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# 1 THE ECOLOGY AND EVOLUTION OF PANGENOMES

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12

## 13 **Abstract**

14 The pangenome is all the genes present in a species and can be subdivided into the accessory  
15 genome, present in only some of the genomes, and the core genome, present in all the  
16 genomes. Pangenomes arise due to gene gain by genomes from other species through  
17 horizontal gene transfer and differential gene loss among genomes. Our current view of  
18 pangenome variation is phenomenological and incomplete. We outline the mechanistic,  
19 ecological and evolutionary drivers of and barriers to horizontal gene transfer that are likely to  
20 structure pangenomes, highlighting the key role of conflict between the host chromosome(s)  
21 and the mobile genetic elements that mediate gene exchange. We identify shortcomings in  
22 our current models of pangenome evolution and suggest directions for future research to allow  
23 a more complete understanding of how and why pangenomes evolve.

24

## 25 **The pangenome concept**

26 The pangenome describes all the genes present in a species and can be subdivided into those  
27 shared by all members of a species—the core genes—and those present in only some  
28 members of a species—the accessory genes [1] (Figure 1). Although a pangenome can be  
29 defined for other taxonomic units (e.g., an ecotype or phylum), we focus here on the single  
30 species level since this is the most commonly used meaning. The pangenome concept  
31 emerged from early comparative studies of bacterial genomes. Comparison of a pathogenic

32 *Escherichia coli* O157 strain with its non-pathogenic relative *E. coli* K12, showed substantial  
33 gene gain in the O157 genome [2]. Shortly afterwards, a three-way comparison of these two  
34 genomes with that of another pathogenic *E. coli* genome, showed that less than 40% of protein  
35 coding sequences were shared between all three strains despite all being members of the *E.*  
36 *coli* species [3], which has proven to have an exceptionally broad pangenome. Even in these  
37 early pangenome studies it was evident that the variation among genomes within a species is  
38 often attributable to horizontal gene transfer (HGT) events. For instance, the difference  
39 between the *E. coli* strains K12 and O157 genomes is largely due to the acquisition of several  
40 large pathogenicity islands by O157 [2]. This variation is part of a wider pattern of variation in  
41 pathogenicity islands seen across *E. coli*, where differential distribution in these genomic  
42 regions is responsible for the classical nomenclature of *E. coli* pathotypes [4]. These range  
43 from chromosomally integrated pathogenicity islands and prophages to independently  
44 replicating plasmids. The advent of next-generation sequencing brought with it an acceleration  
45 in the generation of bacterial genome sequence data, revealing that the size of the pangenome  
46 varies widely among taxa. These studies reveal an overall negative relationship between  
47 pangenome size and the proportion of core genes: “open” pangenomes are larger in size,  
48 have a smaller proportion of core genes, and higher rates of gene gain by HGT, whereas  
49 “closed” pangenomes are smaller in size, have a larger proportion of core genes, and lower  
50 rates of gene gain by HGT (Figure 1) [5]. The concept of a pangenome in eukaryotes is  
51 debated [6, 7], but the available genomic data suggests that the concept is sound, although  
52 the extent of the accessory genome and the processes that drive the evolution of pangenome  
53 content are in many ways different in eukaryotes compared to prokaryotes (Box 1).

54 The current challenge is to move beyond this phenomenological description of pangenomes  
55 to forge an understanding of the mechanisms and processes that determine their structure. A  
56 genome sequence is a snapshot of a strain in time. Some of the genes and mutations in that  
57 snapshot share a long history and are destined to remain associated, while other members  
58 are transient: recent acquisitions in the process of leaving. How do we distinguish between

59 these categories? If a genome is a family photograph, how do we distinguish family members  
60 from the photobombers? A starting point is to understand the processes and mechanisms that  
61 promote or prevent gene gain and loss, and thereby shape the content of the pangenome.  
62 Gene gain by a lineage in the context of the pangenome can be conceptually separated into  
63 two distinct processes, operating on different timescales and affected by different  
64 environmental drivers. The first describes the specific gene acquisition event, which occurs at  
65 the level of individual cells and is effectively instantaneous, while the second represents the  
66 stable assimilation of acquired genes within populations or their non-random elimination from  
67 a lineage, and is on-going, with effects emerging over a longer period and in different ways in  
68 different environments. In this review, we first outline the molecular, ecological and  
69 evolutionary drivers of gene gain and loss which mediate changes in the composition of the  
70 pangenome, and then discuss how evolutionary theory can be applied to understand the  
71 structure of pangenomes.

72

### 73 **Drivers and barriers of gene gain and loss**

74 Gene acquisition introduces variation, and thus provides the raw material upon which selection  
75 can subsequently act [8]. Various mechanisms actively facilitate the movement of genetic  
76 material across membranes. These are particularly well-described in prokaryotes but there is  
77 evidence that equivalent mechanisms may exist in model eukaryotes such as yeast (see Box  
78 1). In recent decades, the canonical processes — conjugation, transduction, and  
79 transformation — have been joined by additional phenomena, including nanotubes [9] and  
80 vesicles [10] that can facilitate nucleotide exchange. These varied mechanisms of gene  
81 exchange offer the potential for gene acquisition, but the likelihood of its occurrence depends  
82 on a range of ecological, mechanistic and evolutionary factors, explored in this section  
83 (summarised in Figure 2).

84

85 *Ecological opportunity for HGT*

86 The proximal environmental triggers activating expression of gene exchange machinery vary  
87 between systems and with different species, but some common themes can be identified. One  
88 of these is stress. For example, the SOS response to DNA damage, triggered by some  
89 antibiotics, reactive oxygen, and UV radiation, activates transfer of the *Vibrio cholerae* STX  
90 element [11], causes integron rearrangement [12], and activates integrated bacteriophage  
91 [13]. Transposons in *E. coli* become active under nutritional stress [14], plasmid conjugation  
92 rates are increased in response to host inflammation in mammalian gut [15], and starvation  
93 conditions activate natural competence [16]. However, different stress responses can have  
94 divergent effects in different species [17], and donors, recipients, and mobile genetic elements  
95 may each have their own cues. For example, some mobile genetic elements, such as the  
96 pheromone-inducible conjugative plasmids of *Enterococcus*, have evolved mechanisms to  
97 detect the presence of recipients [18], and transformation is induced by quorum sensing and  
98 by specific nutrients in some species of *Vibrio* [19].

99 Ecology appears to be a principal determinant of gene-sharing [20], suggesting that the  
100 transfer of genes is to some extent limited by ecological opportunity and occupancy of shared  
101 habitats. Several gene transfer mechanisms including conjugation and nanotubes require  
102 close physical proximity and thus HGT is probabilistically likely to be most efficient between  
103 immediate neighbours [21]. Consequently, the size of the gene pool from which a species can  
104 draw will be dependent on the diversity of environments they occupy as well as the community  
105 diversity these contain. Correspondingly, networks of gene sharing have shown that co-  
106 occurrence of species in a habitat increases the probability of gene sharing [22-25]. Niche  
107 specialists likely to exist in stable environments with very low diversity, such as endosymbionts  
108 [24], have more closed pan-genomes than those that exist in diverse communities and more  
109 variable environments.

110 Among symbionts and pathogens with low rates of gene gain through HGT, variation in gene  
111 loss among lineages can be the primary cause of diversity among clonal lineages, and can  
112 lead to large phenotypic differences [26]. Whereas gene loss can be positively selected in

113 large populations with efficient selection, in intracellular symbionts and pathogens with low  
114 effective population size, gene loss is more likely to be a result of relaxed selection and drift  
115 [27]. How the balance of gene gain and loss contributes to the formation of a pangenome is  
116 well-illustrated by *Yersinia enterocolitica*. The species is composed of five phylogenetically  
117 distinct groups, four of which are pathogenic to humans and have emerged from a non-  
118 pathogenic ancestor, driven by a single acquisition of a large virulence plasmid [28]. Following  
119 plasmid acquisition, the splits between the four pathogenic groups are delineated at a  
120 pangenome level by differential losses of genes present in the ancestor, alongside HGTs  
121 leading to switches in serotype [29].

122

### 123 *Mechanistic drivers and barriers of HGT*

124 Once acquired there are significant barriers to the maintenance of novel genetic material which  
125 shape the patterns of gene sharing among species. Newly acquired DNA must replicate to  
126 ensure it is passed to daughter cells, either by carrying with it replication machinery compatible  
127 with that of the host (in the case of plasmids) or by integrating into a resident replicon (e.g. a  
128 chromosome or already-present plasmid). Integration can occur through general recipient-  
129 encoded processes such as homologous recombination which is dependent on regions of  
130 sequence homology flanking the heterologous gene [30, 31] or by the activity of entities such  
131 as transposons, integrons, and insertion sequences, which can facilitate capture of incoming  
132 DNA (e.g., [32]).

133 Genes must also be transferable and able to function in the host in order to have a phenotypic  
134 effect visible to selection [33], which is dependent on recognition of promoters allowing for  
135 gene expression [34], and comparable GC content, codon usage and compatible genetic  
136 codes allowing for efficient translation [35], and in the case of DNA transfer between eukaryotic  
137 genomes effective splicing of introns. Newly acquired genes evolve faster than older genes in  
138 the same genome, potentially because of adaptation to their new genomic context [36, 37]. As  
139 a general principle, many of these processes become more challenging across larger genetic

140 distances [38]. Correspondingly gene sharing has been shown to be most common between  
141 closer phylogenetic relatives [25], which enhances both the likelihood of the transfer event and  
142 the compatibility of genes between donor and recipient.

143 Mechanistic limitations are also likely to define the types of genes that are more readily shared,  
144 and therefore more likely to contribute to the accessory genome. Incoming DNA can disrupt  
145 cellular processes leading to severe fitness costs, and these genes are likely to be rapidly lost  
146 from the population by purifying selection. Genes encoding core cellular functions, such as  
147 those associated with transcription and translation, can be highly toxic when expressed in  
148 foreign hosts [34, 39] and are poorly represented among horizontally transferred genes [40,  
149 41]. This strong incompatibility may be due to disruption of or failure to maintain the large  
150 number of protein-protein interactions that the protein must engage in to properly function.  
151 Genes embedded within more complex interaction networks are therefore more disruptive and  
152 less likely to maintain the necessary functional interaction network when transferred, a  
153 phenomenon termed the complexity hypothesis [42, 43]. Mobile genetic elements (MGEs)  
154 themselves are often associated with significant fitness costs that are caused by a range of  
155 factors, including the biosynthetic cost of maintaining and expressing additional DNA, toxic  
156 gene products, and epistasis between chromosomal and MGE-encoded genes [44]. This  
157 disruptive effect of HGT is not surprising from an evolutionary perspective: HGT brings  
158 together genes that have different evolutionary histories, and there is no a priori reason to  
159 expect that these genes should function together harmoniously [45].

160

### 161 *Evolutionary conflict and collaboration in the pangenome*

162 Many of the mechanisms for horizontal gene transfer are encoded by infectious MGEs such  
163 as viruses, plasmids, and transposable elements. Therefore, pangenomes are composites of  
164 the host chromosome(s) together with MGEs that may be shared with other species. MGEs  
165 encode accessory genes that may represent adaptive additions to the pangenome (e.g. by  
166 providing a new ecological function or access to an otherwise inaccessible niche), but also

167 encode genes for selfish MGE-directed functions such as replication and transmission, as well  
168 as many genes of unknown function. As semi-autonomous evolving entities we should expect  
169 MGEs to maximise their own fitness through both vertical and horizontal transmission [46].  
170 Encoding beneficial accessory genes can increase MGE fitness through enhanced vertical  
171 transmission as positive selection drives clonal expansion [47]. However, being beneficial is  
172 not necessary for MGE success. Many environmental plasmids do not encode any obvious  
173 accessory genes [48] and are therefore likely to be genetic parasites. Experimental studies  
174 show that high rates of horizontal transmission through conjugation can maintain costly  
175 resistance plasmids in the absence of positive selection [47, 49, 50], and non-beneficial  
176 plasmids can invade biofilm populations [51, 52]. Indeed, experiments with antibiotic  
177 resistance and mercury detoxification plasmids have shown that positive selection for these  
178 functions can limit their horizontal transfer by reducing the availability of recipient cells [47,  
179 53]. Although, in the long run, purely infectious elements would be expected to become  
180 increasingly efficient parasites by shedding their accessory genes, mobile genetic elements  
181 that persist through horizontal transmission are likely to be especially prone to mediating gene  
182 exchange [54]. Higher rates of horizontal transmission expose these MGEs to a wider diversity  
183 of genomic environments, offering greater opportunity for other MGEs (e.g., transposons) to  
184 integrate and hitch a ride. This inherent nestedness of pangenomes means that potentially  
185 conflicting selective pressures may operate at different levels of complexity (e.g., at the level  
186 of the gene, MGE, genome, population, and species etc.).

187 The predominance of gene exchange mediated by MGEs means that this form of gene sharing  
188 is, at least partially, constrained by MGE host range. Phages are believed to have relatively  
189 narrow host ranges, which are often limited to within a species or genus [55, 56]. Plasmid  
190 host ranges can be broader, and are dependent on the diversity of replication genes required  
191 for stable maintenance in different host taxa [57]. Correspondingly, plasmids appear to be  
192 more important mediators of gene exchange across larger genetic distances [58]. However,  
193 interactions between MGEs allow smaller, simpler elements to escape these restrictions.

194 Transposons for example, which are themselves unable to transfer between cells, can hitch a  
195 ride on a conjugative plasmid, as has been observed for plasmid-encoded antibiotic  
196 resistances in hospital outbreaks of Enterobacteriaceae [59, 60]. Further transfer of  
197 transposons between plasmids with different host ranges then expands the range of potential  
198 hosts accessible to these transposon-encoded genes. Plasmids too can be composite  
199 mosaics of other elements, including other plasmids, broadening the range of hosts in which  
200 they can replicate, while transposons can become nested within one another, increasing  
201 opportunities for spread [61, 62]. A consequence of the self-interested activity of MGEs for  
202 genome evolution is that selfish genes encoding MGE-related functions spread between  
203 lineages alongside the MGE-encoded accessory functions that enhance host fitness or niche  
204 adaptation. Indeed, plasmid, phage, and transposon-encoded functions are usually highly  
205 represented in the pangenome and in comparative studies of horizontal gene transfer [5, 63].  
206 Because they can replicate by both vertical and horizontal transmission, MGEs can have  
207 fitness interests that do not necessarily align with those of other parts of the (vertically-  
208 inherited) genome. These 'divided loyalties' manifest in the fitness costs associated with MGE  
209 acquisition and horizontal transmission, and result in intragenomic conflict. For example, while  
210 conjugation provides an efficient mechanism for plasmids to transfer between bacteria, the  
211 expression of conjugative machinery imposes a biosynthetic fitness cost on the donor cell [64],  
212 and leaves the donor cell open to predation by pilus-targeting phage [65]. Resolution of host-  
213 MGE conflict frequently requires compensatory mutation(s) to the MGE or the chromosome to  
214 reduce the fitness costs of the newly acquired genes [46], which is promoted by positive  
215 selection for MGE-encoded functions since this increases the population size and mutation  
216 supply for MGE-carriers [66, 67]. Diverse compensatory mechanisms have been identified to  
217 stabilise plasmids, but two common routes are mutations affecting host gene regulatory  
218 networks [68, 69] or plasmid replication [45, 70]. By stabilising MGEs within bacterial lineages,  
219 compensatory evolution can set the stage for more extensive coevolution between the MGE  
220 and chromosome, driving reciprocal adaptations and counter-adaptations [46]. For example,

221 bacteria-plasmid coevolution rapidly led to the emergence of co-dependence of chromosomal  
222 and plasmid replicons under antibiotic selection, together providing high-level resistance but  
223 separately providing inadequate levels of resistance to persist in the environment they evolved  
224 in [71, 72]. Compensation and coevolution can, in turn, drive the complete domestication of  
225 MGEs and their integration into a more exclusively vertical mode of replication. In practice,  
226 domestication involves downregulation, inactivation, or loss of the machinery involved in  
227 horizontal transmission [73, 74]. For example, bacterial genomes contain numerous  
228 prophages, some of which are incapable of horizontal transmission and now serve their  
229 bacterial hosts as anti-competitor toxins [75]. Alternatively, recombination can relocate mobile  
230 genes to less-mobile parts of the genome, e.g. chromosomal capture of resistance genes from  
231 plasmids, a process rapid enough to be readily observable in the laboratory [50, 69, 76]. In so  
232 doing, the signatures of gene acquisition are gradually lost from the genome sequence,  
233 potentially explaining why many accessory genes originally transferred by an MGE are no  
234 longer obviously associated with MGEs.

235

### 236 *Resisting HGT*

237 Due to the potential for conflict between MGEs and the host chromosome, immunity systems  
238 which actively target incoming foreign DNA are widespread across eukaryotes and  
239 prokaryotes. Systems exist in both eukaryotes (e.g. RNAi [77]) and prokaryotes (e.g. H-NS  
240 [78]) to silence gene expression from foreign DNA. In prokaryotes CRISPR-Cas systems and  
241 restriction-modification (R-M) systems target novel DNA for degradation, and can be an  
242 effective defence against MGEs, potentially reducing HGT [79, 80]. A comparative analysis of  
243 79 prokaryote genomes show that R-M systems structure gene sharing by favouring  
244 exchanges between genomes with similar R-M systems [81]. The relationship between HGT  
245 and CRISPR-Cas systems appears more complex: There are well-described cases where  
246 CRISPR-Cas systems are negatively associated with MGE carriage within a species [82], but  
247 CRISPR-Cas can also promote HGT in some cases [83]. Type-III CRISPR-Cas systems target

248 actively transcribed DNA via spacers derived from RNA transcripts [84] and may therefore be  
249 more effective against phages and plasmids than DNA acquired by transformation [85]. Over  
250 broader taxonomic scales, however, the correlation between CRISPR-Cas systems and the  
251 rate of HGT is less clear and deserves further study [86, 87]. It is likely that additional  
252 mechanisms for resisting gene acquisition will continue to be discovered [88]. Resistance  
253 mechanisms protecting cells against incoming DNA can also be encoded by MGEs  
254 themselves, highlighting how conflict between MGE could act to limit HGT. Both plasmids and  
255 phages defend their host cells against super-infection through self-exclusion mechanisms [89,  
256 90] and can encode their own CRISPR-Cas systems with spacer sequences targeting other  
257 MGEs [91].

258

### 259 **How and why do pangenomes evolve?**

260 The next step is to synthesise these varied drivers of gene gain and loss into a general theory  
261 of pangenome evolution to answer the question: what structures the pangenome? On the one  
262 hand, it is conceivable that the pangenome is dominated by adaptive gene gain and loss, such  
263 that the pangenome is effectively a record of the responses to the myriad selection pressures  
264 that a species faces. At the other extreme, it is possible that the pangenome exists because  
265 selection is unable to prevent the spread of mildly deleterious gene acquisitions and deletions,  
266 and/or that these occur primarily due to the self-interest of MGEs. The key to distinguishing  
267 between these competing models of the pangenome is to disentangle how gene acquisition  
268 and loss, genetic drift, population subdivision and selection interact to shape the pangenome.

269

### 270 *Population genetic approaches to analysing the pangenome*

271 Evolutionary biologists have developed a mature body of population genetic theory to  
272 understand how mutation, selection and genetic drift interact to shape patterns of genetic  
273 variation [92]. A key insight from population genetic theory is that effective population size

274 ( $N_e$ ) shapes patterns of molecular evolution by modulating the efficacy of natural selection  
275 relative to genetic drift [93]. In species with a low  $N_e$ , selection is weak relative to the genetic  
276 drift and evolution is dominated by the stochastic spread of weakly deleterious mutations. In  
277 contrast, selection prevents the spread of weakly deleterious mutations and drives selective  
278 sweeps of beneficial mutations in species with high  $N_e$ . Like spontaneous mutation, both gene  
279 acquisition [38, 44, 94, 95] and loss [96-98] tend to reduce fitness. Therefore, selection should  
280 shape patterns of gene gain and loss in species with high  $N_e$ , whereas the composition of the  
281 pangenome in species with low  $N_e$  will be shaped by underlying rates of gene gain and loss.

282 Genome size increases with  $N_e$  across a wide range of bacteria [99, 100], and this correlation  
283 provides a good starting point for applying population genetic approaches to understand the  
284 pangenome. In part, this correlation is driven by the inability of natural selection to prevent the  
285 spread of weakly deleterious mutations in species with low  $N_e$  [101], such as endosymbiotic  
286 bacteria [102] and intracellular pathogens [103]. Many genes in bacterial genomes only  
287 provide a fitness benefit under very specific environmental conditions [96], and effective  
288 selection for marginally beneficial genes acquired by HGT in species with high  $N_e$  is also likely  
289 to contribute to the positive correlation between  $N_e$  and genome size. Simply put, because  
290 species with large  $N_e$  are likely to occupy wider environment profiles, they are also likely to be  
291 under a wider diversity of environmental conditions driving selection for gene diversity and  
292 therefore larger genome sizes (Figure 1). As such species with high  $N_e$  also have large  
293 pangenomes [5, 100], and McInerney et al. [5] argue that this correlation is evidence that the  
294 pangenome is adaptive. The concept of population structure is key to this argument: in species  
295 with low levels of population structure, adaptive gene acquisition and loss events will sweep  
296 to fixation, and these will therefore not contribute to the pangenome. Population subdivision  
297 provides the opportunity for selection to contribute to increasing the pangenome size of a  
298 species because selective sweeps of locally adaptive gene gain and loss events will affect the  
299 accessory gene complement and thus pangenome size [104]. The point at which ecologically

300 and genetically distinct subpopulations (or ecotypes) become sufficiently diverged to be  
301 considered multiple, different species each with their own pangenome is contentious [33, 105].  
302 Other studies using population genetics have questioned the role of selection in shaping the  
303 pangenome. Comparing levels of synonymous nucleotide diversity, a surrogate measure of  
304  $N_e$ , with a measure pangenome fluidity showed a positive correlation between  $N_e$  and  
305 pangenome fluidity, that could arise because genetic drift leads to the loss of effectively neutral  
306 accessory genes in species with low  $N_e$  [106]. Further support for this idea comes from  
307 comparing the observed distribution of gene frequencies in the pangenome with an expected  
308 distribution generated by a neutral model. This approach, inspired by the infinite alleles model,  
309 assumes that bacteria gain genes from an infinite pool of horizontally transferred genes and  
310 subsequently lose these genes through drift [107, 108]. Accessory genes show a distribution  
311 that is close to the expectations of a neutral model for widely distributed marine bacteria, but  
312 with deviations that are consistent with selection shaping the pangenome [108]. It is unclear,  
313 however, that currently available genomic data provide the necessary breadth and depth of  
314 ecological sampling to adequately test these models.

315

### 316 *The limits of a population genetic approach*

317 Population genetics theory provides some simple guiding principles for understanding the  
318 pangenome, but there are also potential difficulties with applying these models to understand  
319 the pangenome [109]. For example, classical population genetic tests for selection rely on  
320 comparing observed patterns of genetic polymorphisms and divergence with expected  
321 patterns from a neutral model where evolution is driven by mutation and drift, but not selection.  
322 Neutral models in population genetics assume that mutations at different sites in the genome  
323 are not linked. This is a justifiable assumption in eukaryotic species with obligate sexual  
324 reproduction, but the pangenome changes through the gain and loss of blocks of genes, for  
325 example because they are all encoded on a MGE. An important consequence of this is that  
326 strong selection for one gene (e.g. an antibiotic resistance gene) can lead to the spread of

327 linked mildly deleterious genes by co-selection, if there is a net fitness benefit of the MGE.  
328 Similarly, genes that are linked to addiction systems, such as toxin-antitoxin systems, can be  
329 maintained in populations by the toxic effects of MGE loss. In a broader perspective, the strong  
330 linkage disequilibrium observed in clonal bacterial species means that there might be no  
331 effectively neutral variation [109].

332 A second important difficulty is that population genetic models ignore the evolutionary conflicts  
333 of interest that can occur between MGE-encoded accessory genes and chromosomal core  
334 genes in the same genome where selection at the MGE and chromosomal levels are not  
335 aligned. A key concept from evolutionary ecology is that trade-offs exist between the efficacy  
336 of vertical and horizontal transmission [110], preventing the evolution of elements that are to  
337 provide a big benefit to their host and transfer efficiently between hosts. Trade-offs may also  
338 limit the ability of MGEs to maximize the fitness benefit that they provide to different hosts,  
339 further limiting the benefits that hosts gain from acquiring MGEs [72]. All else being equal, we  
340 would therefore expect that MGEs with high mobility, such as broad-host range conjugative  
341 plasmids and lysogenic phage, to impose greater fitness costs than genetic elements with a  
342 low mobility, such as non-transmissible plasmids and defective prophage. This logic is  
343 somewhat counter-intuitive, because many of the pangenome accessory genes with the  
344 clearest ecological functions, such as antibiotic resistance genes, are often found on MGEs  
345 with high mobility [111-113]. These potentially adaptive genes may be rare 'rubies in the  
346 rubbish' from the perspective of their bacterial hosts [8], with the rest of the linked genes being  
347 either merely useless or else functioning solely to promote their own replication and  
348 transmission at the host's expense.

349

## 350 **Perspective**

351 Short-read sequencing technologies have produced a rapid accumulation of sequence data,  
352 revealing the ubiquity and extent of pangenomes, especially in prokaryotes. At present,  
353 however, we lack a unified theory to understand the forces structuring pangenomes, and this

354 will probably require the development of new theory that links together concepts from  
355 evolutionary ecology and population genetics. To achieve this, there are some important  
356 obstacles that need to be overcome:

357 • Defining the concept of pangenome adaptation: Adaptation is the "process of optimisation  
358 of the phenotype under the action of natural selection" [114]. As a pangenome emerges  
359 as an analytical result from comparing multiple genomes, we must take care when  
360 specifying what adaptation means in this context, i.e. who or what is being optimised. While  
361 a pangenome *can* contain adaptive genes that are transferred between species, the  
362 pangenome does not evolve *for the purposes of* maintaining a pool of niche-adaptive  
363 genes. Instead, its contents are defined by selection occurring at lower organisational  
364 levels: the individual bacterial lineage that has acquired locally-beneficial genes, and the  
365 persistent MGE. Neither does a broadly adaptive pangenome imply that the accessory  
366 genes in a given genome are beneficial to that strain. Recent migration or gene acquisition  
367 can result in a strain carrying neutral or deleterious genes which have not yet been lost  
368 [115]. Finally, if the pangenome is defined as the sum-total of all genes in a species,  
369 improved sequencing resolution will increasingly capture transient events which are  
370 unlikely to be adaptive, inflating the size of the pangenome but diluting the signal of  
371 adaptation. Enhanced biological insight into the gene function, as well as bioinformatic  
372 tools that help us distinguish between transient associations and longer-term partnerships,  
373 will guard us from incorrectly inferring adaptation in such instances.

374 • Measuring the rates of HGT in nature: The rate of horizontal gene transfer is key to both  
375 the population genetic and eco-evolutionary perspectives on the pangenome, but our  
376 knowledge of rate of HGT in the wild remains very limited. It might be possible to measure  
377 these rate by using statistical methods to infer rates of HGT from genomic data, and  
378 experimental methods that allow the spread of genes to be measured under natural  
379 communities in real time using for example microcosm experiments [54, 116].

380 • Sampling genomes at ecologically-relevant scales: Microbial genomes are being  
381 sequenced at an incredible rate, but it is very challenging to understand sequence data in  
382 a population genetics context, there are often huge sampling biases in microbial sequence  
383 datasets (intensive sampling of clinical outbreaks is the most extreme example). Given the  
384 vast population size of microbes, we will only ever be able to achieve very sparse sampling  
385 of microbial genomes, even with the most ambitious sequencing projects. We therefore  
386 need to develop approaches to identify and sample ecologically coherent microbial  
387 populations [113] or ecotypes [33]. For example, it is clear that some microbial populations  
388 are structured at an incredibly fine scale, such as individual particles of detritus [117], and  
389 this structuring can play a key role in the evolution of the pangenome [104]. Comparing a  
390 small number of bacterial genomes sampled from many niches is likely to produce an  
391 abundance of rare accessory genes, but these could either represent adaptive accessory  
392 genes that are locally abundant but globally rare, or deleterious accessory genes that are  
393 both locally and globally rare. One key technological development that may help with this  
394 problem is to move from sequencing the genomes of bacterial isolates to single-cell  
395 sequencing of bacteria from environmental samples.

396 • Developing eco-evolutionary models of pangenome evolution: The neutral theory of  
397 molecular evolution has been so useful in revealing the action of natural selection because  
398 it makes quantitative and falsifiable predictions that be tested by comparing datasets.  
399 Given the complexity of forces shaping the pangenome it may be necessary to look outside  
400 genetics for potential approaches: Pangenomes share many characteristics with  
401 metacommunities, most notably the idea that entities (genes or species) are sampled from  
402 a pool to form discrete sets (genomes or communities) that share biological cohesiveness  
403 (pangenome or metacommunity). Metacommunity ecology has a well-developed body of  
404 theory to understand how communities are assembled and structured [118], which may  
405 help to unravel the processes causing the structure of pangenomes.

406



408 **BOX 1: Do eukaryotes have pangenomes?** The existence of pangenomes in eukaryotes is  
409 debated [6, 7]. What is evident is that, unlike the situation in prokaryotes, genome evolution in  
410 eukaryotes is dominated by processes other than HGT, including sexual recombination and  
411 gene duplication [119] often combined with domain reshuffling [120]. Nevertheless, HGT can  
412 and does occur: for example, *Saccharomyces* undergoes transformation under starvation  
413 conditions [121] and can receive DNA by conjugation from bacteria [122], although HGT from  
414 prokaryotes contributes less than 0.5% of the gene repertoire of *Saccharomyces* (reviewed in  
415 [123]). Additionally, a range of other mechanisms introduce genetic material into eukaryotic  
416 cytoplasm offering the potential for HGT, including: viral vectors [124], integration of viral  
417 fragments [125], RNA exchange [126], trophic interactions through phagocytosis of prey cells  
418 [127], and anastomosis of cell structures [123, 128]. The role of HGT in accessory genome  
419 variation is unclear, but likely to be less important than in prokaryotes and a relatively minor  
420 contributor compared to other factors like strain level duplication [129] and differential gene  
421 loss. Pangenome studies in eukaryotes are challenging due to their more complex genome  
422 architectures and a lack of replete genome-level sampling. Analyses of model fungi suggest  
423 core genome fractions of between 80-90% [129], whilst in the marine alga *Emiliania huxleyi*,  
424 17% of genes present in the assembled genome of the model strain CCMP1516 were  
425 absent in four other strains, indicating a putative accessory genome [130]. Consistent with  
426 the complexity of eukaryotic genome architecture, distinct dispensable or supernumerary  
427 chromosomes systems are observed in some fungi that show signs of HGT derivation,  
428 operate to carry an accessory genome, and define the niche and host range of the recipient  
429 lineage [131-133]. Therefore, while the existing studies suggest that the pangenome  
430 concept is well-founded for eukaryotic microbes, the extent of accessory genome variation  
431 is likely to be far lower than in prokaryotes: ~10-15% of genes in eukaryotes compared to  
432 up to ~65% in some prokaryotes.

433

434 **Figure 1: The pangenome concept.** Pangenomes vary extensively in size and the  
435 proportion of core versus accessory gene content. It is likely that species with large, open  
436 pangenomes occupy more varied niches and more complex communities, and have larger  
437 effective population sizes compared to species with smaller pangenomes.

438

439 **Figure 2: The drivers and barriers of horizontal gene transfer.** Horizontal gene transfer  
440 is likely to be affected by a wide range of ecological, evolutionary and mechanistic factors,  
441 which will in turn determine the degree of pangenome fluidity observed in a species.

442

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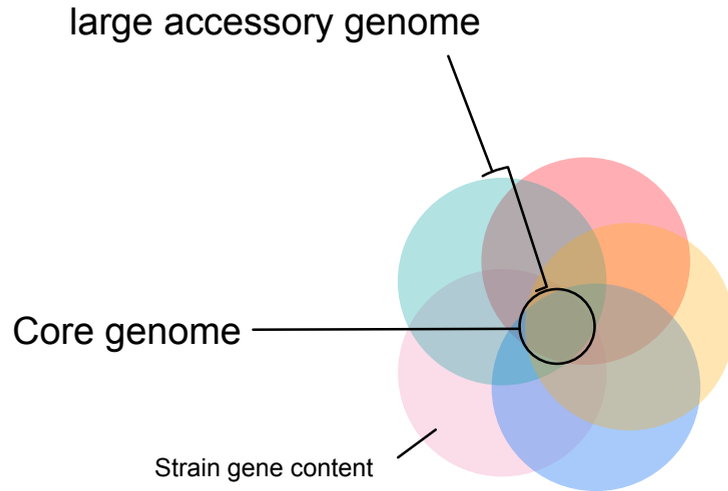
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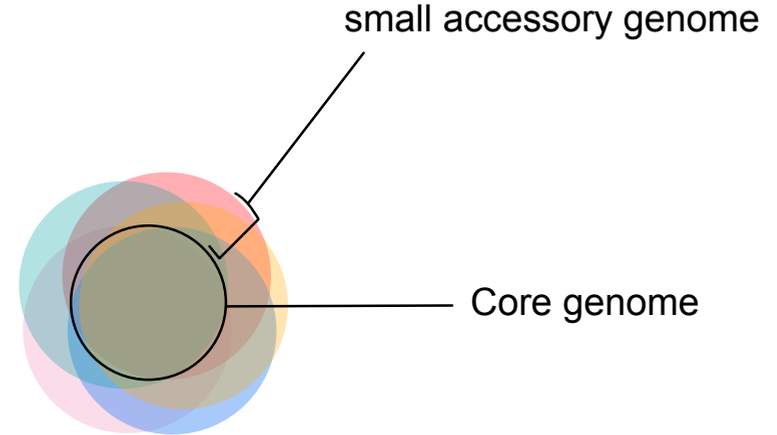
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## Open pangenomes



## Closed pangenomes

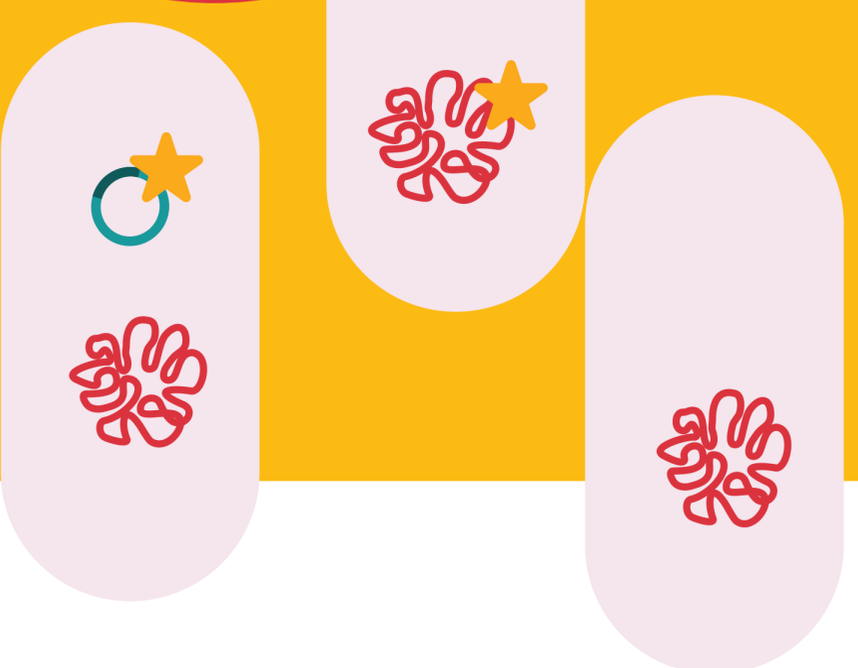
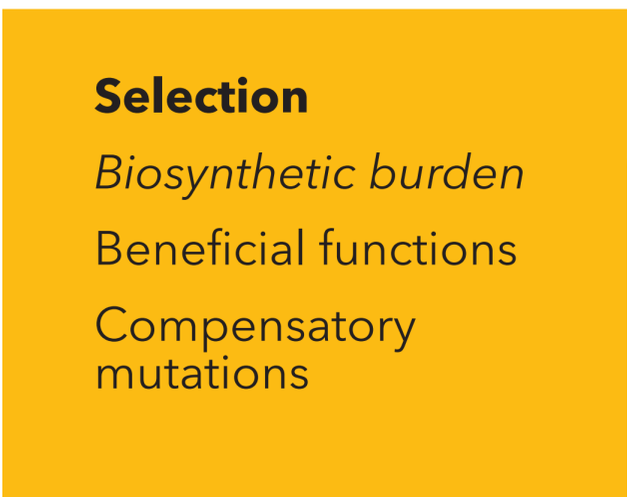
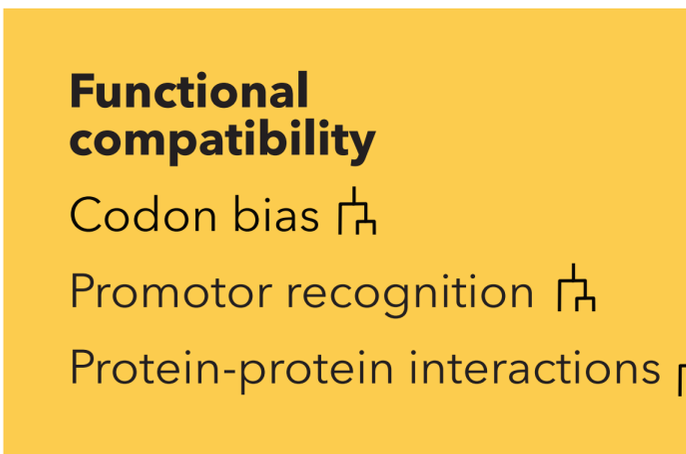
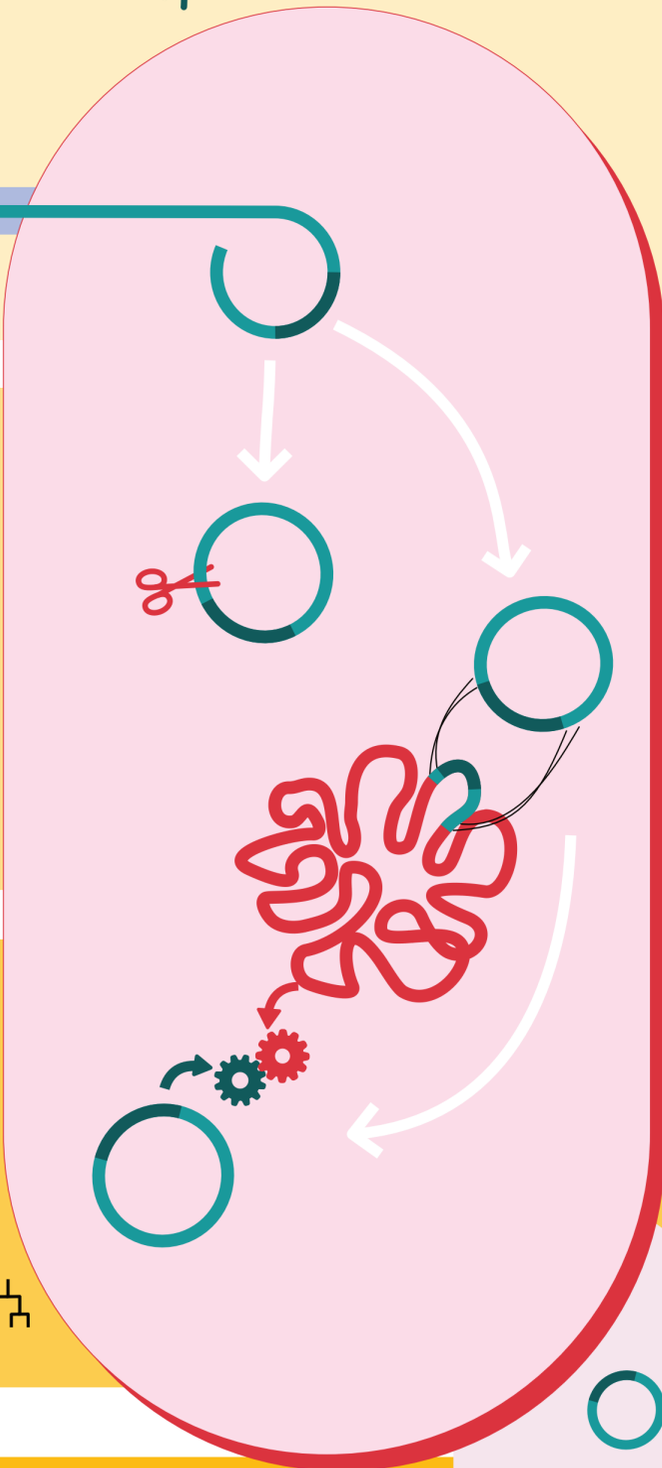
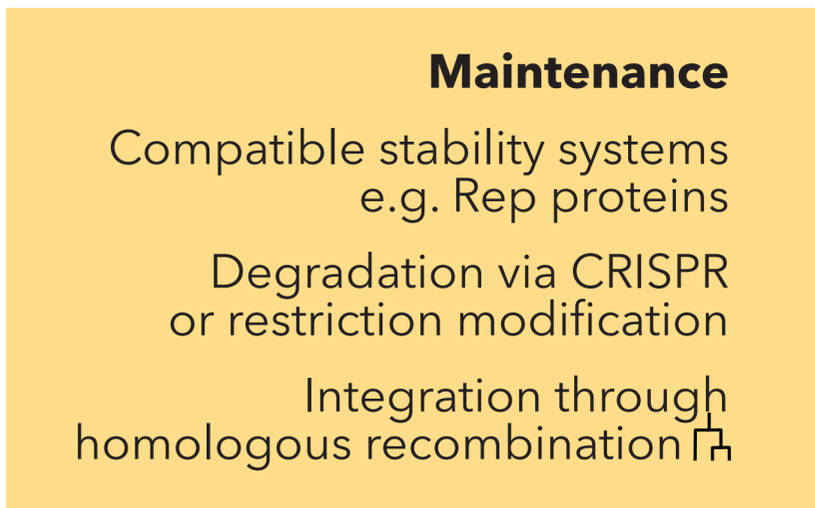
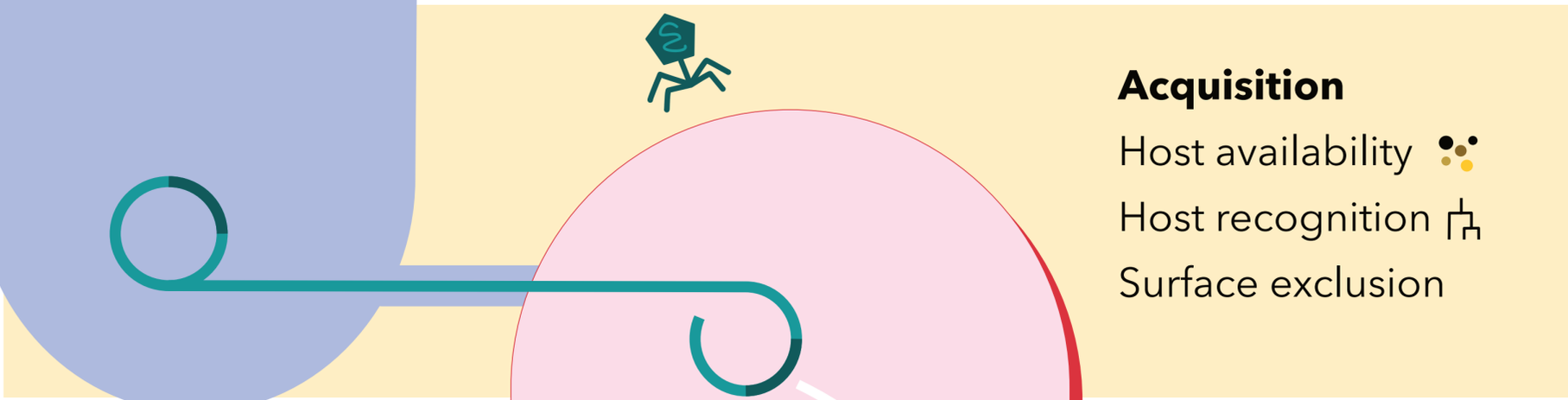


Common among....

niche generalists  
diverse community interactions  
large population size



niche specialists  
limited community interactions  
small population size



●●● likelihood scales with community diversity

⌞ likelihood scales with relatedness