

ArrayOme: a program for estimating the sizes of microarray-visualized bacterial genomes

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ABSTRACT

ArrayOme is a new program that calculates the size of genomes represented by microarray-based probes and facilitates recognition of key bacterial strains carrying large numbers of novel genes. Protein-coding sequences (CDS) that are contiguous on annotated reference templates and classified as 'Present' in the test strain by hybridization to microarrays are merged into ICs (ICs). These ICs are then extended to account for flanking intergenic sequences. Finally, the lengths of all extended ICs are summated to yield the 'microarray-visualized genome (MVG)' size. We tested and validated ArrayOme using both experimental and *in silico*-generated genomic hybridization data. MVG sizing of five sequenced *Escherichia coli* and *Shigella* strains resulted in an accuracy of 97–99%, as compared to true genome sizes, when the comprehensive *ShE.coli* meta-array gene sequences (6239 CDS) were used for *in silico* hybridization analysis. However, the *E.coli* CFT073 genome size was underestimated by 14% as this meta-array lacked probes for many CFT073 CDS. ArrayOme permits rapid recognition of discordances between PFGE-measured genome and MVG sizes, thereby enabling high-throughput identification of strains rich in novel genes. Gene discovery studies focused on these strains will greatly facilitate characterization of the global gene pool accessible to individual bacterial species.

INTRODUCTION

To date, the entire genomic sequence of more than 180 bacterial strains has been determined. Based on comparative analysis of multiple genomes of the same species, it is increasingly apparent that some bacteria possess an extremely plastic genome (1–4). Foreign DNA segments, acquired via horizontal gene transfer, result in a genomic mosaic that reflects the lifestyle of the bacterium, pathogenic traits, adaptation to particular ecological niches and evolutionary history (5). This 'optional' genomic repertoire, which we refer to as the 'mobilome' (mobile genome), includes episomal plasmids, transposons, integrons, prophages and a growing list of genomic islands (GIs) (6,7). Pathogenicity islands, the virulence-associated subset of GIs, have now been identified in many bacterial species and are undoubtedly recognized as major players in the moulding of pathogenic traits. The high cost of genome sequencing has been a barrier to high-throughput prospecting of the mobilome (8). Even costly mega-scale metagenomics projects do not facilitate this process as the derived data are largely skewed towards abundant DNA sequences and low-prevalence mobilome sequences would rarely fit within a wider genomic context (9). A rapid and more cost-effective approach to discovering strain-specific DNA sporadically dispersed among hundred of members of the same species remains a major challenge (10).

Since DNA microarrays were first used to compare the genomes of *Mycobacterium bovis* BCG strains with that of *Mycobacterium tuberculosis* strain H37Rv to reveal several strain-specific deletions (11), comparative genomic hybridization technology has been extensively applied to investigate genome diversity among distinct isolates of many bacterial species including *Bacillus anthracis* (12), *Brucella* spp. (13),

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Campylobacter jejuni (14,15), *Escherichia coli* (8,16–18), *Helicobacter pylori* (19), *Mycobacterium* spp. (20,21), *Porphyromonas gingivalis* (22), *Pseudomonas aeruginosa* (23), *Rickettsia prowazekii* (24), *Salmonella* spp. (25), *Shewanella* spp. (26), *Xylella fastidiosa* (27) and *Yersinia* spp. (28,29). Indeed, microarray-based comparative genomic indexing (CGI) (18) has become a powerful generic tool in the study of bacterial pathogens.

In particular, *E.coli* appears to be a front-runner in the genome diversity stakes comprising isolates with a total genome size of 4.5–5.5 Mb (30). The common *E.coli* chromosomal ‘backbone’ of ~3.6 Mb is punctuated by numerous integrated GIs (1,3,16). The ability of *E.coli* to tap into a large mobile gene pool has probably led to its remarkable success as both an innocent commensal and a virulent pathogen, with its near ubiquitous distribution in the environment and in animal hosts. To date, human pathogenic *E.coli* are classified into at least 10 distinct pathotypes including, for reasons of genetic relatedness, the four species of *Shigella* that are now considered to be subspecies of *E.coli* (31).

CGI data identifies the subset of genes common to both the microarray used and the genome of the strain under investigation. This complement of genes represents an entity, which we define as the microarray-visualized genome (MVG). In this paper, we describe, test and validate a new program, ArrayOme v1.0, which provides an accurate estimate of the size of the MVG of a bacterium based on CGI data alone.

Using an approach we have termed Microarray-Assisted mobilome Prospecting (MAMP), we plan to exploit this tool to screen large numbers of isolates and identify *E.coli* and *Shigella* strains that are rich in novel genetic material for further detailed analyses. The MAMP approach combines CGI, ArrayOme and pulsed-field gel electrophoresis (PFGE) to predict the size of the novel, non-microarray-borne mobilome in a test strain (Figure 1). ArrayOme calculates the sizes of the MVGs of strains based on their possession and presumed reference template-like organization of CGI-defined subsets of microarray-borne genes. The sizes of novel mobilomes borne by individual strains can be estimated by comparing MVG sizes with PFGE-measured chromosome sizes. The degree of size-discordance between PFGE-measured physical genomes and MVGs can be used to identify isolates rich in novel genetic material that are likely to carry unique GIs or prophages. Ochman and Jones (16) had estimated that the amount of unique DNA present in four *E.coli* strains investigated ranged from 65 to 1183 kb based on differences in chromosome length and gene content relative to *E.coli* K-12 MG1655.

Previously reported experimental CGI data and *in silico* hybridization data derived from analysis of various *E.coli* and *Shigella* strains ‘hybridized’ against *E.coli* K-12 or the newly developed *ShE.coli* metagenome microarray are used to test ArrayOme v1.0. This generic bioinformatics program can be easily used to analyse any CGI data set for which a reference annotated template exists.

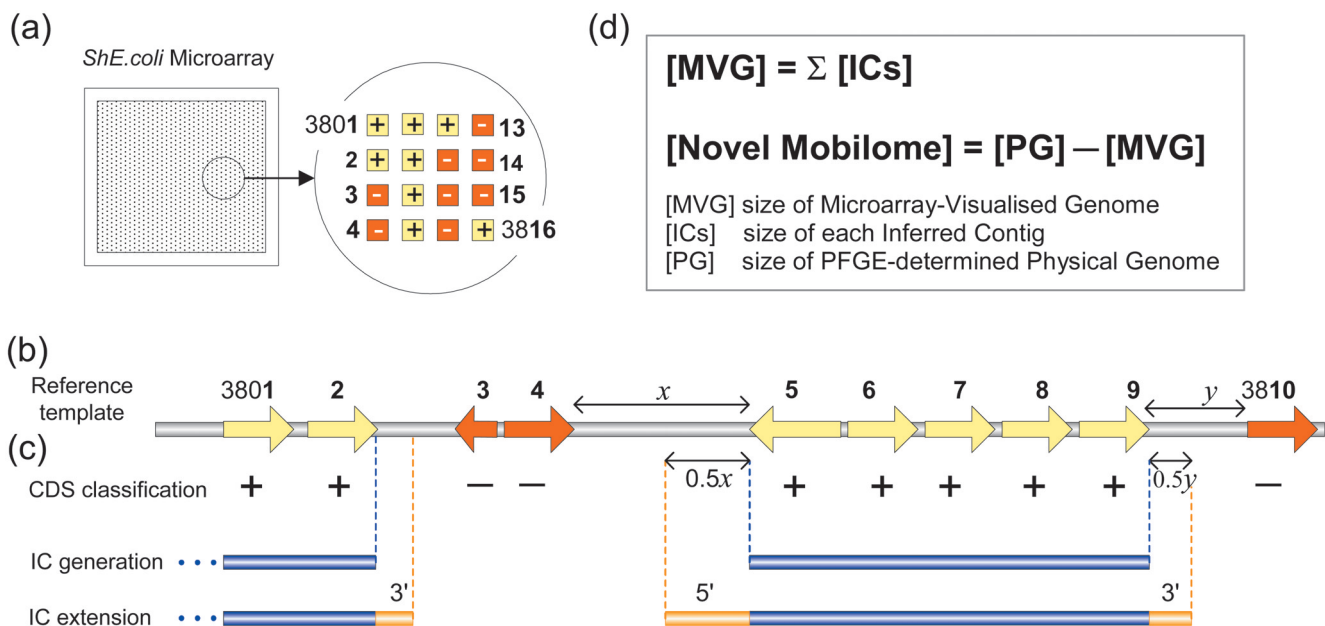


Figure 1. Microarray-Assisted mobilome Prospecting (MAMP): a method for determining the discrepancy between the physical genome size and that accounted for by known genes represented on an expanded species-specific microarray. (a) A schematic representation of a microarray-based CGI output for an hypothetical region of the genome in a strain under investigation by the MAMP technique. Scanned raw data are normalized permitting classification of microarray-represented genes as ‘Present (+)’ or ‘Absent (–)’ in individual test strains. (b) The arrows represent the genetic organization of these CDS within *E.coli* K-12 MG1655, *E.coli* K-12 W3110, *E.coli* O157 EDL933, *S.flexneri* 2a Sf301 or other virulence-associated gene clusters included on the MG1655, W3110 or *ShE.coli* microarrays. (c) Contiguous CDS classified as ‘Present’ are merged into an IC with intergenic non-coding segments between the contiguous CDS included in the corresponding IC. Each IC was then extended in both directions by lengths equal to half the flanking 5’ and 3’ intergenic segments, as indicated by the double-headed arrows of lengths 0.5x and 0.5y in the examples shown. When the *ShE.coli* meta-array sequences were used, each probe was mapped onto a single source reference template to allow for the generation of specific ICs. (d) The size of the MVG was calculated as the sum of all IC lengths. Consequently, the size of the non-microarray-borne novel mobilome was estimated as equal to the discrepancy between the pulsed-field gel electrophoresis-determined physical genome size and the MVG size. Figure not drawn to scale.

MATERIALS AND METHODS

Experimental CGI data

Experimental CGI data versus the *E.coli* K-12 MG1655 or the *E.coli* K-12 W3110 microarrays that had previously been interpreted were taken from the papers of Anjum *et al.* (18) and Fukiya *et al.* (8), respectively. The MG1655 microarray featured 4264 of the 4289 annotated protein-coding sequences (CDS) of MG1655, while the W3110 microarray bore probes specific for 4071 of 4390 W3110-annotated CDS. The microarray-borne DNA probes were amplified by PCR using CDS-specific primers and spotted onto slides as described previously (8,18). Following the reported microarray hybridization and data normalization protocols, the original authors had classed the status of individual CDS in test strains as 'Present' or 'Absent'. Data associated with probes yielding low-reference signals relative to background were excluded from further interpretation. In this study, we have grouped CDS mapped onto these latter probes and CDS that were not represented on the microarray as being of 'Indeterminate' status.

Method to estimate the size of the MVG

A heuristic method was employed to estimate the size of the MG1655 MVG based upon the genetic organization of all 4289 annotated CDS present on the MG1655 reference template. The complete nucleotide sequence and annotation of MG1655 downloaded from GenBank (accession number U00096) served as the reference template. The method used is schematically shown in Figure 1. First, each of the 4289 annotated MG1655 CDS was flagged as 'Present' or 'Absent' in a test strain based on the results of a microarray hybridization experiment. Next, contiguous CDS classified as 'Present' were merged into an inferred contig (IC) with intergenic non-coding segments between these CDS included in the corresponding IC. Each IC was then extended in both directions. The 5' end of each IC was extended by a length equal to half of that of the intergenic segment between the 5' end of the IC and the adjacent 'Absent' CDS. Occasionally, prior to extension, the 5' end of the IC overlapped the flanking 'Absent' CDS. In such cases, the 5' end of the IC was extended by 64 bp, a length approximately equal to half the average intergenic distance in the reference template MG1655. These same steps were employed for the extension of the 3' ends of ICs as well. Finally, the lengths of all extended ICs were summated to yield the MVG size. Details of individual ICs could also be generated as optional output files. A computer program, named ArrayOme v1.0, was encoded using the programming language C++ to implement this algorithm. ArrayOme is freely available on request for academic purposes.

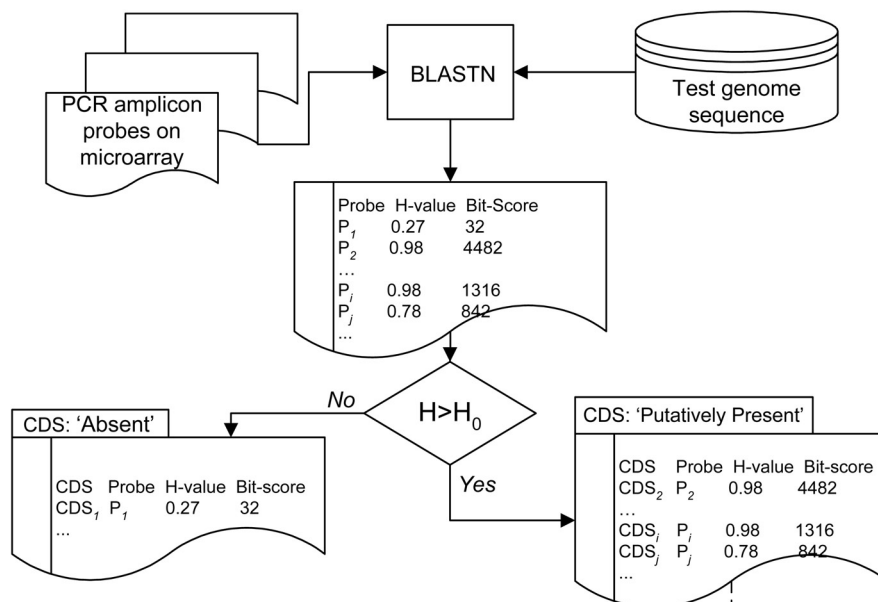
In silico CGI

Simulated CGI data corresponding to *E.coli* and *Shigella* strains with fully sequenced genomes were obtained using an *in silico* DNA-DNA 'hybridization' approach. The DNA sequences of the following genomes were downloaded from GenBank: *E.coli* K-12 MG1655 (accession number U00096) (32), *E.coli* O157:H7 EDL933 (AE005174) (1), *E.coli* O157:H7 Sakai (BA000007) (2), uropathogenic *E.coli* CFT073 (AE014075) (3), *Shigella flexneri* 2a Sf301 (AE005674) (4) and *S.flexneri* 2a 2457T (AE014073) (33).

In a microarray experiment, 'hybridization is a complex process that is not fully understood' (34), in which bound DNA probes on the microarray and labelled targets in solution form heteroduplexes via Watson-Crick base pairing (34). Adjusting the temperature, salt concentrations and the concentration of labelled target molecules can alter the stringency of hybridization. In the *in silico* approach presented here, we utilize the degree and extent of DNA sequence similarity between the probe and target molecules as a surrogate measure for the likelihood of formation of stable heteroduplexes between these entities. The *in silico* CGI procedure applied included the following steps (Figure 2):

- (i) Each of the DNA sequences of the probes was used as a query in a similarity search against a test genome sequence using a locally installed version of BLASTN (35) and default NCBI BLASTN parameters.
- (ii) The stretch of sequence from the test genome with the highest Bit score for each query sequence was retrieved and a homology score (H -value) calculated for each probe in turn. This homology score had been proposed by Fukiya *et al.* (8) and reflected the degree of similarity between the matching test genome sequence and the probe itself in terms of the length of match and the percentage sequence identity at the DNA level. For each query, the H -value was calculated as follows (8): $H = (i \times l) / l_q$, where i is the level of identity of the region with the highest Bit score expressed as a frequency of between 0 and 1, l , the length of the highest scoring matching sequence (including gaps) and l_q , the query length. If there were no matching sequences with a BLASTN E -value < 0.01 , the H -value assigned to that query sequence was defined as zero (8). Therefore, H belonged to the set, $H \in [0, 1]$.
- (iii) For each probe, a threshold value of H (H_0) was used to determine whether the corresponding CDS was to be classed as 'Putatively Present' or 'Absent'. As mentioned above, each probe had been assigned an H -value. If $H > H_0$ for a particular probe, the CDS mapped onto that probe was considered to be 'Putatively Present' and the probe itself further scrutinized to minimize misclassification of CDS as 'Present' due to signals arising from cross-hybridization events. The threshold value, H_0 , was selected to optimize sensitivity and specificity of the *in silico* approach versus true experimental CGI analysis (see Results and Discussion).
- (iv) To increase the specificity of the *in silico* CGI output, cross-hybridization analysis was performed by a BLASTN search of each probe sequence against the remaining probe sequences in turn. For instance, a BLASTN similarity search was performed using the sequence of probe P_i as a query against the sequence of probe P_j ($i \neq j$) and an H -value designated as H' was obtained. If $H' > H_0$, the probes P_i and P_j were considered as a potential cross-hybridization pair and the original BLASTN results linked to these probes further examined. The CDS corresponding to either probe P_i or P_j would be re-classified as 'Absent' if the results of the BLASTN search against the test genome satisfied both of the following conditions: (i) it had a lower Bit score than that associated with the second probe and (ii) it had a H -value less than H_x , a second H threshold value that was also determined following review of the real

(a) Primary similarity



(b) Cross-hybridization filter

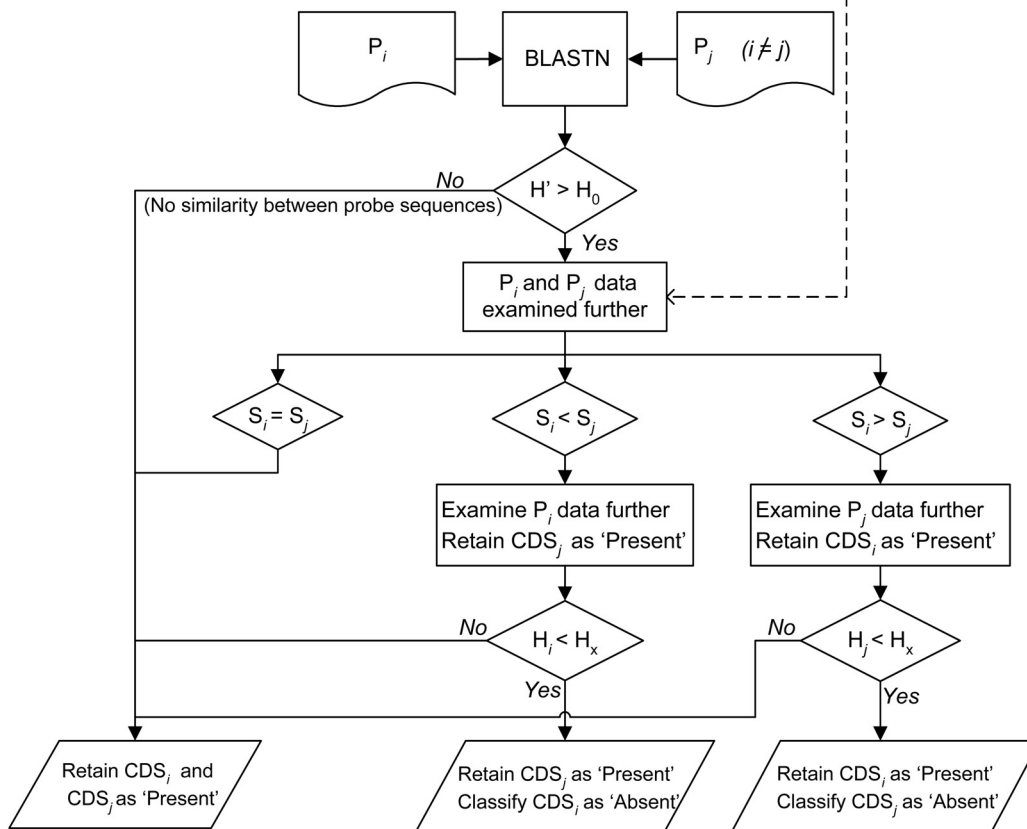


Figure 2. A flow diagram of the logic steps used for *in silico* CGI. CDS, P, H and S denote CDS mapped onto specific probes, individual microarray-borne amplicon probes, H-values and Bit scores, respectively. P_i and P_j denote distinct individual probes, with the subscripts i and j identify matching CDS, H-values and Bit scores. The steps involved in the primary similarity search are shown in (a), while the cross-hybridization algorithm that re-categorizes CDS from 'Putatively Present' to 'Present' or 'Absent' are shown in (b). A direct comparison of *in silico* generated data and true experimental CGI data for *E. coli* EDL933 obtained using the MG1655 microarray was used to validate the algorithm. The key threshold values, $H_0 = 0.40$ and $H_x = 0.96$, were set to optimize the sensitivity and specificity of identifying genes as 'Present' when using the *in silico* as compared to experimental approach (see Figure 3).

hybridization results of *E.coli* EDL933 versus the MG1655 microarray (see Results and Discussion). The remaining 'Putatively Present' CDS were now designated as 'Present' resulting in a final *in silico* CGI data set of CDS classed as 'Present' or 'Absent' only. A Perl program was written to perform the BLASTN-based *in silico* DNA-DNA hybridization steps with a C++ program used to implement the remaining steps and produce the final *in silico* CGI output file.

RESULTS AND DISCUSSION

Analysis of *E.coli* K-12 microarray data using ArrayOme v1.0

In a 'compact' bacterial genome, CDS occupy the majority of the chromosome. For example, 88% of the *E.coli* K-12 MG1655 chromosome contains CDS. The average CDS length in MG1655 is 953 bp, while the mean distance between CDS is 128 bp. We have utilized previously reported CGI data derived from the *E.coli* K-12 MG1655 microarray (18) and the complete chromosomal sequence of MG1655 as a reference template to predict the MVG sizes of a range of pathogenic *E.coli* strains associated with diarrhoeal diseases. The fully sequenced enterohaemorrhagic *E.coli* (EHEC) strain EDL933 was included in this set. Of the 4264 MG1655 CDS represented as probes on the microarray, 3775 were identified as 'Present' in EDL933, while the status of a further 114 was 'Indeterminate' (see Materials and Methods). Inputs of these data into ArrayOme produced 113 or 193 ICs (see Figure 1) scattered along the MG1655 template chromosome depending on whether the 'Indeterminate' CDS were classified as 'Present' or 'Absent', respectively. After 5' and 3' extension of each of these ICs, the MG1655-MVG size of EDL933 was calculated as the sum of all IC lengths resulting in an estimate of 4141–4254 kb. The size of the genome occupied by non-MG1655 genes was then deduced by subtracting this MVG size from its known physical size based on the available EDL933 genome sequence to yield an estimate of 1275–1387 kb for the non-MG1655 mobilome present in EDL933. This number compared well with the previously reported total length (1.34 Mb) of species-specific 'O-islands' identified in the EDL933 deduced following pairwise similarity alignment between the EDL933 and MG1655 genomes (1). Ochman and Jones (16) had proposed that the size of the novel mobilome of a strain could be estimated based on the differences in chromosome lengths and gene contents between a test strain and a reference strain, but had not stipulated in detail the procedure used to calculate the amount of 'missing DNA' relative to the reference strain. Importantly, the ArrayOme approach takes account of likely lost intergenic sequences (Figure 1). As mentioned above, the average length of intergenic sequences in MG1655 was 128 bp. Hence, as there were 400–514 missing CDS in EDL933 the size of its non-MG1655 mobilome would have been underestimated by ~51–66 kb using a method based on the total size of missing CDS alone. To improve prediction accuracy, we used the strategy of creating ICs and then subjecting these virtual entities to rule-defined 5' and 3' extensions to allow the inclusion of likely conserved intergenic stretches within the MVG. Based on this algorithm, reference template-borne

non-coding sequences flanking CDS identified as 'Absent' were also attributed as missing from the final virtual genome.

Similar calculations were performed with the remaining CGI data reported by Anjum *et al.* (18) for the 25 other pathogenic *E.coli* strains. However, because of the large number of CDS classified as 'Indeterminate' for many of these strains, we applied a set of ArrayOme-implemented logic rules in an attempt to better assign these genes based on biological realities. First according to Rule I, all 'Indeterminate' CDS that were conserved at a nucleotide level across all six fully sequenced *E.coli* and *Shigella* genomes were re-classified as 'Present'. In this study, we chose a threshold of $H > 0.64$ as it reflected a degree of nucleotide conservation equivalent to 80% identity over 80% of the CDS length. Selection of H -values ranging from 0.4 to 0.8 resulted only in minor differences in the number of CDS regarded as conserved. Next according to Rule II, single and paired 'Indeterminate' CDS that were directly flanked by genes identified as 'Present' were also deemed to be 'Present', while the reverse was applied to CDS flanked by 'Absent' CDS. When 'Indeterminate' CDS were present in clusters of three or more, the implicated CDS were classified as 'Absent' regardless of the status of the flanking CDS. Indeed, if an intermediate category was used in the interpretation of CGI data, many such CDS would be likely to represent truly divergent genomic sequences. An additional sub-rule based on the G+C-content of the 'Indeterminate' CDS cluster as compared to the genome average could also be applied to clusters of ≥ 3 CDS to vary the final interpretation (see instructions for ArrayOme program). The choice of rules and the order in which these were to be implemented were entirely user-defined. Depending on the interpretation criteria applied, many CDS classed as 'Indeterminate' following experimental CGI analysis might represent examples of gene sequence polymorphism resulting in weaker, poorly interpretable microarray hybridization signals. Indeed, when wide ranges of inferred MVG sizes are obtained following analysis of un-reconciled microarray data these could reflect the extent of genomes occupied by microarray-related but divergent genes. The MVG sizes for all 26 *E.coli* strains predicted using the reconciled CGI data and ArrayOme are available in the Supplementary Material. The estimated number of 'Absent' MG1655 CDS ranged from 222 to 888, while the MVG size estimates spanned 3.7–4.4 Mb. The chromosome sizes of natural *E.coli* strains vary by as much as 1 Mb and are currently recognized to range from 4.5 to 5.5 Mb (30). Based on these limits, we speculate that as much as 1.8 Mb ($5.5 - 3.7 = 1.8$ Mb) of the genome of one of these *E.coli* isolates may differ from that of *E.coli* K-12 and that this mobilome alone could harbour up to 1800 non-MG1655 CDS.

Recently, Fukuya *et al.* (8) reported the CGI results of 22 pathogenic *E.coli* and *Shigella* strains using an *E.coli* K-12 strain W3110 microarray that bore probes for 4071 of 4390 annotated CDS. We have estimated the MVG sizes of these test strains using their reported CGI data. The 4641 433 bp *E.coli* K-12 W3110 chromosome was used as the reference template for this analysis (<http://gib.genes.nig.ac.jp/>). The enteropathogenic *E.coli* strain E2348/69 was found to possess 3568 W3110 CDS (8). CDS categorized as 'Indeterminate' were re-classified based on the logic rules described above. Entry of the reconciled E2348/69 CGI data into ArrayOme

resulted in the generation of 164 or 165 ICs and an estimated MVG size of 4096–4125 kb. Consequently, as the physical chromosome size of *E.coli* E2348/69 had previously been estimated by PFGE as ~ 4.7 Mb (17), it was predicted to possess a non-W3110-related genome complement of ~ 0.6 Mb, suggesting an accessory gene content of about 600 relative to W3110. The soon to be available E2348/69 genome sequence (http://www.sanger.ac.uk/Projects/Escherichia_Shigella/) would allow validation of these predictions. The estimated W3110-MVG sizes for all 22 strains are available as Supplementary Material.

Determination of threshold values of H to be used in *in silico* CGI

To classify CDS as ‘Present’ or ‘Absent’ using a BLASTN-facilitated *in silico* DNA–DNA hybridization approach, suitable threshold values of H (see Materials and Methods) were required. Based on the experimental microarray data of Anjum *et al.* (18) for EDL933, 400 CDS were classified as ‘Absent’, 3775 CDS as ‘Present’ and a further 89 as ‘Indeterminate’ out of a total of 4264 MG1655-CDS probes spotted onto the microarray. BLASTN-based *in silico* hybridization of the 4264 microarray probes versus the *E.coli* EDL933 genome sequence resulted in similarity Bit scores and the H -values for each of the amplicon probe sequences. The distribution of H -values corresponding to probes was bipolar with the average

H -value for the 400 ‘Absent’ CDS (0.087; SD = 0.009) being significantly less than that for the 3775 CDS identified as ‘Present’ (0.962; SD = 0.110) and the 89 microarray-represented CDS of ‘Indeterminate’ status (0.852; SD = 0.309) (Figure 3). The H -values associated with the 89 ‘Indeterminate’ calls were predominantly clustered with those of probes mapped onto CDS categorized as ‘Present’, with only 12 having an H -value of ≤ 0.40 . Based on the observed spread of H -values corresponding to the true experimental data, the threshold value, $H_0 = 0.40$, was selected as the primary discriminant between CDS designated as ‘Putatively Present’ and those classed as ‘Absent’ (Figures 2 and 3). BLASTN data corresponding to CDS classed as ‘Putatively Present’ were then further scrutinized using the cross-hybridization filter described in Materials and Methods. The H_x threshold value was set at a high value of 0.96 to minimize the likelihood of miscalling a CDS as ‘Present’ when sequence similarity existed between its probe and a second microarray-borne probe(s) mapped onto an alternate CDS (Figure 2). With the EDL933 data set, this algorithm correctly predicted 389/400 ‘Absent’ and 3714/3775 ‘Present’ CDS for EDL933, yielding satisfactory sensitivity (98.4%) and specificity rates (99.7%) in the *in silico* CGI classification of CDS as ‘Present’ (Figure 3). By using these same H -value parameters, $\sim 86\%$ (77/89) of the ‘Indeterminate’ CDS were classed as ‘Present’ by *in silico* simulation. This number was similar

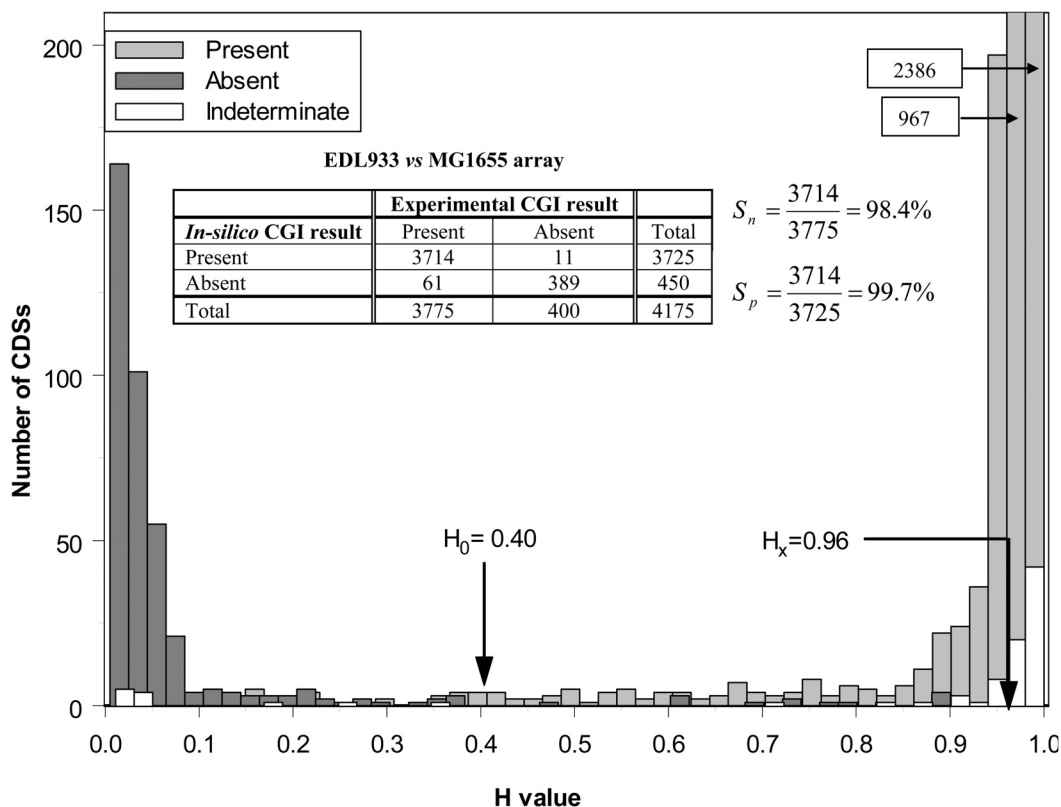


Figure 3. The distribution of H -values corresponding to 4264 amplicon probes spotted onto the MG1655 microarray obtained following a BLASTN similarity search against the EDL933 chromosomal sequence are shown. Interval-grouped H -values are plotted with the data stratified into the experimental CGI categories of ‘Present’ (3775), ‘Absent’ (400) and ‘Indeterminate’ (89). The selected threshold values for *in silico* CGI, $H_0 = 0.40$ and $H_x = 0.96$, are as indicated. The numbers in the boxes at the top right corner correspond to the ‘Number of CDS’ associated with the two bars that extend beyond the limits of the graph. The inset table shows a direct comparison of experimental CGI data derived by Anjum *et al.* (18) and *in silico* CGI data for the CDS classified as ‘Present’ or ‘Absent’ only. The sensitivity (S_n) and specificity (S_p) of identifying genes as ‘Present’ when using the *in silico* as compared to experimental approach, are shown on the right-hand side.

to the percentages of experimentally defined 'Indeterminate' CDS that were re-categorized as 'Present' following application of our logic rules concerning species-wide CDS conservation and the status of flanking CDS and reaffirms the value of these rules for the purpose of MVG sizing. In subsequent *in silico* hybridization simulations, CDS were judged to be 'Present' or 'Absent' using the algorithm shown in Figure 2 with the threshold values set at $H_0 = 0.40$ and $H_x = 0.96$.

Estimated MVG sizes based on *in silico* CGI data generated using the *ShE.coli* meta-array

The recently developed PCR amplicon-based *ShE.coli* microarray contains many more *E.coli* CDS than the first generation *E.coli* K-12 microarrays. The *ShE.coli* meta-array probe sequences used in this study represented 6239 CDS comprising 4264 *E.coli* K-12 MG1655 CDS, 1101 *E.coli* EDL933 CDS, 516 *S.flexneri* 2a Sf301 CDS and a further 358 virulence-associated *E.coli* CDS derived from strains representative of different pathotypes of *E.coli*, particularly enteropathogenic *E.coli* (EPEC) and enterotoxigenic *E.coli* (ETEC). Of the 358 virulence-associated CDS, 132 were located on *E.coli* chromosomes while the remaining 226 mapped onto a range of plasmids based on published literature and GenBank submissions. The 25 MG1655 CDS not represented on the microarray were assigned based on logic rules described in the text with any remaining 'Indeterminate' CDS classed as 'Present' for simplicity. The *ShE.coli* metagenome microarray sequences presently constitute the most comprehensive representation of the global gene pool available to *E.coli* and *Shigella* strains.

In this study, we have used the five completed *E.coli* and *Shigella* genomes, MG1655, Sakai, CFT073, Sf301 and 2457T, to assess the ArrayOme-based estimation of MVG sizes with data derived by *in silico* CGI using sequences of probes present on the MG1655 and *ShE.coli* microarrays. Results of the analysis with EDL933 were also shown. However, it should be noted that as the EDL933 experimental

data were used to train the *in silico* CGI analysis, the simulated output for this strain was of minimal significance. The total sets of MG1655 and *ShE.coli* microarray DNA probes were subjected to BLASTN analysis against each of the six completed genomes and the results interpreted using the logic steps shown in Figure 2. Given the *in silico* CGI algorithm applied, all microarray-represented CDS were classified as either 'Present' or 'Absent' with no 'Indeterminate' category. When the EHEC O157:H7 Sakai genome was analysed using the *ShE.coli* microarray, 5082 CDS were deduced to be 'Present', which comprises 3780 MG1655 CDS, 1083 EDL933 CDS, 164 Sf301 CDS, 50 other *E.coli* chromosomal virulence-associated CDS and 5 plasmid-borne virulence genes (Table 1).

MVG sizes were deduced using *in silico* *ShE.coli* microarray data in a manner similar to that described for the *E.coli* K-12 microarrays. In addition to the MG1655 chromosome, we used the annotated chromosomal sequences of *E.coli* EDL933 and *S.flexneri* Sf301 as reference templates for creating ICs. The genetic organizations of all other *E.coli* genes represented on the microarray were inferred from sequence and annotation details available from GenBank and related publications. When CGI data derived using meta-arrays are analysed, ArrayOme generates a set of linear ICs using as the reference template a single genome or GenBank sequence for each IC. The choice of template for each IC is directed by the contents of the user-generated Microarray Index file that specifies mapping details among probes, CDS and reference templates. Each probe is mapped onto a single CDS, which in turn is mapped onto a single template only. Further details of the algorithm used are available in the Supplementary Material. Comparisons of actual chromosome sizes and ArrayOme-estimated MVG sizes derived following the analysis of *in silico* CGI data are shown in Table 1. For five of the six test genomes, the discrepancy between the physical chromosome size and the MVG size as calculated using *ShE.coli*-derived *in silico* data was <2.8%. However, with CFT073 the margin of error was significantly larger; ArrayOme

Table 1. The ArrayOme-predicted MVG sizes of *E.coli* and *Shigella* strains based on data derived by *in silico* hybridization of the complete genomes against the *ShE.coli* microarray amplicon probe sequences

Strain	Size of the complete chromosome (kb) ^a	No. of CDS classified as 'Present' ^b					Size of the <i>ShE.coli</i> -MVG [MG1655-MVG] (kb) ^c	Discrepancy between the chromosome length and size of the <i>ShE.coli</i> -MVG (kb) (% error) ^d
		MG1655 CDS (n = 4264 + 25)	EDL933-specific CDS (n = 1101)	Sf301-specific CDS (n = 516)	Other chromosomal CDS (n = 132)	Plasmid-borne CDS (n = 226)		
<i>E.coli</i> O157:H7 EDL933	5528	3783	1097	162	50	5	5566 [4192]	+38 (+0.7)
<i>E.coli</i> K-12 MG1655	4639	4288	58	59	17	8	4771 [4639]	+132 (+2.8)
<i>E.coli</i> O157:H7 Sakai	5498	3780	1083	164	50	5	5556 [4191]	+58 (+1.1)
<i>S.flexneri</i> 2a Sf301	4607	3541	122	515	16	14	4519 [3882]	-88 (-1.9)
<i>S.flexneri</i> 2457T	4599	3534	123	499	17	14	4507 [3882]	-92 (-2.0)
<i>E.coli</i> UPEC CFT073	5231	3638	278	174	26	9	4498 [4013]	-733 (-14.0)

^aThe lengths of the complete chromosomes, shown to the nearest kilobase (kb), are based on genome sequences lodged with GenBank.

^bA total of 4264 *E.coli* K-12 MG1655 CDS, 1101 *E.coli* EDL933 CDS, 516 *S.flexneri* 2a Sf301 CDS, 132 other *E.coli* chromosomal virulence-associated CDS and 226 plasmid-borne *E.coli* virulence genes are classified as 'Present' or 'Absent' based on *in silico* CGI analysis. The 25 MG1655 CDS not represented on the microarray were assigned based on logic rules described in the text with any remaining 'Indeterminate' CDS classed as 'Present' for simplicity, leading to the classification of all 4289 MG1655 CDS.

^cThe data corresponding to plasmid-borne genes was omitted when calculating MVG sizes as these CDS would normally be considered to be borne on episomal entities other than the main chromosome given their original identified location. The ArrayOme-predicted sizes of *ShE.coli*-MVGs (left) and MG1655-MVGs (right, square brackets) are shown to the nearest kilobase (kb).

^dThe percentage errors between the reported lengths of the chromosomes and the sizes of the *ShE.coli*-MVGs are shown within parentheses.

Table 2. The MVG sizes of 'virtual genomes' constructed by precise deletion of Islander-defined GIs

Virtual strain designation	Virtual genome			No. of CDS classified as 'Present' ^b				Plasmid-borne CDS (n = 226)	Size of the MVG (kb) ^c	Discrepancy between the sizes of genome and MVG (kb) (% error) ^d
	No. of GIs deleted ^a	Total length of deletion (bp)	Size of the derivative virtual genome (kb)	MG1655 CDS (n = 4264 + 25)	EDL933-specific CDS (n = 1101)	Sf301-specific CDS (n = 516)	Other chromosomal CDS (n = 132)			
EDL933-V9	9 (398)	321 129	5207	3776	905	144	48	3	5351	+144 (+2.8)
MG1655-V3	3 (81)	65 825	4573	4219	53	55	17	5	4705	+132 (+2.9)
Sakai-V8	8 (327)	235 511	5263	3774	888	142	48	3	5333	+70 (+1.3)
Sf301-V3	3 (70)	75 968	4531	3538	112	469	16	11	4445	-86 (-1.9)
2457T-V4	4 (68)	82 476	4517	3527	113	448	17	11	4424	-93 (-2.1)
CFT073-V6	6 (435)	395 611	4836	3606	214	138	20	6	4357	-479 (-9.9)

The ArrayOme-facilitated MVG size prediction was based on data derived by *in silico* CGI hybridization of the complete genomes against the *ShE.coli* microarray amplicon probe sequences.

^aThe GIs that were deleted from the corresponding complete genomes were based on the Islander-derived data of Mantri and Williams (36). The details of these GIs along with the precise boundaries were downloaded from the database of Islander (www.indiana.edu/~islander) (36). The total numbers of CDS contained within the deleted GIs are shown within parentheses.

^bA total of 4264 *E.coli* K-12 MG1655 CDS, 1101 *E.coli* EDL933 CDS, 516 *S.flexneri* 2a Sf301 CDS, 132 other *E.coli* chromosomal virulence-associated CDS and 226 plasmid-borne *E.coli* virulence genes are classified as 'Present' or 'Absent' based on *in silico* CGI analysis. The 25 MG1655 CDS not represented on the microarray were assigned based on logic rules described in the text with any remaining 'Indeterminate' CDS classed as 'Present' for simplicity.

^cThe data corresponding to plasmid-borne genes was omitted when calculating MVG sizes as these CDS would normally be considered to be borne on episomal entities other than the main chromosome given their original identified location. ArrayOme-predicted MVGs are shown to the nearest kilobase (kb).

^dThe percentage errors between the lengths of virtual chromosomes and the sizes of the *ShE.coli*-MVGs are shown within parentheses.

underestimated the true genome size of CFT073 by 733 kb or 14.0% of the complete genome. This was entirely consistent with the fact that the *ShE.coli* microarray lacked probes specific for at least 1501 CFT073-borne genes [(3) and H.-Y. Ou and K. Rajakumar, unpublished data]. As expected, analysis of MG1655 microarray data yielded significantly lower estimates of MVG sizes for all strains except MG1655, consistent with the known incomplete coverage of these genomes on this single-strain microarray (Table 1).

In order to further validate the ArrayOme tool, we generated a set of virtual derivative genomes. These derivatives were constructed from the six intact *E.coli* and *Shigella* genomes analysed above following defined 'deletions' of tRNA gene-borne GIs previously identified by the Islander algorithm (36). These virtual genomes were analysed by *in silico* comparison with the *ShE.coli* microarray sequences and the resulting data processed using ArrayOme (Table 2). Eight archetypal GIs encompassing 235 511 bp were deleted from the EHEC Sakai genome. Taking precise account of likely island boundaries, we constructed a virtual genome of 5 262 939 bp in length. The predicted MVG size of this derivative (Sakai-V8) was 5333 kb; a discrepancy of ~1.3% compared with the known length of its virtual genome. A similar level of precision was obtained with four of the five other virtual derivatives but with CFT073-V6 the margin of error was significantly larger. However, the degree of discordance between the 'physical' length and the MVG size had fallen from 14.0% for the parent strain to 9.9% for the CFT073-V6 hypothetical construct, consistent with the fact that the deletion derivative had lost 311 CFT073-borne genes not represented on the *ShE.coli* microarray [(3) and H.-Y. Ou and K. Rajakumar, unpublished data].

Predicted size of mobilomes in *E.coli* and *Shigella* strains based on *in silico* CGI data generated using the *E.coli* K-12 MG1655 microarray

The sequenced genomes of *E.coli* O157:H7 Sakai, uropathogenic *E.coli* CFT073 and *S.flexneri* Sf301 were used to test the

proposed method of predicting the size of the non-MG1655 mobilome using results obtained via simulated CGI analysis versus the MG1655 microarray (Table 3). For the Sakai, CFT073 and Sf301 genomes, there were 3822, 3662 and 3553 CDS classified as 'Present', respectively. Based on this *in silico* CGI data and the rule-based assignment of the 25 unrepresented MG1655 CDS as described earlier, the size of the MVG was estimated for each strain in turn. The discrepancies between the true chromosome and the MVG lengths were then used to compute the size of individual strain-borne mobilomes. These sizes were then compared with the sizes of strain-specific sequences reported by the authors of the original genome sequences published. For example, in CFT073 there were 191 ICs and the MVG size was estimated to be 4013 kb. Thus, the size of novel mobilome was calculated as ~1218 kb, which was comparable with the total length of non-MG1655 sequences (1306 kb) reported by Welch *et al.* (3). The results of this analysis for all three strains are shown in Table 3.

Contribution of IS elements to MVG size estimates

In all calculations performed to this point, we have included the contribution of IS-associated CDS present in MG1655, EDL933 and Sf301, where these have been identified as 'Present' based on experimental or *in silico* CGI data. The MG1655 genome contains 52 annotated IS-associated CDS, particularly comprising IS1, IS2, IS3 and IS5, contributing to up to 44 kb of additional DNA. In addition, *ShE.coli* probes mapped onto IS-associated CDS on EDL933 and SF301 templates accounted for up to 73 kb of the MVG size. However, ArrayOme could be run using an option that assigned all IS-associated CDS as 'Absent' resulting in reduced MVG size estimates (Supplementary Material). Empirical or experimentally guided adjustments could then be made to account for the estimated IS content of individual strains. ArrayOme also offers the option of accounting for other amplified

Table 3. ArrayOme-predicted MVG sizes of three sequenced *E.coli* and *Shigella* genomes based on *in silico* CGI data derived using the *E.coli* K-12 MG1655 microarray

Strain	Sequenced genome Size of chromosome (bp)	Total size of non-MG1655 sequences (bp) ^a	MVG No. of CDS classified as 'Present' ^b	Size of the MVG (kb)	Size of the non-MG1655 mobilome (kb) (% error) ^c
<i>E.coli</i> O157:H7 Sakai	5 498 450	1 393 070	3822	4191	1307 (−6.2)
<i>E.coli</i> UPEC CFT073	5 231 428	1 306 391	3662	4013	1218 (−6.7)
<i>S.flexneri</i> 2a Sf301	4 607 203	~700 000	3553	3882	~725 (∼+3.6)

^aThe total lengths of strain-specific sequences with respect to *E.coli* K-12 MG1655 genome were reported by the authors of the published sequences. The extent of non-MG1655 sequences present in *S.flexneri* Sf301 was reported by Jin *et al.* (4) as ~0.7 Mb.

^bThe 4264 annotated *E.coli* K-12 MG1655 CDS represented on the microarray were classified as 'Present' or 'Absent' based on *in silico* CGI analysis described in the text. The 25 MG1655 genes not represented on the microarray were assigned based on logic rules described in the text with any remaining 'Indeterminate' CDS classed as 'Present' for simplicity.

^cThe percentage errors between the reported lengths of the non-MG1655 sequences and the sizes of the non-MG1655 mobilomes as determined using ArrayOme are shown within parentheses.

CDS that may be borne on duplicated regions of the genome in its calculation of MVG sizes.

CONCLUSION

In summary, the ArrayOme program that we have developed offers a powerful and simple means of transforming microarray data into an accurate estimate of bacterial genome size. The ArrayOme program is versatile and can process any CGI data set for which a suitable reference genomic template(s) is available, thus offering to add significant value to both pre-existing and newly generated data sets. We have applied ArrayOme to analyse freely available CGI data derived using a two-strain *H.pylori* array (19) and a *Salmonella enterica* array (25), and have included our findings in the Supplementary Material. Coupled with physical genome sizing, researchers will be able to exploit information on MVG sizes for high-throughput identification of strains rich in novel genetic material. Indeed, the ability of ArrayOme to predict the genome size of a strain to within ~2% when CGI data are derived using a comprehensive species-related meta-array, suggests that the accuracy of PFGE-based chromosome sizing would be the major limitation to Microarray-Assisted mobilome Prospecting (MAMP). PFGE-based chromosome sizing typically gives rise to errors of ~5–10% depending on the detailed strategy used and the presence or absence of confounding episomal replicons (37). However, we are currently developing a PFGE strategy based on the sizing of linearized bacterial chromosomes. Together with the use of standard DNA size markers and accurate calibration standards generated from the chromosomes of various sequenced strains, an accuracy of ±2% should be achievable. In addition to the identification of strains for gene prospecting studies, MAMP would be invaluable for selecting strains for future genome sequencing projects, ensuring maximum return of novel, non-redundant data per sequenced genome.

It is clear that even with current high-throughput genomic sequencing facilities, it will not be feasible to sequence hundreds of isolates to identify and decode the global gene pool accessible to a single bacterial species. However, the application of a strategy such as MAMP, underpinned by ArrayOme and CGI technology, followed by either random or targeted gene-discovery studies focused on selected strains bearing mobilome-rich regions would make a major contribution to this effort.

SUPPLEMENTARY MATERIAL

Supplementary Material is available at NAR Online.

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