Diversification of bacterial genome content through distinct mechanisms over different timescales

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Bacterial populations often consist of multiple co-circulating lineages. Determining how such population structures arise requires understanding what drives bacterial diversification. Using 616 systematically sampled genomes, we show that Streptococcus pneumoniae lineages are typically characterized by combinations of infrequently transferred stable genomic islands: those moving primarily through transformation, along with integrative and conjugative elements and phage-related chromosomal islands. The only lineage containing extensive unique sequence corresponds to a set of atypical unencapsulated isolates that may represent a distinct species. However, prophage content is highly variable even within lineages, suggesting frequent horizontal transmission that would necessitate rapidly diversifying anti-phage mechanisms to prevent these viruses sweeping through populations. Correspondingly, two loci encoding Type I restriction-modification systems able to change their specificity over short timescales through intragenomic recombination are ubiquitous across the collection. Hence short-term pneumococcal variation is characterized by movement of phage and intragenomic rearrangements, with the slower transfer of stable loci distinguishing lineages.

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Streptococcus pneumoniae is a human respiratory commensal and pathogen in which extensive genetic diversity underlies phenotypic variation in traits such as antibiotic resistance, virulence and antigenic profile. Alongside considerable allelic variation in the core genome, the species contains many ‘genomic islands’ (GIs); genetic loci only found in a subset of the population. Any GI may be transferred between cells through transformation, as pneumococci possess a competence system. Some GIs, referred to as ‘mobile genetic elements’ (MGEs), encode functions that promote their own transfer between cells. At least three types of MGEs have been characterized in S. pneumoniae: phage, most commonly of the Siphoviridae family; plasmids, of which just two cryptic examples are known in pneumococci; and integrative and conjugative elements (ICEs), which have played an important role in the spread of antibiotic resistance.

Horizontal movement of DNA can be limited by bacterial ‘immunity’ mechanisms. Although pneumococci lack CRISPR elements, they do encode restriction-modification systems (RMSs) that can cleave MGEs when present within the cell as double-stranded DNA. While the pneumococcal competence machinery imports DNA into the cell in restriction-insensitive single-stranded form, the acquisition of novel GIs by this mechanism necessitates the synthesis of the complementary strand of DNA after integration into the chromosome. If the pattern of modification of the imported DNA differs from that of the recipient’s genome, the locus may become vulnerable to endonucleolysis by RMSs that cleave unmodified motifs. Hence, variation in many RMSs affects multiple mechanisms of GI transfer. However, the best-characterized pneumococcal RMSs, DpnI and DpnII, do not inhibit the acquisition of GIs by transformation. DpnI cleaves GATC motifs when fully methylated in double-stranded DNA, while the DpnII RMS cleaves unmethylated GATC motifs, but encodes a methylase that modifies imported single-stranded DNA. Another factor that has been suggested to affect the movement of sequence is the lineage’s ‘pherotype’, determined by the sequence of the competence stimulating peptide (CSP) pheromone it secretes. CSP is a critical signal in triggering competence for transformation, and multiple sets of cognate signalling peptides and receptors are found in the population. It remains controversial as to whether the exchange of sequence is inhibited by differences in pherotype between pneumococci.

A recently published set of 616 draft pneumococcal genomes provides an opportunity to examine the extent of, and processes underlying, variation in gene content within a single population. This systematic collection of isolates carried by children in Massachusetts was previously divided into fifteen monophyletic sequence clusters (SCs), and a sixteenth diverse group of rarer genotypes (SC16), based on variation in the core genome. One monophyletic SC (SC12), composed of atypical unencapsulated pneumococci that have caused outbreaks of conjunctivitis, was found to be a distinct outlier from the rest of the population. The analysis presented here describes the evolutionary processes that generate this population structure. SCs were found to be characterized by their complement of stable GIs, including those transferred primarily through transformation and conjugation, while changes over shorter timescales frequently represented the consequence of phage transmission and intragenomic recombination.

**Results**

SCs have distinct accessory genomes. The original analysis of 616 pneumococcal genome sequences identified 5,442 clusters of orthologous genes (COGs), of which around 1,500 were ‘core’ to almost all isolates and around 3,000 were rare. Applying the ‘power law’ method for quantifying the pangenome suggested

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**Figure 1 | Existence of distinct clusters in the pneumococcal population and the properties of the cCOGs with which they are associated.**

(a) Comparison of pairwise distances between isolates in terms of their core genome divergence, as measured by the cophenetic distance calculated from a maximum likelihood core genome phylogeny, and the difference in their accessory genomes, as measured by the Jaccard distance based on the variation in the COG content of their sequences. Points in red indicate comparisons within monophyletic sequence clusters, while purple points represent comparisons between isolates within the diverse SC16. Points in green indicate comparisons between the atypical unencapsulated isolates of SC12 and other sequence clusters; points in turquoise represent all other comparisons between isolates in different sequence clusters. (b) Properties of the COGs characteristic of SC12. The 78 COGs found in >95% of SC12 isolates, and found at a frequency <5% in the other monophyletic sequence clusters, were classified according to their function or location. (c) The 141 cCOGs of all other sequence clusters classified in the same manner.
the gene pool available to this population was unbounded (Supplementary Fig. 1). However, this was heavily influenced by numerous rare COGs that individually had little impact on the population structure, and were the most likely to represent false-positive gene predictions. An alternative representation (Fig. 1) showed the distribution of variation across the population using pairwise comparisons between isolates. This revealed three distinct groups of points that suggested differences in gene content were approximately proportional to core genome divergence.

The mainly red group of points nearest the origin of the plot demonstrated isolates within the same monophyletic SC were highly similar in their core and accessory genomes, while the set of turquoise points showed the greater level of divergence between representatives of different non-SC12 SCs (purple points represent comparisons between isolates in SC16). The discontinuity between these two sets of points indicated clonal structure in the population, as higher rates of recombination were predicted to generate a more homogenous distribution (Supplementary Fig. 2). Hence the co-circulating lineages that could be distinguished through their core genomes also maintained distinct accessory genomes. The set of green points represented comparisons between isolates in SC12 and those in other SCs, highlighting the divergence of SC12 from the rest of the population.

Potential speciation of atypical genotypes. Genetic loci unique to SC12 seemed likely to explain its distinctive phenotype and disease tropism. To define a set of candidate genes, ‘characteristic COGs’ (cCOGs) were identified in each SC as those COGs found in greater than 95% of genomes in that cluster, while being present in fewer than 5% of genomes in any other monophyletic SC. This identified 78 cCOGs in SC12 (Fig. 1b). Forty-four cCOGs were found within putative MGEs, and a further three were found within the conjugative element-related pneumococcal pathogenicity island 1 (PPI-1)8, although the pit iron transporter operon within this locus implicated in pathogenesis25 was absent from SC12 but present in all other SCs (Supplementary Fig. 3a).

Other non-MGE GIs contributing to SC12 divergence appeared to represent single gain or loss events conserved across the SC. All lacked a functional capsule polysaccharide synthesis (cps) locus and conserved a distinct set of large surface proteins (Supplementary Fig. 3b–e). The SC12 isolates also lacked either of the fucose utilization loci, one of which was evident in all other SCs (Supplementary Fig. 4); the conservation of these sequences across deep-branching clades suggested exchange at this locus was not rapid, and therefore a single deletion in an ancestor of SC12 would account for the observed pattern. Analogously, it seemed likely a single acquisition of genes encoding dihydroxyacetone kinases, in place of the pneumococcal histidine triad protein gene phpA (Supplementary Fig. 3f), would explain their
conservation in SC12 and absence from other isolates in the collection. Hence, the SC12 isolates appeared to be genetically, antigenically and metabolically distinct from the other SCs, and therefore may represent a separate species.

**Clonal association of GI diversity.** Each of the other SCs had a smaller number of cCOGs (Fig. 1c and Supplementary Table 1). In some cases, these corresponded to putative protein antigens or cps genes; for instance, all SC4 representatives expressed capsule type 22F, not found elsewhere in the sample19. The other cCOGs showed little similarity in sequence or putative function, although a substantial number were located in the 3' variable region of PPI-18. Although extensive diversity was observed at this locus across the species, there was little evidence of variation within SCs (Fig. 2), suggesting genes within this locus may underlie lineages' distinctive traits: at least some allelic variation has previously been associated with differences in virulence in a mouse model of disease26. Distinct loci within PPI-1, each ~20-kb long and encoding metabolic genes, were evident in SC1, SC5 and the serotype 3 isolates27. However, not all alleles were unique to a SC: SC2 and SC3 shared an ~10 kb gene cassette, a 3.8 kb allele was common to SC6, SC10, SC13 and SC15, and both complete and incomplete versions of a previously described lantibiotic synthesis gene cluster28 were found in the PPI-1 loci of SC8, SC9 and SC12. In SC12, these genes were accompanied by a putative RMS, which alone constituted the 3' variable region of PPI-1 in SC4 and SC11. The read mapping suggested SC4 also possessed the version of the island found in SC6, but in fact these genes were found on an ICE and appear to exemplify the contribution of MGEs to the diversity of sequences within PPI-18,28. MGEs themselves accounted for almost a quarter of the cCOGs not associated with PPI-1, suggesting such elements did not necessarily exhibit a high level of mobility, and instead may contribute to the stable differences between SCs.

**Diversification driven by MGEs.** The smallest putative mobile sequences previously characterized in pneumococci, three families of short interspersed repeats29, were generally stable in frequency within SCs (except for expansion and contraction of boxB tandem arrays), with small differences between lineages (Supplementary Fig. 5). All three families were evident in SC12 at typical frequencies, in contrast to the related species S. pseudopneumoniae30 and S. mitis31. Similarly, some types of insertion sequences (ISs) were ubiquitous across the sample, while others exhibited stable associations with particular lineages (Supplementary Fig. 6). Acquisition of novel ISs was observed within SC8: IS1202 was gained through serotype switching events twice, while ISSpn5 was imported as part of ICE 'scars'8. An extensive search for longer MGEs (see Supplementary Methods) identified 2,260 putative MGE-derived genetic loci, with a median length of 31 COGs (range 2–91 COGs). On the basis of their distribution around the chromosome, 16 insertion sites could be robustly identified within the core genome (Supplementary Fig. 7 and Supplementary Methods). As in Escherichia coli and Salmonella enterica, the majority of the insertion sites (15 of 16) were in intergenic regions despite the high coding densities of bacterial genomes32. However, in contrast to these enteric bacteria, all but two insertion sites were closer to the origin of

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**Figure 3 | Mobile genetic elements found in the pneumococcal population.** The 2,226 MGE sequences identified in the collection that could be classified as derived from a putative ICE, PRCI or prophage are each represented by a node, coloured according to the sequence cluster of the host in which it was found as displayed in the key. These are linked by vertices based on their similarity in terms of COGs using the Mountford index and classified using functional domains that appear characteristic of different MGE types. Clusters of nodes described in the text are annotated with letters. The grey boxes display the annotation of representative nodes, indicated by the dashed lines, from the main clusters in the network.
replication than the terminus. As the distance of genes from the replichore boundaries is conserved even more strongly than synteny in pneumococci33, this result should apply across the species, although sequence variation prevented the re-identification of three known insertion sites for large conjunctive elements that lie close to the terminus of replication (Supplementary Fig. 7).

A network was constructed in which each putative MGE was represented by a node, coloured according to the SC of the host cell, with vertices linking elements determined as being similar using Mountford’s index34. This allowed all but 34 putative MGEs to be classified into three groups based on the presence of functional domains (Fig. 3, Supplementary Figs 8,9; Supplementary Methods). The most numerous group (1,083 nodes) represented putative ICEs (Supplementary Table 2). These spanned the full range of detected MGE lengths, likely reflecting the efficiency of conjugation in transferring long segments of DNA between streptococci35, permitting modular variation through the insertion or deletion of sequence segments36. Hence, these elements are effective vectors for the import of novel DNA into a species. For instance, all antibiotic resistance genes encoded by MGEs were found on ICEs in the component labelled A and B. These consisted of sequences related to Tn5253, generated through the insertion of Tn916-type elements and other cassettes into Tn5252-type elements8,37 (Supplementary Figs 10,11). Conversely, ICEs in component C did not appear to carry ‘cargo’ genes, but did exhibit extensive similarity to Streptococcus suis MGE ICEsSu22457,38, which contained a cluster of resistance genes not evident in the pneumococcal elements (Supplementary Fig. 12). Hence, in other species, these elements can fulfill the role played by component A and B ICEs in pneumococci.

Component C was one of the six ICE network components predominately associated with SC12, and in this case appeared to represent a conserved GI distinguishing these isolates from the rest of the population. Such sharing of MGEs through recent common ancestry (that is, vertical transmission of the MGE) was indicated by these cliques of highly connected nodes within the same SC. Conservation of ICEs has been observed in multidrug-resistant lineages in which the ICEs encoded resistance genes37,39,40, but many examples identified here, such as those in component C or the Tn5252-type MGE similar to ICEsPuPN128 found in SC6 (Supplementary Fig. 11), lack such obviously beneficial cargo genes. In some cases, the importance of vertical transmission to the spread of some MGEs may reflect the absence of modules encoding the machinery needed for horizontal mobility. Examples were evident in component D (Supplementary Fig. 13), the longest representative of which appeared potentially mobile, whereas the shorter members may have lost some of the machinery for transfer between cells41.

The second group of MGEs, accounting for 471 nodes, likely represents phage-related chromosomal islands (PRCIs), mobilized in cis by ‘helper’ prophage42. First identified as ‘pathogenicity islands’ carrying superantigens in Staphylococcus aureus43, these pneumococcal examples encoded a high proportion of sequences for which no robust functional prediction could be made. Representatives from components E and F exhibited similarity with the Streptococcus pyogenes PRCI SpyC11 (Supplementary Figs 14,15) and were typically between 8 and 15 kb in length with putative integrase and regulatory genes transcribed in one direction and a DNA primase gene transcribed in the opposite direction. Representatives from component G were similar in size and genetic organization, with an integration module that showed limited similarity with the enterococcal PRCI E/CJIV58344 (Supplementary Fig. 16). The most unusual representatives were in component H, in which the putative integrase and primase genes were linked to a central, transposase-flanked portion that closely matched a GI from Streptococcus mutans L J23 (Supplementary Fig. 17). However, there was generally little evidence of the ICE-type modular evolution: PRCIs exhibited less variation in size (Supplementary Fig. 8), and the same core set of functions tended to be conserved between them. Sequence variation was instead mosaic in nature, with the level of sequence divergence between representatives changing at breakpoints that varied between elements, likely representing the consequence of homologous recombination.

Exhibiting a similar mosaic pattern of sequence variation were prophage, the third group of 672 nodes. These generally conserved a distinctive module order, and had a consistent orientation across the five insertion sites containing full-length prophage in which the genes active during MGE replication were aligned with the strong coding bias of the pneumococcal genome, akin to the ‘polarization’ seen in enteric bacteria45. In marked contrast with ICEs and PRCIs, few instances of prophage being stably associated with a lineage were recorded, as implied by the connectivity within component I (Supplementary Table 2). Notable exceptions to this trend were evident: isotypes of prophage oXCI141, independently observed to be stably associated with the serotype 3 genotype predominant in Massachusetts49,27, were identified in the expected hosts (Supplementary Fig. 18). These viruses were within component I, which encompassed the previously described diversity of pneumococcal phages4. Similarly conserved between isolated genomes were two atypical phage: one similar to Enterococcus faecalis V583-pp144 present in all but four members of SC4 (components J and K; Supplementary Fig. 19), and another similar to S. oralis prophage oPH10 present in five SC11 isolates (component L; Supplementary Fig. 20). Yet, the largest set of nodes that showed stable association with host SCs was found within component I; these represented a GI identified in the multidrug-resistant PMEN1 lineage46 that likely represented a ‘prophage remnant’ that has lost its mobilization machinery (Supplementary Fig. 18). Such degradation of an MGE can occur when selection acts to conserve a beneficial cargo gene47; the only candidate in this instance was a coding sequence (CDS) with a functional domain associated with RMSs, suggesting this gene may have been preserved to protect against other MGEs.

**Potential barriers to sequence exchange.** The apparently high rate of phage transmission suggested there would be strong selection for mechanisms that prevented infection with these viruses, which may also inhibit the exchange of other GIs. RMSs seemed likely to play such a role, and 11 candidate RMSs were identified using Pfam domains46 (Supplementary Tables 3 and 4). Three of these were present at the dpn locus, of which two were the previously characterized DpnI and DpnII systems3. The only example of switching between these two systems within a SC occurred on the long branch within SC12 (Fig. 4b). The one other change at the dpn locus within a SC involved replacement of a Type II RMS (designated DpnIII and represented by SPN23818640-18650 in the genome of S. pneumoniae ATCC 700669 (ref. 8)) present in all but one isolate of SC13, in which it had been replaced by DpnII. DpnIII likely targets a different motif to DpnI and DpnII, both sensitive to adenine methylation11, as functional domain information suggested the DpnIII RMS modified cytosine bases. As both DpnI and DpnII do not prevent the uptake of GIs by transformation2, but are likely to be similarly effective against MGEs found as double-stranded DNA forms in the cell, it was unsurprising to find that the accessory genome diversified at
approximately the same rate in isolates carrying either system (Supplementary Fig. 21). However, there was also little difference in the equivalent rate estimated from those isolates carrying DpnIII, which appeared to be a typical Type II restriction system. Furthermore, increasing numbers of non-Dpn accessory RMSs also did not appear to affect the rate of accessory genome diversification, despite these Type I and II systems being potentially able to cleave MGEs or any GIs imported by transformation (Supplementary Fig. 22). Despite their apparent lack of effect on the plasticity of genome content, these accessory RMSs exhibited a similar level of conservation across clades as the Dpn systems (Fig. 4b).

The absence of an observed effect may reflect the influence of other aspects of the transformation mechanism. One candidate was pherotype, which was also conserved across deep-branching clades. All 15 monophyletic SCs were uniformly associated with either CSP-1 or CSP-2, with no isolates having acquired the rarer pherotypes or switched between the more common types (Fig. 4a) despite requiring a change within the range of commonly observed transformation events37. One explanation is that interpherotype exchange of sequence is infrequent38. However, any inhibition of exchange between the pherotypes does not appear to substantially affect their relative rates of recombination. No significant difference was observed in the rate of diversification through homologous recombination relative to point mutation (Wilcoxon rank-sum test of previously calculated r/m values39, N = 15, W = 20, P = 0.46), and no substantial difference in the relative rate of accessory to core genome diversification could be identified between them (Supplementary Fig. 23).

Figure 4 | Mechanisms potentially affecting GI transfers. (a) Maximum likelihood phylogeny based on the core genome annotated according to the distribution of sequence clusters. The branches of the phylogeny are coloured according to a maximally parsimonious reconstruction of CSP pherotype. The ‘CSP-3-like’ sequence was identical to the previously described CSP-3 (ref. 16) but lacking an FNIFNF peptide. (b) Variation in accessory RMSs. The columns to the left indicate which of the three RMSs is present at the dpn locus by black bars in the appropriate rows. The eight columns to the right indicate the presence of other putative accessory RMSs, as inferred from the distribution of the relevant methylase COGs. Columns are labelled with the accession code of the sequence in Supplementary Table 4, with black bars again indicating the presence of an RMS in the corresponding isolate. (c) Variation in the ivr locus. The left columns show reads corresponding to the 5’ part of the full-length spnIVRhsdS gene assigned to the two alternative 5’ TRD-encoding sequences A or B. The heatmap indicates the proportion of reads corresponding to the spnIVRhsdS gene that matched each allele, with red indicating a higher proportion and blue a lower proportion. The right columns show reads likely corresponding to the 3’ part of spnIVRhsdS assigned to the three alternative TRD-encoding sequences a, b or c. (d) Variation in the tvr locus. Eleven different spnTVRhsdS TRD-encoding sequences were identified across the population. When the TRD-encoding sequence was present as part of a full-length CDS, the cell is coloured red, if the TRD was found in the 5’ half (these are labelled with uppercase Roman numerals), and orange, if found in the 3’ half (these are labelled with lowercase Roman numerals). Where the TRD-encoding sequence was present as a lone fragment, the corresponding cell in the grid is coloured blue. Empty cells indicate the TRD-encoding sequence was absent from the corresponding isolate.
Rapid RMS variation through DNA inversion. The lack of a detectable impact of either accessory RMSs or phenotype on the rate of genome content diversification indicated there may be an alternative mechanism inhibiting the spread of GIs. Two candidates were ‘core’ RMSs that were ubiquitous in the sampled population (Figs 4 and 5). Both of these were Type I RMS loci containing multiple sequences encoding different DNA-binding target recognition domains (TRDs) of specificity subunits together with a recombinease. One of these loci encoded TRDs on both strands of the genome (Fig. 5a), and had previously been demonstrated to undergo rearrangements through sequence inversion in S. pneumoniae TIGR4 (ref. 48). This phase variation resulted in five TRD-encoding sequences being combined into up to six different full-length genes, each encoding a putatively functional Type I RMS specificity subunit formed of two TRDs. This region was denoted the ‘inverting variable restriction’ locus (ivr locus), with the specificity subunit encoded by the spnIVRhsdS gene. The rapid variation in the composition of spnIVRhsdS across the sequenced collection (Fig. 4c) was hypothesized to be driven by intragenomic recombinations catalysed by the recombinease encoded by ivrR within the ivr locus48. Hence, ivrR was disrupted using an antibiotic resistance marker to stabilize three different versions of spnIVRhsdS generated by intragenomic recombination during routine culturing of S. pneumoniae R6 (ref. 49).

The mutants were characterized by SMRT sequencing, with de novo assemblies confirming that each had a different spnIVRhsdS allele (Supplementary Fig. 24). The mutant with the same spnIVRhsdS sequence as the R6 genome (composed of the TRDs denoted Aa) was found to have adenines methylated at the N6 position in three motifs. Two of these, TCGAG and TCTAGA (underlined adenines were methylated; Supplementary Table 5), likely represented the activity of two Type II RMSs. Twenty CDSs in the S. pneumoniae R6 genome matched the RMS-associated functional domains listed in Supplementary Table 3. The most likely candidates for causing these methylation patterns were SpnIM, an ‘orphan’ methyltransferase encoded by a CDS adjacent to an endonuclease pseudogene, predicted to target the TCTAGA motif49, and Spr1102, which appears to form a functional Type II RMS with Spn1103 (orthologous with the accessory RMS with accession code LK020705 in Fig. 4). The third motif, CAG(N)₆TTYG, was bipartite and likely to represent the activity of the Type I RMS encoded by the ivr locus. SMRT sequencing of the second mutant, in which the 3’ region of the
spnIVRhsdS gene had been switched by an inversion such that it was composed of TRDs Ab, found the same Type II RMS motifs and an altered Type I motif, GAA(N1)TYYG. The maintenance of the 5’ half of the spnIVRhsdS allele was consistent with the conservation of the TTYG component of the original motif. Correspondingly, SMRT sequencing of a third mutant with a Ba spnIVRhsdS allele, in which only the 5’ region of spnIVRhsdS differed from allele Aa, identified a bipartite methylated motif of CAG(N2)-GTG; this preserved the CAG component of the original motif, while nevertheless again altering the system’s overall specificity.

Rapid RMS variation through DNA translocation. In contrast with the ivr locus described above, the TRD-encoding sequences at the second ‘core’ RMS (the SP_0886-SP_0892 region of the S. pneumoniae TIGR4 genome) were all on the same strand (Fig. 5b). In many isolates, apparently functional specificity subunit genes were formed through the combination of two TRDs, as at the ivr locus. Alignments of the locus in closely related members of sequence type 3280 (ref. 19) suggested that ‘shuffling’ of TRDs occurred through lateral translocation of DNA (Supplementary Fig. 25); PCR amplification confirmed this was genuine variation and not an assembly artefact (Supplementary Fig. 26). This was unlikely to represent spontaneous, irreversible mutation because isolates in SC2 and SC3 apparently alternated between two different forms (Fig. 4d and Supplementary Fig. 27). Rather, the changes were likely catalysed by the putative recombinase, TvrR, encoded by this locus, henceforth termed the ‘translocating variable restriction’ (ivr) locus. These alterations would likely involve excision and re-integration of DNA; the putative TvrTA toxin–antitoxin system may select against failure to re-insert the gene cassette during rearrangements, as these systems are likely to be effective in stabilizing such dynamic genetic loci. To test whether variation in this locus could occur through intragenomic recombination, individual colonies from three different isolates, each of which had a different full-length spnTVRhsdS gene (Fig. 5b), were serially passaged in broth three times. A PCR was designed to amplify an ~3 kb product from the ‘native’ version of the locus, which could also detect rearrangements through the amplification of shorter products as the consequence of a primer binding site within a TRD-encoding sequence being shuffled closer to the 3’ edge of the locus (Fig. 5c). In the case of CH2060, only the ~3 kb product was clearly observed, suggesting any rearrangements were rare in this isolate; with BR1109, a shorter band became prominent over the time course, suggesting infrequent rearrangement; whereas the variant locus was easily detectable in ND6010 after a single night’s growth. None of these shorter bands were observed following the replacement of the 3’ end of the locus, including tvrR, with a kanamycin resistance marker (Supplementary Fig. 28). Again, the high rate of this mechanism was reflected by the extensive variation in spnTVRhsdS observed across the population (Fig. 4d). These data indicated that the ‘shuffling’ of spnTVRhsdS configurations was rapid, commonly occurring within SCs, whereas the horizontal acquisition of new spnTVRhsdS TRD-encoding sequences was much less frequent.

S. pneumoniae R6 lacks a full-length tvr locus specificity subunit gene (spnTVRhsdS), hence the absence of any corresponding signal from the previous SMRT sequencing data. To determine whether the system was active when a full-length spnTVRhsdS allele was present, the three tvr loci in which the 3’ end had been replaced with a kanamycin resistance marker were introduced into the S. pneumoniae R6 Aa mutant. When the tvr locus from isolate BR1109 was introduced, the same methylation motifs were detected as in the original Aa mutant, with no evidence of another RMS being active. This may be the consequence of a small truncation of the methylase subunit, or a low specificity, or efficiency, of methylation. However, introducing the loci from isolates CH2060 and ND6010 resulted in the detection of both the Type II RMS and ivr locus-associated motifs, along with an additional Type I methylation motif (Supplementary Table 5): GATA(N2)RTC in CH2060 and GAGA(N2)-TGA in ND6010. Hence, the tvr locus encodes an RMS with a specificity apparently determined by the sequence of spnTVRhsdS, a gene that can vary through intragenomic recombination.

Discussion

The observation of distinct co-circulating lineages, as defined by the core genome, is often assumed to mirror selectively important differences in gene content. While pneumococci belonging to the same lineage are more likely to share accessory genome loci, this is generally not the result of lineages maintaining large numbers of unique genes; instead, they are characterized by combinations of MGEs, individually common GIs. The SC12 isolates were an exception in this population. They appear to be adapted to a different, although likely overlapping, ecological niche and may merit recognition as a novel species.

Not all GIs were stable; different types demonstrated different dynamics across the population, as illustrated by Fig. 6. The black line traces the general decline in gene content similarity from focusing only on near-identical isolates to comparing the entire collection. This partly reflects the low rate at which GIs primarily depending on transformation for their mobility were exchanged, as exemplified by the conservation of PPI-1 alleles (Fig. 2), genes involved in sugar metabolism (Supplementary Fig. 4) and capsule type19 within SCs. Other lines trace the divergence assessed using only the subset of COGs associated with the three large MGE types (Supplementary Methods). These show that PRCIs and ICEs were stable within SCs, but diverged between them. Such conservation may reflect these MGEs providing an advantage to their host, although few examples of potentially beneficial cargo were identified. Nevertheless, these MGEs appear to be reliant on vertical transmission for their success, and therefore the clonal dissemination of ICE-associated antibiotic resistance37,39,40 can also be thought of not only as hosts maintaining selectively advantageous MGEs, but also as MGEs providing a benefit for their long-term host.

In contrast to such a ‘symbiotic’ long-term association between MGEs and hosts, prophage were much less stable (Fig. 6). The typically destructive transmission mechanism of phage imposes a strong selective cost on host cells, meaning isolates carrying prophage are likely to be selected out of the population comparatively quickly. This lowered vertical transmission through inheritance necessitates that phage have an elevated rate of horizontal movement between cells51. As RMSs are effective at inhibiting horizontal transmission, the benefit they confer on their host cell is likely to be high if cells are frequently being infected by phage, as in this population. Yet, this inhibition is contingent on the donor and potential recipient harbouring different RMSs. Hence, RMSs can be highly effective at preventing the spread of MGEs if diverse across the population; however, this did not appear to be the case for the accessory RMSs in this collection, as typified by the dpr locus only having two common alleles. This contrasts with the extensive variety of prophage from which variants may emerge able to evade such defences. Furthermore, the observed stability of RMSs on non-MGE GIs (Fig. 6) implied they were exchanged less frequently than phage. This makes it difficult to conceive how they might co-evolve at the same rate as phage, and also makes them unlikely to
be effective in preventing transmission between isolates of the same SC. Hence, the maintenance of these RMSs may partly reflect persistence as ‘selfish’ elements.

Unlike GI exchange by transformation, the speed of intragenomic recombination is not limited by the need to encounter a suitable donor cell, and hence it can facilitate adaptation over short timescales. Rapid intragenomic changes affecting RMSs have previously been described in other species, through phase variation produced by DNA inversion, variation in the length of tandem repeat arrays, and homopolymeric tracts. The rate

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**Figure 6 | Dynamics of the accessory genome across the population.** (a) Rate of change in different components of the accessory genome. The horizontal axis represents a threshold maximum cophenetic distance separating isolates, based on the core genome maximum likelihood phylogeny; the vertical axis represents the similarity observed in different aspects of the genome when considering all pairwise comparisons below this cophenetic distance threshold. The black line uses the median Jaccard similarity metric to trace the change in overall COG content between isolates. The blue and purple lines represent the median value of a similar metric calculated only using the subset of COGs characteristic of different MGE classes. Prophage-associated COG content (excluding the prophage remnant GI) diverged considerably within sequence clusters, indicating these MGEs are relatively transiently associated with pneumococcal hosts. By contrast, PRCI and ICE content (excluding the ICE ‘scars’) were stable within sequence clusters, but varied substantially between them. When considering the distribution of RMSs, each pairwise comparison was coded one where both isolates shared the same profile (as calculated from the data in Fig. 4), and therefore the system could not be effective in preventing an MGE transmission, and zero otherwise. In the case of the ivr locus, the TRDs most commonly predicted to form the spnIVRhsdS gene were used to calculate this metric; in the case of the tvr locus, the profile of all TRDs at this locus, including whether or not they were present in a full-length spnTVRhsdS gene, was used. The plotted lines show the proportion of pairwise comparisons in which isolates had identical profiles based on the same core genome cophenetic distance thresholds as for the similarities in terms of COGs. This found the dpn locus and other accessory RMS to be conserved over relatively long evolutionary timescales, whereas the ivr and tvr loci were divergent between even very closely related isolates. (b) The distribution of pairwise cophenetic distances, calculated from a maximum likelihood core genome phylogeny, represented as a histogram.
of diversification facilitated by the *ivr* and *tvr* loci means they are likely to be effective at preventing phage transmission in clonally related cell populations, as is evident from the high levels of diversity seen at these loci even between closely related isolates (Fig. 6). Consistent with such an anti-phage activity, of the three motifs associated with the *ivr* locus, the most frequent was found at a density of one site per 2.3 kb in the subset of COGs associated with prophage, with the least frequent was present at a density of one per 16 kb. The motifs associated with the *tvr* locus were found at densities of one per 3.8 kb and one per 5.4 kb in the same sequences. As complete prophage are typically over 30 kb in length, these systems should frequently be effective at preventing viral infection.

Hence, closely related pneumococci are most likely to be distinguished by shuffling of these variable RMSs and their complement of prophage. However, the reversible nature of the RMS alterations means that once an equilibrium level of diversity is reached through intragenomic recombination, it does not tend to increase (Fig. 6), with the exception of transfer of *RMSs*.

RMS alterations means that once an equilibrium level of diversity is reached through intragenomic recombination, it does not tend to increase (Fig. 6), with the exception of transfer of *RMSs*. If a Type I system). These have been submitted to the ENA with accession codes listed in Supplementary Table 7.

Detection and classification of MGEs. Short interspersed repeat sequences were detected using Hidden Markov models as described previously. Analysis of the distribution of ISs involved constructing a reference sequence from the ISs found to be associated with pneumococci in the ISFinder database and analysing the distribution of sequence by read mapping as for Gls. The identification and classification of multi-gene MGEs is described in Supplementary Methods and Supplementary Table 6. BLAST was used for alignment of translated nucleotide sequences; these were displayed using ACT. Accession codes for MGE assemblies are listed in Supplementary Table 7.

Detection of RMSs. Fifty-eight COGs were identified through searching the overall set for those containing one of the Pfam domains listed in Supplementary Table 3. Representatives of each example were then manually investigated using ACT, and functional loci selected on the criteria that they contain apparently full-length endonuclease and methylase genes (as well as a specificity subunit gene, if a Type I system). These have been submitted to the ENA with accession codes listed in Supplementary Table 4.

Ascertaining the arrangement of *ivr* loci in silico. For each isolate, the Illumina reads were mapped to the reverse complement of the region defined by coordinates 454708–456366 in the genome of *S. pneumoniae* R6 [EMBL accession code: AE007317].49, corresponding to the complete spr0449 *sprVRLhsdS* CDS and invariant 5′ region of the spr0448 *sprVRLhsdS* CDS. The unmapped pairs of those reads mapped in the correct orientation for providing information on the downstream region should correspond to the 5′ variable region of *sprVRLhsdS*. Consequently, the number of uniquely mapping reads with at least 95% similarity along their full length to each of the two alternative 5′ sequences (A and B in Fig. 5a) found in the locus were tallied, and their relative frequencies displayed as a heatmap in the first pair of columns in Fig. 4c. Seven isolates had ten or fewer reads matching the two *ivr* locus 5′ TRD sequences in total; *de novo* assemblies confirmed these isolates had complete, or partial, deletions of the *ivr* locus, and consequently all *sprVRLhsdS* TRD sequences were set as having zero coverage in these isolates.

The 5′ TRD-encoding sequence most frequently found immediately downstream of the *sprVRLhsdS* CDS was then used as the reference sequence for a second round of mapping, as appropriate for each isolate. In this case, the unmapped read pairs downstream of the mapped reads should correspond to the 3′ TRD-encoding sequence. The relative frequencies of these reads with at least 95% similarity along their whole length to each of the three 3′ TRD-encoding sequence were displayed as the three adjacent columns in Fig. 4c. Ten isolates had fewer than ten reads mapping to the 3′ TRD-encoding sequences. *De novo* assemblies confirmed these corresponded to the clade of seven isolates within SC12, which shared a deletion that eliminated all of the 3′ *ivr* locus TRD-encoding sequences, and a further three isolates across the collection that suffered from deletions or rearrangements at the locus that explained this lack of matches. All 3′ TRD-encoding sequences of *sprVRLhsdS* were set as having zero coverage in these isolates.

Ascertaining the arrangement of *tvr* loci in silico. A single COG (CL500804) corresponded to the majority of the *tvr* locus TRDs. Every member of this COG was scanned for the Pfam domain Methylyase _S_ (PF14020), found in single copy in *sprVRLhsdS* TRD-encoding sequences and in two copies in putatively functional *sprVRLhsdS* genes. The amino-acid sequences corresponding to this domain were extracted from each CDS, aligned using MUSCLE, and clustered using BAPS. This identified 11 different groups of sequences, which could then be classified as corresponding to 5′ or 3′ TRD-encoding sequences based on the order of domains within putatively functional CDSs. The distribution of these sequences across the population is shown in Fig. 4d.

Ascertaining *ivr* and *tvr* orientations by PCR. To confirm the orientations of the *ivr* and *tvr* loci by PCR, isolates were cultured overnight in THY broth (Todd–Hewitt broth containing 0.5% yeast extract), and their DNA extracted using DNEasy kits (Qiagen). In each reaction, 50 ng of genomic DNA was used as the template for PCR amplification with the specified primers (Supplementary Table 8) independently acquired a mutation at a single core locus with a probability of 0.1, or a single accessory locus, also with a probability of 0.1. In addition, with a Hobbitt brood (r), a given individual underwent recombination with a randomly selected donor. This involved each locus of the core and accessory genome being independently acquired from the donor with a fixed probability of 0.025 in this simulation (equating to an import of around 50 kb in a pneumococcus). The next generation of 1,000 sequences were then selected from the population at random with replacement, with each simulation run for 10,000 generations. A neighbour-joining tree was generated from the core loci using APEX, and the cohenetic distances plotted against the accessory genome dissimilarity (calculated as a jaccard distance) as in Fig. 1. Qualitatively similar results were obtained for different mutation rates and numbers of generations.
using the OneTag DNA polymerase and appropriate buffer (NEB). Product elongation was performed at 48°C for a time commensurate with the expected product size. Reactions were supplemented with 0.25 mU T4 DNA ligase (NEB) at room temperature for 1 h, or with BamHI (NEB) at 37°C for 1 h, as appropriate. The three digestion products were purified with a DNA Purification Kit (Qiagen) and mixed in equimolar proportions for ligation with T4 ligase (NEB) at 4°C for 24 h. Full-length ligation products were then amplified using primers R6hsdSL and Lint, which added an ApaI site onto the 3′ end of the agarose gel electrophoresis. The extract DNA was reamplified using primers LUpVL and RDwnVL, and then used to transform the primer pairs R6hsdSL and Lint, which added an ApaI site onto the 3′ end of the agarose gel electrophoresis. This construct was then reamplified with the same primer pair and used to transform thawed S. pneumoniae R6 cells in the presence of 10 ng of CSP-1 and 5 µl 500 µm calcium chloride. After 2 h incubation, cells were spread on blood agar plates supplemented with 5 mg/l erythromycin, and multiple colonies picked for screening using PCR to identify mutants with different spnTVRhsdS genes, resulting in the isolation of the three patterns found in isolates S. pneumoniae R6 Aa, Ab and Ba.

Construction of S. pneumoniae R6 tvrR knockouts. Three isolates with different putatively functional alleles of the spnTVRhsdS genes were identified from the collection19: CH2060, BR1109 and ND610. For each of these, the region upstream of tvrR in the tvr locus was amplified through PCR using the primers LUpVL and LDwnVL, and the region downstream of the tvr locus amplified using RUPLV and RDwnVL. These primers added an Apal site onto the 3′ end of the upstream product, and a BamHI site onto the 5′ end of the downstream product. An aph3′ gene was amplified from the Janus cassette20 using the primers kanL and kanR, which generated a DNA fragment containing the resistance marker flanked by the primer pairs R6hsdSL and Lint, which added an ApaI site onto the 3′ end of the agarose gel electrophoresis. This construct was then reamplified with the same primer pair and used to transform thawed S. pneumoniae R6 cells in the presence of 10 ng of CSP-1 and 5 µl 500 µm calcium chloride. After 2 h incubation, cells were spread on blood agar plates supplemented with 5 mg/l erythromycin, and multiple colonies picked for screening using PCR to identify mutants with different spnTVRhsdS genes, resulting in the isolation of the three patterns found in isolates S. pneumoniae R6 Aa, Ab and Ba.

SMRT sequencing of samples. Initially 2–4 µg genomic DNA was converted into ~20 kb fragments through hydrodynamic shearing using a MegaRuptor (Diagenode). The DNA fragments were separated on 2% agarose gels, excised, and resuspended in water. The extracted DNA was reamplified using primers LUpVL and RDwnVL, and then used to transform the parental isolate, using the appropriate CSP as determined from the genome sequence, and S. pneumoniae R6 Aa, using CSP-1, as described above.

Analysis of the data was conducted with using smrtanalysis version 2.1.0. De novo assembly, using PacBio data exclusively, was performed using the Hierarchical Genome Assembly Process, protocol RS_HGAP_Assembly2. Base modification and motif analysis was performed using protocol RS_Modification_and_Motif_Analysis1n. All data have been submitted to the ENA under the study accession code ERP005506.

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Author contributions

N.J.C., S.D.B. and W.P.H. conceived and designed the study. N.J.C., P.G.C., A.E.S. and W.P.H. performed the experiments. N.J.C. was supported by an AXA Foundation post-doctoral fellowship. This work was approved by all authors.

Additional information

Accession codes: The putative mobile genetic element and RMS sequences have been deposited in the European Nucleotide Archive under the accession codes LK020676 to LK020715. The SMRT sequence data have been deposited in the European Nucleotide Archive under the study code ERP005506.

Supplementary Information

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