) encoded by that transcript (OC_{protein}) (Equations 3 and 4). This 224 measure, which we named the transcript energy score (TES), couples transcript levels in the 225 cell with the encoded protein costs (see Material and Methods). Higher TES values reflect 226 greater impact, while lower TES values indicate a smaller impact of a given transcript on the 227 total energy budget of the cell. We performed the TES-based analysis in two ways: by including 228 all genes expressed per phase and by considering only those which were exclusively expressed 229 230 in one phase (Figure 5).

The attack phase is generally characterized by a much smaller number of expressed genes, 231 which are more evenly distributed across the narrower range of TES values (Figure 5). In 232 233 contrast, the TES values of the growth phase are predominantly grouped at the lower end of the TES range. This suggests that the attack phase of predatory *B. bacteriovorus* is underpinned by 234 high transcription of a relatively small number of genes many of which encode expensive 235 236 proteins. In contrast, the growth phase is characterized by relatively low transcription of many genes that encode cheaper proteins. These differences are even more apparent in the analysis of 237 phase-specific genes (Figure 5B). Together, this suggests that the active predation in B. 238 bacteriovorus requires proteins which are composed of expensive AAs. As an analogy, we 239 240 previously speculated that similar phenotypes might exist in the context of animal predation [1]. 241



Figure 5. The comparison of transcriptome energy scores (TES) between two phases in the life 243 cycles of *B. bacteriovorus*. Calculations were performed (a) for all genes that are expressed per 244 phase and (b) only for phase-specific genes. The TES value represents the average opportunity 245 246 cost of a gene product multiplied by its frequency in the transcriptome at a given life cycle phase. The significance of the difference between the two phases was tested by the Mann-247 248 Whitney U test with continuity correction. We depicted the corresponding W-value, p-value, and effect size (r). Outliers were removed from the graph for the clarity of presentation, but 249 250 were included in the calculation of statistics. The X symbol represents the mean, and the 251 horizontal line within a violin-plot represents the median. Transcription data was retrieved from Karunker et al. 2013 (Karunker et al., 2013). 252

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254 Discussion

Functional outsourcing presumes that genes that support essential functions can be lost from 255 the genome if their activity can be substituted through environmental interactions [8]. We have 256 successfully applied this concept to the evolution of AA auxotrophies in animals and bacteria 257 [1,5]. We found that all animals and some bacterial groups which are capable of harvesting a 258 sufficient amount of AAs from their ecosystem lose the ability to produce expensive AAs on 259 their own [1,5,8]. This AA outsourcing is at least partially driven by energy-optimizing 260 selection, which not only favors the loss of the ability to synthesize expensive AAs but also 261 allows for more frequent usage of expensive AAs in the proteome [1,5]. We proposed that these 262 phenotypes could have been triggered in animals by predation and aerobic respiration [1]. If 263

these processes are indeed selection-driven, it would be expected that they occur convergentlyunder similar ecological pressures across the tree of life.

Unlike most other bacteria, Bdellovibrionota and Myxococcota are aerobic predators; they hunt and consume their bacterial prey under aerobic or microaerobic conditions [11–13,23]. As an independent evolutionary event, these behaviors were likely crucial for the outsourcing of AA production in animals [1]. We have shown here that obligate aerobic predation left an astonishingly similar metabolic impact on Bdellovibrionota, resulting in the largely overlapping set of AA auxotrophies compared to animals. In contrast, very few auxotrophies can be observed in Myxococcota, which are facultative predators [18,20].

There are likely multiple factors that influenced the vast differences in AA biosynthesis 273 capabilities between Bdellovibrionota and Myxococcota. However, the most apparent and 274 potentially vital factor is that Myxococcota are only facultatively predatory [18], making them 275 276 unable to consistently obtain AAs in large quantities. This is supported by the observation that 277 Bdellovibrionota grow and assimilate carbon at higher rates than Myxococcota [18]. Furthermore, it has been observed that Bdellovibrionota are significantly more abundant in 278 279 aerobic environments than in anaerobic ones, in contrast to Myxococcota, which show no apparent preference [23]. This might indicate that the metabolism of Bdellovibrionota is more 280 281 dependent on efficient respiration, which could also drive them toward heightened AA auxotrophy levels [1]. Another important factor might be feeding efficiency-Bdellovibrionota 282 283 are always physically connected to their prey, while Myxococcota secrete hydrolytic enzymes 284 around their prey, which carries the risk of diffusion, leading to a lower return of the energy expended on predation [24]. 285

The reduction in the ability to synthesize expensive AAs is directly correlated with the increased usage of expensive AAs in the proteomes [1,5]. Animals use more frequently expensive AAs in their proteomes than their sister group, choanoflagellates, and here we showed that the same is true for Bdellovibrionota when compared to their related group, Myxococcota [15]. Although Myxococcota have larger proteomes than Bdellovibrionota [11,24], their encoded AAs are on average cheaper by a large margin, which suggests that their complex lifestyle requires a wide range of functions that do not require proteins with very expensive AAs.

On the other hand, the ecology of Bdellovibrionota is relatively simple, consisting of two distinct phases: the attack phase, during which cells actively hunt, and the growth phase, during which cells consume their prey and subsequently divide [11,12,14]. A previous study produced

transcriptome data for these two phases in B. bacteriovorus [14], providing a unique 296 opportunity to examine the effects of energy-related selection on the *B. bacteriovorus* life cycle. 297 We used a novel measure, the transcript energy score (TES), which combines the energy cost 298 of a coding gene with its level of transcription, producing higher scores for more expensive and 299 highly expressed transcripts. Using this metric, we found that the growth phase is underlined 300 by a much larger set of transcripts with relatively similar expression levels that encode for 301 302 cheaper AAs, while the attack phase consists of a relatively small subset of transcripts with a 303 wide range of expression levels that encode for more expensive AAs. This supports the possibility that the energy saved through the outsourcing of AAs was invested in the attack 304 phase, enabling more efficient predatory behavior and thus ensuring a more consistent influx 305 of AAs from the environment, creating a positive feedback loop. 306

In conclusion, it is evident that every level of biological organization-from metabolism and 307 proteome composition to the regulation of gene transcription-is intimately tied to the 308 organism's energy budget. The repeated convergent evolution of similar sets of AA 309 310 auxotrophies across eukaryotes and prokaryotes can be attributed to obligate predation under aerobic conditions, which itself goes hand in hand with the evolution of novel and expensive 311 predation-related functions. Deeper understanding of the evolutionary pressures leading to 312 predatory behavior in bacteria is especially important, as it has implications not only for 313 understanding the regulation of ecological networks [18] but also for potential medical 314 315 applications [25].

316

317 Materials and Methods

All proteomes used in this study were retrieved from the NCBI GenBank. We acquired the highest-quality Bdellovibrionota (89) and Myxococcota (203) proteomes by using the NCBI filter to exclude atypical genomes, metagenome-assembled genomes, and genomes from large multi-isolate projects.

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We conducted the detection of amino acid (AA) biosynthesis pathway completeness using the protocol described in our earlier publication [5]. Briefly, we prepared a reference database of 387,892 enzyme sequences across 2,095 bacterial species, with each enzyme annotated with the biosynthetic pathway it is involved in [5]. We combined the reference database with our proteomes and clustered the sequences using MMseqs2 [21] with the following parameters: cluster-mode 0, -cov-mode 0, -c 0.8, and -e 0.001. We then functionally annotated all members
of a cluster based on the presence of enzymes from the reference database in that cluster.

For each AA biosynthesis pathway and species in the database, we calculated a pathway completeness score (i.e., prototrophy index) by dividing the number of detected enzymes by the total number of enzymes in that pathway,s resulting in values ranging from 0 to 1. If a species contained alternative biosynthetic pathways for an AA, the pathway with the highest completeness score was selected. We then calculated the AA auxotrophy index of the i-th amino acid (AI_i) by subtracting the completeness score from 1 [5]:

$$AI_i = 1 - completness \, score \tag{1}$$

We used the energy costs of AA biosynthesis as described in our earlier publications [1,5]. The opportunity cost reflects the impact of AA synthesis on the cell's energy budget and is calculated as the sum of the energy lost in the synthesis of AAs (direct cost) and the energy that would have been produced if a cell catabolized precursors instead of making AAs. Using the AA opportunity cost, we also calculated the opportunity cost of an average AA in each proteome $(OC_{proteome})$ using the following equation:

343
$$OC_{proteome} = \frac{\sum_{i=1}^{n=20} OC_i \times N_i}{\sum_{i=1}^{n=20} N_i} = \sum_{i=1}^{n=20} OC_i \times F_i$$
(2)

In this equation, OC_i represents the opportunity cost of the i-th AA, N_i denotes the total number of occurrences of this AA in the entire proteome, and F_i represents the frequency of the AA in the proteome.

The permutation analyses were performed separately for each species by first determining the 347 number of auxotrophic AAs and then generating all possible permutations for that number of 348 auxotrophic AAs. For each permutation, we then calculated the average opportunity cost of the 349 auxotrophic AAs. For example, if a species was found to be auxotrophic for 10 AAs, we found 350 all possible combinations of 10 AAs and calculated the average opportunity cost of each. Since 351 there is a limited number of possible average values, each value was treated as a bin. We 352 calculated the proportion of permutations within a bin by dividing the number of elements in 353 that bin by the total number of permutations. The obtained distribution represents empirical 354 probability mass function (PMF) which was then used to calculate the probability that the 355 observed set of auxotrophies in a given species is a result of a random process. We calculated 356

p-values by summing the proportions of permutation in the range from the actual value observedin nature to the most extreme value at the closest distribution tail [1].

359 For transcriptome analysis, we obtained data on differential gene expression in the two distinct life cycle phases of *Bdellovibrio bacteriovorus* from an earlier study [14]. For each gene (i), we 360 361 calculated its proportion in a given transcriptome phase by dividing its transcription value (t_i) by the sum of all transcription values in that phase. This proportion in essence represents the 362 frequency of each transcript in the transcriptome phase ($f_{\text{transcript}}$). We then multiplied this 363 number by the opportunity cost (high respiration mode) of the protein encoded by that gene 364 (OC_{protein}) to obtain a transcript energy score (TES). This score is meant to represent the energy 365 impact of each transcript on the total energy budget of a transcriptome phase. TES was 366 calculated using the following equation: 367

368
$$TES_i = \frac{t_i}{\sum_i^n t_i} \times OC_{protein} = f_{transcript} \times OC_{protein}$$
(3)

369 The opportunity cost of proteins was calculated using the following equation:

370
$$OC_{protein} = \frac{\sum_{i=1}^{n=20} OC_i \times n_i}{\sum_{i=1}^{n=20} n_i} = \sum_{i=1}^{n=20} OC_i \times f_i$$
(4)

In this equation, OC_i represents the opportunity cost of the i-th AA, n_i denotes the total number 371 of occurrences of this AA in the protein, and f_i represents the frequency of the AA in the protein. 372 We used the Mann-Whitney U test with continuity correction to compare the opportunity costs 373 of an average AA in proteomes (OCproteome) of Bdellovibrionota and Myxococcota using the 374 package rcompanion (https://CRAN.R-project.org/package=rcompanion). We used the same 375 test to compare transcript energy scores (TES) of different life cycle phases in *B. bacteriovorus*. 376 To calculate correlations, we used the cor.test() function in the R stats (v. 3.6.2) package. The 377 heatmap was visualized using the ComplexHeatmap package [26]. 378

Supplementary Materials: File S1: Full pathway completeness heatmap; File S2: Per-species
statistics on binary encoded auxotrophies; Table S1: Database with pathway completeness
detection results.xlsx; Table S2: Summary of statistics based on binary encoding of
auxotrophies

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