

The *Thermosynechococcus* Genus: Wide Environmental Distribution, but a Highly Conserved Genomic Core

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Cyanobacteria thrive in diverse environments. However, questions remain about possible growth limitations in ancient environmental conditions. As a single genus, the *Thermosynechococcus* are cosmopolitan and live in chemically diverse habitats. To understand the genetic basis for this, we compared the protein coding component of *Thermosynechococcus* genomes. Supplementing the known genetic diversity of *Thermosynechococcus*, we report draft metagenome-assembled genomes of two *Thermosynechococcus* recovered from ferrous carbonate hot springs in Japan. We find that as a genus, *Thermosynechococcus* is genomically conserved, having a small pan-genome with few accessory genes per individual strain as well as few genes that are unique to the genus. Furthermore, by comparing orthologous protein groups, including an analysis of genes encoding proteins with an iron related function (uptake, storage or utilization), no clear differences in genetic content, or adaptive mechanisms could be detected between genus members, despite the range of environments they inhabit. Overall, our results highlight a seemingly innate ability for *Thermosynechococcus* to inhabit diverse habitats without having undergone substantial genomic adaptation to accommodate this. The finding of *Thermosynechococcus* in both hot and high iron environments without adaptation recognizable from the perspective of the proteome has implications for understanding the basis of thermophily within this clade, and also for understanding the possible genetic basis for high iron tolerance in cyanobacteria on early Earth. The conserved core genome may be indicative of an allopatric lifestyle—or reduced genetic complexity of hot spring habitats relative to other environments.

Key words: cyanobacteria, great oxygenation event, hot springs, comparative genomics, pan-genome, *Thermosynechococcus*

Water oxidizing cyanobacteria fundamentally altered the distribution of carbon and electrons on Earth (Canfield, 1998; Raven, 2009; Ward and Shih, 2019). A marker of this reorganization is in the Great Oxygenation Event (GOE), which marks a major transition in the evolution of life on Earth ~2.3 billion years ago (Lyons *et al.*, 2014; Fischer *et al.*, 2016). While the GOE is widely accepted to have been driven by the production of O₂ by oxygenic photosynthesis performed by members of the cyanobacteria, the timing and proximal trigger for the GOE is debated. At least six hypotheses for the timing of the GOE have been considered: i) the evolution of oxygenic photosynthesis by cyanobacteria just before the GOE (Fischer *et al.*, 2016; Shih *et al.*, 2017), ii) an earlier evolution of cyanobacteria with O₂ accumulation delayed due to the transition of cyanobacteria from small-scale freshwater to large-scale marine environments (Sánchez-Baracaldo, 2015), iii) the transition from

unicellular to multicellular organisms for increased evolutionary success (Schirmer *et al.*, 2015), iv) the inhibition of early cyanobacteria due to high iron concentrations (Swanner *et al.*, 2015a; 2015b), v) a possible nitrogen throttle on cyanobacterial growth (Fennel *et al.*, 2005; Shi and Falkowski, 2008; Allen *et al.*, 2019), vi) or depressed Archean productivity due to phosphate availability (Hao *et al.*, 2020).

When and where the last common ancestor of cyanobacteria emerged is a matter of debate (Sánchez-Baracaldo, 2015; Shih *et al.*, 2017; Tria *et al.*, 2017) and cyanobacterial taxonomy continues to be refined (Knoll, 2008; Tomitani *et al.*, 2006; Shih *et al.*, 2013; Soo *et al.*, 2019; Parks *et al.*, 2020). Although uncertainty exists as to how much can be learned about past ecology from contemporary biology, an increased understanding of today's organisms may help us generate, or reject hypotheses about former evolutionary states.

Cyanobacteria are found in a wide range of environments—the knowledge of which continues to expand (Whitton, 1992; Puente-Sánchez *et al.*, 2018; Callieri *et al.*, 2019). As a single phylogenetic group, members of the *Thermosynechococcus* genus have been documented to inhabit a range of chemical environments, some of which can be seen as analogous to the Proterozoic environments on the early Earth (Ward *et al.*, 2019). Phylogenetic analysis of the *Thermosynechococcus* indicates they are a relatively recent divergence (Sánchez-Baracaldo, 2015; Shih *et al.*,

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2017). As a coherent group which spans a limited evolutionary distance, they provide a test case for querying how much environmental adaptation can be achieved within a limited time frame; that *Thermosynechococcus* inhabit a wide array of environments implies that rapid adaptation is possible, but the basis for this is unknown. Those *Thermosynechococcus* with genomes available are from hot springs which vary in temperature from 44–98°C, pH ranging from 5.4–9.3, sulfate concentrations between 0.06 mM and 17.4 mM, and iron concentrations between 0.4 µM and 261 µM (Table 1) at their source (however it must be emphasized that the source waters are not where the organisms reside unless specifically noted and the source water chemistry only gives a starting point of a gradient).

The *Ca. T. nakabusensis* OHK43 (OHK43) genome was recovered from material at the edge of the Okuoku-hachikurou Onsen (OHK) source pool where iron concentrations were measured to be 114 µM, and the *Ca. T. nakabusensis* Jinata (J003) genome was recovered at Jinata hot spring along a gradient where iron concentrations were 100 µM and greater (Ward *et al.*, 2017; 2019). In contrast, the other springs where *Thermosynechococcus* genomes originate are from hot springs with iron concentrations below a toxicity threshold of tens of micromolar Fe(II) suggested previously (Swanner *et al.*, 2015a). Furthermore, Nakabusa hot spring is sulfidic with 0.1 mM sulfide, while sulfide concentrations for Jinata and OHK were below the detection limit of a Cline assay (unpublished data), showing additional differences in geochemistry, which together are shown in Table 1.

Seminal work by Papke *et al.* (2003) on phylotype:geographical relationships of *Thermosynechococcus* posed questions as to how these organisms could be so widely distributed: in their analysis, the distribution of *Thermosynechococcus* could not be explained by measured geochemical parameters from their respective environments of origin. *Thermosynechococcus* thus appear to be cosmopolitan, but the basis for this remains unresolved. Motivated by the finding of *Thermosynechococcus* members in ferrous iron carbonate hot springs that we have been studying (Ward *et al.*, 2017; 2019), we sought to test the hypothesis that *Thermosynechococcus* members in high iron springs may have genetically resolvable traits associated with iron adaptation, by comparing to strains which are from lower iron environments.

Materials and Methods

Genome recovery

The OHK43 genome was recovered from genome-resolved metagenomic sequencing of samples from Okuoku-hachikurou Onsen (OHK) in Akita Prefecture, Japan, following methods described previously (Ward *et al.*, 2019; 2020) and described briefly here. Samples of thin biofilms in the outflow of the hot spring were sampled for metagenomic sequencing in September 2016. DNA was preserved in the field with a Zymo Terralyzer BashingBead Matrix and Xpedition Lysis Buffer (Zymo Research) after disruption of cells in polyethylene sample tubes via attachment to and operation of a cordless reciprocating saw (Makita JR101DZ). Microbial DNA was extracted and purified after return to the lab with a Zymo Soil/Fecal DNA extraction kit (Zymo Research). Quantifica-

Table 1. Geochemical parameters for hot spring source waters. Values in parentheses indicate the geochemical values of the sites where *Thermosynechococcus* sequences were observed, if known. Other values indicate the source water geochemistry of each spring, which can be used as a reference point for the start of a gradient in cases in which the explicit site where *Thermosynechococcus* sequences were observed is unknown. ^aConcentrations were derived from geochemical modelling in Mei-hua *et al.* (2015). ^bIron for Jinata and Okuoku-Hachikurou hot springs is ferrous, all others are totals of ferrous plus ferric iron. References are related to the first publication of the strains or the geochemistry of the hot spring. *information available online through local governments, last accessed in 11/2019.

Name	Strain name (<i>Thermosynechococcus</i>)	Max. Temperature [°C]	pH	Sulfate [mM]	Total Iron [µM]	References
Kangding Geothermal Field Lianhua Lake hot spring (Sichuan, China)	<i>T. elongatus</i> PKUAC-SCTE542 (<i>elongatus</i> PKUAC)	94 (67.2)	6.35–8.84 (7.95)	1.2	10.4 ^a	Mei-hua <i>et al.</i> , 2015; Guo <i>et al.</i> , 2017; Tang <i>et al.</i> , 2018
Okuoku-hachikurou Hot Spring (Akita, Japan)	<i>Ca. T. nakabusensis</i> OHK43 (OHK)	44	6.8 (6.8)	6.5	114 ^b (>100)	Ward <i>et al.</i> , 2017
Yunomine Hot Spring (Wakayama, Japan)	<i>T. vulcanus</i> NIES2134/1-2178 (<i>vulcanus</i>)	91	8	0.06–0.229	<1.8	Ichikuni and Kikuchi, 1972 Onsen information sheet*
Nakabusa Hot Spring (Nagano, Japan)	<i>T. nakabusensis</i> NK55a (NK55a)	76	8.5–9	0.218–0.246	0.4	Nakagawa and Fukui, 2002; Nakamura <i>et al.</i> , 2002; Stolyar <i>et al.</i> , 2014 Onsen information sheet*
Jinata Hot Spring (Shikinejima, Tokyo, Japan)	<i>Ca. T. nakabusensis</i> Jinata (J003) <i>T. sp.</i> M3746_W2019_013 (Jinata 2)	63 (37–46)	5.4 (6.7)	17.4	261 ^b (>100)	Ward <i>et al.</i> , 2019, Alcorta <i>et al.</i> , 2020
Chung-Lun Hot Spring (Taiwan)	<i>T. sp.</i> CL1/1-2178 (CL1)	62	9.3	1.35–1.39	0.6	Yeh <i>et al.</i> , 2005; Hsueh <i>et al.</i> , 2007; Maity <i>et al.</i> , 2011; Cheng <i>et al.</i> , 2020
Beppu Hot Spring Kamegawa Shinoyu (Oita, Japan)	<i>T. elongatus</i> BP1/1-2178 (<i>elongatus</i> BP1)	78	6.8	1.09	3.6	Onsen information sheet*
Shivlinga hot spring, Ladhak, India	<i>T. sp.</i> M46_R2017_013 (Shivlinga) *excluded in genome comparison due to low completeness	70 (46)	7 (8)	1	No data	Alcorta <i>et al.</i> , 2020; Roy <i>et al.</i> , 2020
Tattapani, India	<i>T. sp.</i> M55_K2018_012 (Tattapani 1) <i>T. sp.</i> M98_K2018_005 (Tattapani 2)	98 (55)	7.7 (7.9)	No data	0.49	Saxena <i>et al.</i> , 2017; Kaushal <i>et al.</i> , 2018; Alcorta <i>et al.</i> , 2020

tion of DNA was performed with a Qubit 3.0 fluorimeter (Life Technologies). DNA was submitted to SeqMatic LLC for library preparation using an Illumina Nextera XT DNA Library Preparation Kit prior to 2×100 bp paired-end sequencing via Illumina HiSeq 4000 technology. Raw sequence reads were quality controlled with BBTools (Bushnell, 2014) and assembled with MegaHit v. 1.02 (Li *et al.*, 2016). The OHK43 genome bin was recovered via differential coverage binning with MetaBAT (Kang *et al.*, 2015). Completeness and contamination/redundancy were determined with CheckM v1.1.2 (Parks *et al.*, 2015). The genome was uploaded to RAST v2.0 for annotation and characterization (Aziz *et al.*, 2008). Presence or absence of metabolic pathways of interest was predicted using MetaPOAP v1.0 (Ward *et al.*, 2018). Taxonomic assignment was determined with GTDB-Tk v1.2 (Parks *et al.*, 2018, 2020; Chaumeil *et al.*, 2019).

Organismal phylogenies

Concatenated ribosomal phylogenies were constructed following methods from Hug *et al.* (2016). Members of the *Thermosynechococcaceae* and outgroups were identified using GTDB (Chaumeil *et al.*, 2019) and their genomes downloaded from the NCBI WGS and Genbank databases. Ribosomal protein sequences were extracted using the tblastn function of BLAST+ (Camacho *et al.*, 2009) and aligned with MUSCLE (Edgar, 2004). Trees were built with RAxML v.8.2.12 (Stamatakis, 2014) on the Cipres science gateway (Miller *et al.*, 2010). Transfer bootstrap support values were determined with BOOSTER (Lemoine *et al.*, 2018). Visualization of trees was performed with the Interactive Tree of Life Viewer (Letunic and Bork, 2016).

Genome comparison

We compared the core- and pangenomes of *Thermosynechococcus* at the genus level, family level and with a sub-sample of organisms across the GTDB defined class Cyanobacteria. ProteinOrtho (Lechner *et al.*, 2011) was used for the identification of Conserved Likely Orthologous Groups (CLOG) and analysis.

At the genus level, ten *Thermosynechococcus* strains from varying hot spring environments (Table 1 and 2) were compared: The genomic data from ten currently available sequences of *Thermosynechococcus* strains *T. sp.* CL1/1-2178 (CL1), *T. elongatus* BP1/1-2178 (*elongatus* BP1), *T. vulcanus* NIES2134/1-2178 (*vulcanus*), *T. sp.* NK55a/1-2022 (NK55a), *T. elongatus* PKUAC-SCTE542 (*elongatus* PKUAC), *T. sp.* M55_K2018_012 (Tattapani 1), *T. sp.* M98_K2018_005 (Tattapani 2), *Ca. T. sp.* J003 (J003), *T. sp.* M3746_W2019_013 (Jinata 2) and *Ca. T. sp.* OHK43 (OHK43) were compared using ProteinOrtho, BLAST (Altschul *et al.*, 1990), and FeGenie (Garber *et al.* 2020). Due to low completeness of the genome, we excluded the eleventh available species (*T. sp.* M46_R2017_013). Phylogenetic relationships between the strains were established using concatenated ribosomal protein phylogenies following Hug *et al.* (2016), taxonomic classifications with GTDB-tk (Chaumeil *et al.*, 2019) and average nucleotide identities (Rodriguez-R and Konstantinidis, 2014).

For the analysis at the family level, we included 6 more species that appear as representative strains at the *Thermosynechococcaceae* family level in the GTDB. The representative GTDB strains include *Acaryochloris marina* MBIC11017, *Cyanothece* sp. PCC 7425, *Acaryochloris* sp. CCMEE 5410, *Synechococcus* sp. PCC 6312, *Synechococcus lividus* PCC 6715 and *Acaryochloris* sp. RCC1774. This resulted in a total of 16 strains for family level analysis.

At the class level we used the ten genus level strains and additionally 16 species, 15 after (Beck *et al.*, 2012) and in addition one *Gloeobacter* species (*G. kilauensis* JS-1). The class level analysis includes all of the genus level strains, and some family and class level strains. Here the goal was to test the coherence of our analysis parameters when compared to previous studies of cyanobacterial core- and pangenomes (Beck *et al.*, 2012, 2018). We acknowledge that the distribution of species within the families of the class is not even, however this comparison was used to verify our methods and the stability of the class level core. This resulted in a total of 26 genomes for the class level analysis.

To compare the appearance of genes related to iron uptake and

Table 2. Genus level average nucleotide identities (ANI) and genome sizes (diagonal). Additionally, genomes sizes for family level species are shown in the lower left.

Genome Size	<i>Ca. T. nakabusesensis</i> Jinata	<i>Ca. T. nakabusesensis</i> OHK	<i>T. elongatus</i> BP-1	<i>T. elongatus</i> PKUAC-SCTE542	<i>T. sp.</i> CL1	<i>T. nakabusesensis</i> NK55a	<i>T. vulcanus</i> NIES2134	<i>T. sp.</i> M3746_W2019_013	<i>T. sp.</i> M46_R2017_013	<i>T. sp.</i> M55_K2018_012	<i>T. sp.</i> M98_K2018_005
<i>Ca. T. nakabusesensis</i> Jinata	2.31 MB	99.70	92.65	86.82	87.53	99.68	92.59	99.76	83.92	87.85	87.85
<i>Ca. T. nakabusesensis</i> OHK		2.25 MB	92.58	86.85	87.52	99.84	92.55	99.76	83.95	87.88	87.87
<i>T. elongatus</i> BP-1			2.59 MB	86.65	87.34	92.53	99.14	92.56	83.86	87.63	87.61
<i>T. elongatus</i> PKUAC-SCTE542				2.64 MB	89.95	86.81	86.63	86.88	85.45	90.54	90.58
<i>T. sp.</i> CL1					2.64 MB	87.50	87.32	87.55	85.51	92.27	92.26
<i>T. nakabusesensis</i> NK55a						2.51 MB	92.45	99.69	83.92	87.84	87.82
<i>T. vulcanus</i> NIES2134							2.57 MB	92.51	83.79	87.55	87.56
<i>T. sp.</i> M3746_W2019_013								2.38 MB	83.99	87.87	87.90
<i>T. sp.</i> M46_R2017_013									2.39 MB	86.03	86.06
<i>T. sp.</i> M55_K2018_012										2.39 MB	99.92
<i>T. sp.</i> M98_K2018_005											2.37 MB

Genome Sizes for GTDB representative family level species:

<i>Acaryochloris marina</i> MBIC11017	8.36 MB
<i>Acaryochloris</i> sp. CCMEE 5410	7.87 MB
<i>Cyanothece</i> sp. PCC 7425	5.78 MB
<i>Synechococcus lividus</i> PCC 6715	2.65 MB
<i>Synechococcus</i> sp. PCC 6312	3.72 MB
<i>Acaryochloris</i> RCC 1774	5.93 MB

regulation we used FeGenie with standard parameters. We used results from ProteinOrtho for further analysis of the core, shared, unique, TS (*Thermosynechococcus*) core and TS shared clusters. Proteinortho was applied such that the output also included singleton clusters (only containing a single protein) with an algebraic connectivity of 0 as a measure for the structure of the orthologous clusters. We did not obtain many differences when running ProteinOrtho on our data using a value of 0 or 0.1 (default) for the algebraic connectivity, however, a value of 0 resulted in slightly larger clusters for the core and a few less singletons. A comparison of results from FeGenie and ProteinOrtho resulted in similar gene clusters for iron related genes such that our results here are based on both program outputs.

Results and Discussion

Phylogeny of the *Thermosynechococcus* and the species proposal *T. nakabusensis*

The *Thermosynechococcus* genus is phylogenetically coherent within the Cyanobacteria (Fig. 1) and the genome sizes of genus members are similar to one another (Table 2). Based on similarity observed with ANI and GTDB-tk (Table 2), 5 species are present within the *Thermosynechococcus* genus: *T. elongatus* BP1 and *T. vulcanus* belonging to one species, *T. NK55a*, *Ca. T. sp. J003*, *T. sp. M3746_W2019_013* (Jinata 2) and *Ca. T. sp. OHK43* (OHK43) genomes belonging to a second species, Tattapani 1 and Tattapani 2 belonging to a third species, and *T. CL1* and *T. elongatus* PKUAC-SCTE542 correspond to one species each. For *T. sp. M46_R2017_013* (Shivlinga) the species designation remains unresolved due to low completeness. For the species including *T. NK55a*, J003, with Jinata 2,

and OHK43, we propose the name *Thermosynechococcus nakabusensis* after the first and so far only isolated species member which originates from Nakabusa hot spring in Nagano Prefecture, Japan.

Genus and family level comparison of the genus *Thermosynechococcus* and the family *Thermosynechococcaceae*

Comparing conserved likely orthologous groups (CLOGs), we analyzed i) the core-genomes: those CLOGs shared by all genomes in an analysis, ii) the shared CLOGs: those shared by at least 2 but not all of the genomes in the analysis, and iii) unique CLOGs: those CLOGs that are unique to a single genome (Table 3). The *Thermosynechococcus* genus specific core (core TS) comprises CLOGs shared by all ten genus level genomes that are not present in any other species, and the *Thermosynechococcus* genus-specific shared CLOGs (shared TS) corresponds to CLOGs that are shared by at least two and at most nine genus level genomes.

Comparing the genomes of the ten genus members, the protein core is made up of 1,737 CLOGs and contains 66 to 75% of the putative protein coding genes in a genome. This value is different from a recent analysis (Cheng *et al.*, 2020), who compared 5 genomes available at that time and found a core of 1,264 CLOGs. The higher value observed here is attributed to the analysis of a revised *T. elongatus* PKUAC-SCTE542 genome which had a much lower quantity of pseudogenes in comparison to the previously available version.

The percentage of the *Thermosynechococcus* genomes which belongs to the genus core is higher than in other com-

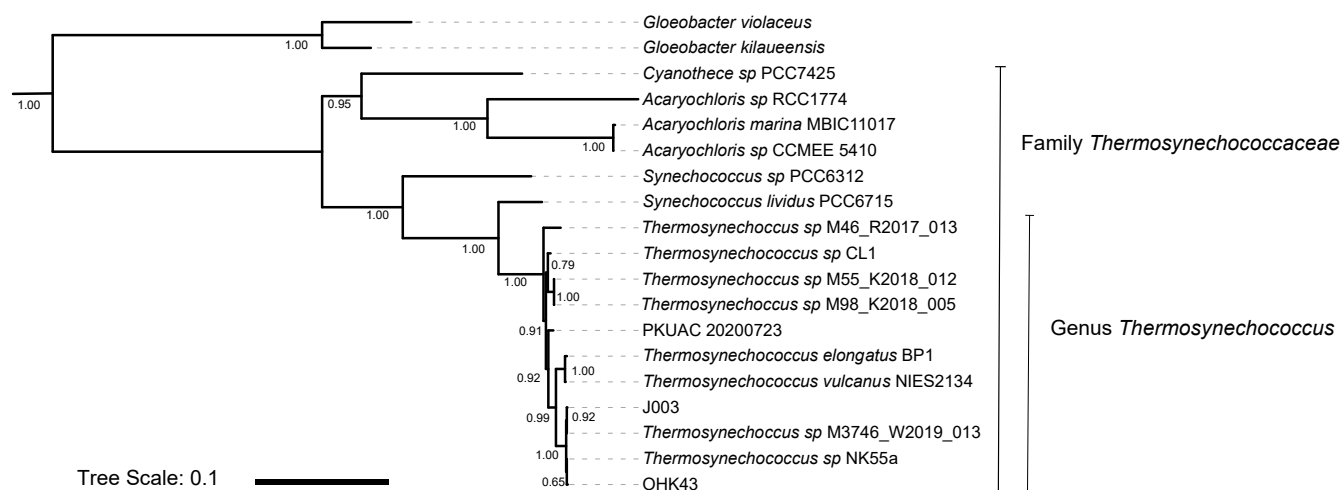


Fig. 1. *Thermosynechococcaceae* phylogeny built with concatenated ribosomal proteins. Branch supports are derived from bootstrapping with BOOSTER and the tree scale bar indicates substitutions per nucleotide site.

Table 3. Numbers of CLOGs per grouping and phylogenetic level. *note that the number of genus specific core CLOGs increases due to the exclusion of some family level genomes at the class level.

	Genus (10 strains)	Family (16 strains)	Class (26 strains)
Core of all genomes analyzed	1737	1225	723
<i>Thermosynechococcus</i> genus specific core	—	14	67
Uniquely shared between 7 <i>Thermosynechococcus</i> (not including J003/Jinata2/OHK43)	1	0	0
Uniquely shared between J003, Jinata 2 and OHK43 only (high iron organisms)	1	1	1

parisons of organisms analyzed at this taxonomic level, for example Wu *et al.* (2018) found that the core genes of species belonging to the genus *Comamonas* account for 18–33% of all genes, and Barajas *et al.* (2019) reported that the core genome within the genus *Streptococcus* ranges in size from 9.6% to 24%. At the species level, Reno *et al.* (2009) found that 69–79% of genes made up the core in *S. islandicus* strains. The high proportion of CLOGs which make up the core leaves few unique CLOGs for each genome, and only slight variations in genome content between genomes are observed: focusing on the organisms from high ferrous iron springs compared to low iron springs for example, only one CLOG is specific to the seven genomes that do not include J003, Jinata 2, and OHK43 and 1 CLOG is specific to J003, Jinata 2 and OHK43 (Table 3, Supplementary Fig. S1).

Golicz *et al.* (2020) suggested that the size of a pangenome is related to organismal lifestyle, with sympatric organisms having open pangenomes with many accessory genes and allopatric organisms having more closed and conserved pangenomes. The *Thermosynechococcus* genus could thus be considered as allopatric as they have a comparatively large core and few shared and unique genes. Since all *Thermosynechococcus* in this study are found in hot springs—which typically have reduced microbial diversity in comparison to other environments such as soils (Ward *et al.*, 1998)—the conserved core and small pangenome of the *Thermosynechococcus* genus may reflect a more limited opportunity for lateral gene transfer, which in turn could lead to less opportunity for lateral gene transfer and smaller genomes.

Chen *et al.* (2020) also noted that differences in horizontal gene transfer (HGT) are related to genome size, with smaller genomes showing less HGT and larger genomes having a greater HGT probability. They also found that hot spring cyanobacteria specifically have smaller genome sizes and less HGT into the genome. Excluding massive gene loss within the *Thermosynechococcus* genus, it is tempting to speculate that the proportionally large *Thermosynechococcus* genus core might be indicative of a more ancient gene repertoire of hot spring cyanobacteria, with other cyanobacteria gaining functionality through HGT over evolutionary timescales. Although still tentative, the observation that thermotolerance is phylogenetically scattered across the cyanobacterial tree of life and occurs mostly in organisms comprising smaller genomes is in line with this hypothesis of substantial gene gain by HGT in many cyanobacterial lineages and a small ancestral core.

In contrast to the genus level where genome size varies from 2.25 MB to 2.64 MB, at the family level it varies up to 8.36 MB (*Acaryochloris marina* MBIC11017, Table 2). Running Proteinortho analyses with the ten genus level *Thermosynechococcus* and the six family level sequences, the shared genes, not core genes, comprise a larger percentage of the genomes for smaller genomes, while unique genes are abundant in larger genomes (Fig. 2a). The overall core is reduced by about 30%, from 1,737 to 1,225 CLOGs.

At the family level, the number of CLOGs shared between the seven *Thermosynechococcus* genus members that do not include J003, Jinata 2 and OHK43 (the strains

from higher iron environments) is reduced from one (found when analyzing at the genus level) to zero, showing that this CLOG is found in other closely related members at the family level. However, the number of CLOGs found only in J003 and OHK43 changes from 23 to 20 CLOGs, highlighting that these are unique even at the family level (Supplementary Table S2).

Genomic differentiation of the Thermosynechococcus genus from other cyanobacteria

At the class level the number of *Thermosynechococcus* genus level core CLOGs increases (67) when compared to the family level (14), and this is due to the exclusion of some family level species at this level (in our analysis the class level comparison was made specifically in reference to a previous study (Beck *et al.*, 2012) for comparative purposes; see Materials and Methods and Table 3). Consistent with this former study, the overall core decreases by almost 60% when class level representatives are added (from 1,737 core CLOGs at the genus level to 723 core CLOGs at the class level). This is expected and in line with larger analyses of microbial genomes which have shown that the continued addition of taxonomic diversity in an analysis leads to increasingly smaller cores (Charlebois and Doolittle, 2004; Lapierre and Gogarten, 2009). Moreover, Beck *et al.* (2018) suggested that the clustering of CLOGs depends on the data analyzed including variability in genome size as well as phylogenetic distance between the analyzed genomes. Overall these results conform with Beck *et al.* (2012), with slight variations attributed to the different genomes used and the different orthology methods and parameters used for each analysis. Finally, we confirm that the class level core is stable when adding a higher diversity of organisms—similar to Beck *et al.* (2018)—as the number of core family CLOGs only slightly decreases when adding more species at the class level (660 core CLOGs were obtained with 16 strains in Beck *et al.* [2012] and 621 core CLOGs with 77 strains in Beck *et al.* [2018]).

Investigating the genes which may differentiate the *Thermosynechococcus* genus from other cyanobacteria, we found that 67 CLOGs appear in all *Thermosynechococcus* but not in other cyanobacteria in our class level analysis (Table 3). However, taking into account that 14 core CLOGs were observed when analyzing with the full compliment of family members, only 8 CLOGs are found to be unique to the *Thermosynechococcus* genus when considering all cyanobacteria analyzed in this paper (see Supplementary Table S1). Amongst the genus CORE proteins observed at the different levels, a number are of unknown function and could be of interest to further biochemical studies. The 8 genes may code for proteins that make the *Thermosynechococcus* unique from other cyanobacteria. Since all the *Thermosynechococcus* analyzed are from hot springs, these genes potentially provide some basis for that lifestyle.

Adaptation of Thermosynechococcus to their respective environments

It is important to note that although the spring source water properties differ, the source water is not necessarily

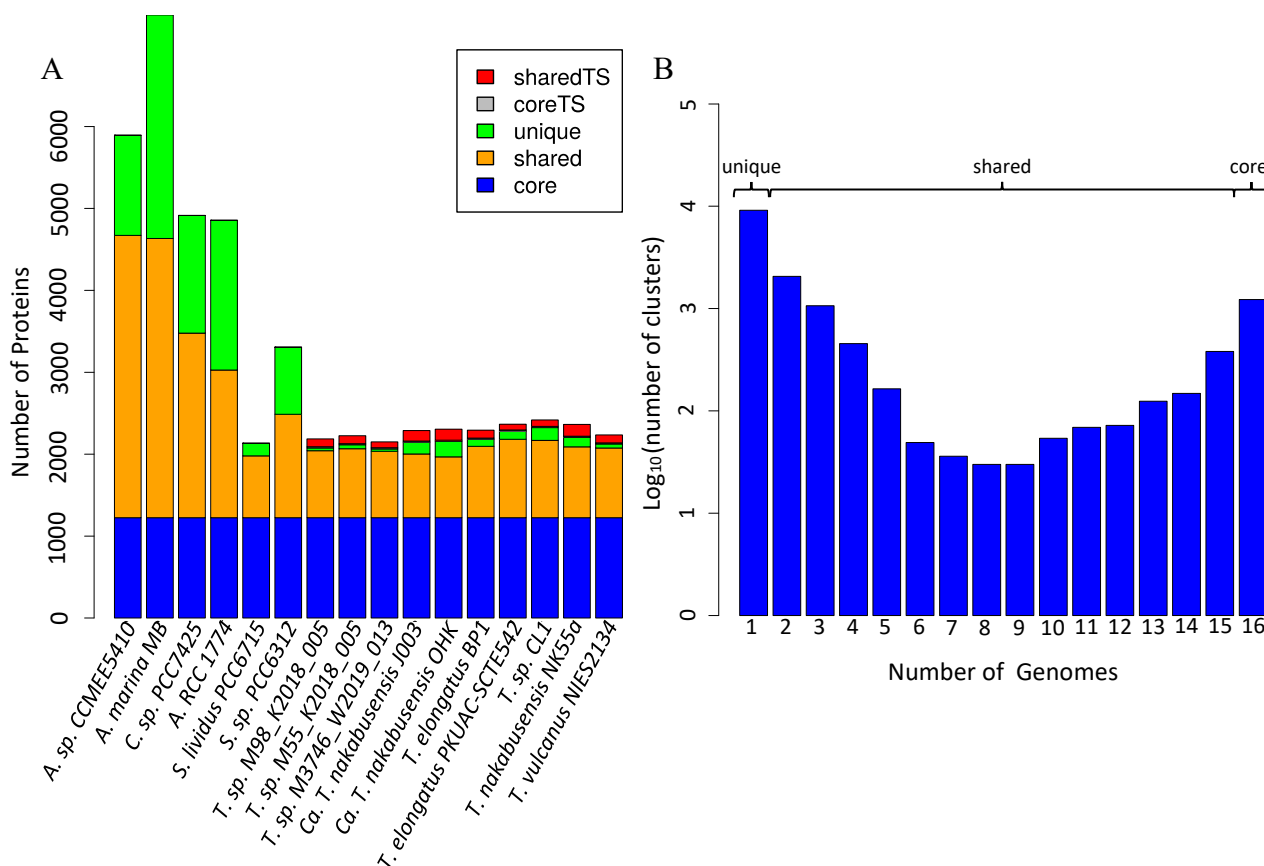


Fig. 2. (A) Thermosynechococcaceae family level comparison of core- and pangenomes. CoreTS indicates CLOGs found in all genus level genomes, sharedTS are CLOGs found in at least 2 but not all genus level genomes and no other genomes. Full species names as mentioned above. (B) number of CLOGs observed at the family level in relationship to the number of genomes in the analysis. The core at the family level is made of the proteins found in all 16 genomes. Shared are those found in more than one but not all genomes, and unique genes is the group of proteins present in single genomes.

where the DNA or the isolated organism originated. Additionally, hot spring flow paths frequently change course, so even if an organism is isolated at one time from one location, it may have been previously growing under a different set of conditions, leading to confusion between the relationship between genotype and phenotype. Acknowledging this, the gross qualities of the hot springs analyzed here differ significantly in pH and the type of reductant present (*e.g.* iron vs. sulfide) (Table 1) implying that organisms inhabiting them experience different environments. We were especially interested in those CLOGs that are shared between the strains from environments with elevated iron concentrations (the genomes J003, Jinata 2 and OHK43) but which are not present in any other cyanobacteria. Previous studies have shown that some cyanobacteria express higher levels of genes involved in iron homeostasis in iron limiting conditions (Cheng and He, 2014), and we investigated the presence or absence of iron related gene products in *Thermosynechococcus* compared to other cyanobacteria using FeGenie and BLAST comparisons. Only one CLOG is uniquely shared between J003, Jinata 2 and OHK43 amongst the genus members. Analyzed at the class level, 19 CLOGs are uniquely shared between Jinata 1 and OHK43. It is notable that this number is higher than the CLOGs uniquely shared between all three strains, and that these 19 CLOGs do not appear in the second strain from Jinata

(Jinata 2) or in the fourth strain of the same species (NK55a). Two CLOGs are uniquely shared between OHK43 and Jinata 2, and six CLOGs are uniquely shared between J003 and Jinata 2. Of these genes, some show high partial identity but low coverage matches with genes from other *Thermosynechococcus*, especially *T. sp.* NK55a (Supplementary Table S2). With our current understandings after considering results from BLAST and FeGenie (Supplementary Table S2) none of the CLOGs uniquely shared by the organisms from high iron hot springs comprises genes that could explain adaptation to elevated iron concentrations. From these analyses we conclude that there is no protein sequence resolvable genomic signature specific to strains from Jinata and OHK hot springs related to iron tolerance or oxidative stress response.

Thermosynechococcus lack genes coding for ferrous iron transport and uptake proteins EfeB, EfeO and EfeU, the metal transport gene ZupT, the cellular iron storage protein Bfr, and the iron regulator active under iron limiting conditions PfsR (Table 4). In all cases the same genes encoding proteins related to ferrous iron uptake (FeoA, FeoB, YfeA and YfeB), ferric iron uptake or transport (ExbD, FutA, FutB and FutC), siderophore iron acquisition (FpvD), metal ion binding (Ho1 and Ho2) and iron starvation acclimation (IsiA) are present (Table 4).

In the absence of a resolvable genetic signature for iron

Table 4. Genes known to be involved in iron regulation within the class Cyanobacteria and their presence in the *Thermosynechococcus* genus. Presence or absence of genes was confirmed with BLAST searches and FeGenie.

Protein	Pfam identifier	Function	Present in <i>Thermosynechococcus</i>	Reference
FeoA	PF04023	ferrous iron uptake	Yes, all	Lau <i>et al.</i> , 2016
FeoB	PF07664	ferrous iron uptake	Yes, all	Lau <i>et al.</i> , 2016
YfeA	PF01297	ferrous iron uptake	Yes, all	Toulza <i>et al.</i> , 2012
YfeB	PF00005	ferrous iron uptake	Yes, all	Toulza <i>et al.</i> , 2012
FpvD	PF00005	siderophore iron acquisition	Yes, all	Brillet <i>et al.</i> , 2012
ExbD	PF02472	ferric iron uptake	Yes, all	Jiang <i>et al.</i> , 2015
PfsR	PF00440	iron regulator under iron limiting conditions	No, but other cyanobacteria	Cheng and He, 2014
FutA	PF13416	ABC-type ferric iron transport	Yes, all	Katoh <i>et al.</i> , 2001
FutB	PF00528	ABC-type ferric iron transport	Yes, all	Katoh <i>et al.</i> , 2001
FutC	PF00005	ABC-type ferric iron transport	Yes, all	Katoh <i>et al.</i> , 2001
EfeB	PF04261	ferrous iron transport	No, but other cyanobacteria	Lau <i>et al.</i> , 2016
EfeO	PF13473	ferrous iron transport	No, but other cyanobacteria	Lau <i>et al.</i> , 2016
EfeU	PF03239	ferrous iron uptake	No, but other cyanobacteria	Lau <i>et al.</i> , 2016
ZupT	PF02535	metal transport (including ferrous iron)	No, but other cyanobacteria	Lau <i>et al.</i> , 2016
Bfr	PF00210	cellular iron storage	No, but other cyanobacteria	Keren <i>et al.</i> , 2004; Cheng and He, 2014
IsiA	PF00421	iron starvation acclimation	Yes, all	Cheng and He, 2014
Ho1	PF01126	metal ion binding	Yes, all	Cheng and He, 2014
Ho2	PF01126	metal ion binding	Yes, all	Cheng and He, 2014

tolerance, we recall that Ionescu *et al.* (2014) proposed that by simply increasing photosynthetic rate and oxygen production, cyanobacteria might protect themselves from ferrous iron by promoting its precipitation at some distance from the cell (Ionescu *et al.*, 2014). In line with this observation, is worthwhile to note that the biomass accumulation in high iron environments like Jinata hot spring is appreciable, with co-occurring visible biomass and dissolved oxygen concentrations that are elevated above what is expected from atmospheric solubility, indicating active water oxidizing photosynthesis (Ward *et al.*, 2019).

The conserved genomic core of Thermosynechococcus in relationship to environmental distribution is unique

In addition to uncovering a highly conserved genome core in a group of organisms with significant environmental distribution, our work is also relevant to historical proliferation of cyanobacteria, since some modern-day hot springs and their biogeochemistries can be used as historical process analogues (Brown *et al.*, 2005; 2007; Ward *et al.*, 2019). Considering contemporary environments, the analysis of *Thermosynechococcus* also provides insight into island biogeography of microbes. Ionescu *et al.* (2010) observed that the speciation patterns of microorganisms are shaped by local community structures and environmental influences, and Bahl *et al.* (2011) additionally suggest a positive correlation between geographic and genetic distance. Papke *et al.* (2003) found that isolated environments such as geothermal springs may lead to evolutionary divergence of closely related *Thermosynechococcus* strains due to island effects, similar to analyses by Whitaker *et al.* (2003), who found similar trends of divergence in hyperthermophilic archaea. Our analysis suggests that geographically widespread organisms belonging to the genus inhabit hot springs with varying geochemistries without genomically recognizable adaptations specific to their site of origin. Instead, the finding of highly conserved genomes within the genus, and furthermore, that the genetic content of the genus is not markedly

different from other cyanobacteria, implies that the genus is inherently flexible and able to grow in the geochemical regimes studied. The large portion of shared genes within the genus provides a genetic basis for the lack of correlation between geographic and genetic distances within the genus found by Papke *et al.* (2003). Apparently, *Thermosynechococcus* is environmentally promiscuous, and have fewer restrictive requirements concerning their distribution. This is in contrast to other groups of organisms, for example the analyzed by Reno *et al.* (2009) who found that the obligately thermoacidophilic *S. islandicus* archaea show a core and pangenome shaped by their geographical distribution.

Outlook

Based on the genome comparisons presented here, a viability test of isolates in environments other than those of their origin is suggested as future work. For example, genus level iron tolerance experiments are proposed to test if strains from low-iron environments can withstand elevated iron. In a similar way, thermotolerance of these organisms could also be investigated. This could help us understand if *Thermosynechococcus* is indeed less restrictive with respect to their geochemical requirements or to identify mechanisms unresolvable by the CLOG approach that account for the geographical distribution.

Data availability for newly described strains

The Whole Genome Shotgun project for OHK43 has been deposited at DDBJ/ENA/GenBank under the accession JACOMP000000000. The version described in this paper is version JACOMP010000000.

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