1	Title: Temperature-induced transcriptional responses of the deep-biosphere bacterium,
2	Kosmotoga olearia, illuminate its adaptation to growth from 20°C to 79°C
3	Running title: Transcriptional response to temperature in K. olearia
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18 Abstract

19 Temperature is one of the defining parameters of an ecological niche, and ambient temperature 20 change is a physiological challenge faced by all living cells. Most organisms are adapted to 21 growing within a temperature range that rarely exceeds $\sim 30^{\circ}$ C, but the anaerobic thermophilic 22 bacterium Kosmotoga olearia is capable of growing over an extremely wide temperature range (20°C - 79°C). To pinpoint genomic determinants of this flexible phenotype, we compared 23 24 transcriptomes of K. olearia cultures grown at its optimal 65°C to those at 30°C, 40°C, and 25 77° C. We found that changes in temperature significantly affect expression of 573 of 2,224 K. 26 *olearia* genes. At different temperatures K. *olearia* remodels its metabolism dramatically, with increased expression of genes involved in energy and carbohydrate metabolism at high 27 temperatures and up-regulation of amino acid metabolism at lower temperatures. At sub-optimal 28 29 temperatures, many transcriptional changes were similar to those observed in mesophilic bacteria 30 at physiologically low temperatures, including up-regulation of genes encoding enzymes for fatty 31 acid synthesis, typical cold stress genes, and ribosomal proteins. In comparison to other 32 Thermotogae, K. olearia has multiple copies of some cold-associated genes, suggesting that an increase in gene copy number is a strategy for cold adaptation. Many of these cold response 33 34 genes are predicted to be laterally acquired, highlighting the role of gene exchange in bacterial 35 thermoadaptation. Notably, at 77°C one third of the up-regulated genes encode proteins with 36 hypothetical functions, indicating that many features of adaptations to high temperature growth 37 are still unknown.

39 Importance

40 The subsurface is arguably the largest habitat on Earth, and insights into the composition, 41 adaptation, and evolution of its microbial communities is likely to advance our knowledge of the 42 biosphere and of global element cycling. Some subsurface systems, such as oil reservoirs, are 43 thought to be so stable that any changes in environmental conditions, including temperature, occur over thousands of years. Yet a member of the oil reservoir microbial communities, the 44 45 bacterium Kosmotoga olearia, is capable of growing over a range of 59°C. Our finding of 46 coordinated temperature-specific gene expression patterns, and by extension temperature specific 47 metabolism, suggests that *Kosmotoga* populations encounter variable environments, probably 48 through migration. This raises the question: are deep subsurface microbial communities more dynamic than currently perceived? The large number of identified temperature-responsive genes 49 50 also indicates that temperature response is a complex polygenic trait.

51

53 Introduction

54 Microorganisms are capable of growing over an impressive temperature range, at least from -55 15° C to 122° C (1, 2), and temperature is one of the most important physical factors determining 56 their distribution, diversity, and abundance (3). However, individual microbial species grow only 57 within a much narrower temperature interval. For example, Escherichia coli O157:H7 thrives in 58 the laboratory between 19° C and 41° C (4), while *Geobacillus thermoleovorans* has a growth 59 range of 37° C to 70° C (5). Microorganisms with temperature ranges >50°C are rare, and research into the few with ranges >40°C has focused on psychrophiles (e.g., (2)). Kosmotoga 60 61 *olearia* TBF 19.5.1 (hereafter referred to as K. *olearia*) is an anaerobic thermophile from the 62 bacterial phylum Thermotogae with a growth range that spans almost 60° C (6). How does this lineage achieve such physiological flexibility and what are the evolutionary advantages and 63 64 implications of having this capability?

Fluctuations in temperature induce broad physiological changes in cells, including 65 alterations to cell wall and membrane composition, translation, and energy metabolism (3, 7, 8). 66 67 These physiological changes can be classified into two broad types of cellular response. Cold or 68 heat *shock* designates the changes observed *immediately* after the shift of a culture to a lower or 69 higher temperature, while *prolonged growth* at a specific lower or higher temperature elicits an 70 acclimated low- or high-temperature response (7). Most studies on prokaryotes have focused on 71 temperature shock responses. Among the Thermotogae, responses to both heat shock and prolonged growth at high temperatures have been studied in the hyperthermophile Thermotoga 72 73 maritima, which can grow between 55°C and 90°C (9, 10). During prolonged high temperature 74 growth T. maritima strongly up-regulates central carbohydrate metabolism genes and expresses a 75 few typical heat shock protein genes (10). Little is known about how T. maritima responds to

76 sub-optimal temperatures, although it encodes some genes implicated in cold shock response. 77 For example, its family of cold shock proteins (Csp), which are nucleic acid chaperones known 78 to be induced during cold shock and cold acclimation in mesophilic bacteria (7, 11), exhibits 79 nucleic acid melting activity at physiologically low temperatures (12). Similarly, responses to 80 cold shock in a few other thermophiles involve many of the genes implicated in mesophiles' cold 81 shock response (e.g., (13, 14)). In this study we systematically assess bacterial physiological 82 changes associated with response to prolonged growth at both high and low temperature using K. olearia as a model system. 83

84 The K. olearia genome (NC_012785) has 2,302,126 bp and is predicted to encode 2,224 85 genes (15). Within the Thermotogae, genome size, intergenic region size, and number of 86 predicted coding regions correlate with the optimal growth temperature of an isolate (16), with 87 hyperthermophilic Thermotogae genomes being the most compact. Phylogenetically, the 88 Thermotogae order Kosmotogales comprises the genera Kosmotoga and Mesotoga spp., the latter being the only described mesophilic Thermotogae lineage (8). Assuming a hyperthermophilic 89 90 last common ancestor of the Thermotogae (17), the Kosmotogales can be hypothesized to have 91 acquired wide growth temperature tolerance secondarily by expanding its gene repertoire. 92 Moreover, it is likely that the ability of the Kosmotogales common ancestor to grow at low 93 temperatures made the evolution of mesophily in Mesotoga possible (8).

Such adaptations of lineages to new environments can be greatly facilitated by lateral gene transfer (LGT), since genes already "adapted" to the new conditions are readily available in the microbial communities of the new environment (18). For instance, LGT has been implicated in adaptation to high temperature growth in hyperthermophilic bacteria, including *Thermotoga* spp., and to low temperature growth in Archaea (8, 18, 19). Genome analysis of the mesophilic 99 Mesotoga prima revealed that it laterally acquired 32% of its genes after it diverged from other 100 Thermotogae lineages (16). Many of the predicted gene donors are mesophiles, supporting the 101 importance of lateral acquisition of genes already adapted to mesophilic conditions in the 102 evolution of Mesotoga.

103 To further gain insights into mechanisms of bacterial temperature response we sequenced 104 17 transcriptomes from isothermal and temperature-shifted cultures of K. olearia and examined 105 transcriptional differences at temperatures spanning its wide growth range. Additionally, through 106 comparative genomic and phylogenetic analyses of identified temperature responsive genes and 107 their homologs in two newly sequenced Kosmotoga isolates, as well as in genomes of other 108 thermophilic and mesophilic Thermotogae, we investigated the importance of within-lineage 109 evolution through LGT and gene duplication for adaptation of K. olearia to growth over a wide 110 temperature range.

111

112 **Results and Discussion**

113 Temperature shifts and isothermic conditions elicit different growth patterns in K. olearia. 114 Under laboratory conditions in liquid anaerobic medium we observed growth of K. olearia at 115 temperatures as low as 25°C and as high as 79°C, with optimal growth at 65°C (Fig. 1 and Fig. 116 S1 in the supplemental material). Using a non-linear regression model (20) we estimate a 117 growth-permissive temperature range of 20.2 - 79.3 °C, consistent with the previously reported 118 wide growth range of this isolate (6). Interestingly, we were not able to cultivate K. olearia at 119 temperatures near its range boundaries (30° C and 77° C) by direct transfer from 65°C cultures. 120 Instead, the growth temperature had to be changed sequentially in $\leq 10^{\circ}$ C increments. Particularly

121	at the extremes, even small temperature shifts caused both a longer lag phase and a slower
122	growth rate compared to isothermal cultures (Fig. 1 and Fig. S1 in the supplemental material).
123	This phenomenon has also been noted for mesophilic bacteria, especially for transitions from
124	high to low temperature (21). Our observations suggest that cells shifted to a new temperature
125	need to undergo large physiological changes that require time (i.e. an 'acclimation' period (7))
126	and that these physiological challenges are too great to overcome when temperature changes are
127	large. To illuminate K. olearia's transcriptional responses to changes in temperature we
128	sequenced 17 transcriptomes of replicate mid- to late-log cultures grown isothermally at 30°C,
129	40°C, 65°C, and 77°C, and of two 30°C cultures shifted to 25°C and 4°C (see Table S1 in
130	Dataset S1 and Text S1 in the supplemental material).

Architecture of the K. olearia transcriptome. Analysis of transcription start and stop 131 132 sites predicted a minimum of 916 transcriptional units (TU) in K. olearia (Text S1 and Table S2 133 in Dataset S1 in the supplemental material), 52% of which consist of a single gene. This fraction 134 of single-gene TUs lies between the 43% recorded for T. maritima, which has been shown to 135 have a streamlined genome and a low-complexity transcriptome (22), and the 65% reported for E. coli (23). The average TU length of ~2.39 genes in K. olearia is less than the 3.3 genes per 136 transcript of *T. maritima* (22) but closer to 2.2 genes per transcript in the mesophilic firmicute 137 Geobacter sulfurreducens (24) and 1-2 genes per transcript in Bacteria in general (e.g. (23)). 138 139 Given that the K. olearia genome has more intergenic DNA than T. maritima's genome (the ratio 140 of the nucleotides located in non-coding vs. coding regions is 0.13 in K. olearia and 0.06 in T. 141 maritima), the shorter TU lengths in K. olearia may point to more flexible transcriptional regulation and may be linked to *Kosmotoga*'s ability to grow under more variable conditions. 142

143	Consistent energy generation across different temperature conditions. K. olearia
144	produces ATP from pyruvate using a biochemically well-understood fermentation pathway that
145	generates hydrogen, carbon dioxide and acetate ((6); Fig. 2 and data not shown). Given that
146	pyruvate was the carbon and energy source in all experiments, we surveyed 51 genes predicted to
147	be involved in pyruvate catabolism and identified 15 genes with consistently high expression in
148	all temperature treatments (Fig. 2 Table S3 in Dataset S1 in the supplemental material). In
149	addition to indirectly validating the previously known functional annotations of these genes, we
150	furthermore propose that genes Kole_1509 – 1513 encode a pyruvate ABC transporter (Fig. 2).
151	Their current annotation as a peptide ABC transporter may be erroneous since most of the
152	peptide ABC transporters predicted in T. maritima using bioinformatics have been shown instead
153	to bind and transport sugars (25). Our findings also indicate that the enzymes involved in the
154	pyruvate fermentation pathway are versatile enzymes that are expressed and are capable of
155	functioning across an extremely wide temperature range.

156

Identification of temperature-related transcriptional responses in K. olearia. Based

157 on hierarchical clustering, transcriptome replicates at the same temperature group together (Fig.

158 S2 and Text S1 in the supplemental material), suggesting that the observed changes in

transcription are due to the culture growth temperature. Principal Component Analysis (PCA)

160 clearly separated the transcriptomes into quadrants corresponding to optimal (65° C),

161 intermediate (40°C), low (30°C, 25°C and 4°C) and high (77°C) growth temperatures (Fig. 3).

162 Several genes with a high correlation between their expression level and a specific growth

temperature (vectors in Fig. 3, Table S4 in Dataset S1 in the supplemental material) are known to

- be involved in temperature response. For example, expression of the protease Kole_1599
- 165 positively correlated with the 77°C transcriptomes, where high expression of proteases was

166 expected based on their involvement in heat shock response in T. maritima (9). Similarly, 167 expression of the cold shock protein genes Kole_0109 and Kole_2064 positively correlated with 168 low temperature growth. Lastly, some observed changes presumably were due to the expected 169 decreased metabolic activity of the culture at a non-optimal temperature. This can be exemplified 170 by the high expression and strong correlation of the central carbon metabolism gene 171 glyceraldehyde-3-phosphate dehydrogenase (Kole_2020) with the 65°C transcriptomes. Overall, 172 the observed differential expression of known temperature-responsive genes implies that the 173 remaining detected transcriptional changes also likely reflect temperature-related trends in gene 174 expression.

175 Detailed analysis of changes in gene expression in response to prolonged growth at 176 different temperatures. Putative temperature-responsive genes were identified by pairwise 177 comparisons of each isothermic temperature treatment to the optimal growth at 65°C (i.e., 30°C 178 vs 65°C, 40°C vs 65°C, and 77°C vs 65°C). Across all comparisons 573 genes fulfilled our 179 criteria for temperature responsiveness (i.e., ≥ 2 -fold difference in expression, > 20 reads per 180 transcript, False Discovery Rate < 0.05) with 430, 115, and 169 genes detected in the 30°C vs 181 65°C, 40°C vs 65°C, and 77°C vs 65°C comparisons respectively (Table S5 in Dataset S1 in the 182 supplemental material). Most of these genes were up-regulated (Fig. S3 in the supplemental material) with the exception of down-regulation of many genes involved in carbohydrate and 183 energy metabolism at 30°C (Clusters of Orthologous Groups [COG] categories G and C). The 184 185 latter probably reflects the very slow growth rate at this temperature. Curiously, despite the 186 slower growth rate at 77°C, genes from COG C and G categories were over-represented and up-187 regulated at this temperature (Fig. 4 and Fig. S3 in the supplemental material; discussed in detail 188 below).

189 In all transcriptomes the list of putative temperature-responsive genes was depleted of 190 genes involved in translation (COG category J) and nucleotide metabolism (COG category F) 191 (Fig. 4 and Fig. S3 in the supplemental material) and enriched in genes involved in replication, 192 recombination and repair (COG category L, particularly at 30°C), and signal transduction (COG 193 category T). Most of the identified COG category L genes are either mobile elements or CRISPR-associated proteins. Movement of mobile genetic elements is a common feature of 194 195 stress responses (26) and the up-regulation of CRISPR-associated genes could therefore be a 196 response to the proliferation of these elements. Differential expression of the signal transduction 197 genes suggests the importance of these systems for regulating cellular responses at all tested 198 temperatures. Additionally, at both 30°C and 77°C many genes encoding transcription regulators (COG category K, transcription) are up-regulated, suggesting that prolonged growth at sub- and 199 200 supra-optimal temperatures results in detectable changes in transcriptional gene regulation in K. 201 olearia. Below we discuss the identified temperature-responsive gene expression patterns in more detail. 202

203 i) At 40°C there are pronounced differences in membrane fatty acid composition 204 but no signs of cold stress. Although the growth rate of K. olearia at 40°C is only one-third of 205 that at the optimum $65^{\circ}C$ (Fig. 1 and Fig. S1 in the supplemental material), clustering analysis 206 suggested that the 40°C transcriptome was most similar to that at 65°C (Fig. 3 and Fig. S2 in the 207 supplemental material). The slower growth rate was reflected by the four most highly expressed 208 temperature responsive genes at 40° C showing significantly lower expression than at 65° C. 209 including growth-related genes like the toga protein Kole 1501 (Table S5 in Dataset S1 in the 210 supplemental material). Yet, 94 of 115 putative temperature responsive genes were up-regulated

211	(Table S5 in Dataset S1 in the supplemental material), suggesting that slower metabolism is not
212	the only explanation for the observed transcriptional response to growth at 40°C.

213 Lipid metabolism (COG category I) appears to be particularly important at 40°C. For 214 instance, all of the predicted fatty acid synthesis genes showed the highest expression at 40°C 215 (Table S5 in Dataset S1 and Fig. S4 in the supplemental material), with two genes involved in 216 synthesis of unsaturated fatty acids (Kole_0968) and initiation of fatty acid synthesis 217 (Kole_0969) having significantly higher expression. Biochemical analyses of total fatty acids at 218 40°C and 65°C showed a much greater diversity of fatty acids at 40°C (Table S6 in Dataset S1 in 219 the supplemental material), which may explain the higher demand for these genes at lower 220 temperatures. Interestingly, at 40°C in particular there was increased expression of a phosphate 221 ABC transporter (Kole_0707 – Kole_0711, Table S5 in Dataset S1 in the supplemental material), 222 which may be linked to the increased production of polar membrane lipids at moderately low 223 temperatures. Maintenance of a functional cell membrane is crucial for survival, and bacteria 224 respond to changes in temperature by altering the membrane's fatty acid composition (27). The 225 observation that lipid metabolism genes were among the highly expressed genes at low 226 temperature, despite the lower growth rate, suggests that changes to the cell membrane 227 composition are one of the most important adaptations for survival of K. olearia at lower 228 temperatures.

Proper protein folding at a lower temperature is another physiological challenge that may require enzymatic assistance. For example, proline isomerization happens spontaneously at high temperatures, but at lower temperatures (e.g., 37°C) the reaction needs a catalyzing enzyme peptidylprolyl isomerase (PPIase) (28). Not surprisingly, *K. olearia* has three temperatureresponsive PPIase genes: two PpiC-type genes (Kole_1682 and Kole_0383) that are both highly

234	expressed at 40°C, and one FKBP-type gene (Kole_1745), which shows high expression at all
235	temperatures except 77°C (Table S5 in Dataset S1 in the supplemental material). These
236	expression patterns suggest PPIase is particularly important at moderately low temperatures
237	where the cells are still relatively active. However, the enzymes known to assist protein folding
238	in cellular stress responses, chaperones (e.g., GroEL and Hsp) and protease Do, were
239	significantly down-regulated at 40°C (Table S5 in Dataset S1 in the supplemental material).
240	Among other typical cold stress related proteins, only one of K. olearia's three cold shock
241	proteins (Kole_0109) showed significantly higher expression at 40°C and its up-regulation was
242	merely moderate when compared to its expression levels at 30°C (Table S5 in Dataset S1 in the
243	supplemental material). This overall lack of induction of typical stress-related genes, especially
244	when compared to 30°C and 77°C (see below), suggests that 40°C is still within the "Goldilocks"
245	temperature range for K. olearia.

ii) K. olearia is in cold stress at 30°C. Transcriptomes from 30°C, 25°C, and 4°C 246 247 cultures were very similar to each other (Fig. 3 and Fig. S2 in the supplemental material). 248 Overall, the gene expression differences observed at 30°C were even more pronounced at 25°C 249 and 4°C (Table S5 in Dataset S1 in the supplemental material). However, due to adjustments in 250 culture handling required to obtain enough biomass at lower temperatures (see Text S1 in the 251 supplemental material), some gene expression patterns at 25°C and 4°C may be due to the cells 252 either responding to fresh medium or displaying an immediate cold shock response. Therefore, 253 we focused our further analyses on genes differentially expressed at 30°C, while the 25°C and 254 4°C transcriptomes were used to confirm the patterns observed at 30°C.

Two of three Csp-encoding genes in *K. olearia* (Kole_0109 and Kole_2064, Table S5 in Dataset S1 in the supplemental material) were among the three most highly expressed up-

257	regulated genes at low temperatures, suggesting that the cells were in a cold-stressed state during
258	growth at ≤30°C. Further support for this hypothesis comes from significant up-regulation of
259	genes linked to bacterial cold response (7): a DEAD/DEAH -box RNA helicase (Kole_0922),
260	RbfA (Kole_2103), and NusA (Kole_1529). Hence, the thermophile K. olearia uses homologs of
261	the cold response genes employed by mesophilic bacteria at physiologically low temperatures.

262 With decreasing temperature, we observed up-regulation of several ribosomal proteins 263 (Fig. 3). Some (L10 (Kole 1840) and L7/L12 (Kole 1839)) have already been linked to both cold shock and prolonged low temperature growth responses in bacteria (e.g., (29)). The most 264 265 dramatic differential expression, however, was observed for a ribosomal protein gene not yet 266 connected to cold response (L34; Kole_0258). L34, a bacteria-specific ribosomal protein 267 hypothesized to be a relatively recent addition to the evolving ribosome (30), is required for proper ribosome formation (31). A Bacillus subtilis mutant lacking the L34 gene showed slow 268 269 growth at low temperature (32), suggesting a role for L34 in this condition. Many ribosomal 270 proteins are recruited for extra-ribosomal functions (33), hence some of the up-regulated 271 ribosomal proteins may have alternative roles in response to low temperature that are unrelated to the ribosome itself. However, genes encoding ribosomal RNA (rRNA) methyltransferases. 272 273 rmlH (Kole 1718) and rmlL (Kole 0897), were also significantly up-regulated, and methylation 274 of rRNAs has been associated with responses to environmental stress, including temperature 275 (34). Combined with observations that ribosomes need to be fine-tuned to function properly at low temperature (7), we hypothesize that K. olearia modifies its ribosome by changing 276 stoichiometry of its components and by methylating rRNA. Time required for such ribosomal 277 278 adjustments could also explain the longer lag phase following temperature shifts (Fig. S1 in the 279 supplemental material).

280 To detect a decrease in environmental temperature and elicit an appropriate regulatory 281 response, some bacteria have evolved two-component cold sensors (27). These signal 282 transduction systems consist of a sensor, a membrane-integrated protein with a kinase domain 283 that detects changes in the fluidity of the cell membrane, and the cytoplasmic response regulator, 284 a protein that induces expression of cold-responsive genes. In K. olearia, a histidine kinase with 285 two predicted transmembrane domains (Kole_1017) and two response regulators (Kole_1015) 286 and Kole_1016) showed a steady increase in expression as temperatures decreased from 65°C, but no significant change in expression at 77°C (Table S5 in Dataset S1 in the supplemental 287 288 material), leading us to hypothesize that these genes encode a cold-sensing two-component system. 289

290 iii) Increased amino acid metabolism at sub-optimal temperatures. At lower growth 291 temperatures (and especially at and below 30°C) we observed an over-representation of genes 292 involved in amino acid metabolism (COG category E). At 30°C, and to a lesser extent at 40°C, a 293 peptide ABC transporter gene (Kole_2046 – Kole_2050) and several genes in the arginine 294 (Kole_0092 – Kole_0097) and lysine (Kole_0104 – Kole_0107, 30°C only) biosynthesis 295 pathways were up-regulated, suggesting the potential for accumulation of peptides and amino 296 acids (or their intermediates) at lower temperatures. At 30°C there was also significant up-297 regulation of a citrate synthase gene (Kole_1230). Intriguingly, in *Staphylococcus aureus* citrate 298 was shown to accumulate during prolonged cold stress (29), which could also be the case for K. 299 *olearia*. Alternatively, citrate synthase, together with isocitrate dehydrogenase (Kole_1227), may 300 be involved in converting pyruvate or acetyl-CoA to 2-oxoglutarate, a precursor for several 301 amino acids including arginine. Accumulation of both arginine and lysine was observed during 302 low temperature growth of *Clostridium botulinum*, where these amino acids were suggested to

act as compatible solutes (35). Interestingly, while the cells may accumulate peptides at 30°C, at 40°C there was increased expression of an oligo-peptidase (Kole_1190) and genes involved in lysine degradation (Kole_0958, Kole_0963 – Kole_0966). Such distinguishably different metabolic responses to moderately low (40°C) and low (\leq 30°C) temperatures suggest a finetuned temperature-dependent peptide turnover.

Two paralogs of ornithine carbamoyl-transferase (ArgF; Kole_1433 and Kole_2071) showed significantly lower expression at both 40°C and 30°C. The amino acid ornithine is an intermediate of arginine synthesis and lower expression of ArgF suggests that ornithine, rather than arginine, may accumulate at sub-optimal temperatures. However, ornithine has also been implicated in biofilm formation and species cross-talk (36), suggesting a possible alternative role of this amino acid in cellular responses to low temperature. Our unpublished observation of increased clumping of *K. olearia* cells grown at 30°C indirectly supports this hypothesis.

315 Re-modelling of amino acid metabolism at low temperatures has also been observed in 316 other bacteria (e.g., (35, 37)). Interestingly, the genome of *M. prima* encodes more genes 317 involved in amino acid metabolism than the genomes of K. olearia and other Thermotogae (16). 318 Perhaps the mesophilic *Mesotoga* spp. have adapted to the increased need for peptides by 319 expanding existing and acquiring new amino acid metabolism gene families. Amino acid 320 metabolism genes are also among the most numerous bacterial genes laterally acquired by 321 mesophilic Archaea, which was hypothesized to reflect their adaptation to low temperature growth (19). 322

iv) *K. olearia* is in heat stress at 77°C. Both the multivariate (Fig. 3) and clustering (Fig.
S2 in the supplemental material) analyses showed that the 65°C and 77°C transcriptomes are
distinct. Since 77°C is almost the upper limit for *K. olearia* growth, we hypothesize that the

326 observed differences in expression profiles at 77°C reflect a cell-wide heat stress response. Of 327 the 169 differentially expressed genes, 119 showed increased expression at 77°C (Table S5 in 328 Dataset S1 in the supplemental material). Hypothetical proteins made up a sizeable fraction (41 329 genes; 34%) of the 119 genes, indicating that adaptation to growth at sustained high temperature 330 remains largely uncharacterized. Only one of the known heat shock response genes (9), the 331 extreme heat stress sigma factor-24 (rpoE, Kole_2150), was up-regulated. Among the most 332 highly expressed genes were the structural RNAs ffs (Kole R0010), ssrA (Kole R0006), and 333 rnpB (Kole R0049) (Fig. 3), suggesting an increased rate of RNA turnover at supra-optimal 334 temperature. As mentioned earlier, carbohydrate and energy metabolism genes (COG category C 335 and G) were also up-regulated. It is unclear, however, if the underlying cause is the increased turnover of enzymes at elevated temperatures, or a demand for more enzymes due to increased 336 337 carbohydrate catabolism. Notably, two genes predicted to produce amino sugars for cell surface 338 polysaccharides (Kole_1281 and Kole_1282) were also moderately up-regulated at 40°C, 339 suggesting that cell surface polysaccharides may contribute to temperature adaptation at both 340 sub- and supra-optimal temperatures. Increased carbohydrate metabolism in response to 341 prolonged growth at supra-optimal temperature has been observed previously in T. maritima (10) 342 and therefore may be a common adaptation to high temperature growth in the Thermotogae. The 343 prolonged supra-optimal temperature growth of T. maritima also did not involve up-regulation of typical heat-shock response proteins (10). This highlights the difference between cellular 344 345 response to an immediate heat-shock and to prolonged growth at supra-optimal temperature, and 346 in general justifies classifying the cellular response to temperature into these two distinct 347 categories.

348 v) General stress response genes. Since we hypothesize that at 77° C and 30° C K. 349 *olearia* cells are under stress, genes that are significantly up-regulated at both temperatures are 350 candidates for a general temperature-stress response. There are 25 such genes, three of which 351 were also significantly up-regulated at 40°C (Table S5 in Dataset S1 in the supplemental 352 material). Among the most highly expressed of the 25 genes were Kole_2091, a gene with a 353 distantly related homolog in only two other Thermotogae, and Kole_0418, a gene that within the 354 Thermotogae has homologs only in *Mesotoga* spp. Both genes encode proteins of unknown 355 function. Given such limited distribution within the Thermotogae, these genes may be involved 356 in *Kosmotoga*-specific adaptation to a wide growth temperature range. 357 Conservation of K. olearia's temperature-responsive genes across Kosmotogales. All 358 genes that are required for adaptation and response of K. olearia to a wide range of growth 359 temperatures are expected to be present in other K. olearia isolates, whereas some may be absent 360 from Kosmotoga species having a narrower spectrum of growth temperature. Therefore, we 361 compared the K. olearia genome to the genomes of Kosmotoga sp. DU53 and Kosmotoga 362 arenicorallina (38). Kosmotoga sp. DU53 has a similar growth temperature range (observed 363 range 25°C - 79°C, Table S7 in Dataset S1 in the supplemental material) and >99% average 364 nucleotide identity (ANI) when compared to K. olearia, while K. arenicorallina exhibits a 365 narrower growth temperature range (observed range 35°C - 70°C, Table S7 in Dataset S1 in the 366 supplemental material) and has only 84% ANI when compared to K. olearia. Indeed, the *Kosmotoga* sp. DU53 genome lacks only 10 of the 573 K. olearia putative 367

Indeed, the *Kosmotoga* sp. DU53 genome lacks only 10 of the 573 *K. olearia* putative temperature-responsive genes (BLASTP and TBLASTN searches, E-value $< 10^{-3}$, Table S5 in Dataset S1 in the supplemental material). All 10 genes were expressed in *K. olearia* at relatively low levels (the highest average expression value of 453 is for Kole_0200 at 77°C), suggesting

that they are unlikely to be essential for high or low temperature growth. On the other hand, the *K. arenicorallina* genome does not have detectable homologs of 103 of the 573 putative temperature-responsive genes in *K. olearia* (BLASTP and TBLASTN searches, E-value < 10^{-3}) (Table S5 in Dataset S1 in the supplemental material). The list of absent genes includes several of the arginine and lysine biosynthesis genes that are up-regulated in *K. olearia* during growth at <30°C, and seven of the hypothetical proteins up-regulated at 77°C. Therefore, we hypothesize that these 103 genes play a role in extending the growth range of *K. olearia* to <35°C and ≥70°C.

378 Role of lateral gene transfer in thermoadaptation of *K. olearia*. Obtaining "pre-

379 adapted" genes from other genomes is one way prokaryotes adjust to new environmental 380 conditions (18). Using HGTector (39) we predicted that 354 of K. olearia's 2,118 protein coding 381 genes have been acquired laterally by K. olearia or the Kosmotogales (i.e., Kosmotoga and *Mesotoga*), presumably representing LGT events occurring after the divergence of Kosmotogales 382 from other Thermotogae (Table S8 in Dataset S1 in the supplemental material). Eighty-eight of 383 384 the 354 genes were temperature responsive (Table S5 in Dataset S1, Fig. S5A and S5B in the 385 supplemental material), including several already discussed highly expressed genes (Table 1 and 386 Table S5 in Dataset S1 in the supplemental material). Notably, LGT appears to be especially 387 important in K. olearia's adaptation to the lower growth temperatures. Thirty-eight of the 88 388 temperature-responsive laterally acquired genes are shared with the strictly mesophilic 389 Mesotoga, and most of them were highly expressed at lower temperatures, 30°C in particular 390 (Fig. S5C in the supplemental material). Among these are the previously discussed rRNA 391 methyltransferases (Kole_1718 and Kole_0897). The fatty acid synthesis genes (Kole_0969-392 Kole_0973) that are up-regulated at 40°C, as well as their Kosmotogales and *Mesotoga* homologs, form a distantly related sister clade to other Thermotogae lineages (Fig. S6A in the 393

supplemental material), suggesting that these genes may have been acquired from an un-sampled
lineage. Similarly, the Csp-encoding gene highly expressed at 30°C (Kole_0109) is placed
outside of the Thermotogae clade (Fig. S6B in the supplemental material). Predicted acquisition
of the fatty acid synthesis and Csp genes by (now mesophilic) Archaea (19) additionally argues
for the importance of these genes in adaptation to low temperature growth.

It is notable that some putative lateral acquisitions by *K. olearia* do not have homologs in *Mesotoga*. These include genes encoding the predicted cold temperature sensor (Kole_1015 –
Kole_1017), one of the PPIase genes (Kole_1745), as well as the canonical cold response
enzyme DEAD/DEAH box RNA helicase (Kole_0922). Lack of these genes in *Mesotoga*suggests their potential importance for *K. olearia*'s ability to grow over a wide temperature
range.

405

Role of gene family expansion and lineage-specific gene evolution in

thermoadaptation. Expansion of cold-responsive gene families may represent a common 406 407 strategy for low temperature adaptation, as has been noted in many bacteria, especially in 408 psychrophiles (e.g. (40)). K. olearia exhibits the same trend. For example, when compared to 409 other Thermotogae, all three analyzed Kosmotoga genomes harboured more copies of Csp-410 encoding genes (Table S9 in Dataset S1 in the supplemental material). Additionally, K. olearia 411 has extra homologs (Kole_0111 and Kole_0110) of the putative cold sensor system discussed above. The observed gene family expansions might be important not only for low temperature 412 413 growth, but also for growth over a wide temperature interval. For example, Mesotoga functions 414 with only a single Csp gene, demonstrating that having more copies of this gene is not required 415 for low temperature growth. Having several copies of these genes, however, allow K. olearia the 416 opportunity to regulate them differently at different temperatures. Similarly, the additional cold

417 sensor homologs do not show co-ordinated temperature response: Kole_0110 is up-regulated at
418 40°C, while Kole_0111 is up-regulated at 77°C (Table 1). Therefore, these additional homologs
419 may represent sensors tuned to different temperatures.

420 Gene family expansions can be achieved via within-lineage gene duplication or through 421 LGT. A combination of these mechanisms appears to be at work in K. olearia, as demonstrated 422 by the phylogenetic analyses of Csp genes (Fig. S6B in the supplemental material). Similarly, 423 even though several Thermotogae genomes contain as many copies of PPIase genes as do 424 Kosmotoga genomes (Table S9 in Dataset S1 in the supplemental material), phylogenetic 425 analysis suggests that in the Kosmotogales this gene family has only recently been expanded by 426 both LGT (the FKBP-type, Table 1) and duplication (the PpiC-type, Fig. S6C in the 427 supplemental material).

However, the role of within-lineage evolution of specific genes in response to changing environmental conditions should not be neglected. For example, typical cold response genes RbfA (Kole_2103) and NusA (Kole_1529) were not laterally acquired, but nevertheless show high expression only at 30°C. Deciphering adaptive changes that occurred in such genes compared to thermophilic homologs may elucidate molecular mechanisms of low temperature adaptation.

Why maintain the capacity for growth over such a wide temperature range? Most
bacteria are under selection to eradicate extraneous DNA (and genes) from their genomes (41),
and among free-living bacteria Thermotogae in general have very compact genomes.

437 Kosmotogales, however, have notably larger genomes than other thermophilic Thermotogae (8,

438 16), raising the possibility that expanded genomes are advantageous in *K. olearia*'s habitat. As

discussed above, many of the genes in *K. olearia*, such as the cold-sensor system, were

440 expressed only at specific sub- or supra-optimal temperatures, but do not seem to be important 441 for growth at other temperatures (Table 1 and Table S5 in Dataset S1 in the supplemental 442 material). The regulated response to low temperatures and the preservation of the laterally 443 acquired genes specifically expressed at 40° C and 30° C suggest that K. olearia encounters environments with very different temperatures frequently enough to maintain these genes in its 444 445 genome. Such environments may include oil reservoirs located at different depths, as well as 446 marine sediments influenced by the mixing of cold deep sea water and hydrothermal fluids (42). 447 As a result, this lineage was likely selected to become a temperature generalist. This conjecture 448 is supported by the environmental conditions of the subsurface environments and marine 449 hydrothermal vents from which Kosmotoga spp. have been isolated (6, 43, 44). K. olearia was isolated from a deep subsurface oil reservoir with in situ temperature of 68°C (6), but its 16S 450 451 rRNA sequences also have been detected in many oil fields having *in situ* temperatures of 20°C– 452 50°C (45). Kosmotoga sp. DU53, which is most similar to K. olearia, was isolated from an oil 453 reservoir with an *in situ* temperature of \sim 50°C, while *K. arenicorallina* was isolated from 454 hydrothermal sediments with a temperature of $\sim 40^{\circ}$ C (43). Notably, K. olearia was also identified as a major constituent in a metagenome from a deep subsurface oil reservoir with in 455 456 situ temperature of 85°C and pressure of 25MPa (46). While the reservoir temperature is higher 457 than the maximum K. olearia growth temperature reported here, elevated pressure could extend 458 K. olearia's temperature maximum, as has been demonstrated for some Archaea (e.g. (1)). 459 Therefore, K. olearia's growth temperature range under natural conditions may be even broader 460 than 20-79°C.

461 Concluding Remarks. The present study demonstrates that a bacterium with a relatively
 462 small genome can use transcriptional changes to respond effectively to large changes in

temperature. We showed that *K. olearia*'s response to sustained exposure to a non-optimal
temperature includes up-regulation of hundreds of genes. A substantial fraction of these genes
have been acquired laterally, suggesting that LGT is an evolutionarily successful strategy for
expansion of temperature tolerance; however, gene duplication and subsequent sub-

467 functionalization of the paralogs also plays an important adaptive role.

468 The ability of K. olearia to inhabit both high and low temperature environments suggests 469 that members of this lineage encounter environments with large temperature fluctuations and/or 470 frequently migrate across ecological niches within the deep biosphere (e.g., between deep and 471 shallow subsurface oil reservoirs). Therefore, the subsurface environments, as well as their 472 microbial populations, might be viewed as a connected archipelago instead of isolated islands. 473 As a corollary, we speculate that K. olearia-like ecological generalists could also facilitate LGT 474 among seemingly isolated deep biosphere microbial communities adapted to a narrower 475 ecological niche. For example, we have previously demonstrated high levels of gene flow among 476 hyperthermophilic *Thermotoga* populations in subsurface oil reservoirs and marine hydrothermal 477 vents (47), environments that are separated by non-thermophilic surroundings. The mechanism 478 of such gene flow is not yet known, but K. olearia-like Thermotogae capable of growing both in 479 subsurface oil reservoirs and adjacent marine sediments could serve as mediators of gene 480 exchange.

Although some of the identified 573 temperature-responsive genes are already known to be expressed in Bacteria and Archaea grown at high or low temperatures, most of the upregulated genes have not previously been implicated in temperature response and are in need of better functional and biochemical characterization. For example, the majority of the *K. olearia* genes responsive to elevated temperature encode proteins of unknown functions. Versatile

486	proteins that work across a broad range of temperatures also warrant further biochemical and
487	evolutionary analyses, as understanding of their enzymatic flexibility can aid the design of
488	commercially important thermostable proteins.

489

490 Materials and Methods

- 491 Bacterial culturing, and RNA and DNA isolation. K. olearia was grown at different
- temperatures (4°C, 25°C, 30°C, 40°C, 65°C, and 77°C), but otherwise optimal conditions, as
- described in (6) and Text S1 in the supplemental material.

For each temperature treatment, RNA was extracted in either mid-log phase or late-log 494 495 phase, using the Zymo Research Fungal/Bacterial RNA MiniPrep Kit (Cedarlane Laboratories, 496 Ltd.; Burlington, Ontario) and following the manufacturer's protocols (Table S1 in Dataset S1 in the supplemental material). The extracted RNA was sequenced on either an Ion Torrent PGM 497 (RNA-Seq kit V2) or an Illumina MiSeq (TruSeq RNASeq v2 2x100 bp) from the libraries 498 499 constructed following the manufacturer's instructions (Table S1 in Dataset S1 in the supplemental material). The transcriptomes are available in the Sequence Read Archive 500 501 (http://www.ncbi.nlm.nih.gov/sra) under the accession number SRP075860. 502 **RNA-Seq analysis.** For each transcriptome, sequenced reads were analyzed using the 503 RNA-Seq module in CLC Genomics Workbench version 7.0.4 (http://www.clcbio.com/, CLC 504 bio, Århus, Denmark), resulting in RPKM (Reads Per Kilobase of transcript per Million mapped 505 reads) values for each gene, as described in SI Materials and Methods. RPKM values for all

- genes are listed in Table S4 in Dataset S1 in the supplemental material. Differentially expressed
- 507 genes were identified by doing pairwise comparisons of the transcriptomes of the isothermically
- grown cultures at 30°C, 40°C, and 77°C to the cultures grown at the optimal temperature of

509	65°C. The analyses used the "Empirical Analysis of DGE" function, which employs the "Exact
510	Test" for two-group comparisons (48). A gene was considered differentially expressed in a
511	pairwise comparison if it had $(i) > 20$ reads in at least one of the two transcriptomes, (ii) a
512	statistically significant difference in the RPKM values (corrected for multiple testing using False
513	Discovery Rate [FDR] < 0.05), and (<i>iii</i>) a difference in RPKM values at least two-fold in
514	magnitude. Principal Component Analysis (PCA) and biplot visualization were performed using
515	R packages ade4 and bpca respectively (49, 50). Transcription start and stop sites and number of
516	transcripts were predicted using Rockhopper (51). For detailed descriptions see Text S1 in the
517	supplemental material.
518	Comparative analyses of three Kosmotoga spp. genomes. The genome of K. olearia
519	(accession number CP001634, (15)) was compared to genomes of Kosmotoga sp. DU53
520	(accession number JFHK0000000) and K. arenicorallina (accession number JGCK0000000)
521	(38) using the IMG portal (52) and Geneious v.9. Pairwise Average Nucleotide Identity (ANI)
522	(53) was calculated using the Enveomics Toolbox (54). Protein-coding genes in each genome
523	
	were classified as putatively laterally transferred using a customized version of HGTector (39).
524	were classified as putatively laterally transferred using a customized version of HGTector (39). For detailed descriptions see Text S1 in the supplemental material.

- 526 Archive (SRA) database under the accession number SRP075860.
- 527

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540	
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710

712 Figure Legends:

713 FIG 1. Growth rate of *K. olearia* as a function of temperature. Isothermic growth curves were 714 generated at each temperature from an inoculum grown at that temperature for at least three 715 transfers (except for 25°C and 80°C, for which an inoculum from the same temperature could not 716 be generated; see Text S1 in the supplemental material.). Up-shifted and down-shifted growth 717 curves were generated from an inoculum that was grown at lower and higher temperatures, respectively. Red squares, growth temperature up-shifted from 65°C to 77°C or from 40°C to 718 719 65°C; Blue circles, growth temperature down-shifted from 77°C to 65°C, 65°C to 40°C, or 40°C 720 to 30°C. Data points represent the mean of replicate cultures (see Text S1 in the supplemental 721 material); error bars represent standard error.

722

723 FIG 2. Model of energy generation pathway in K. olearia during growth on pyruvate. The 724 model includes genes likely involved in conversion of pyruvate to acetate, CO₂, H₂, and ATP. 725 The genes were selected from the list of genes highly expressed across all temperature conditions 726 (Table S3 in Dataset S1 in the supplemental material). Acetate transport is not shown. The 727 dashed box indicates hydrogenase activity. The two highly expressed hydrogenases are shown, 728 but their potential interactions with each other or with the membrane are not known. Increased 729 expression of citrate synthase at low temperature, which could redirect acetyl-CoA away from 730 acetate production, is shown in grey. The model also explains the observed lower ratio of carbon 731 dioxide to hydrogen produced by growth on maltose vs. pyruvate (not shown), as during growth 732 on maltose reduced electron carriers would be generated from the conversions of maltose to 733 pyruvate.

734

735	FIG 3. Biplot of the principal component analysis of 12 transcriptomes. Each transcriptome
736	is denoted by a point, while genes are represented by vectors. Genes that point into a specific
737	"temperature quadrant" are up-regulated at the growth temperature(s) of that quadrant, and the
738	five longest (i.e., most highly correlated) gene vectors pointing to each quadrant are shown. Co-
739	ordinates and vector length for all genes can be found in Table S4 in Dataset S1 in the
740	supplemental material. It should be noted that the ffs (Kole_R0010) transcript is only 115 nt, and
741	may not have been fully represented in every transcriptome due to our isolation protocol which
742	selects against small RNA (<200 nucleotides). Also, the high expression of the alcohol
743	dehydrogenase (Kole_0742) is probably due to the RNA isolation method (see Text S1 in the
744	supplemental material.).

745

746 FIG 4. Difference between observed and expected number of temperature responsive genes 747 across functional categories. Functional categories were assigned using the Clusters of 748 Orthologous Groups (COG) database as implemented in IMG (52) and are denoted by one-letter 749 abbreviations along the X-axis (see Fig. S3 legend for notations). NC, for "no category", denotes 750 genes not assigned to a functional category. For each temperature treatment (30°C, 40°C and 751 77°C) only the temperature-responsive fraction of the K. olearia genome was considered. If the 752 temperature-responsive genes were randomly distributed across functional categories we would 753 expect the same fraction of temperature-responsive genes in each COG category. The difference 754 (in percent) between the observed and expected number of temperature responsive genes is 755 plotted on the Y-axis with positive and negative values referring to over- and under-

- representation of the temperature-responsive genes, respectively. For actual number of genes in
- each COG category see Fig. S3 in the supplemental material.

758

760 **TABLE 1. Gene expression in selected laterally acquired temperature-responsive genes**. At each

temperature, the listed RPKM values represent the average expression levels across replicates. Values that

re significantly different from 65°C are shown in bold font.

Locus Tag	Functional annotation	30°C ^a	40°C	65°C	77°C	Identified by
Kole_0109	Cold shock protein	5602	892	222	119	Phylogenetic
						analysis
Kole_0110	Histidine kinase	175	333	144	312	Phylogenetic
						analysis
Kole_0111	Response regulator	166	204	173	446	HGTector
Kole_0505	Glycerol dehydrogenase	721	2668	752	1242	HGTector
Kole_0506	Hypothetical protein	559	2037	461	783	Phylogenetic
						analysis
Kole_0507	Hypothetical protein	555	2193	521	809	HGTector
Kole_0508	Poly (3-	212	423	200	314	HGTector
	hydroxybutyrate)					
	depolymerase-like					
	protein					
Kole_0897	Ribosomal RNA	503	498	228	232	HGTector
	methyltransferase, rmlL					
Kole_0922	DEAD/DEAH box	755	288	89	102	HGTector
	helicase					
Kole_0969	3-oxoacyl-ACP synthase	2386	3063	939	1424	HGTector
	III , FabH					
Kole_0970	enoyl-ACP reductase II,	2226	3243	1486	1641	HGTector
	fabK					

Kole_0971	malonyl CoA-acyl	2304	4211	2303	2647	HGTector
	carrier protein					
	transacylase, fabD					
Kole_0972	acyl carrier protein	6531	12601	4850	4241	HGTector
Kole_0973	3-oxoacyl-ACP synthase	4815	9257	4753	4498	HGTector
	II, fabF					
Kole_1015	Response regulator	1289	515	95	130	HGTector
Kole_1016	Response regulator	783	280	54	72	HGTector
Kole_1017	Histidine kinase	697	275	59	90	Phylogenetic
						analysis
Kole_1281	N-acylneuraminate-9-	482	699	315	840	HGTector
	phosphate synthase					
Kole_1282	N-acylneuraminate	244	283	128	315	Phylogenetic
	cytidylyltransferase					analysis
Kole_1718	Ribosomal RNA	531	332	211	203	HGTector
	methyltransferase, rmlH					
Kole_1745	PPIase FKBP-type	2783	2382	1541	430	HGTector

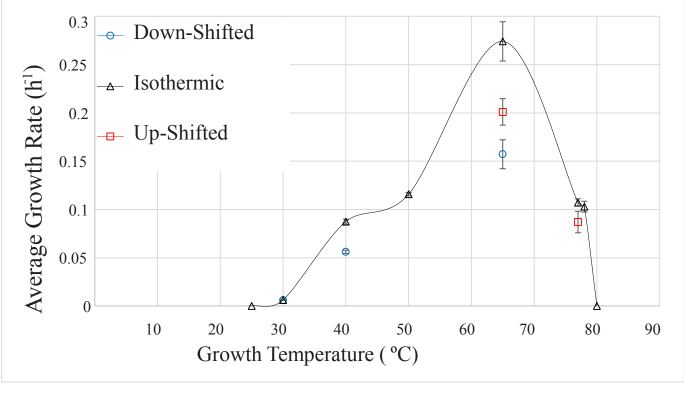


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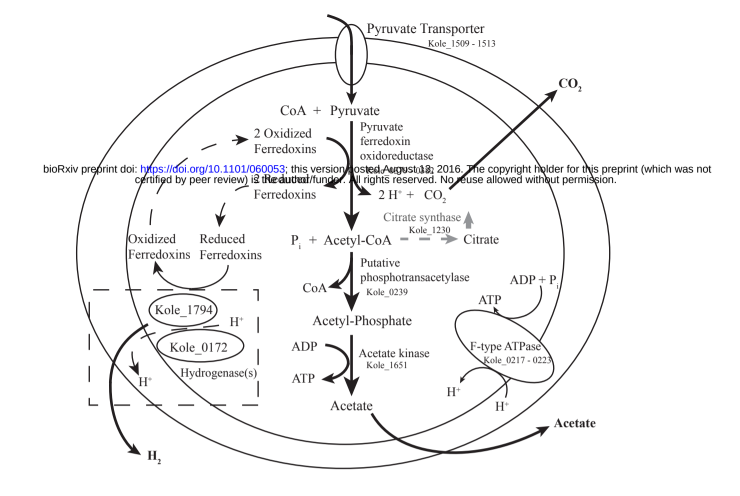


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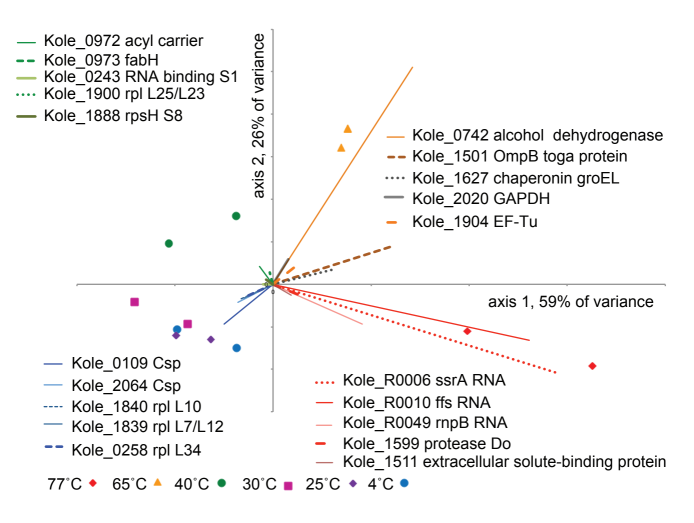


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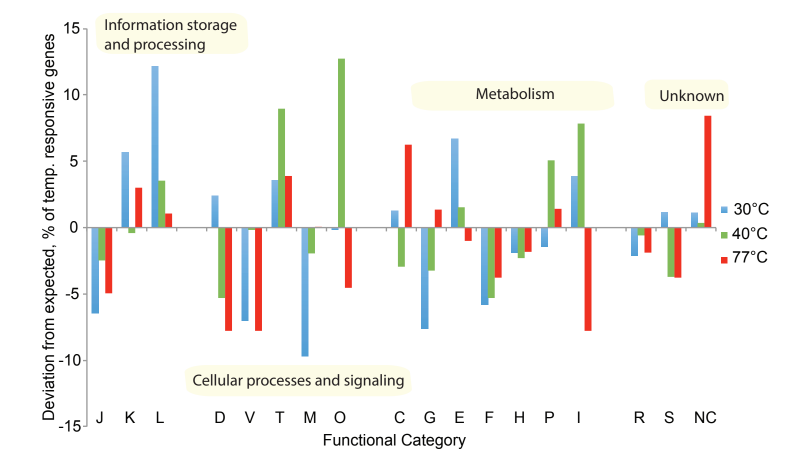


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