The *Pseudoalteromonas* multipartite genome: Distribution and expression of pangenome categories, and a hypothesis for the origin and evolution of the chromid

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Running title: The *Pseudoalteromonas* bipartite genome

Key words: *Pseudoalteromonas*, pangenome, multipartite, chromid, Alteromonadales, secondary replicons.

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Abstract
Bacterial genomes typically consist of one large chromosome, but can also include secondary replicons. These so-called multipartite genomes are scattered on the bacterial tree of life with the majority of cases belonging to Proteobacteria. Within the class gamma-proteobacteria, multipartite genomes are restricted to the two families *Vibrionaceae* and *Pseudoalteromonadaceae*. Whereas the genome of vibrios is well studied, information on the *Pseudoalteromonadaceae* genome is much scarcer. We have studied *Pseudoalteromonadaceae* with respect to the origin of the chromid, how pangene categories are distributed, how genes are expressed relative to their genomic location, and identified chromid hallmark genes. We calculated the *Pseudoalteromonadaceae* pangenome based on 25 complete genomes and found that core/softcore are significantly overrepresented in late replicating sectors of the chromid, regardless of how the chromid is replicated. On the chromosome, core/softcore and shell/cloud genes are only weakly overrepresented at the chromosomal replication origin and termination sequences, respectively. Gene expression is trending downwards with increasing distance from the chromosomal oriC, whereas the chromidal expression pattern is more complex. Moreover, we identified 78 chromid hallmark genes, and BLASTp searches suggest that the majority of them were acquired from the ancestral gene pool of Alteromonadales. Finally, our data strongly suggest that the chromid originates from a plasmid that was acquired in a relatively recent event. In summary, this work extends our knowledge on multipartite genomes, and helps us understand how and why secondary replicons are acquired, why they are maintained, and how they are shaped by evolution.

Introduction
Multipartite genomes are recognized by the concurrent presence of multiple replicons, i.e., cells contain one or more large replicon in addition to the chromosome (Harrison et al. 2010). The majority of bacteria with multipartite genomes are associated with high tolerance to abiotic stresses, or are associated with animals, human or plants as pathogens or symbionts (Misra et al. 2018). This observation, in addition to other data, have prompted scientists to hypothesize that multipartite genomes play crucial roles in the successful spread and establishment of bacteria into a broad range of ecological niches (Heidelberg et al. 2000). A striking and well-studied example is the bacterium *Vibrio fischeri*. Some strains colonize the light-emitting organ of squid (e.g., the Hawaiian bobtail squid *Euprymna scolopes*), and produce bioluminescent light that enables the host to evade predators by counter-illumination (Soto and Nishiguchi 2014). Other strains are in contrast pathogens, which
apparently is made possible due to the presence of a gene capture system, a superintegron and pathogenicity islands located on the chromid (Soto and Nishiguchi 2014; diCenzo et al. 2019). Therefore, a link between the two-replicon genome architecture and the bacteria’s lifestyle has been suggested. The reality is however, that although carrying one or more extra replicon may promote new opportunities for a bacterium to move into new niches, other bacteria thrive in the same environment without additional large replicons, thus demonstrating that multipartite genomes are probably not necessary to succeed in that environment. Our general understanding of the origin, evolution and functional roles of multipartite genomes remains fragmented, and multiple other equally likely hypotheses have been proposed to explain their existence. For example, carrying genes on more than one large replicon allows for replicon-specific gene dosage, and consequently replicon-specific gene expression regulation (e.g., Couturier and Rocha 2006; Dryselius et al. 2008). Also, it has been suggested that the presence of multiple replicons allows bacteria to contain larger genomes, and may reduce the time required to complete replication thus allowing for rapid cell growth and division (diCenzo et al. 2014). Finally, we recently proposed a hypothesis that the presence of two large replicons allows for intracellular spatial separation of different categories of genes, and that there is a link between the skewed gene placement and their function (Sonnenberg et al. 2020). The underlying reason for the separated distribution of gene categories in 3D is however likely very complex, and not easy to dissect.

Given that one or more of the hypotheses above are correct, then it is not surprising that bacteria with multipartite genomes are indeed widely distributed. They are as of today found scattered across the bacterial kingdom, into six of 81 phyla listed in the NCBI taxonomy system (i.e., Proteobacteria, Bacteriodetes, Actinobacteria, Firmicutes, Deinococcus-Thermus and Spirochaetes) (diCenzo and Finan 2017). It is highly likely however that more examples of multipartite genomes remain to be discovered, especially when considering that the number of complete genomes in the databases is still relatively low (approx. 23,000), and dominated by Proteobacteria (57%) and Terrabacteria (34%). One hundred one of 127 multipartite genomes group within the phylum Proteobacteria (diCenzo and Finan 2017), which means that the remaining 26 genomes are distributed among five other phyla. Inside the class gamma-proteobacteria, multipartite genomes are restricted to Vibrionaceae and Pseudoalteromadaceae. The multipartite genome of Vibrionaceae consists of one large circular chromosome (4,1 – 2,7 Mb) known as Chromosome 1 (Chr 1), and one smaller circular replicon (2,3—0,7 Mb) knows as Chromosome 2 (Chr 2), hereafter referred to as the Vibrionaceae
chromid, in accordance with the nomenclature by Harrison et al. (2010). Replication of the *Vibrionaceae* chromosome and chromid is precisely coordinated by a mechanism that is partly understood. Briefly, when the replication fork approaches *crtS* (chromid replication triggering site) in Chr 1, a hitherto unknown mechanism triggers replication of the chromid (Val et al. 2016; Kemter et al. 2018), there is a brief pause in Chr 1 replication before the cycle ends in a synchronized termination of replication. In fast-growing bacteria, such as *Vibrio cholerae* and *Vibrio natriegens*, replication results in higher gene copy numbers of genes surrounding the origin of replication of Chr 1 (*ori1*) (this is known as “gene dosage effect”). Consequently, expression of genes typically decreases with increasing distance from *ori1* (Dryselius et al. 2008; Toffano-Nioche et al. 2012). This correlation does not necessarily apply to slow growing bacteria, or fast-dividing bacteria grown under poor (sub-optimal) conditions. We recently published a study where we calculated the *Vibrionaceae* pangenome based on 124 genomes to study how the four pangene categories (core, softcore, shell and cloud) are distributed on the genome (Sonnenberg et al. 2020). The analysis showed that core/softcore genes are typically found clustered around *ori1*, whereas shell and cloud genes densely populate terminus-proximate regions on Chr 1. On the chromid, genes are more randomly distributed, with no strong distribution pattern. On Chr 1, gene expression levels strongly correlate with distance to *ori1*, with higher expression levels around *ori1*. Interestingly, under slow-growing conditions all categories, except core genes, contributes to this pattern. This prompted us to question whether the observed gene distribution and expression patterns are specific to *Vibrionaceae*, or represent a general trend among bacteria with multipartite genomes.

The family *Pseudoalteromonadaceae* represents an excellent opportunity to study multipartite genomes, e.g., how the genes are distributed and expressed, and its origin and evolution. As of March 2021 the Refseq database contains 53 complete *Pseudoalteromonadaceae* genomes. All genomes are bipartite and consist of one chromosome (3.1–4.9 Mb) and one chromid (0.6–1.8 Mb). Bosi et al (2017) calculated the *Pseudoalteromonas* pangenome based on 38 genomes (Bosi et al. 2017). Briefly, they described the pangenome as open and with a large percentage (80%) unique genes. Further, they estimated the last common ancestor (LCA) of *Pseudoalteromonas* to contain an estimated 2,999 genes, compared to an average of 4,245 genes in the present-day genomes, which support that the genome has undergone a considerable expansion. More recently, Liao et al (2019) studied the evolution of the *Pseudoalteromonas* genome (Liao et al. 2019). Using a phylogenetic approach and timescale analysis, they showed
that the chromosome and chromid have coexisted, probably since *Pseudoalteromonas* diverged from the putative LCA 500 million years ago. The chromid apparently originates from a megaplasmid that over time obtained essential genes (Médigue *et al.* 2005; Rong *et al.* 2016; Liao *et al.* 2019; Xie *et al.* 2021).

In summary, *Vibrionaceae* and *Pseudoalteromadaceae* represent the only two families from the gamma-proteobacteria class with multipartite genomes. Whereas the *Vibrionaceae* genome is well studied, the information on *Pseudoalteromadaceae* is scarce. In this work we set out to gain insight into how pangene categories are distributed on *Pseudoalteromonadaceae* chromosomes and chromids, how genes are expressed relative to their genomic location, which genes can be regarded as hallmark genes of the chromid, and the origin and evolution of the chromid. We present data that support observations on gene distribution and global expression from other bacterial chromosomes, as well as data showing chromid-specific patterns that suggest specific roles of secondary replicons. Several pieces of evidence suggest the likely source of the chromid and its hallmark genes.

**Materials and Methods**

**Genome retrieval and gene annotation**

A total of 25 *Pseudoalteromonas* genomes that were available at the onset of this project (mid 2019) were publicly available at the National Center for Biotechnology Information (NCBI) and downloaded from the RefSeq database at NCBI (O’leary *et al.* 2015) (see File S1 for a complete list). The following genomes were excluded from the analysis: *Pseudoalteromonas atlantica* T6 (GCF_000014225.1) misplaced into *Pseudoalteromonas*, and later re-classified and renamed to *Paraglaciecola atlantica* T6. *Pseudoalteromonas atlantica* ECSMB14104 and *Pseudoalteromonas marina* ECSMB141043 are assembled into one contig, and the nature of their chromids could not be reliably resolved using Mauve. All genome sequences were re-annotated using RAST (Rapid Annotation using Subsystem Technology) version 2.0 (Aziz *et al.* 2008) with default settings. Mauve (Darling *et al.* 2004) was used to align genomes that were annotated with only one replicon.

**Phylogenetic analysis**

Phylogenetic relationships between Alteromonadales genomes were inferred using the nucleotide sequences *gyrB, recA, rpoD, recN* and *topA* as described earlier (Busch *et al.* 2019), and included the seven families *Alteromonadaceae, Colwelliaceae, Idiomarinaceae, Moritellaceae,*
Pseudoalteromonadaceae, Psychromonadaceae and Shewanellaceae (see Figure S1 for complete phylogeny). The nucleotide sequences were aligned using MUSCLE (Edgar 2004). Only unambiguously aligned positions were kept using BioEdit (Hall 1999), which resulted in a 9,216 nt sequence alignment. MEGAX was used to generate a Maximum Likelihood (ML) tree, with the settings GTR (General Time Reversible) model, Gamma Distributed with Invariant (G+I), and Bootstrap with 1,000 pseudoreplicates (Kumar et al. 2018; Stecher et al. 2020). An ML-phylogeny of Pseudoalteromonas was based on 469 single copy marker genes identified by EzTree (Wu 2018). The robustness of nodes was tested with a bootstrap analysis inferred from ML-GTR+G+I.

Pangenome calculations

To classify the annotated Pseudoalteromonas protein sequences into four categories (core, softcore, shell and cloud genes) we performed pangenome analysis using the software package GET_HOMOLOGUES (v3.1.0 (20180103)) (Contreras-Moreira and Vinuesa 2013). The clustering algorithm MCL was used to cluster homologous protein sequences. The parameter “minimum percent sequence identity” was set to 50 and “minimum percent coverage in BLAST query/subj pairs” was set to 75 (default).

Mapping of core, softcore, shell and cloud genes on the Pseudoalteromonas genome

To study the distribution of core, softcore, shell and cloud genes of Pseudoalteromonas, the chromosome and chromid sequences were divided into 4, 6, 8, 10 and 12 equally sized sections, with sector one starting at origin of replication (gidA on the chromosome and parA on the chromid). For each sector, the number of core, softcore, shell and cloud genes were counted. The number of genes in each sector was then divided on the total gene number per replicon (probability of a gene belonging to a sector). The probability of a gene belonging to a sector given equal distribution between sectors was calculated for each of the 4, 6, 8, 10 and 12 sized sectors (1 divided on numbers of sectors). Then, the log10 ratio was calculated of the probability of a gene belonging to a sector divided by the probability given an equal distribution of genes between all sectors. Only a summary of the results when chromosomes and chromids are divided into six sectors is presented in the paper itself. The summary was made by calculating log10 ratio of: The probability of a gene belonging to a sector on average (Average #genes in a sector /Average total #genes) / The probability given an equal distribution of
genes between all sectors (1/#sectors). See File S2 for data. Kruskal-Wallis test and Dunn’s test were used to perform pairwise comparisons of number of genes between all sections (see File S3 for data).

Gene expression analyses
RNA-seq datasets from *P. fuliginea* BSW20308 grown at three different temperatures, i.e., 32° (BioSample accession no. SRR11593421, SRR11593421, SRR11593422), 15° (sample no. SRR11593423, SRR11593424, SRR11593425) and 4° (sample no. SRR11593426, SRR11593427, SRR11593428) (Liao et al. 2019) were downloaded from the NCBI Sequence Read Archive (Leinonen et al. 2011) and analyzed. The quality of the reads was checked using FastQC (Andrews 2010). EDGE-pro v1.0.1 (Estimated Degree of Gene Expression in Prokaryotes) (Magoc et al. 2013) in Galaxy was used to align cDNA reads to the genome assembly (no. GCF_000310105.2) and estimate gene expression as reads per kilobase per million (RPKM) for all protein coding sequences (CDS). The RPKM values were then used to calculate the \( \log_2 \) ratio RPKM CDS:RPKM median to make global expression maps for each of the three datasets. To identify which pangene categories contribute to the gene expression pattern, the chromosome was divided into “upper” and “lower” halves, and the chromid were divided into “upper” and “lower” halves, as well as “right” and “left” halves, and the RPKM median value for each pangene category was calculated (File S4).

BLASTp searches
Homologs of chromid hallmark genes were identified by BLASTp when using the non-redundant database, and excluding the *Pseudoalteromonadaceae* family (taxid: 267888), with the thresholds: e-value < 1e-15, sequence identity > 30% and sequence coverage >70%. (see File S6).

Statistical analysis
Statistical analysis was performed using R in RStudio (RStudio Team 2021). Kruskal-Wallis test and Dunn’s test were used to perform pairwise comparisons of number of genes in replicon sections. The tests were chosen because the data did not follow a normal distribution, and sample sizes were low. R’s kruskal.test() function for the rank-based non-parametric Kruskal-Wallis test and the dunn.test() function for post hoc Dunn’s test were used (see File S2 and S3 for data). Significant difference of gene expression between replicon halves and replicons was performed using R’s wilcox.test().
function for unpaired Wilcoxon signed-rank tests (see File S4 for data). For all analyses \( P \)-values were Bonferroni corrected for multiple comparisons using R’s `p.adjust()` function.

Results

**Pseudoalteromonadaceae branches off from families with multipartite genomes**

Figure 1 shows the overall phylogenetic relationships between bacterial families and genera that form the order Alteromonadales (see Figure S1 for complete phylogeny). The ML-tree (GTR+G+I model) was based on the concatenated nucleotide sequences of *gyrB*, *recA*, *rpoD*, *recN* and *topA* from selected bacteria from the seven families *Alteromonadaceae*, *Colwelliaceae*, *Idiomarinaceae*, *Moritellaceae*, *Pseudoalteromonadaceae*, *Psychromonadaceae* and *Shewanellaceae*. The analysis shows that each genera and family forms well supported monophyletic groups, similar to previous studies (Williams *et al.* 2010; Martin *et al.* 2015). Notably, the family *Pseudoalteromonadaceae* (includes only the genera *Pseudoalteromonas*), which exclusively contains bacteria with multipartite genomes, branches off from the monopartite genome-containing clades as a crown group together with its sister *Alteromonadaceae*. None of the bacteria outside of *Pseudoalteromonadaceae* contain multipartite genomes. These two observations strongly support that the chromid was acquired by the most recent LCA of *Pseudoalteromonadaceae*, likely in a single event (indicated with an arrow in Figure 1). A single origin of the chromid is supported by a phylogenetic analysis that showed congruent phylogenies between the chromosome and chromid (Liao *et al.* 2019). Finally, two separate estimates of time since divergence suggest that *Pseudoalteromonadaceae* branched off approximately 500, and 502-378 million years ago (Liao *et al.* 2019; Xie *et al.* 2021). Compared to *Vibrionaceae*, which also exclusively contains multipartite genomes, the birth of *Pseudoalteromonadaceae* is relatively recent. The time since divergence of *Vibrionaceae* is estimated to approximately 1,100-900 million years ago (Xie *et al.* 2021).

**The Pseudoalteromonadaceae pangenome contains 1,399 core genes**

The definite point of origin of the *Pseudoalteromonadaceae* chromid prompted us to study the multipartite genome in more detail, e.g., to identify where the chromid replicon was acquired from, and how it has evolved after its acquisition. We used a pangenome approach as previously described...
(Sonnenberg et al. 2020). Briefly, available complete genomes were downloaded and re-annotated using RAST (Aziz et al. 2008). Genome datasets were then used to cluster orthologous groups of protein sequences based on the MCL algorithm (van Dongen 2000) in GET_HOMOLOGUES (Contreras-Moreira and Vinuesa 2013). By using 25 available complete Pseudoalteromonadaceae genomes, which were available by the onset of the calculations (see File S1 for complete list), we found a total of 24,991 clusters. The clusters were sub-categorized into 1,399 core (encoded by all 25 genomes), 1,606 softcore (encoded by ≥23 genomes), 7,688 shell (encoded by ≤22 and ≥3 genomes), and finally 15,697 cloud (encoded by ≤2 genomes). This result is comparable to the calculations reported by Bosi et al. (2017), based on 38 Pseudoalteronas genomes (mostly draft genomes), which identified a total of 22,530 clusters, sub-divided into 1,571 core (encoded by all 38 genomes), 2,901 shell (encoded ≤3 and ≥2 genomes) and 18,058 cloud (encoded by one strain) (Bosi et al. 2017).

The distribution of core/softcore genes on the Pseudoalteronas chromid strongly correlates with the direction of replication

To establish the distribution pattern of Pseudoalteronas pangenes, we mapped all representative genes from the four pangene categories core, softcore, shell and cloud to their chromosomal or chromidal locations. First, chromosome and chromid sequences were divided into 4, 6, 8, 10 or 12 equally sized sectors (or bins), with sector one starting from the origin of replication and proceeding clockwise. For each sector, the number of genes from each category were counted. At least for primary replicons, previous data from other bacterial families (Comandatore et al. 2019; Kopejtka et al. 2019; Sonnenberg et al. 2020), have shown that core/softcore genes densely populate regions that are replicated early in the replication cycle, and we hypothesized that Pseudoalteronas would generate a similar distribution pattern. Notably, a recent study showed that most Pseudoalteronas chromids are replicated unidirectionally, except for Pseudoalteronas spongiae and Pseudoalteronas piratica chromids, which are replicated bidirectional (Xie et al. 2021).

Figure 2A shows the result mapped onto a Pseudoalteronas ML-phylogeny based on 469 single copy marker genes identified by ezTree (Wu 2018). Heatmaps summarize the result for Clade 1 (unidirectional replication of chromid), and for Clade 2 (bidirectional replication of chromid). The heatmaps are based on average values from the 25 analyzed genomes (see Materials and methods). Only data for chromids divided into 6 sectors are shown (see File S2 for all datasets). The Kruskal-
Wallis and the Dunn’s post hoc tests were used to identify significant over- or underrepresentation of gene numbers between all pairs of sectors (see File S3). The main finding is that core/softcore genes densely populates late replicating chromidal sectors, regardless of if chromids are replicated uni- or bidirectionally, which is surprising and opposite of what we expected. Specifically, for unidirectionally replicated chromids (Clade 1), core/softcore genes are strongly overrepresented in sector 6, and underrepresented in sectors 2 and 3. For bidirectionally replicated chromids (Clade 2), core/softcore genes are strongly overrepresented in sectors 3 and 4, and underrepresented in sectors 2 and 6.

None of the pangene categories are, in contrast, significantly over- or underrepresented in specific regions of the chromosome. Instead, core/softcore genes are only weakly overrepresented in sectors 1, 5 and 6 (near origin of replication), shell genes are weakly overrepresented in sectors 3 and 4 (near terminus of replication), and cloud genes are weakly overrepresented in sector 3. The general pattern is therefore similar to, but less pronounced than what has been reported for e.g., *Vibrionaceae* (Sonnenberg et al. 2020) *Klebsiella pneumonia* (Comandatore et al. 2019) and *Rhodobacteraceae* (Kopejtka et al. 2019).

In summary, by dividing the two *Pseudoalteromonas* replicons into 4–12 sectors and calculating the log$_{10}$ ratio of the probability of a gene belonging to a sector divided by the probability given an equal distribution, we showed that core/softcore genes are significantly overrepresented in late replicating sectors of the chromid, regardless of how the chromid is replicated, i.e., unidirectionally (Clade 1 strains) or bidirectionally (Clade 2 strains). Chromosomal genes are instead more evenly distributed into each sector of the replicon.

**Gene dosage is in effect on the Pseudoalteromonas fuliginea BSW20308 chromosome, but not on the chromid**

It is well established that the copy number of ori-proximate genes can increase during rapid growth due to the formation of multiple concurrent replication forks, which in turn result in multiple copies of the replicon (e.g., a chromosome), and increased gene expression. This is known as the “gene dosage effect”. To date, this has been described for the *Vibrionaceae* Chr 1 (Rasmussen et al. 2007; Srivastava and Chatteraj 2007; Dryselius et al. 2008; Toffano-Nioche et al. 2012), *Escherichia coli*, *Bacillus subtilis* and *Streptomyces* (Couturier and Rocha 2006; Lato and Golding 2020). To establish if a gene dosage effect is in play in *Pseudoalteromonas* (for the chromosome, the chromid or both),
we downloaded data from one of two available RNA-seq experiments stored at the NCBI Sequence Read Archive (Leinonen et al. 2011). In the selected experiment, P. fuliginea BSW20308 was grown in Difco marine broth 2216 and harvested at 4°C (lowest temperature with growth), 15°C (optimal growth), or 32°C (maximum temperatures with growth) (Liao et al. 2019). These datasets therefore provide an excellent chance to test gene dosage effects at fast and slow growth, which is highly relevant because gene dosage has been reported to be particularly strong at rapid growth (and therefore rapid replication). The three RNA-seq datasets (each in triplicates) were analyzed as previously described (Sonnenberg et al. 2020). Briefly, cDNA reads were mapped onto the P. fuliginea BSW20308 genome (assembly no. GCF_000310105.2) and reads per kilobase per million (RPKM) was calculated for all protein coding sequences (CDS).

Figure 3 shows global expression maps of the chromosome and chromid when P. fuliginea BSW20308 was grown at 4°C, 15°C, or 32°C. Data points (log₂ ratio RPKM CDS:RPKM median) are centered around the RPKM median. Moreover, for each plot a trend line averaged over a sliding window of 100 data points was added to show the overall direction of the data. Expression from the primary replicon (i.e., the chromosome) is trending downwards starting from ori1 and ending at ter1, with a low point at position 1,734,472. In other words, RPKM values are typically higher on the upper half compared to the lower half, which is expected if gene dosage is in effect on a bidirectionally replicated chromosome. This finding is strongly supported by the Wilcoxon signed-rank test (P-adj ≤ 0.05) (see File S4). Expression from the chromid is elevated in the intermediate and late replicating regions. This expression pattern is opposite of what is expected if gene dosage is in effect on a unidirectionally replicated chromid, as in this case. If gene dosage was in effect then overlapping replication cycles would increase the number of DNA copies on the chromidal half (i.e., the “right” half) which is replicated first. The Wilcoxon test does not, however, support significant difference in gene expression neither between upper and lower halves, or left and right halves (see File S4).

In summary, we found that gene dosage appears to be in effect on the Pseudoalteromonas chromosome, but not on the chromid. This applies for all three tested temperatures, 4°C, 15°C, or 32°C, which can be regarded as the minimum, optimum or maximum growth temperatures, respectively.

All pangene categories contribute to higher gene expression on the upper half of the Pseudoalteromonas chromosome under optimal growth temperature

To establish which pangene categories contribute to the gene dosage effect on the P. fuliginea BSW20308 chromosome, we calculated the RPKM median value for each pangene category (Table
1). The Wilcoxon signed-rank test strongly support ($P$-adj $\leq$ 0.05) that all four pangene categories contribute, when *P. fulginea* BSW20308 is cultured at optimal conditions (15°). Interestingly, when grown at sub-optimal conditions (4° and 32°), the same test identifies only shell genes as significant contributors (see Table 1). The RNA-seq data further shows that RPKM median values of core and softcore genes are generally higher than that of shell and cloud genes (see File S4), and this is valid for all three datasets except for the chromosome when grown under 32°. As expected, RPKM values are generally highest when grown at optimal temperature (15°), slightly lower at 4° and lowest at 32°. Overall, expression from chromosomal genes is higher compared to chromidal genes at 15° (RPKM median = 45 and 31, $P$-adj = 0.0), 4° (RPKM median = 30 and 20, $P$-adj = 0.0) and 32° (RPKM median 22 and 20.5 = , $P$-adj = 0.043).

To summarize, under optimal growth conditions all four pangene categories contribute to higher gene expression on the upper part of the chromosome, whereas only shell genes contribute under sub-optimal conditions. As expected, absolute RPKM values are generally highest for core and softcore genes, and the median RPKM value for the chromosome is significantly higher than that of the chromid.

The *Pseudoalteromonas* chromid originates from an ancestral plasmid similar to those found in extant species of Alteromonadales

To investigate where the *Pseudoalteromonas* chromid originates from, we performed BLASTp searches using the chromid ParA, ParB and RepA proteins as queries against the nr. database. The tripartite ParA-ParB-parS system consists of a ParA ATPase, a ParB CTPase and DNA-binding protein, and a centromere-like parS site, and is responsible for faithful segregation of replicons during cell growth and division in approximately three quarters of bacteria (Jalal *et al.* 2020). RepA is the replication initiator protein in *Pseudoalteromonas* chromids (Xie *et al.* 2021). The fundamental function of the partitioning system and the replication initiator protein, together with their widespread distribution in Bacteria and Archaea, make ParA, ParB and RepA excellent candidates for finding clues to the origin of the chromid.

ParA and ParB BLASTp searches both identified homologs from draft genomes of *Rheinheimera* and *Catenovulum* as best hits (e-values=0/0, Identities=39%/41%; see top 20 list in File S5). We believe that these hits represent auxiliary chromosomal ParA and ParB sequences originating from integrated plasmids. Following the top hits are a number of high scoring matches against
plasmids from *Pseudoalteromonas*, *Shewanella*, *Vibrio*, *Catenovulum*, *Alteromonas* and *Glaciecola*. RepA BLASTp identified *Rheinheimera* and *Shewanella* draft genomes and *Catenovulum sediminis* plasmid as best hits, followed by *Aeromonas* plasmids. The BLASTp results therefore strongly suggest that the *Pseudoalteromonas* chromid originates from an ancestral plasmid, or possibly a megaplasmid, similar to those found in extant Alteromonadales species. Moreover, the relatively large size of today’s *Pseudoalteromonas* chromids suggests that the acquired plasmid or megaplasmid has accumulated a vast number of genes that over time evolved into an in-dispensable replicon. A similar origin has been suggested for the *Vibrionaceae* chromid (i.e., Chr 2) (Heidelberg et al. 2000; Fournes et al. 2018) and other chromids (Harrison et al. 2010; diCenzo et al. 2019).

More than half of the chromid hallmark genes in *Pseudoalteromonas* originates from the ancestral gene pool of Alteromonadales

Given that the *Pseudoalteromonas* chromid originates from an ancestral plasmid, then new questions emerge. For example, which type of genes are associated with chromids? Potential genetic sources could be genes from the *Pseudoalteromonas* chromosome, and/or genes from chromosome, chromid or plasmid DNA from closely or distantly related bacteria. To address this, we used the results from our pangenome analysis of *Pseudoalteromonas* genomes, and calculated the number of genes from each pangene category that are associated with the chromid. Any gene that was found at least once on a chromid was included. We found 164 core, 746 softcore, 2,097 shell and 4,790 cloud genes, in total 7,633 genes.

To find the genetic source of chromid genes we carefully selected a set of proteins and used them as queries in BLASTp searches. Of the 164 core genes only 78 are always located on the chromid (see File S6 for complete list of genes). These are hereafter referred to as “chromid hallmark genes”. Their ubiquitous presence on chromids support that they were acquired by the LCA, before diversification of *Pseudoalteromonas* took place (see arrow in Figure 1). Interestingly, about half (31) of the chromid hallmark genes are found clustered close to the replication terminus, and include genes and operons involved in histidine biosynthesis (*hisIEFAHBCDG*), DNA binding protein (*hupB*), acetolactate synthase (*ilvBH*), biopolymer transport system (*tonB-exbB-exbD*) and cell division (*minCDE*) (see File S6 for more information). For the ancestral plasmid to be maintained and become part of the stable genome we see today, the chromid hallmark genes probably provided a selective advantage. We therefore regard these genes as great candidates for studying the origin of early
chromid genes. All chromid hallmark proteins were used as queries in BLASTp searches (File S6). In total, 42 (58%) of the proteins produced highest scoring matches to sequences from Alteromonadales (after excluding matches from *Pseudoalteromonas*), followed by Chromatiales (11%), Vibrionales (10%) and Oceanospirillales (8%). This suggests that more than half of the current hallmark genes originates from the ancestral gene pool of Alteromonadales, whereas the remaining genes were acquired from diverse sources, mostly other gamma-proteobacteria.

**The *Pseudoalteromonas* chromid contains a large number of genes with roles in iron uptake and homeostasis**

A surprisingly large number of genes associated with iron-acquisition and homeostasis are located on the *Pseudoalteromonas* chromid. For example, in all 25 genomes, two *bfr* genes that encode bacterioferritin are located on the chromid, often flanked by *bdf* (encodes bacterioferritin-associated ferredoxin) and *iutA* (aerobactin siderophore receptor gene). Moreover, two complete *tonB-exbB-exbD* systems are found in all genomes, one on the chromid and one on the chromosome. And, in addition to *iutA*, six other TonB-dependent siderophore receptor genes are associated with the chromid, including *fhuA* (ferrichrome), *fhuE* (coprogen, rhodoturulate), *viuA* (vibriobactin), *fepA* (enterobactin), *desA* (deferoxamine B) and *vctA* (enterobactin). As previously reported for *Pseudoalteromonas tunicata* D2 (Thomas *et al.* 2008), a complete siderophore biosynthesis gene cluster is yet to be found in any of the *Pseudoalteromonas* genomes, even though they carry a relatively large number of siderophore receptor genes. This suggests that *Pseudoalteromonas* are “cheaters” i.e., they have siderophore receptors on their surface with affinity to compounds produced by other bacteria (Payne *et al.* 2016). This mechanism is used as a strategy to avoid being discriminated against by other bacteria in the constant struggle between microorganisms to acquire iron.

**Discussion**

The number of studies on multipartite bacterial genomes has steadily increased along with the number of available finished genomes in public databases. As of March 16th 2021, there are 306,881 bacterial genome assemblies listed in the NCBI genome database, of which 22,910 are denoted as “complete” (7.5%). However, whereas some phyla are well represented, with e.g., 57% of complete genomes belonging to Proteobacteria, and 34% belonging to Terrabacteria, most groups of bacteria are poorly
represented, or not represented at all. Opportunities for doing studies on many gap-free multipartite
genomes from single families are therefore rare. *Pseudoalteromonas* represents one of these rare
cases. The chromid appears to originate from a relatively recent event that can be placed at a specific
branch on the evolutionary tree with high confidence. We have therefore taken the opportunity to
study the *Pseudoalteromonas* genome, and mapped how different gene categories are distributed and
expressed in order to shed light on possible mechanisms that have shaped the chromosome and
chromid.

We found that the *Pseudoalteromonas* genome partly confirms observations from other
families, e.g., that core/softcore genes appear more frequent around *ori*1, and shell (accessory) genes
occur more frequent around *ter*1 (Figure 2). We recently reported a similar strong correlation for
*Vibrionaceae* Chr 1 (Sonnenberg *et al.* 2020). Using a slightly different pangenomic approach
Comandatore *et al.* (2019) found a similar distribution pattern in *Klebsiella pneumonia*, whereas
Kopejtka *et al.* (2019) reported a more complex picture with twenty-two species from
*Rhodobacteraceae* showing clustering of core genes close to *ori*C, and eight species showing
clustering around *ter* (Comandatore *et al.* 2019; Kopejtka *et al.* 2019). One plausible hypothesis is
that core/softcore genes, which are associated with essential cell processes, are typically
overrepresented around *ori*C because their gene products are of high demand during fast growth
(Slager and Veening 2016). The rationale is that several concurrent initiations of replication from *ori*C
results, on average, in higher “doses” of *ori*C-proximate genes. In turn, this leads to increased gene
expression (the “gene dosage” effect) (Couturier and Rocha 2006). Analyses of *V. natriegens* and *A.*
salmonicida (Sonnenberg *et al.* 2020), *Salmonella enterica* (Garmendia *et al.* 2018), and eleven
bacterial data sets of diverse origin (Lato and Golding 2020) all confirmed that overall expression
decreases with increasing distance to *ori*C. Our current analysis of a *P. fuliginea* BSW20308 RNA-
seq data replicates a similar pattern (Figure 3).

The distribution pattern for chromid genes is in contrast very different, and perhaps more
difficult to explain. For hitherto unknown reasons, the presence of core genes strongly correlates with
distance to *ter*2. Interestingly, a recent study concluded that chromids belonging to the *P. spongiae*
group are replicated bidirectionally, whereas chromids in all other *Pseudoalteromonas* are replicated
unidirectionally (Xie *et al.* 2021). Accordingly, in bidirectionally replicated chromids *ter*2 is located
at 6 o’clock, and in unidirectionally replicated chromids *ter*2 is located at 12 o’clock. The fact that
core genes are overrepresented at *ter*2 suggests that the genes are typically found in chromid sections
that are replicated in the final part of the replication cycle, a situation that is opposite to that of e.g., the *Pseudoalteromonas* and *Vibrionaceae* chromosomes where the gene dosage effect is in play. Gene dosage is apparently not in effect in *Pseudoalteromonas* chromids which suggests that we need another explanation for the clustering of core genes.

We can only speculate on why core/softcore genes tend to be located at ter2, but an intriguing possibility that we recently introduced for *Vibrionaceae* (Sonnenberg et al. 2020), is that the genomic distribution of gene categories is directly linked with how genes are organized into subcellular locations. In *V. cholerae*, Chr 1 and Chr 2 are longitudinally organized, with ori1 located at the old pole, ter1 and ter2 located at the new pole, and ori2 placed at the cell center (Fogel and Waldor 2005; Srivastava and Chattoraj 2007; David et al. 2014). Together, the data from *V. cholerae* suggests to us that core/softcore and shell/cloud genes are enriched at two separate cellular locations, i.e., at the old and new poles, respectively. Given that this hypothesis is correct, then it is plausible that a similar pattern/mechanism is in play in *Pseudoalteromonas*. It should be stressed that this remains per today a hypothesis, although the cytoplasmic position of individual gene loci have previously been successfully predicted based on the spatially organization of chromosomes (reviewed in Surovtsev and Jacobs-Wagner 2018). Moreover, for the hypothesis to be valid for *Pseudoalteromonas* there is an additional prerequisite that must be fulfilled: ter2 is located at 6 or 12 o’clock (relative to ori2) depending on the replication mechanism that is in play (uni- or bidirectional). If ter2 is deciding the subcellular destination of ter2-proximate core genes then they should, in principle, be located at the same subcellular compartment regardless of the replication mechanism. If, however, ori2 is the decisive genetic component for intracellular positioning of the chromid, then ter2-associated core genes will be located at different spatial places depending on the replication mechanism (and positioning of ter2 relative to ori2). Unfortunately, there is currently no evidence to suggest how *Pseudoalteromonas* cells are spatially organized intracellularly with regards to their chromatin. We note that the Min system (*minCDE*), which represents one of the best studied proteins involved in cellular self-organization (reviewed by Wettmann and Kruse 2018), is located in the vicinity of ter2, but the significance of this is currently unclear to us.

Our results suggest that today’s chromid in *Pseudoalteromonas* originated from a plasmid that was acquired in a single event in the LCA of this family. By comparing the chromid ParAB with database sequences we found that the best hits belong to plasmids found within today’s representatives of Alteromonadales (File S5). An early acquisition of chromid is further supported by
congruent phylogenies of the chromosome and chromid, which support that the two replicons have coexisted since the LCA of *Pseudoalteromonas* (Liao *et al.* 2019; Xie *et al.* 2021). Given an early acquisition of a plasmid or megaplasmid, what then were the main driving forces for retaining and expanding the replicon size into a relatively large chromid? diCenzo *et al.* (2019) recently proposed that the main advantage with secondary replicons, is that they enable increased genetic flexibility and potential to acquire new genetic material (diCenzo *et al.* 2019). As a result, the bacterium is better suited to take advantage of new niche opportunities. It is an appealing concept, and several pieces of evidence from our study support the hypothesis. Perhaps the most compelling evidence comes from our pangenome calculations that identifies the *Pseudoalteromonas* chromid as open and extremely flexible. A total of 7,633 genes are associated with the chromid, which is approximately 10x greater that the number of genes encoded by individual chromids (553-1567 genes; median=781). Moreover, chromid genes are generally expressed at a lower level, which have been suggested to increase the likelihood of newly acquired genes to be retained in the genome (Park and Zhang 2012; diCenzo *et al.* 2019). This is likely because highly expressed and mostly more critical genes on the chromosome are not disrupted, which then leads to less fitness cost for the bacterium. As a final piece of the puzzle, the vast majority of chromid genes in *Pseudoalteromonas* belong to the categories shell or cloud (see File S2), which provides further support for the hypothesis that new genes are preferentially maintained on the chromid and thus increases the genetic plasticity of the *Pseudoalteromonas* genome.

To summarize, we provide data showing that *Pseudoalteromonas* core/softcore genes are weakly overrepresented at oriC-proximate regions, whereas shell/unique genes are weakly overrepresented around ter1. This distribution fits with patterns reported earlier for other bacteria (Comandatore *et al.* 2019; Kopejtka *et al.* 2019; Sonnenberg *et al.* 2020). Similarly, we found that gene expression is trending downwards with increasing distance to oriC, which also fits a general pattern among many bacteria (Garmendia *et al.* 2018; Lato and Golding 2020; Sonnenberg *et al.* 2020). For secondary replicons the situation appears more complex. Here, the distribution pattern for pangene categories, as well as global expression maps, vary greatly among the studied bacteria. Perhaps the reason for the apparent lack of general trends is a direct result of the specialized roles of chromids, which have been shaped by the acquired and retained set of (mostly shell/unique) genes. Finally, we hypothesize that the gene distribution patterns reported by us and others is directly linked to how the DNAs are organized intracellularly, such that different pangene categories are enriched at separate subcellular locations based on their specialized biological functions.
Data availability
Supplemental data are available in Figure S1 and Files S1-6. Figure S1 shows phylogenetic relationships between Alteromonadales families. A list of all 25 Pseudoalteromonas genomes used in this study are available in File S1. File S2 contains distribution data of core, softcore, shell and cloud on Pseudoalteromonas chromosome and chromid divided into 4, 6, 8, 10 and 12 sectors. File S3 contains statistical analysis of pairwise comparisons of number of genes between sectors (Kruskal-Wallis and Dunn’s test). Statistical analysis of gene expression of P. fuliginea BSW20306 chromosome and chromid using Wilcoxon signed-rank test is available in File S4. File S5 contains BLASTp results with chromid ParA, ParB and RepA as queries. A list of chromid hallmark genes and BLASTp results is available in File S6. Supplemental material is available at figshare: https://doi.org/10.25387/g3.14900463.

Funding
This work was supported by the UiT The Arctic University of Norway. The publication charges for this article have been funded by a grant from the publication fund of UiT The Arctic University of Norway. The funder had no role in study design, data collection and interpretation, or the decision to submit the work for publication.

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**Figure legends**

**Figure 1** Summary of an ML-phylogenetic tree showing evolutionary relationships between Alteromonadales families. See Figure S1 for complete tree. Multipartite genomes are restricted to *Pseudoalteromonadaceae*, which is placed at the crown of the tree and branches off from families containing monopartite genomes. The arrow highlights the last common ancestor of *Pseudoalteromonadaceae*, and the likely point of origin of the *Pseudoalteromonadaceae* chromid. The color scheme shows family affiliation of genera. Numbers of strains included in collapsed nodes are shown in parentheses. Bootstrap values at the nodes were calculated using the ML method, and the GTR+G+I model, with 1,000 replicates.

**Figure 2** Heatmaps of distribution of core, softcore, shell and cloud genes in *Pseudoalteromonas* genomes. Genes were placed into one of six equally sized sectors of chromids (A) or
chromosomes (B), with sector 1 starting at the origin of replication (12 o’clock). Unidirectionally replicated chromids are found in species that belong to Clade 1, as shown in the ML-phylogeny (GTR+G+I model, 1,000 replicates), whereas bidirectionally replicated chromids belong to representatives of Clade 2. Heatmaps are based on log_{10} ratio values of the probability of a gene belonging to a sector on average divided by the probability given an equal distribution of genes among all sectors. Positive values (shades of orange) suggest that gene categories are overrepresented, whereas negative values (shades of blue) suggest underrepresentation. Asterisks indicate significant over- or underrepresentation of gene categories using Dunn’s test P-value ≤ 0.05 (see File S2 and S3 for more details). The phylogenomic tree is based on 469 single marker genes identified by EzTree (Wu 2018). Bootstrap values at the internal nodes were inferred from a ML–G+I analysis.

Figure 3 Global expression maps of *P. fuliginea* BSW20306 chromosomal and chromid genes centered on the median. Data points (log_{2} ratio RPKM CDS:RPKM median) for each CDS are shown, as well as a trend line averaged over a sliding window of 100 data points. The temperatures 4° and 32° corresponds to sub-optimal growth conditions and 15° corresponds to optimal growth conditions.
Table 1 Comparison of gene expression levels for pangenes located on the upper or lower halves of the *P. fuliginea* BSW20308 chromosome

| Optimal growth conditions (15°) | Sub-optimal growth conditions (4°) | Sub-optimal growth conditions (32°) |
|---------------------------------|-----------------------------------|-----------------------------------|
| core softcore shell cloud       | core softcore shell cloud         | core softcore shell cloud         |
| **Upper half**                  |                                   |                                   |
| Q₁                              | 41 39 26 20                        | 30 29 11 7                        | 20 19 10 8                        |
| Q₂                              | 84 82 43 37                        | 77 74 24 15                       | 41 39 18 15                       |
| Q₃                              | 253 245 94 70                      | 290 267 67 30                     | 115 107 43 34                     |
| Max                             | 11 063 11 063 134 285 37 786       | 48 320 48 320 169 723 53 846      | 5 208 5 208 35 549 8 083          |
| **Lower half**                  |                                   |                                   |
| Q₁                              | 29 28 16 13                        | 24 21 7 5                         | 18 17 8 6                        |
| Q₂                              | 65 64 28 26                        | 66 65 16 12                       | 40 38 14 13                       |
| Q₃                              | 185 181 59 55                      | 267 232 46 32                     | 97 94 32 27                       |
| Max                             | 11 172 11 172 19 840 1 635         | 6 927 6 927 17 176 578            | 2 047 2 047 4 884 523            |
| *P*-value Q₂<sup>b</sup>        | 0.00 0.00 0.00 0.00                 | 0.08 0.07 0.00 0.15               | 0.40 0.40 0.00 0.20               |

<sup>a</sup> Q₁ is the RPKM value at the first quartile. Q₁ is defined as the middle number between the smallest number and the median (i.e., the second quartile Q₂), if the data numbers (in this case RPKM values) are ordered from smallest to largest. The third quartile (Q₃) is the middle value between the median (Q₂) and the maximum (Max) value.

<sup>b</sup> Adjusted *P*-values from Wilcoxon signed-rank test, to test if Q₂ values (median) of genes located on the upper half of the chromosome are significantly different from Q₂ values of genes located on the lower half. Values below 0.05 are considered significant.
