TITLE: Soil, ocean, hot spring, and host-associated environments reveal unique selection pressures on genomic features of bacteria in microbial communities

AUTHORS:

Peter F. Chuckran¹
Bruce Hungate¹
Egbert Schwartz¹
Paul Dijkstra¹

¹Center for Ecosystem Science and Society (ECOSS), Northern Arizona University, Flagstaff, AZ, USA

*Corresponding author: pfc25@nau.edu, pfchuckran@gmail.com
Free-living bacteria in nutrient limited environments often exhibit small genomes which curb the cost of reproduction – a phenomenon known as genomic streamlining. Streamlining has been associated with a suite of traits such as reduced GC content, fewer 16S rRNA copies, and a lower abundance of regulatory genes, such as sigma (σ) factors. Here, we analyzed these traits from 116 publicly available metagenomes derived from marine, soil, host associated, and thermophilic communities. In marine and thermophilic communities, genome size and GC content declined in parallel, but GC content was higher in thermophilic communities. In soils, the relationship between genome size and GC content was negative, suggesting a different selection pressure on genome size and GC content in soil bacteria. The abundance of σ-factors varied with average genome size, ecosystem type, and the specific functions regulated by the sigma factor. In marine environments, housekeeping and heat-shock σ-factor genes (rpoD and rpoH respectively) increased as genome size declined, and σ-factor responsible for flagella biosynthesis (fliA) decreased, suggesting a trade-off between nutrient conservation and chemotaxis. In soils, a high abundance of fliA and the stress response σ-factor gene (rpoS) was associated with smaller average genome size and often located in harsh and/or carbon-limited environments such as deserts or agricultural fields – suggesting an increased capacity for stress response and mobility in nutrient-poor soils. This work showcases how ecosystem-specific environmental constraints force trade-offs which are then embedded in the genomic features of bacteria in microbial communities, specifically genome size, GC content, and regulatory genes, and further highlights the importance of considering these features in microbial community analysis.
BACKGROUND

Assessing microbial communities through a trait-based framework highlights important relationships between microbes and their environment which may not be detectable through taxonomic analyses alone [1–6]. Notably, genomic characteristics such as genome size, GC content, number of regulatory genes, and number of 16S rRNA gene copies, have been shown to be indicators for growth rates [7], life history strategies [8] and population dynamics [9] of bacteria. Relationships between genomic features and environmental factors such as nutrient usage [9–11], aboveground cover [12, 13], temperature [14], and precipitation [15] have demonstrated the potential utility of genomic traits for assessing the relationship between bacteria and their environment.

The genome size of free-living bacteria may be reduced by a process called genomic streamlining, wherein nutrient limitation selects for smaller genomes as a way to reduce the cost of reproduction [16]. Streamlined genomes are associated with a number of traits which also reduce reproductive costs, most notably a lower GC content, fewer regulatory genes (specifically those encoding σ-factors), smaller intergenic spacer regions, and fewer 16S rRNA gene copies [11]. Consequently, bacteria with streamlined genomes tend to have a higher resource use efficiency and lower growth rates compared to bacteria with larger genomes and more rRNA gene copies [17]. Although experimental genome reduction has shown mixed relationships between genome size and growth rate [18, 19], many studies have found evidence supporting this relationship [7, 20–22]. In this way, genome size could be a functionally useful trait in predicting ecological phenomena such as growth rate. The prevalence of small genomes has long been recognized in marine systems [23] where the streamlined SAR11 clade, with a genome of only ~1.3 Mbp, makes up 25% of all planktonic bacteria [24]. Streamlining is also prevalent in soils,
as represented by the recently described *Candidatus Udaeobacter copiosus*, a ubiquitous taxon in soils, with a genome size of 2.81 Mbp [25].

Temperature can also influence genome size due to increased fitness of small cells at high temperatures [14]. Accordingly, small cells and smaller genomes are typically associated with higher optimal growth temperatures. This relationship is most pronounced in thermophilic communities [26], but has also been demonstrated in marine systems [27–29] and more recently in soils [30]. These patterns between genome size, GC content, and number of 16S rRNA gene copies as a result of temperature-induced genome reduction often resemble patterns in streamlined genomes [14].

Small genomes are also prevalent in host-associated bacteria, however the processes underpinning the reduction in genome size involve several mechanisms, including drift, rapid mutation rate, or other mechanisms, which could be more important than streamlining [9]. In environments where nutrients are abundant but population sizes small, deletions in bacterial genomes are more likely to become fixed in a population [9, 31], a process particularly common in host-associated gut microbiota, where population sizes are small due to isolation [32]. Bacteria subject to higher levels of mutation are more likely to be AT-rich since there is a mutational bias from GC \( \rightarrow \) AT [9, 33–35]. Since the mechanisms driving the evolution of host-associated bacteria often stray from streamlining, genome reduction in host-associated bacteria may yield different patterns in genome reduction. Specifically, streamlining, which is more a directional rather than stochastic process, will often select for specific genes [9].

Much of our knowledge concerning bacterial genomic traits has been derived from cultures or isolates. This present us with substantial bias in our understanding of these relationships [36], especially for genomic traits of bacteria in complex microbial communities [37]. An alternative
approach is to examine genome size on a community level *in situ*. While metagenome-assembled genomes can provide astounding insights into evolutionary processes [38], assembly can be difficult and computationally expensive, making this approach difficult to apply across a large number of communities. Fortunately, genomic traits like GC content, and average genome size, often can be easily estimated, and do not require an extensive knowledge of the taxa within the community. Here we present a comparison of genomic traits from 118 metagenomes from soil, marine, host-associated, and thermophilic systems. We hypothesize that the average genome size in soil microbial communities will be larger than in marine, host-associated, or thermophilic communities, consistent with findings from isolates [14]. We also predict that average genome size and GC content will be positively correlated in free-living soil, marine, and thermophilic communities, consistent with streamlining. Finally, we predict that while both free-living and host-associated communities with small average genome sizes will demonstrate a low GC content, free-living communities will also exhibit additional streamlined traits such as a reduced number of σ-factor and rRNA gene copies.
METHODS

Dataset Curation

Metagenomes from soil, marine, thermophilic, and host-associated communities were downloaded from the Integrated Microbial Genomes & Microbiomes (IMG/M) [39] system. We searched for soil and marine samples that were untreated and from natural systems (i.e. not an incubation or microcosm). For thermophilic samples we searched for communities derived from natural hot-springs, and for host-associated samples we included communities associated with an animal symbiont. We then selected samples which were both sequenced and assembled by the Joint Genome Institute (JGI) and where > 35 Mbp were assembled. Replicates appearing to be derived from a single sample (i.e. identical metadata and sample name) were discarded. In order to limit potential bias introduced by a specific study site or set of protocols of a given study, no more than 4 samples were used from any single geographical location and no more than 14 samples were selected from a single study. Ecosystem type was determined for soil samples using the available metadata and study description. Samples from non-published metagenomes were included with the consent of the primary investigator of the study. In total, 116 samples from 30 different studies were used in this analysis (Supplemental Fig. 1; Supplemental Table 1; [15, 40-62]).

Collection of Genomic Traits

Average genome size for each metagenome was estimated using the program MicrobeCensus (parameters -n 50000000) [63] on QC filtered reads accessed through the JGI Genome Portal [64]. MicrobeCensus uses the abundance of single-copy genes to estimate the
number of individuals in a population, which is then divided by the total number of read base-pairs to provide an estimate of the average genome size in a metagenome.

From IMG/M, we accessed the size of the metagenomic sample (bp), GC-%, total number of 16S rRNA gene copies, and the total number of $\sigma$ factors identified by the KEGG Orthology database (Table 1 [65]). We estimated the number of genomes per metagenome by dividing the total base pair count of the metagenome by the estimated average genome size from MicrobeCensus. The average number of 16S rRNA gene copies per genome and the number of $\sigma$-factors gene copies per genome was then determined by dividing the total number of 16S rRNA or $\sigma$-factor gene copies by the estimated number of genomes.

Data Filtering

To ensure that any observed trends were not heavily influenced by the abundance of nonbacterial genomes, such as large eukaryotic genomes, we assessed the relationship between average genome size and the relative abundance of assembled bacterial reads. The phylogenetic distribution of assigned reads for each metagenome were downloaded from IMG/M and grouped by domain. The relationship between the relative abundance of bacteria and average genome size of the community was then calculated for each ecosystem to assign a cutoff which demonstrated the least amount of bias (as determined by linear regression). As a result, samples where bacteria made up less than < 95% of the assembled reads were discarded.

Since archaeal abundance in thermophilic microbial communities is often high, filtering samples with < 95% bacterial reads discarded a large number of thermophilic samples. Post filtering, only 5 thermophilic samples were left for analysis – a sample size ultimately too small to generate conclusions. Rather than omitting the thermophilic environments from our analysis
entirely, and because small archaeal genomes abundance have been shown to be correlated with higher optimum growth temperatures [14], we decided to include thermophilic samples with > 5% archaeal abundance in several of the comparisons. Although these data do not examine bacterial streamlining specifically, we find that they still provide valuable insight into how genomic traits are distributed in these communities. Mixed thermophilic samples (those including > 5% archaea) are shown separately in figures and analyses. In comparisons of genome size versus bacteria-specific traits, such as 16S rRNA gene copies or abundance of sigma factors, we only report samples where bacteria comprise > 95% of annotated reads.

Analysis

Multiple regression was used to determine the relationship between genome size and genomic characteristics – specifically, GC content, 16S rRNA gene relative abundance, the relative abundance of $\sigma$-factor genes, and the relative abundance of specific $\sigma$-factor variants. Models were constructed with the command lm or lmer from the R (v3.6.1 [66]) package lme4 [67]. For each response variable, we constructed multiple models considering all parameters and interactions. Final models were selected using Akaike information criterion (AIC) values. The addition of a new parameter resulting in a reduction of the AIC value by at least 4 indicated a significantly better fit with increased model complexity.

To assess the abundance of $\sigma$-factor genes between different ecosystems, we used both the multi-response permutation procedure (MRPP) as well as the permutational multivariate analysis of variance (PERMANOVA). The MRPP was conducted using all samples while PERMANOVA was conducted using 11 randomly selected genomes from each ecosystem to
ensure balanced design. Both analyses were conducted using Bray-Curtis dissimilarity matrices constructed from the relative abundance of each \( \sigma \)-factor. To visualize differences in the distribution of different types \( \sigma \)-factors between ecosystems we used nonmetric multidimensional scaling (NMDS) on Bray-Curtis distances. MRPP, PERMANOVA and NMDS were done using the \textit{vegan} package [68] in R (v3.6.1).

\textit{Isolates}

To compare relationships between genomic characteristics of a microbial community with characteristics of isolates, we accessed over 6,000 isolates of bacteria, archaea, and fungi from the IMG/M system in June of 2020. Isolates were selected if they were (1) publicly available; (2) previously published; (3) sequenced by JGI. Metadata was used to group samples into one of three ecosystem types: soil, marine, thermophilic, or host-associated. To avoid potential bias introduced by large studies selecting for specific taxa, we randomly selected no more than 20 isolates from a single study. Relationships between genomic characteristics were analyzed using multiple regression analyses as described above for the analysis of community-level traits. ANOVA was used to assess differences in the distribution of genomic characteristics between isolates and metagenomic averages.
RESULTS

Average Genome Size and GC Content

Average genome size was significantly different between ecosystems (ANOVA; $F_{4,111} = 135.9$, $p < 0.01$). Specifically, average genome size was higher in soils compared to marine, host-associated, or thermophilic communities (Fig. 1a, Tukey’s HSD $p < 0.01$). GC content (%) varied between each ecosystem (ANOVA; $F_{4,111} = 140.3$, $p < 0.01$), and was highest in soil, followed by thermophilic, host-associated, and then marine communities (Fig. 1b). Overall, GC content and average genome size were positively correlated; however, this trend varied between ecosystems (Fig. 1c). A comparison of multiple models, using AIC values as selection criteria, indicated that GC content was best predicted by average genome size, ecosystem, and their interaction ($F_{9,106} = 136.1$, $p < 0.01$, Supplemental Table 2). Specifically, GC content was positively correlated with average genome size in marine and thermophilic communities, negatively correlated in soil communities, and not significantly related in host-associated communities (Fig. 1c). The relationship between average genome size and GC content was offset between marine and thermophilic communities: thermophilic communities had a higher GC content than marine communities with the same average genome size (Fig. 1c). The relationship between GC content and average genome size was strongly driven by the abundance of archaea in the mixed thermophilic samples (Supplemental Fig. 2). In soils, average genome size and GC content were significantly different between ecosystem types (ex. Deserts, grasslands, forests; ANOVA: Mbp - $F_{7,38} = 24.35$, $p < 0.01$; GC-% - $F_{7,38} = 4.986$, $p < 0.01$; Fig. 2).

The average genome size and GC content of the metagenomes fell within the range of isolates from each ecosystem (Supplemental Fig. 3). However, the mean genome size and GC
content derived from metagenomes varied from isolates in both soil and thermophilic environments (ANOVA; p < 0.05), but not in marine environments.

16S rRNA gene copies and Sigma factors

Host-associated communities had the highest number of 16S rRNA gene copies per genome, followed by soils and then thermophilic and marine communities (Supplemental Fig. 4). A comparison of AIC values indicated that ecosystem type alone was the best predictor of 16S rRNA gene copies per genome (Supplemental Fig. 4, Supplemental Table 2).

The relative abundance of σ-factors genes per metagenome changed with estimates of average genome size, however this relationship varied significantly between ecosystems (Fig. 4a; Supplemental Table 2). Average genome size was significantly correlated with the relative abundance of σ-factors in thermophilic environments ($R^2 = 0.49$), but not in soil, marine, or host associated environments ($R^2 < 0.2$; Fig. 4a). The distribution of σ-factor types within a metagenome varied more between ecosystems than within (Fig 3; Fig. 4b; MMRP, $A = 0.34$, $p < 0.01$), and ecosystems differed significantly (Fig 3; Fig. 4b; PERMANOVA, $R^2 = 0.50$, $p < 0.01$).

The relationship between average genome size and the relative abundance of individual σ-factors was dependent on both ecosystem type and the type of σ-factor (Fig. 4c, Supplemental Table 3). In host-associated communities, the relative abundance of only one σ-factor, $\sigma H$, was significantly ($P = 0.018$) negative correlated with average genome size. Abundance of all other sigma factors were unchanged with genome size in host-associated communities (Supplemental Table 3). In soil communities the relative abundance of $rpoH$ per metagenome significantly increased ($P < 0.01$) with larger average genome size, while the relative abundance per
metagenome of rpoS, sigH, sigB, and fliA decreased (P < 0.01). In marine communities, we
found that the relative abundance of fliA, rpoE, and sigH significantly increased (P < 0.01) with
genome size, and the abundance of rpoH, and rpoD significantly decreased (P < 0.01). Due to
the small samples size of thermophilic communities, we did not include the relationships
between σ-factors and average genome size for thermophilic environments; however, correlation
coefficients and statistics for all linear regressions between average genome size and σ-factor
abundance for each ecosystem can be found in Supplemental Table 3. A visualization of average
σ-factor copies per genome can be found in Supplemental Fig. 5.
Our hypotheses were that: 1) average genome size in soil microbial communities would be higher than in marine, thermophilic, and host-associated microbial communities; 2) GC content would be positively correlated with the average genome size of free-living (not host-associated) microbial communities; and 3) host-associated microbial communities exhibiting smaller average genome sizes, although AT-rich, will lack other traits associated with streamlining, such as a reduction in regulatory $\sigma$-factors. The first and third hypotheses were generally supported, however the second hypothesis was rejected since trends between genomic traits varied between environments.

As expected, microbial communities in marine, host, and thermophilic environments had a smaller average genome size than those in soil, consistent with previous findings from studies using bacterial isolates and single-amplified genomes [8, 11, 69]. Since smaller genomes tend to have lower GC content [70], we expected to find a positive correlation between GC content and average genome size for each ecosystem. Surprisingly we only found this relationship in marine and thermophilic communities. In thermophilic communities, this relationship appeared confounded with the presence of archaea, thus making it impossible to distinguish between archaeal abundance or streamlining as a driver for smaller genome size in these extreme environments. It is worth noting that the relationship between genome size and GC content in thermophilic communities was offset higher from marine systems, even for bacterial dominated thermophilic communities. This offset is likely the result of a requirement for thermal stability in hot environments which is provided by the GC triple-hydrogen bonds versus the AT double-bond [71, 72].
Both GC content and average genome size in host-associated communities were low, a common feature of symbiotic bacteria [32]. Although host-associated bacteria in small populations often have AT-rich genomes [9], the relationship between GC content and average genome size was not significant for host-associated communities. Reduced genetic flow in these communities could mean that changes in nucleotide frequency and genome size develop independently in populations. Therefore, these trends might exist within, but not between, communities. In other words, host-associated environments might produce small AT-rich genomes, but these two traits do not covary between communities.

Soil communities exhibited a negative relationship between average genome size and GC content. This does not necessarily exclude streamlining as a driver of genome size in soils but suggests other drivers of genome size and GC content. For instance, fungal reads may reduce the overall GC content of a metagenome while raising estimates of average genome size. We found that fungal isolates generally have a lower GC content than bacteria with a similarly sized genome (Supplemental Fig. 6). Although we attempted to avoid the influence of fungal genomes by limiting our dataset to metagenomes which were dominated by bacteria, it is possible that even a low abundance of large fungal genomes affected our estimates.

Another explanation is that soil microbial communities in nutrient limited environments select for smaller genomes with a higher GC content, which may be advantageous when carbon is limited [73]. A GC basepair has carbon to nitrogen ratio of 9:8 while an AT basepair has a ratio of 10:7. A reduction in GC content therefore decreases the amount nitrogen required for DNA synthesis, which has been suggested as an explanation of the low GC content that is commonly exhibited in marine systems, where nitrogen is limiting [74]. Similarly, small genomes in soil, where C is generally limiting [75, 76], might preferentially select for GC rich
DNA. In this dataset, communities from deserts, agricultural fields, and grasslands had a smaller average genome size and higher GC content (Fig. 2). These environments tend to have lower soil and microbial carbon to nitrogen ratios than forests [77]. Similarly, bacterial communities in forests tended to have larger average genome sizes and lower GC content. Other environmental factors fall along this gradient which might also influence GC content, such as temperature and moisture, which has been shown to influence nucleotide composition in terrestrial plants [78].

Our work shows a potential connection between nucleotide frequency and ecosystem properties, emphasizing the need to develop a more complete understanding of genomic features across soil microbial communities.

We did not find that the relative abundance of $\sigma$-factors was associated with average genome size. However, we did observe that marine communities maintained a lower abundance of $\sigma$-factor gene copies in comparison to other ecosystems, even when average genome size was comparable. One explanation is that the reduction of $\sigma$-factor gene copies is particularly effective in reducing reproductive costs in marine systems. Marine systems are considered to be nutrient poor relative to soils and a general reduction in the proportion of $\sigma$-factors in bacterial genomes may function as an adaptation to nutrient constraints. We also found many trends between average genome size and the abundance of specific $\sigma$-factor genes in marine communities. In marine metagenomes, the relative abundance per genome of $rpoD$ and $rpoH$, which encode for $\sigma^D$ and $\sigma^H$ respectively, was negatively correlated with average genome size. These trends are perhaps caused by the abundance of the streamlined SAR11 clade, which only contain $\sigma^D$ and $\sigma^H$ [24]. Conversely, the abundance of the gene $fliA$, which encodes for the $\sigma^{28}$ and regulates flagella biosynthesis [79], increased with average genome size. This relationship tracks a copiotroph-oligotroph framework in marine systems, wherein nutrient scarcity selects for smaller, more
streamlined, individuals while increased nutrient availability selects for larger individuals capable of chemotaxis [17, 80].

In soils, the relative abundance of many $\sigma$-factors were negatively correlated with estimates of average genome size. Most notably, we observed a decrease in the relative abundance of rpoS ($\sigma^S$) but no significant change in the abundance of rpoD ($\sigma^D$) with increasing average genome size. The balance between rpoS and rpoD may be a trade-off between stress tolerance and growth [81, 82]. A higher ratio of rpoS to rpoD has been shown to increase the cell’s capacity to cope with stress but limit its ability to grow on a variety of carbon sources [82–84]. We see this reflected in the environments from which the metagenomes were samples, with microbial communities from high stress environments, such as deserts, having a higher abundance of rpoS compared to lower-stress carbon-rich environments, such as forests (Supplemental Fig. 7).

Surprisingly, we found a high abundance of fliA gene copies in soil communities with smaller genomes, several of which were sourced from desert environments. Motility may be more valuable in nutrient limited environments, whereas in environments with high nutrient inputs, nutritional competency may be more paramount. However, these results contrast with the commonly held notion that chemotaxis is most prevalent in mesic soils. One explanation is that motility may be especially important when water availability is ephemeral. A greater number of regulatory mechanisms would therefore be advantageous as it would allow for a rapid response to periodic pulses of moisture. Another possibility is that bacteria utilize biofilms surrounding fungal hyphae, or “fungal highways” [85], which could explain the persistence of flagellated bacteria even in xeric environments [86].
Conclusion

We found a number of compelling relationships between ecosystem parameters and genomic traits of a microbial community, most notably with genome size, GC content and the distribution of $\sigma$-factors. Several of these relationships align with evolutionary mechanisms which relate to known drivers in these environments, such as streamlining in oceans and drift in endosymbionts. Still, we observed trends in soils which were not in-line with known mechanisms of genome reduction, emphasizing the need to develop an understanding of the controls of genomic features in soils. Finally, this work highlights the utility of understanding microbial communities through the lens of genomic traits in addition to genetic potential.
Availability of data and materials

All data used in this analysis were accessed from the IMG/M database (https://img.jgi.doe.gov/cgi-bin/m/main.cgi). Study names and GOLD project IDs can be found in Supplementary Table 1. Data which was not both publicly available and previously published was used with permission from the listed authors on IMG/M. This publication does not act as a primary publication for included studies and use of these data requires consent from the team which generated these data.

Competing interest

The authors declare that they have no competing interests

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Authors’ contributions
PC gathered and analyzed the data and wrote the manuscript. PD assisted in study design, writing, and editing. BH and ES contributed to writing and editing the manuscript. All authors read and approved the final manuscript.

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Table 1: Gene name, description, and KEGG ortholog identifier (K numbers) for each σ-factor used in the analysis.

| σ-factor gene | Functions regulated by σ-factor | K Number  |
|---------------|---------------------------------|-----------|
| rpoD          | Primary sigma factor, "Housekeeping" [87] | KO:K03086 |
| rpoE          | Envelope stress [88]             | KO:K03088 |
| fliA          | Flagella biosynthesis [79]       | KO:K02405 |
| rpoH          | Heat shock [89]                  | KO:K03089 |
| sigI          | Heat shock [90]                  | KO:K03093 |
| sigH          | Heat shock, oxidative stress [91] | KO:K03091 |
| rpoN          | Nitrogen assimilation [92, 93], Motility [93], Quorum sensing [94] | KO:K03092 |
| rpoS          | Stress response [95, 96], Stationary phase [97] | KO:K03087 |
| sigB          | Stress response [98], Stationary phase [99] | KO:K03090 |

Figure 1: Average genome size and GC-content calculated from environmental metagenomes. (A) Violin-plots and boxplots of the average genome size (Mbp) of microbial communities in different ecosystems. (B) Violin-plots and boxplots showing GC-% between systems. (C) GC-% as a function of average genome size (Mbp) of a metagenome, separated by system. Point shape and outline represent source system; point fill represents system including thermophilic samples with archaea.

Figure 2:
GC content (%) as a function of average genome size (Mbp) in soils, with color indicating source environment.

Figure 3:
The relative abundance $\sigma$-factors in a metagenome separated by ecosystem. Each bar represents the abundance of $\sigma$-factors in a single metagenome, and metagenomes are ordered from smallest to largest (left to right) for each ecosystem.

Figure 4:
The relative abundance of $\sigma$-factors ($\sigma$-factor count / gene count) as a function of average genome size and system. (A) The relative abundance of all $\sigma$-factors (total $\sigma$-factor count / gene count) in a metagenome against average genome size. Source environment indicated by color for host associated (red), soil (green), thermophilic (orange) and marine (blue) communities. (B) NMDS of Bray-Curtis distance of the relative abundance of $\sigma$-factors ($\sigma$-factor count / total gene count) from a metagenome. (C) The relative abundance ($\sigma$-factor count / total gene count) of 9 $\sigma$-factors (rows) versus average genome size, separated by environment (columns).
Figure 1:

A. Box plot showing the average genome size (Mbp) for different systems: Thermophilic including archaea, Thermophilic, Soil, Marine, and Host associated.

B. Box plot showing the GC% for different systems: Thermophilic including archaea, Thermophilic, Soil, Marine, and Host associated.

C. Scatter plot showing the relationship between GC% and average genome size (Mbp) for different systems. The equations for the regression lines are:

- $R^2 = 0.078$
- $R^2 = 0.67$
- $R^2 = 0.36$
- $R^2 = 0.53$

The system community types are:
- Host associated
- Marine
- Soil
- Thermophilic
- Thermophilic including archaea
Figure 2:
Figure 3:
Figure 4: