A domain sequence approach to pangenomics: applications to *Escherichia coli* [version 1; peer review: 2 approved]

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**Abstract**

The study of microbial pangenomes relies on the computation of gene families, i.e. the clustering of coding sequences into groups of essentially similar genes. There is no standard approach to obtain such gene families. Ideally, the gene family computations should be robust against errors in the annotation of genes in various genomes. In an attempt to achieve this robustness, we propose to cluster sequences by their domain sequence, i.e. the ordered sequence of domains in their protein sequence. In a study of 347 genomes from *Escherichia coli* we find on average around 4500 proteins having hits in Pfam-A in every genome, clustering into around 2500 distinct domain sequence families in each genome. Across all genomes we find a total of 5724 such families. A binomial mixture model approach indicates this is around 95% of all domain sequences we would expect to see in *E. coli* in the future. A Heaps law analysis indicates the population of domain sequences is larger, but this analysis is also very sensitive to smaller changes in the computation procedure. The resolution between strains is good despite the coarse grouping obtained by domain sequence families. Clustering sequences by their ordered domain content give us domain sequence families, who are robust to errors in the gene prediction step. The computational load of the procedure scales linearly with the number of genomes, which is needed for the future explosion in the number of re-sequenced strains. The use of domain sequence families for a functional classification of strains clearly has some potential to be explored.

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Introduction

Microbial pangenomics has attracted interest over recent years, stimulated by the availability of sequence data from whole-genome re-sequencing projects\(^1\)-\(^7\). The pangenome of a prokaryotic species is the collection of gene families for the entire species, as opposed to a single genome, which is the set of genes in a functional organism. The pangenome diversity can be huge, which is also reflected in the span of phenotypes. An example of this is found in the model organism *Escherichia coli*\(^8\)-\(^9\). Currently there are more than a thousand *E. coli* genomic projects listed\(^10\) and this number will grow in the near future, along with genomes for many other bacteria; it is reasonable to assume that pangenomics will attract more attention.

The fundamental step in any pangenome analysis is the computation of gene families. Several approaches to computing gene families have been used in previous pangenome analyses\(^11\)-\(^12\), but this part of the analysis has received little attention. A pangenome analysis typically involves the estimation of the size of the core and the pangenome, measured by the number of gene families, and several methods have been proposed for doing this\(^13\)-\(^14\). The core is the set of gene families present in all genomes of the population. The remaining gene families are more or less abundant among the genomes. The sample pangenome size is the total number of gene families found in the currently available genomes, while the population pangenome size is the number of gene families we expect to see if every single strain was sequenced. It is the latter which is of scientific interest, but it must be estimated from the former. A fraction of the gene families will be found only in a small number of genomes, and those observed in only a single genome are called ORFans.

The first step of a gene family computation is to obtain a list of protein coding genes for each genome under study. Completed genomes will have a set of annotated genes, but the quality of these annotations may vary between projects. A pangenome analysis will often include draft genomes as well, where annotations are lacking. For these reasons it is convenient to start the analysis by a gene-finding step, treating all genomes the same way, and minimizing variability due to different annotation procedures. Even if prokaryote genes are in most cases simpler to detect than eukaryote counterparts, there are still problematic cases\(^15\). In case of the draft genomes, where the genome sequence is spread out on a (large) number of contigs, the gene-finding problem is even more difficult. Ideally, we would like to compute gene families in a way that minimizes the effect of various gene finding algorithms.

The second step is to group proteins into gene families. A gene family is basically the set of orthologs and in-paralogs collected from the various genomes. The most common approach so far is based on all-against-all alignments to compute some kind of similarity/distance between all proteins, and then finally cluster these. This approach poses some problems. First, the all-against-all approach is not computationally feasible in the long run since the number of genomes grows rapidly. Second, the clustering procedures always involve some granularity threshold implicitly defining the size/number of gene families. Some kind of thresholding seems impossible to avoid, but it would be desirable to allow it some variability over gene families, since some gene families are expected to be more conserved than others. Finally, false predicted ‘genes’ will not align well to any other, and produce singleton clusters adding, sometimes dramatically, to the total number of gene families found.

When grouping protein coding genes, we find it natural to consider the presence of common protein domains. Domains are closely linked to gene function\(^16\). Previous work has demonstrated that the functional repertoire of a genome can be predicted by the knowledge of domain content\(^17\)-\(^18\). In addition, focusing on the combination of domains in each protein, instead of the frequency of each domain separately, we come even closer to the protein function\(^19\)-\(^20\). A grouping of proteins by function rather than by evolutionary distance seems like a good alternative, since pangenomics is in many cases about the study of functional diversity.

The Pfam-A database\(^21\) is a comprehensive collection of domains, curated and of high quality. Each entry is characterized by a profile hidden Markov model, and until recently it has been quite time consuming to search against such a database. However, with the launching of the HMMER3 software\(^22\)-\(^23\), it is now possible to search with all proteins within a genome against the entire Pfam-A database in reasonable time (few minutes), even on a laptop. The idea of using domain information in comparative genomics is not new, and Yang *et al.* used genome-wide domain frequencies to reconstruct phylogenetic trees\(^24\). Their study considered diverse genomes from both prokaryotes, eukaryotes and archaea, quite opposite of pangenomics, where we focus only on genomes from the same species. An argument for using domains was that structure is more conserved than sequence, and that domain information provides a view into ancient history\(^25\). In this perspective one may question the use of domain information for pangenomics studies, since they may provide too low resolution.

In this paper we consider domain sequence families defined by the domain sequence of each protein as an alternative to gene families for pangenome studies. The domain sequence is the ordered sequence of domains in each protein, instead of the frequency of each domain separately, we come even closer to the protein function. We describe the procedure of computing such gene families, and consider the effect of gene prediction and of the use of draft genomes in the analysis. We use this approach in a pangenome study of the model organism *Escherichia coli*, the prokaryote species with the largest set of available genomes.

Materials and methods

The methods we have used for gene prediction and database scanning are all standard and free software, see below. We acknowledge that the gene finders have tuning possibilities, and that slightly different results may be obtained by using this. We have used only default settings, or setting suggested in accompanying manuals, in order to make the approach as universal as possible. The data are from the model organism *Escherichia coli* only, and many of the more specific results cannot be extended to other species. However, the procedures are applicable to all prokaryotes where we have genomes from many strains available.

Data

Genome sequences for *Escherichia coli* were downloaded from NCBI\(^26\) on June 15 2012. At that time, there were 56 completed and 291 draft genomes available as contigs.
Gene finders
In a pan-genome study involving draft genomes some automated gene finding is needed. For each genome we predicted genes using the three popular gene finders Prodigal\textsuperscript{27}, GeneMark\textsuperscript{28} and Glimmer\textsuperscript{29}.

The Prodigal gene finder (v2.60) was run by the command line

\texttt{prodigal -i gnom.fsa -o pred.txt -q}

where \textit{gnom.fsa} was a FASTA-formatted input file with the genome sequences, and \textit{pred.txt} was the output file.

The GeneMark predictions were obtained by first training a Hidden Markov Model (HMM) using GeneMark.S (v4.6b):

\texttt{gmsn.pl -combine -clean -gm gnom.fsa}

which produced a model file \texttt{GeneMark.hmm.combined.mod}. Next, this was used to predict genes by \texttt{GeneMark.hmm}, prokaryote version 2.6:

\texttt{gmmhmp -m GeneMark.hmm.combined.mod -o pred.txt gnom.fsa}

The Glimmer predictions were obtained by the tigr-glimmer wrapper software for Linux. First, the long.orfs software was used to extract training sequences from the genome sequences:

\texttt{tigr-glimmer long-orfs -n -1 -t 1.15 -g 30 seq.fasta coord.txt tigr-glimmer extract -t seq.fasta coord.txt >> train.fsa}

where \textit{seq.fasta} was the FASTA file with a single genome sequence. This was repeated by looping through all genome sequences, producing the file \textit{train.fsa} with training sequence data. Next, an Interpolated Context Model (ICM) was trained:

\texttt{tigr-glimmer build-icm -r glim.icm < train.fsa}

where \textit{glim.icm} was the model file output. This was finally used to make gene predictions:

\texttt{tigr-glimmer glimmer3 -o50 -g110 -t30 -linear -extend gnom.fsa glim.icm pred.txt}

In all cases the gene predictions were stored as a table with one row for each predicted gene, and with columns GenomeSequence (name of genome sequence where it is found), Strand (1 or -1), Left (smallest coordinate), Right (largest coordinate), Partial (logical indicating if the gene runs over the start/end of the genome sequence). This format makes comparison of gene predictions straightforward.

Effect of gene prediction
We wanted to examine the effect of various gene finders on the list of proteins obtained from each genome. For this study we only focused on the 54 completed genomes with annotated proteins in the RefSeq database\textsuperscript{30}. Given the curation of the RefSeq database, and the fact that \textit{E. coli} is the most well studied prokaryote genome, we expect these RefSeq annotations to be as close to the truth as one can expect for any prokaryote organism. Thus, by using the three gene finders described above we obtained four sets of proteins for each of the 54 genomes. We compared these sets using the Jaccard distance defined as

\[ J(S_a, S_b) = 1 - \frac{|S_a \cap S_b|}{|S_a \cup S_b|} \]  

where \( S_a \) and \( S_b \) are two sets of proteins and \(|| \) indicates cardinality (number of elements in the set). A distance of 0.0 means the two sets \( S_a \) and \( S_b \) are identical, while the maximum distance of 1.0 means they are non-overlapping. In this comparison, two proteins were considered equal only if they were exactly identical (identical amino acid sequence). After extraction of domain sequences from each protein (see below) we again computed Jaccard distances. This time two proteins were considered equal if they had identical domain sequence.

Final gene prediction
For the sake of completeness all genomes, both complete and draft, were subjected to the same analysis. Since we subsequently filtered all results through the Pfam-A database (see below), we were less concerned about false positive gene predictions at this step. To obtain a comprehensive gene-list, we ran all three gene-finders on each genome, and compiled the union of the results into one long list of ORFs for each genome. In the cases where the same stop-codon had alternative starts, we always chose the longer ORF, again due to the later Pfam-A filtering. All partial ORFs (lacking start and/or stop codon) were eliminated before the analysis, but we made no attempt to resolve overlaps between different ORFs. Finally, all remaining ORFs were translated to protein.

Pfam search
FASTA-formatted files of protein sequences were queried against the Pfam-A database version 26.0\textsuperscript{21} using the HMMER 3.0 software\textsuperscript{22,23}. The search was done using the command line

\texttt{hmmscan -o out.txt -cut_ga -noali -cpu 0 -domtblout res.txt Pfam-A.hmm prot.fsa}

where \textit{res.txt} is the name of the output file, \texttt{Pfam-A.hmm} is the HMMER3 database and \textit{prot.fsa} is the FASTA-formatted input file of protein sequences to scan. We used the option \texttt{-cut_ga} in order to invoke the curated inclusion thresholds inherent in the Pfam-A models. The e-value (independent E-value) was used as the criterion in the cases where we wanted to overrun these thresholds, using a stricter level of significance.

Domain sequences
For every protein sequence the significant hits were listed in their order of appearance along the protein. In the cases where two or more domains were overlapping, the one with largest E-value was eliminated, and this was repeated in a recursive procedure until no more overlaps between hits were found. There is a small number of nested domains (domains inside domains) in the Pfam database, but we
chose to eliminate all occurrences of overlaps. The domain sequence of a protein is the ordered list of Pfam accession numbers, and multiple hits of the same domain may occur as long as they are non-overlapping. It is a high-level alternative to the amino acid sequence representation of a protein. Proteins giving no hits in Pfam-A were discarded from the downstream analysis. This step presumably filtered out the majority of the false positive gene predictions from the gene-finding step. Finally, we grouped into domain sequence families all proteins having identical domain sequence.

Pan-matrix

The fundamental data structure for a pangenome analysis is the pan-matrix. This matrix has one row for each genome and one column for each domain sequence or gene family, and cell \((i,j)\) holds the number of copies of domain sequence family \(j\) in genome \(i\), alternatively just 0 or 1 to indicate absence/presence, the latter being used in this paper. Let \(x_i\) be the number of genomes in which family \(j\) is present. It is natural to view \(x_i\) as a random variable, the randomness mainly due to our random selection of genomes since there are obviously many genomes we could have, but have not yet, looked into. The variable \(x_i\) can take any integer value from 1 to \(G\), where \(G = 347\) is the number of genomes in this study.

Each row of the presence/absence pan-matrix is a coordinate vector indicating the location of the corresponding genome in a domain sequence space. This domain sequence space is closely related to a functional space in the sense that neighbors in this space is likely to display similar functions. This space is high-dimensional (many different functions), but in order to visualize the genomes relative location in this space we used a principal component decomposition to extract the most dominant directions. The genomes location along these dominant directions give us some indications of their relative location in the functional space.

Heaps law and population closedness

Using a Heaps law type of regression model, Tettelin et al. made an estimate of pangenome closedness\(^5\). A closed pangenome means the gene families are sampled from a stable and finite set, as opposed to an open pangenome where gene families ‘migrate’ in and out at unknown rates. If we arrange the \(G = 347\) genomes in some fixed order, we can define \(n_g\) as the number of new families seen in genome \(g\) compared to the previous \(g – 1\) genomes. The Heaps law says that

\[
E(n_g) = \beta g^\alpha
\]

where \(\beta\) is some intercept and \(\alpha\) is the parameter of interest. An \(\alpha\) below 1 indicates the number of new families does not decrease fast enough for the pangenome to be limited, i.e. if the number of genomes \(G\) grows towards infinity the number of new families observed will never reach zero. Parameters can be estimated from data, by repeatedly permuting the order of the genomes and counting \(n_g\) for \(g = 2, ..., G\), after each re-ordering.

Binomial mixture model analysis

As explained above, let \(x_i\) be the number of genomes in which we observe domain sequence family \(j\). Let \(y_j\) be the number of families found in \(g\) genomes (number of \(x_i’s\) with value \(g\)). Then \(y_j\) is also a random variable. The probability density of this variable, e.g. the one depicted in Figure 3, can be described by a \(K\) component binomial mixture model

\[
f_y = \sum_{k=1}^{K} \pi_k b(\rho_k)
\]

where \(b(\rho_k)\) is the binomial density with probability \(\rho_k\) and \(\pi_k\) is the mixing proportion (the \(\pi_k’s\) sum to 1). The left pie of Figure 7 is a visualization of a \(K = 12\) component model, where the size of each sector is the corresponding \(\pi_k\) and the color of each sector indicates \(\rho_k\), for \(k = 1, ..., 12\). Summing \(y_j\), we get the number of domain sequence families seen so far, i.e. the sample pangenome size. From the binomial mixture model we can also predict \(y_j\) the number of families not yet seen, and in this way we can estimate the population pangenome size\(^14\). The probabilities \(\rho_k\) we refer to as the selection probabilities. A domain sequence family with selection probability 0.1 will on average be present (selected) in 1 out of 10 genomes.

Genome diversity can also be expressed as the overlap between genomes, and for this we could use the mean Jaccard distance defined above, where \(S_j\) and \(S_j\) are two sets of domain sequences (two genomes). Genome fluidity, introduced in\(^3\), is an almost identical measure. A small Jaccard distance-genome fluidity indicates a small degree of uniqueness in each genome, i.e. large overlap. The expected overlap between two genomes can also be computed directly from a fitted binomial mixture model in an elegant way. The expected number of domain sequence families in a genome is

\[
E_1 = N \sum_{k=1}^{K} \rho_k \pi_k
\]

where \(N\) is the population pangenome size. The expected number of domain sequences found in two independent genomes simultaneously is

\[
E_2 = N \sum_{k=1}^{K} \rho_k^2 \pi_k
\]

since the probability of selecting a domain sequence family twice is \(\rho_k \cdot \rho_k\), given that \(\rho_k\) is its selection probability. The expected overlap between two genomes can be expressed as \(E_2/E_1\). Note that the pangenome size \(N\) cancels out, and the result only depends on the selection probabilities in combination with the mixing proportions. The expected overlap between 3 and more genomes can be computed along the same lines.

Results and discussion

Gene prediction

Before we commenced the full scale analysis, we wanted to investigate the effect of gene finding algorithms. To this end, we considered only the 54 complete \(E. coli\) genomes with annotations in the RefSeq database\(^39\) at the time of this study. In Table 1 we show how the sets of genes differ between the official RefSeq annotation and the three most commonly used prokaryote gene finders. From the upper triangle of the table we see that the Jaccard distance to the RefSeq annotations is large in all cases. Even for Prodigal, which comes closest to the RefSeq annotations, the distance is 0.26. For a list of 5000 proteins, this corresponds to around 750 proteins on
Table 1. Gene finding For each of the 54 RefSeq-annotated E. coli genomes we compared the set of annotated genes and the set of predicted genes from Prodigal, Glimmer and GeneMark. The Jaccard distance between all pairs of sets was computed, see Method section. First, two genes were identical only if they have identical amino acid sequence. The upper triangle (above the diagonal) shows the mean Jaccard distance over the 54 genomes in this case. Next, we computed the Pfam-A domain sequence of each predicted protein, discarding all proteins without Pfam-A hits, and again computed the Jaccard distance between the sets. The lower triangle (below the diagonal) shows the mean Jaccard distance over the 54 genomes in this case.

<table>
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<th>Prodigal</th>
<th>GeneMark</th>
<th>Glimmer</th>
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<tr>
<td>Glimmer</td>
<td>0.35</td>
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</tbody>
</table>

each list not being found in the other. Also the differences between the gene finders are large, and there is a severe uncertainty in any list of genes obtained by any single gene finder. In the lower triangle of Table 1 we see how the differences are reduced dramatically once we only consider domain sequences. This means all proteins without Pfam-A hits are discarded, and the remaining proteins are only compared by the domain sequence. The difference in start codon predictions, causing the majority of differences in the first comparison, no longer have any impact. If two proteins have the same domains, it does not matter how they compare outside these domains.

In the RefSeq annotated genomes we found on average around 2750 unique domain sequence families (from 2555 to 2957). Using the union of the three gene finders, we found on average 2725 of the RefSeq annotated domain sequence families among these, i.e. around 25 families (1%) in each genome is not found by any of the three gene finders. On the other hand, the three gene finders find on average close to 100 extra domain sequences families in each genome. This means that either the RefSeq annotated genomes have missed quite a number of domain-containing protein sequences, or the built-in inclusion thresholds in the Pfam-A models are too liberal, resulting in a substantial number of false positive hits. To investigate the latter, we tried out stricter significance thresholds in the HMMER3 software. Even for an E-value of $E = 10^{-10}$ we found around 70 extra domain sequence families per genome in the union of the three gene finders compared to the RefSeq annotation. We believe this illustrates that annotated genomes still contain several ‘holes’, which is also the conclusion of others. The fact that gene finding in prokaryotes is simpler than in eukaryotes may have created an illusion that it is straightforward and nearly 100% reliable, when careful studies have shown that typical bacterial genomes have around 10% genes that are false positive, and another 10% of the proteins discovered by proteomics are missing. Angiuoli et al. suggested an annotation procedure based on the whole-genome alignment of all genomes in a pangenome study, and found a disturbing amount of inconsistencies in public annotations. For E. coli less than 50% of the gene-structures were consistently annotated across 30 genomes, and the majority of the inconsistencies were due to difference in start codons positions.

E. coli domain sequences

The input to the analysis was a set of 347 genomes of E. coli, 56 of these were completed and 291 were available as contigs, all public data downloaded from NCBI. The left panel of Figure 1 shows the size distribution for the completed and the draft genomes. The first step was to predict genes in all genomes by three different gene finders, as explained in the Methods section. As seen in the center panel of Figure 1, this resulted in a list of roughly 7000–8000 predicted genes in each genome, which is far more than the 4500–5500 genes usually found annotated for E. coli genomes. This reflects the disagreement between the gene finders, and we expect a large proportion of false positives in this set. The second step was to scan all predicted proteins against the Pfam-A database. Protein sequences producing no hits in Pfam-A were discarded from the downstream analysis. Roughly 4000–5000 proteins from every genome produced significant hits in Pfam-A, see Figure 1, right panel. We also note that the difference between completed and draft genomes is more or less nonexistent after this step. This justifies the use of draft genomes in pangenome studies like this.

Considering all sequences from the 347 genomes, a total of 3679 unique Pfam-A domains/families produced significant hits, which is approximately 30% of the entire database. This produced a total of 5724 unique domain sequence families for the 347 E. coli genomes, with around 2500 in any single genome. This is a small number compared to the number of gene families previously found for E. coli. The domain sequence families are expected to be fewer and larger clusters compared to the previously computed gene families. First, not all genes have domains listed in Pfam-A, and these are discarded here. This significantly shortens the list of clusters, presumably also eliminating a large majority of the false positive gene predictions. Second, some domain sequence families are large, containing proteins sharing perhaps only a single domain. In these cases a domain sequence family may contain more than one gene family.

The most common domain sequence family in E. coli is the single-domain protein containing the major facilitator superfamily (MFS) transporter with accession PF07690.11. This is found in around 50 copies in every single genome. Other very frequently occurring single-domain proteins are the binding-protein-dependent transport system inner membrane component (PF00528.17), and the ATP-binding domain of ABC-transporters (PF00055.22). Among the multi-domain proteins the sequence PF00126.22, PF03466.15 is the most common. This is a domain sequence characteristic of HTH-type transcriptional regulators, and it occurs in more than 30 copies in each of the 347 E. coli genomes investigated here. Figure 2 shows that more than 3000 of the domain sequence families are defined by a single domain, while gradually fewer families are defined by multi-domain-sequences. Single-domain proteins also clearly outnumber the multi-domain proteins in the genomes, contrary to what is sometimes claimed. The longest domain sequence contains 25 non-overlapping Pfam-A hits in the same protein, mainly multiple copies of PF05662.9 and PF05658.9, both short repeats associated with haemaglutinins. This protein is found in 4 of the 347 genomes.

In Figure 3 we show the distribution (presence/absence) of domain sequences over the E. coli population. The leftmost bar is the
Figure 1. Complete and draft genomes. The box and whisker plots illustrate the differences between completed and draft genomes in this study. The left panel shows that the 56 complete genomes are somewhat smaller in size measured in megabases. This is most likely due to unresolved overlaps between the contigs in the draft genomes. The middle panel contains the number of unique genes predicted by the three gene finders after the elimination of all partial predictions (lacking start or stop codon). Notice the large number of predicted genes in virtually all cases, annotated *E. coli* genomes usually have 4500–5500 genes. Among the draft genomes some genomes have very few predicted genes, seen as circles. The rightmost panel shows the number of predicted gene with at least one Pfam-A hit. Except from four draft genomes with extremely few genes, the differences between complete and draft genomes are now ignorable.

The most remarkable result is the large number of ORFans. More than 15% of the domain sequence families found in *E. coli* are seen in only 1 out of 347 genomes. The Pfam-A models are curated and have a built-in threshold for assigning significant hits. However, given our approach, where we scan a list of presumably many false positive protein predictions, we may be extra restrictive when considering the hits. In Figure 4 we show how a systematic change in E-value cutoff changed the number of core families and ORFan families. As we lower the E-value cutoff, little happens to the number of core families, but the number of ORFans drops markedly. This demonstrates that the ORFan are, as usual, the most uncertain families. In the downstream analysis we made a parallel analysis using the E-value cutoff 10^{-10} as a strict alternative to the built-in thresholds.
Figure 2. Domain sequence lengths. The bar chart shows how many domain sequences have 1 domain, 2 domains,... etc up to 25 domains. These are all non-overlapping hits in the protein. Single-domain proteins makes up more than half of the total number.

Figure 3. Gene family distribution. The distributions of how many domain sequence families are found in 1, 2,..., 347 genomes. There are 909 ORFan families (leftmost bar), 479 core families (rightmost bar) and in total there are 5724 unique domain sequence families (sum of all bars).
The 347 text files show the results from scanning each of the fasta-files against the Pfam-A database

347 Data Files
http://dx.doi.org/10.6084/m9.figshare.95928

Functional space
Domains are usually well conserved, and may show too few differences between strains within a species, i.e. all genomes contain more or less the same domain sequences. To examine this, we computed the Manhattan distance between every pair of genomes, which is simply the number of domain sequences where the two genomes differ in presence/absence status (ignoring copy numbers). Figure 5 shows a histogram of all pairwise distances. The median distance between two genomes is around 500, meaning there are 500 distinct domain sequences contained in one of the genomes but not the other. Only two genomes are identical, and this is actually the same genome, BL21(DE), sequenced at two different locations (Korea and Austria), but stored under unique accession numbers in the database. All other cases show at least 3 or more differences, and the median shows around 500 differences. Hence, even genomes of the same serotype have plenty of differences in domain sequence absence/presence. Using the strict E-value cutoff reduced the median distance to around 450, but did not change the shape of the histogram. The small ‘bump’ on the right hand side is due to a few genomes being quite different from the rest. These are all draft genomes with a large number of contigs, producing a smallish number of domain sequences.

We may think of each domain sequence as representing a small repertoire of functions the genome can inhabit. For each genome the vector of presence/absence of the set of domain sequences will correspond to a location in a functional space, and the Manhattan distance between two genomes in this space is a functional distance. A pangenome tree can be made from these Manhattan distances, as described in \(^\text{36}\). However, any tree visualization will suffer when we have as many as 347 leaf nodes. Supplementary figure 1 illustrates a pangenome tree for the 347 \textit{E. coli} genomes.

This sample of \textit{E. coli} genomes contains several potential subsamples, i.e. strains who have more in common than just being of the same species. It is natural to see where these are located in this functional space, and here we have looked at the four largest subsamples in the data set. In \(^\text{37}\) it was found that ETEC type of strains shared some genomic features distinguishing them from other \textit{E. coli}, and it is reasonable to expect that certain strain subsets will cluster within the total collection of \textit{E. coli} genomes. We used a principal component analysis on the present/absent pan-matrix as described in the Methods section. The functional space is high-dimensional, and plotting each genome as a dot in the first two dimensions...
principal component directions as in Figure 6 give us only a rather rough overview of the relations. Still, the clustering of the enterohemorrhagic serotypes (O157:H7, light blue dots and 0104:H4, dark blue dots) is clearly visible. The collection of diarrheagenic strains (red dots) are clumped at several locations, indicating a mixture of functional content. The Human Microbiome Project strains (green dots) are all located to the right hand side of Figure 6. Using the strict E-value cutoff hardly changed the picture in Figure 6 at all. This is due to the extreme stability of a multivariate analysis like PCA. Since we know from Figure 4 that this strictness reduce the number of ORFans, it means the picture in Figure 6 is marginally influenced by ORFan gene families.

We also conducted a Heaps law analysis as suggested by 13. From this it seems the population of E. coli domain sequences is open. We fitted the Heaps law model to these data, using 100 random permutation of genome ordering, and came up with the estimate 0.94 for the parameter α in the model. A value of α < 1.0 is consistent with an open population. This means the population size is unbounded when extrapolating the Heaps law trend into many more sequenced genomes. However, unbounded is a mathematical term, and an unbounded function can still grow very slowly. The model predicts that after 1 million sequenced E. coli genomes(!) we still have not seen more than 9442 domain sequences. As such, the result is not that different from the mixture model estimate.

We repeated the computations using the strict E-value cutoff in the HMMER3 software. The sample pangenome then reduces to 4745 (from 5724) observed domain sequence families, since only very significant (E < 10^{-10}) Pfam-A hits are now considered. The binominal mixture model gives a pangenome size estimate of 4973, which still means a coverage around 95%. The Heaps law analysis, however, results in a closed population, with α = 1.01 and an estimate of its size at 4876 domain sequences. This illustrates how the choice of cutoffs in the sequence clustering may change a result completely. The Heaps law analysis is extremely sensitive to the number of ORFans in the data set.

The sample core size is 479 domain sequences, and the binominal mixture model predict the population core to be 462. However, the notion of the core is difficult, since we require a core domain sequence to be present in every single genome. Due to the uncertainties

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**Figure 5. Functional distances.** A histogram over all pairwise Manhattan distances between the genomes. The distance between two genomes is defined as the number of domain sequence families they differ in presence/absence status, i.e. a distance of 500 means there are 500 different families contained in one but not the other genome.

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**Pangenome analysis**

In this sample of 347 E. coli genomes we find 5724 unique domain sequences. Using the binominal mixture model suggested in 4, the total pangenome size is estimated to 6040. Employing the suggested bagging procedure, we find that in 90% of the re-sampled cases the estimate is between 5998 and 6136, reflecting the uncertainty in the data. This should be regarded as a lower bound estimate, but indicates we have already seen the majority of domain sequence families in E. coli. The observed set of domain sequences covers around 95% of the predicted pangenome.
in gene prediction and computation of sequence clusters, such a crisp limitation of the core is unfortunate. From the binomial mixture model we get a much better and more robust picture by considering the estimated selection probabilities behind every mixture component. In Figure 7 we have visualized the *E. coli* pangenome as a pie chart, where the colors indicate the selection probabilities. Using the Bayesian Information Criterion (BIC) we found that 12 components was optimal for the current data set. This means *E. coli* domain sequences can be grouped into 12 distinct sectors with respect to how often they appear in the genomes. The core genes is one of these types, having selection probability 1.0 since they occur in every genome (darkest blue sector). Notice also the large sector of domain sequences with a selection probability of 0.988. Even the third darker blue sector has a selection probability of 0.966. The large number of domain sequence families in these sectors are also highly conserved, and could be seen as core families for all practical purposes.

The coloring in Figure 7 reflects the commonly used division of the pangenome into three types of genes; the core (dark blue), the shell (light blue/green) and the cloud (orange/pink). By allowing the mixture model to have many more components, we found 12 was optimal here, we can cope with the fact that genomes are not uniformly distributed within the population. The distribution in Figure 3 is affected by this, e.g. since we have 31 genomes of serotype O157:H7 in the sample we expect there will be a small ‘bump’ in the distribution at 31 genomes, reflecting the domain sequence families common to these closely related genomes. The binomial mixture approach can be illustrated by a lunch buffet table, with each single genome as a plate to be filled with content from the pangenome (the buffet table). Some families on the buffet is always selected, and turns up on every plate. These are the core families. The other families are more or less popular, and have different probabilities of being selected. Some families are very unlikely to be selected, but if there is a large number of them, some of them
will end up in almost every genome. However, these ORFans are so unlikely to appear they are never seen twice.

The right hand pie in Figure 7 shows the expected distribution of domain sequence families within an average E. coli genome. Clearly, the darker blue domain sequences makes up the majority. The size of the pangenome is largely due to the big number of ORFan domain sequences (pink sectors in left pie), but in any average genome these makes up a small minority (pink sectors in right pie). The expected overlap between two genomes is 0.90, i.e. on average 90% of the domain sequence families in one genome is also found in another. We also computed the genome fluidity to 0.11 and the mean Jaccard distance between two genomes to 0.18, both indicating a small uniqueness (lack-of-overlap) in each genome.

Conclusion
Any study of pangenomes involves the clustering of sequences into gene families, and the results obtained will invariably depend on the way gene families are computed. In this paper we have proposed an alternative based on protein domains to obtain stable sequence clusters. This means we loose some information from every genome, since only proteins with known domains are considered. In this study we used only the Pfam-A database, and by extending it to also include other databases (e.g. Pfam-B, CDD, InterPro), some more hits would probably be found. Also, the resolution is on the lower end since all proteins having identical sequence of domains are clustered into the same family. However, the gain is a robustness with respect to gene prediction, eliminating many false positive ‘proteins’ from the analysis as well as the potential effect of the inconsistent annotation of start codons. Computing gene families based on the alignment of full protein sequences is more sensitive, but given the substantial uncertainty in gene prediction and annotation revealed here and by others, the domain sequence information may be a fruitful approach for pangenome studies. More attention should be given to improve gene prediction for prokaryotes, and the recent approach taken by Angiuoli et al., considering all genomes simultaneously to improve consistency, is a good idea for pangenome data.

Another advantage of the domain sequence approach is the fact that Pfam-A domains have a built-in significance threshold optimized for each individual HMM, and by using this we obtain families with variable heterogeneity reflecting the various degree of conservation of different types of proteins. The procedure for computing such domain sequence based gene families is straightforward and highly reproducible, using only publicly available software. Finally, each genome is only scanned once, which means the computational load scales linearly in the number of genomes.

Despite the lower resolution in the domain sequence families, we find plenty of differences between strains of E. coli. We also find meaningful groupings of strains, indicating that the differences are not just random ‘noise’ but have plenty of biological foundation. The use of domain sequences for classification of strains clearly has some potential, and this should be pursued further.

With as many as 347 genomes, and more than 1000 soon available, we would expect the E. coli pangenome to be fairly well covered.
Using the domain sequence families, this seems to be the case. Our lower bound estimate of pangenome size indicates we have seen perhaps as much as 95% of all domain sequences ever to be found in this species. Domain databases, like Pfam-A, will still grow but the number of single domain proteins is leveling out, and the future growth will mainly be due to new domain sequence combinations. There is an endless number of possible domain combinations even with todays databases, and for E. coli we must expect some new combinations in future sequenced genomes. However, with an overlap of 90% between any two E. coli genomes the time of big surprises is gone. The Heaps law analysis indicates the population is open, but we believe this analysis is too sensitive to a smallish number of ORFan families in the data set. Also, the entire concept seems to build on the assumption of a uniform sampling of strains, which is clearly not the case for E. coli.

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Authors contributions

LS and DWU conceived of the study. LS carried out the programming. LS and DWU drafted the manuscript. Both authors read and approved the final manuscript.

Competing interests

No competing interests were disclosed.

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The authors provide an excellent-outstanding computational analyses of the *e. coli* pangenome with respect to protein coding across more than 50 complete genomes and 350 complete and whole genome shotgun (incomplete) genomes. Importantly, they evaluate the proteins and protein families by recomputing/predicting families, rather than relying on GenBank/RefSeq annotations, which can vary depending upon source. Their work is well-documented and should serve as a gold standard for pangenome analyses of protein families for many bacteria.

There are however, some minor improvements that can be made to this paper:

1. On top right column of page 7, the authors explicitly state that “Only two genomes are identical…BL21(DE3). Since several non-identical K-12 strains are included in their Data File(MG1655, W3110, DH10A, and a couple others), authors should simply state how many K-12 strains were included in their analyses.

2. It would be especially helpful to the community, if the authors can post data files of their analyses, e.g., the raw data used to generate their pie charts in Figure 7. I am sure users would more widely cite their efforts if such results can be made easily accessible.

**Competing Interests:** No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.
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The authors provide a very interesting insight into how one can reliably identify genes and classify them into gene families in fully sequenced bacterial genomes and apply this approach to 347 fully sequenced *Escherichia coli* genomes.

They avoid annotation problems by first running a very crude gene prediction program, that returns a large number of genes with a large proportion of false positive open reading frames (ORFs). For these ORFs they then predict Pfam-A domains. All ORFs for which no domain could be predicted were discarded. They then show that the remaining set open reading frames closely resembles the number of ORFs expected from manually curated genome annotations. They then go on and analyse the domain sequence composition of all remaining ORFs and provide a comprehensive pan genome analysis. In conclusion, the paper presents a very interesting approach to solve gene classification problems in pan-genome studies, a field that rapidly gains significance as whole genome sequencing becomes cheaper and cheaper.

The paper is well written and very comprehensive. The data is presented clearly and is directly downloadable from within the paper. The author even explain complex problems with interesting similes, which I thought was a nice idea and helped me to understand the presented ideas.

There are only a few minor problems that should be addressed:

1. One thing needs clarifying. The authors mention in paragraph two on page five that within the RefSeq annotated genomes they can identify 2725 of the 2750 unique domain sequence families for genes that were automatically identified. Also they say that they identify 100 extra unique domain sequence families in automatically identified genes. Does that mean that on average 2825 unique domain sequence families were identified? If this is so then I do not understand why in the second to last paragraph the authors write that in every single genome there are about 2500 unique sequence domain sequences. Especially since Figure 1 shows that on average there are more proteins with Pfam hits for draft genomes, and draft genomes are on average longer. Does that have to do with the fragmentation of the data that leads to splitting of domains? If so, how would that affect functional predictions?

2. How do the authors deal with gene predictions where one gene is predicted on the top strand and another on the bottom strand or generally genes predicted in different reading frames by different gene prediction programs? Did the authors test whether the data that was missed by/additional to the refseq annotation were overlapping genes but in different reading frames and hence could not have been picked up by Pfam no matter what the threshold is?

3. I think it would be interesting to estimate the effect that errors in gene prediction have on prediction of metabolic function etc. would have in theory. For example, what were the annotations of the unique domain sequence families that were missed by refseq or the gene finders. Were those
mostly proteins of unknown function or did it include proteins which could potentially be predictive for the environment a particular bacterium could survive in?

4. The paper does need some proof reading, there are quite a few grammatical mistakes.

5. Throughout the paper the authors confuse the word “that” or “which” with “who”. For example in the abstract it says: “Clustering sequences by their ordered domain content give us domain sequence families, who are robust to errors in the gene prediction step.” In this case “who” refers to “domain sequence families” and hence should be substituted with “that” or “which”.

6. The authors frequently use the words “eukaryote” and “prokaryote” as adjectives (e.g. third paragraph of the introduction: “Even if prokaryote genes are in most cases simpler to detect than eukaryote counterparts, there are still problematic cases.”) The proper adjective should be “eukaryotic” or “prokaryotic” (and in the example it should probably also say “their eukaryotic counterparts”).

7. On page four, in the Methods part of the paper, the authors present a formula for the binomial mixture model. In this formula the running variable is k but changes to kappa in a few places (in the formula as well as in the text). This is a bit confusing and needs fixing.

8. In a few places references are given in the following format: “as described in36” in those cases either the “as described in” part should be left out or it should say as described by Snipen and Ussery36” or “as described in one of our earlier papers36”.

9. There are quite a few cases where plural and singular are confused, for example in the last paragraph of page 7 it says “in the first two principal component” it should say “first two principal components”.

10. The figure caption of Figure 7 requires some revision.

11. In the first paragraph of page 11 the reference for Angiouli et al. is missing. The authors also mention in this sentence the recent approach of Angioula et al without explaining what this approach is and how it compares to the presented approach. After reading the manuscript again I noticed that this paper was mentioned also in the beginning of the Results and Discussion section, but I still think it would be helpful for the reader if the authors briefly outlined what Angiouli et al did to improve consistency and in particular how this could be integrated into the presented approach.

12. In general the Conclusions section is harder to read and less clear than the rest of the paper, maybe it would help to spend a little bit of time revising this section.

**Competing Interests:** No competing interests were disclosed.
I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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