1	Gene Evolutionary Trajectories in <i>M. tuberculosis</i> Reveal Temporal Signs of Selection
2	Álvaro Chiner-Oms ^{1,*} , Mariana G. López ¹ , Iñaki Comas ^{1,2,*}
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4	1. Instituto de Biomedicina de Valencia, IBV-CSIC, Valencia, Spain
5	2. CIBER en Epidemiología y Salud Pública, Valencia, Spain
6	*Correspondence to achiner@ibv.csic.es, icomas@ibv.csic.es
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24 Abstract

Genetic differences between different Mycobacterium tuberculosis complex (MTBC) strains determine their ability to transmit within different host populations, their latency times, and their drug-resistance profiles. Said differences usually emerge through de novo mutations and are maintained or discarded by the balance of evolutionary forces. Using a dataset of ~5,000 strains representing global MTBC diversity, we determined the past and present selective forces that have shaped the current variability observed in the pathogen population. We identified regions that have evolved under changing types of selection since the time of the MTBC common ancestor. Our approach highlighted striking differences in the genome regions relevant for host-pathogen interaction and, in particular, suggested an adaptive role for the sensor protein of two-component systems. In addition, we applied our approach to successfully identify potential determinants of resistance to drugs administered as second-line tuberculosis treatments.

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51 Introduction

The Mycobacterium tuberculosis complex (MTBC) is a genetically monomorphic group of bacteria 52 53 ^{1,2} whose members cause tuberculosis in humans and animals. The MTBC comprises both 54 human-associated (L1, L2, L3, L4, L5, L6, L7, L8, and L9) and animal-associated (A1, A2, A3, 55 and A4) clades ^{3–7}. Due to the absence of mobile genetic elements and measurable recombination 56 among strains and other species ^{8–10}, chromosomal mutations represent the source of MTBC 57 genetic diversity. The maximum genetic distance between any two MTBC strains is around 2,500 58 single nucleotide polymorphisms (SNPs). Strikingly, studies have highlighted large phenotypic 59 differences between strains involving traits like gene expression, drug resistance, transmissibility, 60 and immune response despite this limited variation. In some cases, the mutations driving 61 phenotypic differences have been identified - for example, non-synonymous variants in genes 62 such as rpoB, katG, or embB cause drug-resistant phenotypes ¹¹⁻¹³. Furthermore, single mutations in regulatory elements can induce alterations to downstream gene expression, which 63 can foster differential virulence characteristics ^{14,15}. Finally, specific gene mutations may affect 64 transmission ⁹, host tropism within the complex ¹⁶, and the host immune response ¹⁷. However, 65 66 many of the genomic determinants of these phenotypes remain elusive despite robust evidence that they are driven by genetic differences between strains ^{18,19}. 67

Several types of evolutionary forces play crucial roles in the fixation of mutations in bacterial populations. Previous research has provided evidence for the ongoing positive selection of specific genes and regions ^{9,20–23}, while other studies have reported ongoing purifying selection of specific genomic regions, especially in epitopes and essential genes ²⁴. Additionally, there exists some evidence that genetic drift may have significant functional and evolutionary consequences ²⁵.

74 Detecting selection in MTBC at the genome-wide level remains a challenging task due to limited 75 genetic diversity. The significant accumulation of non-synonymous substitutions has been 76 previously used to characterize patterns of mutation accumulation in large categories of genes 77 ^{24,26}; however, these studies employed a limited number of strains. Of note, the number of MTBC 78 sequences has undergone a recent and rapid expansion, with studies involving hundreds to 79 thousands of strains. The large number of available sequences has allowed, for example, the 80 estimation of the ratio of non-synonymous to synonymous substitutions (dN/dS) signatures in 81 more than 10,000 strains ²⁷, thereby allowing the identification of novel targets of selection with 82 some probably related to host-pathogen interactions. Host-pathogen interaction signals are 83 specially challenging as they are likely obscured by the force exerted by antimicrobial therapies.

Weaker signals are also expected in genes related to second-line drugs related to the relative under-use of related treatments and the low abundance of associated resistant strains in genome databases ²⁸.

87 We reasoned that to detect signs of selection, we should focus on when and/or where they 88 occurred in the phylogenetic tree instead of averaging signs across the phylogeny. In this new 89 study, we developed a methodology to study temporal signs of selection in MTBC genes and 90 identified positive selection in a larger number of genes than previously described. This allowed 91 the identification of past and currently unknown players in MTBC evolution, particularly two-92 component systems, related to host adaptation and second-line drug resistance. This new 93 methodology can be applied to other tuberculosis settings to explore signs of selection associated 94 with changing selective pressures and could be extremely useful to unravel hidden details in the 95 evolution of other human pathogens.

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97 Results

98 Dataset Preparation

We downloaded all samples described in Brites et al.⁴, Coll et al.²⁹, Stucki et al.³⁰, Guerra-99 Assunçao et al.³¹, Zignol et al.³², Bos et al.³³, Ates et al.³⁴, Comas et al.¹⁰, Comas et al.³⁵, 100 Borrell *et al.* ³⁶, and Cancino-Muñoz *et al.* ³⁷ and obtained whole-genome sequencing data from 101 102 9,240 samples comprising the primary human- and animal-adapted MTBC lineages. We mapped 103 Fastg files for each sample against the inferred ancestor of the MTBC and extracted genomic 104 variants (Methods), from which we derived a multiple sequence alignment and a phylogeny. The 105 huge size of the phylogeny and the multifasta file obtained made unaffordable certain parts of the 106 planned subsequent computational analyses, hence we used Treemer to prune the tree down to 107 4,958 leaves (**Table S1**) while maintaining 95% of the original genetic diversity. With this final set 108 of selected samples, we reconstructed a multiple sequence alignment and a phylogeny (Figure 109 S1a).

We mapped each genomic variant to the inferred phylogeny using PAUP (Phylogenetic Analysis Using Parsimony). This step provides information regarding the branch in which every mutation appeared, which allows the identification of homoplastic variants - those that appeared multiple times in different branches of the phylogeny - and the relative 'age' of every mutation, calculated as the node-to-root genetic distance.

116 Scars of Past Selection and Drift in Almost Half of the MTBC Genome

As a first step, we calculated the pN/pS values for genes that possessed up to ten identified variants (n=3,690). A previous study stated a mean pN/pS value for the complete MTBC genome considerably under 1 ³⁸. In agreement with this result, we found that 90% of the genes evaluated possess a pN/pS value less than 1 (**Figure 1a**, pN/pS IQR 0.477-0.804), suggesting ongoing evolution under purifying selection. A high pN/pS may reflect the recent origin of the MTBC, given the time-dependent nature of the accumulation of non-synonymous variants ³⁹.

- 123 Of note, the pN/pS value for a gene results from the pN and pS values calculated with all gene 124 mutations found across the phylogeny (what we term the 'overall pN/pS' in Figure 1b). This value 125 does not reflect changes in selective pressures over time and lineages as the pathogen has 126 potentially faced different environmental "challenges." As we estimated the genetic distance to 127 the root for each mutation as a relative measure of time, we calculated temporal trajectories for 128 pN/pS for each gene during MTBC evolution (Methods, Figure 1b). By doing so, we classified 129 all genes according to their pN and pS trajectories over time into five different categories (Figure 130 **1c**, **Figure S2a**, **Table S2**: (i) pS almost always higher than pN (n=2,032); (ii) pN almost always 131 higher than pS (n=154); (iii) pS > pN but inverts to pN > pS at a certain point (n=35); (iv) pN > pS 132 but inverts to pS > pN at a certain point (n=370); and (v) complex pN and pS trajectories with 133 multiple cross-points, which don't support proper categorization (n=1,099). If our classification 134 reflects differences in the selection pattern over time, we expect that those genes with stable 135 trajectories ('always higher'/'always lower') will have accumulated low variances in pN/pS when 136 pooling timepoints. Conversely, we expect changing trajectories to display high variance between 137 timepoints (Methods, Figure S2b). As predicted, we failed to observe significant differences in 138 variance (Welch t-test, p-value > 0.05) in genes belonging to the 'pN almost always higher' or 'pS 139 almost always higher' categories. In both cases, the pN/pS cumulative variation has a value 140 around zero. However, categories with changing trajectories displayed significant differences 141 (Welch t-test, p-value << 0.01), using 'pS almost always higher category' as the reference 142 category.
- In summary, and in contrast with the observation that 90% of genes possess an overall pN/pS <144 1, only 55% of genes (n=2,032) maintained a pN/pS value below a value of 1 since divergence 145 from the MTBC common ancestor. This set of 2,032 genes is overrepresented for experimentally 146 confirmed essential genes in both *in vivo* and *in vitro* conditions (chi-square test, p-values 0.003 147 and <2.2E-16, respectively). In contrast, 45% of the genes (n = 1,658), mainly those initially

148 classified as being under purifying selection, may have faced other types of selective pressures

149 or genetic drift.

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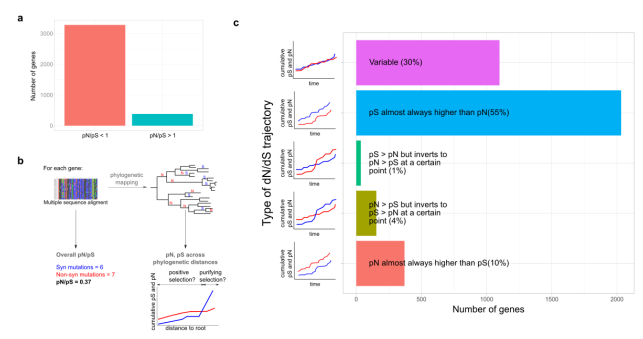


Figure 1. Gene-by-Gene Calculation of pN/pS Over Phylogenetic Time. a. Bar plot showing the number of genes currently displaying a pN/pS > 1 and a pN/pS < 1. b. From the alignment, we inferred the current pN/pS; however, when mapping different mutations onto the phylogeny, we inferred how the pN and pS rates changed over time. c. Five categories grouping studied genes according to their trajectories.

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These results suggest that many genes have been subjected to periods of non-synonymous 157 158 substitution accumulation. Distinguishing between genetic drift and positive selection at a 159 particular time point remains challenging. We expect founder effects to play a crucial role during 160 the early evolution of MTBC, and they may drive a number of the unstable trajectories observed. 161 However, given that MTBC is clonal, positive selection and genetic drift are both expected to have 162 a functional impact. Our analysis identifies a set of genes that shows a pN/pS > 1 near the root 163 but changed to pN/pS < 1 near the leaves (n=370), suggesting that selection and/or founder 164 effects favored the fixation of non-synonymous mutations at early times but that the gene 165 functionality remained conserved at later times. We found that this gene category was enriched 166 for conserved hypotheticals (fisher test, p-value = 0.02) and protein and peptide secretion (fisher-167 test, p-value = 0.05). Intriguingly, we also discovered that certain genes that fell into this category 168 encode known MTBC epitopes (which we will explore below). Of particular note, the presence of

169 154 genes almost always exhibiting a pN higher than pS. This gene category is enriched for non-170 essential *in vitro* genes (chi-square test, p-value=0.005) from three main categories; antibiotic 171 production and resistance (fisher-test, p-value=0.02), conserved hypotheticals (fisher-test, p-172 value=0.02), and unknown functions (fisher-test, p-value=0.03). The mix of genes with a clearly 173 identified function and hypothetical genes suggests that, in some cases, positive selection has 174 been acting through the evolutionary story of some genes while others are likely under genetic 175 drift.

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Evolutionary Trajectories Identify Sensor Proteins of Two-Component Systems Under Positive Selection

179 An increasing number of non-synonymous mutations that start to grow near the leaves may 180 indicate the action of more recent selective forces and suggest unpurged transitory 181 polymorphisms. To distinguish between the two possibilities, we examined the group of genes 182 with a pS > pN in the internal branches but a pN > pS near the leaves (n=58, **Table S2**). Antibiotic 183 resistance genes represent a clear instance of recent positive selection, and we hypothesized 184 that their initial trajectory should reflect conservation of gene function, as they usually perform 185 relevant biological functions and only recently started to diversify due to antibiotic selective 186 pressure. Encouragingly, data for the antimicrobial resistance genes such as *rpoB*, *katG*, *embB*, 187 gidB, and rpsL supported this hypothesis. The genetic distance from the root at which we detected 188 a change in the selective pressures correlates with the time at which each antibiotic became a 189 treatment for tuberculosis. Genes related to resistance to the most recently employed drugs 190 began to accumulate non-synonymous variants at a higher genetic distance to the root than those 191 used in early periods. This point is placed at 1.566483e-04 for gidB (streptomycin, first antibiotic 192 used in tuberculosis treatment in 1946), 1.637692e-04 for katG (isoniazid, use began in 1952), 193 and 1.774088E-04 for both embB (ethambutol, 1966) and rpoB (rifampicin, 1972). These results 194 suggest that our approach possesses sufficient sensitivity to detect recent instances of positive 195 selection.

Among those genes unrelated to antimicrobial resistance, we found several components of toxinantitoxin systems, including *vapC29*, *vapB3*, *vapC35*, *vapB40*, *vapC22*, and *vapC47*, which are critical for the adaptation of bacteria to different stressful conditions. For example, VapC22 has a significant role in virulence and innate immune responses in particular ⁴⁰. Other significant virulence regulators in MTBC are the two-component systems (2CS), which are critical players in extended transcriptional networks. 2CSs comprise a sensor protein coupled to a transcription 202 factor - the sensor protein activates the transcription factor in response to a specific stimuli to 203 trigger a regulatory cascade. We have previously described *phoR*, which encodes the sensor 204 component of the PhoPR 2CS, as an important player in MTBC evolution ⁹ as illustrated by the 205 high levels of accumulation of non-synonymous variants over time. Our data shows that kdpD, a 206 gene that encodes the sensor component of the KdpDE 2CS, displays a similar pattern, with a 207 dN/dS value that reached ~2 at some points during MTBC evolution. In both 2CSs, the genes 208 encoding the regulatory protein (*phoP* and *kdpE*) display high conservation at the functional level, 209 with the pS values consistently higher than the pN values. For the NarLS 2CS, both the regulatory 210 protein (narL) and the sensor protein (narS) exhibit changing patterns towards recent positive 211 selection; however, as for the other described 2CSs, the sensor domain of narS accumulates 212 more non-synonymous variants (fisher test, p-value = 0.036). Our analysis suggests that sensor 213 proteins of 2CSs allow MTBC strains to adapt to varying environments during host-pathogen 214 interaction.

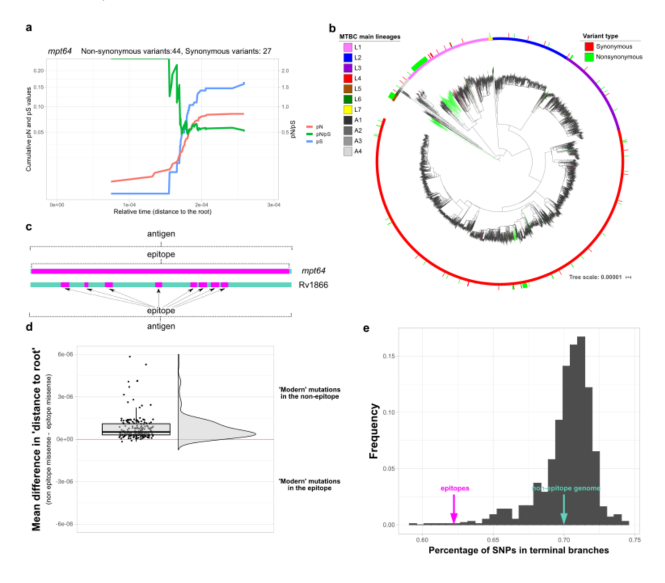
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Epitope Mutations are Older and Show Divergent Evolutionary Trajectories Compared to the Rest of the Antigen

Contrary to many other pathogens, the *M. tuberculosis* genome regions recognized by the host tend to be conserved, albeit with some exceptions ^{24,41}. Given our new results revealing past "scars" of selection in MTBC genes, we analyzed the pN/pS trajectory of a total of 179 antigens harboring 1,556 epitopes ⁴². Specifically, we aimed to evaluate a hypothesis that epitope and nonepitope regions of the antigen experience different selective pressures and that the former most likely reflect interactions with the immune system while the latter reflects the evolution of gene function.

225 Our results revealed that $\sim 60\%$ of the antigens analyzed exhibited a pN/pS value of < 1 across 226 phylogenetic history, providing evidence for their conservation since their diversification of the 227 MTBC from a common ancestor (Table S3). Of note, a relevant proportion of antigens (11%) 228 accumulated a high number of non-synonymous variants in internal branches, which now appear 229 to be conserved (**Table S3**). For example, the *mpt64* gene encodes for a known antigen employed 230 in diagnostic tests. When mapping the genetic variants in the MTBC phylogeny, most non-231 synonymous mutations map to the L5 ancestral branch in a large clade of the L1.2.2 sublineage 232 and a group of L4.10 strains (Figure 2a, b). Other antigens, such as eccD2, Rv1866, fadD21, or 233 Rv2575, exhibited a similar pattern. Apart from human-adapted clades, specific antigens 234 accumulated non-synonymous mutations in deep branches of the animal-adapted lineages, such

as Rv2575 or *IlvB1*. This suggests that these antigens were under positive selection or genetic
drift driven by founder effects when MTBC diversified.





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239 Figure 2. Specific Antigenic Genes Show Signs of Early Positive Selection. a. Cumulative pN, pS, and 240 pN/pS trajectories over time for the mpt64 antigen (Rv1980c). The x-axis represents the genetic distance 241 of each node to the root. The left y-axis represents the cumulative pN (red line) and pS (blue line) values. 242 The right y-axis represents the pN/pS. b. Maximum-likelihood MTBC phylogeny with mapped mpt64 243 variants. The sticks in the outer circle mark the strains with variants identified (red synonymous, green non-244 synonymous). Deep non-synonymous mutations can be found in deep nodes of L1 and L5. c. Some 245 epitopes comprise the entire antigen (such as in mpt64), while in genes such as Rv1866, the epitope 246 represents a small subset of regions embedded in the antigen. d. Raincloud plot of the mean differences 247 in the distance (to root) value between the non-epitope and the epitope mutations for each antigen. e.

Distribution of SNPs found in terminal branches for 1,000 randomly selected sets of non-epitope fragments (grey bars). The percentage of SNPs observed in the epitopes differs from this distribution (\sim 62%, z-score = -4,28, pink arrow), while the percentage of SNPs found in the rest of the genome remains similar to the distribution (\sim 70%, green arrow).

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253 For another group of antigens (27%), the pN/pS value failed to show a definitive trajectory (**Table** 254 **S3**). Specific antigens showed a pattern of pN/pS value of ~ 1 since the diversification of the 255 MTBC from a common ancestor. This pattern could reflect two different causes: genetic drift or 256 differential selective pressures in different MTBC clades/lineages, which could be masked when 257 calculating a common pN/pS for all lineages. The second option is defined by an accumulation of 258 non-synonymous mutations in specific MTBC clades and synonymous mutations in other clades. 259 As a result, the overall pN/pS value would be ~ 1. We observed this scenario, for example, in the 260 *lpqL*, *mce2A*, and *esxH* genes; in these cases, we found an elevated accumulation of non-261 synonymous mutations in deep branches of the L1, L2, and *M. africanum* lineages, although they 262 are highly conserved in modern lineages. Other genes exhibited a similar pattern (**Table S3**), 263 while others could have evolved under the effect of genetic drift.

In general, the evolution of antigens does not essentially differ from other genes in their respective
functional categories. When we compared the trajectories of the antigens against such genes, we
failed to encounter statistical differences between the distributions (Fisher test, BH adjusted pvalue > 0.05).

268 Of note, antigens have a myriad of distinct functions, but the immune system only recognizes 269 specific regions of the antigens - the epitopes. In some cases, epitope regions cover the entire 270 antigen (as for *mpt64*), so selection acts on the antigen and epitope equally. In other cases, 271 epitopes represent only a small fraction of the antigen and may be subject to different selective 272 pressures than the rest of the gene (Figure 2c). When exploring whether selection at the epitope 273 level drives different antigen trajectories, we encountered the Rv1866 locus as a clear example. 274 This antigen has a pN/pS value of >1 near the root, but its value changes to <1 near the leaves, 275 suggesting the action of distinct types of selection across the phylogeny; however, the epitopes 276 contained are highly conserved with a pN/pS value of <1 during the complete trajectory.

In most cases (**Table S3**), the evolutionary trajectories of epitopes seem to be unlinked to the rest
of the antigen, with most epitopes being conserved. We hypothesized that epitopes might reflect
past selection events to adapt to different populations during the initial expansion of the MTBC.
In general, the mean relative phylogenetic age (measured as the genetic distance to the root) of

281 the non-synonymous variants present in the epitopes is older than the non-synonymous variants 282 of the non-epitope regions of the antigen. This phenomenon can be observed when pooling all epitope vs. non-epitope variants (Welch t-test, p-value = 8e-07) and when splitting by different 283 284 genes (Figure 2d) (although with considerable overlap, as expected). Consequently, we expect 285 fewer mutations to accumulate at phylogeny tips if epitope conservation becomes more important 286 at a later stage. The proportion of mutations in epitopes falling in terminal branches (62%) is 287 significantly lower than in sets of regions of the same size randomly selected from the non-epitope 288 genome (70%, z-score = -4.28, P(x < Z) = 0.00001, **Figure 2e**). This suggests the more robust 289 nature of negative selection on epitopes than the rest of the genome in circulating strains.

Thus our results provide further evidence for the generally unlinked nature of gene and epitope evolution, which had been previously established in smaller sets of samples ^{24,38}. In addition, we demonstrate that interaction with the immune system likely drives epitope conservation (as it is the only function in common among epitopes), while non-epitope regions reflect the selection of the gene's biological function. Finally, mutations in epitopes mainly reflect older fixation events while the rest of the genome accumulates mutations more rapidly.

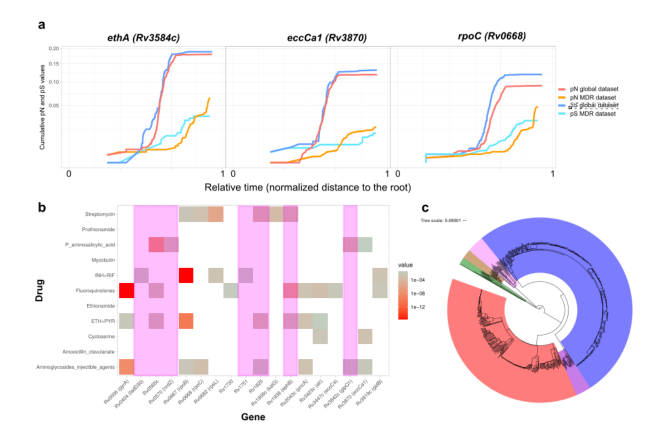
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Novel Candidate Drug Resistance Regions Revealed by a Dataset Enriched for MDR/XDR TB-associated Strains

299 Identifying genes involved in resistance to second- and third-line drugs and new and repurpose 300 drugs remains challenging. We reasoned that if our approach was powerful enough to identify 301 changing selective pressures due to the introduction of first-line antibiotics, we should detect 302 changes in genes associated with the treatment of multidrug-resistant (MDR)- and 303 extensively drug-resistant (XDR)-tuberculosis patients. We assembled and compared a dataset 304 enriched for MDR (n=312) and XDR (n=132) strains and additional sensitive controls to our global 305 dataset (Figure S1b, Table S1). Our analysis revealed instances of genes with an increased pN 306 value towards the leaves of the tree for the MDR/XDR dataset compared to the global dataset. 307 Our approach correctly identified genes associated with MDR, such as gyrA (quinolones), ethA 308 (ethionamide), and *rpoC*, which compensates for the fitness cost of MDR strains (**Figure 3a**). 309 Importantly, we also identified less-well-studied genes with a similar profile, including Rv0552, 310 Rv1730c, alr (Rv3423c), eccC4 (Rv3447), eccCa1 (Rv3870) (Figure 3a), and Rv3883c (mycP1). 311 To formally evaluate their association to different drugs, we generated computational models 312 (Methods, Figure S1b) to link the observed drug-resistant phenotypes with mutations in genes 313 with a changing pN/pS pattern. Well-known resistance-conferring genes such as rpoB, katG, or

rpsL exhibit a strong statistical association with drug-resistant phenotypes, as expected (**Table S4**, **Figure 3b**). Corroborating our observations, the identified less-well-studied genes displayed a significant association with resistant phenotypes for second-line drugs. For example, Rv1730 weakly associated with fluoroquinolone-resistant phenotypes (Wald test, p-value = 0.02), *alr* with D-cycloserine and fluoroquinolones (Wald test, p-value = 0.007 and p-value = 0.001), *eccC4* with fluoroquinolones (Wald test, p-value = 0.01), and *eccCa1* with D-cycloserine and aminoglycoside injectable agents (Wald test, p-value = 0.04 and p-value = 0.002).

321 Of note, our analysis did have certain limitations; for example, given the combined therapy 322 administered in tuberculosis treatment, the same gene may correlate with several antibiotics. 323 Likewise, given the enrichment of this dataset with L4 and L2 strains (Figure 3c), non-324 synonymous phylogenetic variants in genes such as fadD30 (Rv0404), Rv0565c, nrdZ (Rv0570), 325 Rv1751, Rv1825, ephB (Rv1938), and glpQ1 (Rv3842c) appear to be associated with drug-326 resistant phenotypes but are likely neutral markers, a previously reported phenomenon ⁴³. The 327 identification of previously uncharacterized genes represents the overall value of the analysis, 328 with results requiring corroboration by fine-grain *in vitro* experiments.



330 Figure 3. Identification of Genes Related to Second-line Antibiotic Resistance. a. Three genes 331 showing signs of ongoing positive selection in the MDR-enriched dataset but ongoing purifying selection in 332 the global dataset. The x-axis represents the node-to-root genetic distance normalized in the 0-1 range to 333 merge data from both trees as a measure of relative time. The y-axis represents the cumulative dN and dS 334 values. b. A computational model has been constructed for each antituberculosis drug to identify specific 335 gene mutations associated with resistance. In the matrix, rows represent antibiotics and columns represent 336 genes suspected to be under positive selection in the MDR-enriched dataset. Colored cells (from gray to 337 red) indicate a statistically significant association between non-synonymous mutations found in the genes 338 and resistant phenotypes. Genes marked in pink show a strong association with drug-resistant phenotypes 339 due to phylogenetic variants, suggesting that the association may be spurious. c. Maximum-likelihood 340 phylogeny constructed with the MDR-enriched dataset showing an overrepresentation of L2 (blue) and L4 341 (red) strains.

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343 Selection Also Acts in Non-coding Regions

344 Beyond mutations affecting coding regions, we (and others) have established the importance of 345 mutations in intergenic regions in shaping the pathogen's phenotype, as they can alter gene 346 regulation. Hence, natural selection can also target these positions. Using a Poisson distribution, 347 we identified 290 intergenic regions possessing more mutations than expected by chance (BH 348 adjusted p-value < 0.05). 270 of the intergenic regions harbor homoplastic mutations, 349 representing a good correlate of positive selection in MTBC. Certain mutations had been 350 previously categorized as resistance-conferring variants, including 1673425C>T (upstream 351 fabG1), 4243221C>T (between embC-embA), or 2715342C>G (upstream eis) (Table S5). We 352 found other mutations in intergenic regions suspected of being related to drug resistance; 353 however, the exact mutations were not present in the PhyReSse and ReseqTB catalogs.

354 We also calculated the ratio of intergenic variants per intergenic site compared to the ratio of 355 synonymous variants per synonymous site of the flanking genes (pl/pS) for each intergenic region 356 as a measure of selective pressure, as previously proposed by Thorpe et al. ⁴⁴. We found a mean 357 pl/pS value of 1.03 (95% CI: 0.98 - 1.07), near the expected value of 1 when under no selection; 358 however, 123 intergenic regions appeared as outliers of this distribution (**Table S6**) as they exhibit 359 pl/pS values greater than 2.058 (calculated as Q3 + 1.5*IQR ⁴⁵). A gene set enrichment analysis 360 (GSEA) of gene ontology (GO) functions of flanking genes of these intergenic regions 361 demonstrated that the most overrepresented functions (Hypergeometric test, BH corrected p-362 value < 0.05) are responses to acid chemicals, REDOX processes, and regulation of DNA 363 templated transcription. The identification of REDOX is in agreement with oxidative metabolism

playing a role in macrophage survival and drug resistance ^{46–48}. A previous study reported that changes in regulatory regions (mostly intergenic) could significantly affect the transcription rates of downstream genes ⁴⁹. Therefore, the positive selection of these regions may not be surprising.

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368 Discussion

369 Pathogen diversity reflects a balance between evolutionary forces. In the case of the virtually 370 clonal MTBC (⁹, highly diverse and highly conserved genes can be identified despite low genetic 371 diversity¹, thereby suggesting the activity of distinct evolutionary forces. While metrics such as pN/pS present with certain limitations ³⁹, they allow the identification of the footprints of 372 evolutionary forces. pN/pS has the power to identify selection at the genome-wide level ^{26,27}, 373 374 including traces of positive selection in specific genes, gene categories, and/or lineages ^{23,50,51}. 375 Analyses revealed an average pN/pS value across the MTBC genome of around 0.7, well below 376 the value of 1 expected for any organism but high compared to others. This likely reflects the 377 recent emergence of MTBC with the presence of many transitory polymorphisms ³⁹ and the impact 378 of genetic drift in the form of bottlenecks and founder effects (Herbergh 2008). However, the 379 balance of evolutionary forces shaping genetic diversity is dynamic, and what was under positive 380 selection or drift in the past may be under negative selection in the present and vice versa. This 381 idea is illustrated in our work by the striking discovery of scars of elevated non-synonymous rates 382 in almost half of the MTBC genome, contrasting with previous reports (Coscolla et al. 2015; 383 Pepperell et al. 2013).

384 Our analyses identified different temporal evolutionary dynamics in *M. tuberculosis* genes. In one important category, genes are subjected to positive selection or genetic drift early in MTBC 385 386 evolution but to purifying selection near the leaves. A prominent example of this phenomenon is 387 the accumulation of early non-synonymous variants in epitopes such as mpt64. Deep mutations 388 may reflect past events such as founder effects or drift, but our analysis suggests that mutations 389 in epitopes are older when compared to other regions of the genome and that epitope evolution 390 is not linked to the evolution of the rest of the antigen and functional category. These observations 391 are compatible with scenarios suggesting early co-evolution of host and pathogen populations ⁵.

We also identified genes subjected to purifying selection in the past but to current positive selection. The abrupt shift in the pN/pS values in resistance-conferring genes illustrates the impact of antibiotic treatments on MTBC evolution. While our novel approach detected an increase in the pN/pS in a set of genes in MDR and XDR strains, we did not observe this increase

396 in strains not exposed to second-line drugs. This finding allowed the proposal of a set of candidate 397 genes that confer resistance to second-line antitubercular drugs. Previous reports have suggested that genes such as *alr* or *eccCa1* can confer resistance to MDR treatments ^{15,52–56}; 398 399 however, novel genes identified in this study highlight our incomplete understanding of the genetic 400 basis of resistance, in particular for second-line and new drugs. Our approach also detected 401 genes unrelated to antibiotic resistance that have been subjected to recent positive selection, a 402 finding missed when applying averaged pN/pS ratios. We commonly encountered the sensor 403 component of 2CSs in this gene-set, and our previous data established robust signs of recent 404 positive selection in *phoR*, the sensor component of the PhoPR 2CS ^{9,57}. This finding suggested 405 that non-synonymous mutations in *phoR* participate in host adaptation by regulating *PhoP*, a 406 major regulator of MTBC physiology and virulence. We now show a similar occurrence in two 407 other sensor proteins - KdpD and NarS. Thus, the accumulation of non-synonymous mutations in 408 sensor proteins may represent a common strategy used by mycobacteria to adapt to the changing 409 environment during infection.

410 In addition to coding regions, we also found traces of selection in non-coding sequences, which 411 agrees with previous findings ⁴⁴. While identifying selection pressures on intergenic regions 412 remains challenging, given the problematic interpretation of the functional effect of variants that 413 fall in these areas, homoplastic mutations and the comparison of variants against surrounding 414 genes provide a good framework. Variants accumulation in these regions can impact the 415 regulation of nearby gene expression ^{16,49}. Again, drug resistance appears to represent the 416 strongest selective force; however, variants found in these regions also impact transcription factor 417 activity and oxidative metabolism

418 We are aware of the limitations of our current study. The study of past traces of selection in MTBC 419 members remains challenging due to the low genetic diversity present; however, we attempted to 420 maximize genetic diversity to gain resolution by including a broad representation of the main 421 MTBC lineages. Unfortunately, subtle traces of selection affecting small subclades or groups of 422 strains can be masked using this strategy - indeed, this is illustrated by our study when lineage-423 positive signs of selection fail to appear in our analysis. For example, Menardo et al. have 424 described a high number of non-synonymous mutations in the epitopes of esxH⁴¹. This finding is 425 not reflected when considering all lineages but only when we search lineage by lineage (Table 426 S3). Further analysis focusing on specific subclades may illuminate differential evolutionary 427 pressures within the MTBC. Furthermore, we only analyze mutations fixed in the phylogeny, so 428 we only infer an approximate picture of the evolutionary forces that have shaped complex

evolution in the past. In addition, the low variability present in the MTBC, strain subsampling, and
lack of metadata/dates for most deposited genomes make absolute dating for some studied
mutations extremely challenging. We are also aware that, in some cases, genetic drift may be
mistaken with other selection forces; however, this does not preclude those changes from having
a functional effect ²⁵.

434 Finally, we note that our approach can be used as a blueprint to study the evolution of several 435 bacterial species. For example, the Salmonella genus includes strains exhibiting high host-436 specificity and those with the general ability to infect many hosts ⁵⁸. The gene-by-gene evaluation 437 of past and current selective pressures could shed light on the genomic determinants that drive 438 differing specificity. The same approach could be valid with *Helicobacter pylori*, a pathogenic 439 bacteria that causes gastric infections and is highly specialized at infecting human hosts ^{59,60}. 440 MTBC displays virtually no recombination or ongoing horizontal gene transfer (which is not the 441 case of *H. Pylori* or *Salmonella*), making the interpretation of the results more straightforward; 442 however, we anticipate that, taking into account population structure, our approach can be 443 adapted to answer a range of evolutionary questions in pathogen evolution.

444

445 Methods

446 Variant Analysis Pipeline and Phylogenetic Reconstruction

447 All samples were analyzed using our variant analysis pipeline, which has been extensively 448 described in a previous publication ⁶¹. Briefly, FASTQ files were trimmed to remove low-quality 449 reads using fastp ⁶² (version 0.12.5, arguments --cut by quality3, --cut window size=10, --450 cut mean guality=20, --length required=50, --correction) and aligned to the most likely inferred 451 ancestor of MTBC²⁴ using the BWA-MEM algorithm ⁶³. Potential optical and PCR duplicates were 452 removed with Picard tools ⁶⁴, while reads with a MAPQ value < 60 were also discarded. Variant calling was performed using SAMtools ⁶⁵, VarScan ⁶⁶, and GATK ⁶⁷. A pileup file was created with 453 454 SAMtools from the BAM files, and VarScan was then used to extract variant positions from this 455 pileup file (version 2.3.7, arguments --p-value 0.01 --min-coverage 20 --min-reads2 20 --min-avgqual 25 --min-strands2 2 --min-var-freq 0.90), while GATK was used to extract INDELS (version 456 457 3.8-1-0-gf15c1c3ef, HaplotypeCaller and SelectVariants functions). To remove mapping errors, 458 detected variants were discarded if found in INDEL areas or areas of high variant accumulation 459 (more than three variants in a 10-bp defined window). Variants were then annotated using SnpEff (version 4.2) ⁶⁸. Variants associated with proline-glutamate/proline-proline-glutamate (PE/PPE) 460

461 genes, phages, or repeated sequences were also filtered out (**Table S7**) as they tend to 462 accumulate false-positive SNPs owing to mapping errors. Finally, with the selected high-quality 463 variant calls, a non-redundant variant list was created and used to retrieve the most likely allele 464 at each genomic sequence to generate a variant alignment.

465 The first phylogeny was constructed with all samples that passed a minimum depth coverage 466 threshold (median 25x) and had no mixed infections (n=9,240). This initial phylogeny was constructed using MEGA-CC⁶⁹ and the Neighbor-Joining algorithm. Later, we pruned the 467 468 phylogeny with Treemer ⁷⁰ to obtain a smaller tree for subsequent computational analyses. A 469 reduction of just 5% of the initial genetic diversity led to the selection of 4,958 samples. With these 470 selected samples, a maximum likelihood phylogeny was constructed using IQTREE ⁷¹ (version 471 1.6.10) with the GTR model of evolution, taking into account the invariant sites and with an 472 ultrafast bootstrap ⁷² of 1.000 replicates.

473

474 Phylogenetic Variant Mapping and pN/pS Trajectories

475 After phylogenetic reconstruction, the mutations called in the 4,958 samples (n=368,719) were 476 mapped onto the phylogeny. For his purpose, the ancestral state of each polymorphism in each node was reconstructed using PAUP ⁷³ with a weight matrix that punished reversions with a 10X 477 478 multiplier. From this information, the phylogenetic node at which each variant appeared was 479 obtained. Later, a relative age derived from the branch length information for each variant was 480 assigned for each variant. This relative age is the genetic distance from the ancestral node to the 481 bisection point of the target branch on which the variant appears. Finally, the cumulative pN and 482 pS trajectories were calculated for each gene using the potential synonymous and nonsynonymous sites inferred using the SNAP tool ⁷⁴ and plotting the pN and pS values at each 483 484 timepoint, taking into account the variants that appeared before this timepoint.

- 485 Initially, we classified the genes according to their pN/pS trajectories with the following criteria:
- 486 I. genes with a cumulative pN/pS < 1 at more than 95% of the sampled points were classified
- 487 as 'pS almost always higher than pN'
- 488 II. genes with a cumulative pN/pS > 1 at more than 95% of the sampled times were classified
 489 as 'pN almost always higher than pS'
- 490 III. genes in which the cumulative pN/pS changed from >1 to <1 or vice versa more than three
 491 times were classified as 'variable'

492 IV. genes in which the cumulative pN/pS changed from >1 to <1 or vice versa less than four
493 times and that the cumulative pN/pS started being <1 but ended >1 were classified as 'pS
494 > pN but inverts to pN > pS at a certain point'

V. genes in which the cumulative pN/pS changed from >1 to <1 or vice versa less than four
times and that the cumulative pN/pS started being >1 but ended <1 were classified as 'pN
pS but inverts to pS > pN at a certain point'.

This classification was reviewed manually at a later stage. Genes with less than ten mutations were not considered for subsequent analyses.

500 The cumulative pN/pS variation for each gene was calculated as:

501

502
$$pN/pS var = \sum_{i=4}^{n} x_i - x_{i-1}$$

503

with *x* the cumulative pN/pS value at each of the sampled *i* points. The first three values of each gene's cumulative pN/pS value were not considered, as the initial values can show significant differences due to a low number of mutations.

507

508 Epitope and Antigen Analysis

All linear epitopes (n=1,556) found in the IEDB database ⁴² that belong to *M. tuberculosis* in August 2019 were downloaded. All linear epitopes with overlapping coordinates with regards to the H37Rv reference strain were merged into unique non-overlapping 'contigs' (n=718). The potential synonymous and non-synonymous sites were inferred using the SNAP tool ⁷⁴. All genes containing such epitopes were considered antigens, except those genes not considered in the variant calling step, as explained above (PE/PPE, phages).

515 The percentage of SNPs that occur in these 718 regions that appear in terminal branches of the 516 phylogeny were determined using the information derived from PAUP. The percentage of SNPs 517 in the rest of the genome (not considering these 718 regions) that fall in terminal branches were 518 also determined. To evaluate if the difference between these values was statistically significant, 519 718 segments of the non-epitope genome with the same length as the epitope regions set, 1,000 520 times, were selected. For each iteration, the percentage of SNPs found in terminal branches was

 $\gamma - \mu$

521 calculated and plotted in a distribution. Finally, a z-score (see below) between the distribution and522 the value observed for the epitopes was calculated.

523

524
$$z$$
-score = $\frac{x - \mu}{\sigma}$

525

526 Gene Set Enrichment Analysis

527 Several approaches for functional category enrichment were performed to compare genes 528 present in our sets of interest against other genes. For the essentiality enrichment, the *in vivo* ⁷⁵ 529 and *in vitro* ⁷⁶ classification of genes was used, and the enrichment in these categories tested 530 with Fisher tests. For GO enrichment, the Bingo tool ⁷⁷ was used with a hypergeometric test 531 (sampling without replacement) and the Benjamini-Hochberg correction for multiple testing 532 comparisons. Finally, the enrichment of the functional categories was also evaluated ⁷⁸ employing 533 Fisher tests corrected with the Benjamini-Hochberg procedure.

534

535 Drug-resistant Dataset Preparation and Analysis

536 Drug-resistant strains were downloaded from the TBportals database ⁷⁹ on October 22, 2019 537 (n=656). Samples were classified according to their drug-resistant phenotype and then passed 538 through the variant analysis pipeline described above. A maximum-likelihood phylogeny was 539 constructed using IQTREE with the previously described options, including samples from the 540 Comas *et al.*, 2013 study to achieve nodes from lineages underrepresented in the TBportals 541 database.

542 The pN/pS trajectories were calculated and classified as explained for the other dataset.

A matrix was next created that included phenotypic information for each tested drug (resistant/susceptible) and the presence/absence of non-synonymous mutations in the gene set classified as having a trajectory in which the pS > pN but inverts to pN > pS at a certain point for each sample. A set of binomial logistic regression models was constructed with this data, explaining the observed phenotypes based on the presence of non-synonymous mutations on selected genes. These models were trimmed *a posteriori* following a backward stepwise methodology, selecting the set of regressors that show the best Akaike Information Criterion.

550

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738

739 Author Contributions

I.C. conceived this work. A.C.O. and M.G.L. analyzed the data. A.C.O. wrote the first version of
 the draft. A.C.O., I.C., M.G.L. critically reviewed and contributed to the final version of the paper.

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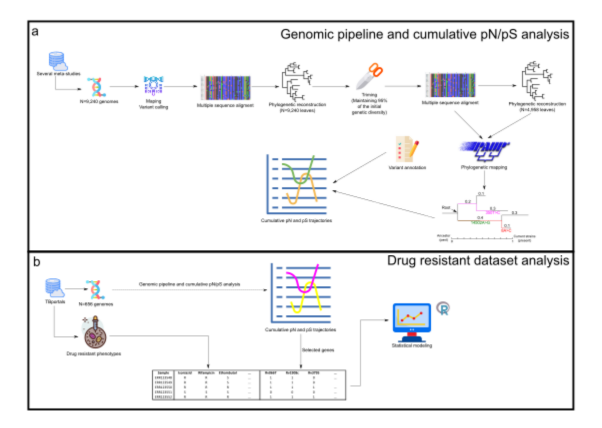
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753 Supplementary Material



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756 Figure S1. Workflow Followed in Different Analyses. a. From public repositories, we downloaded more 757 than 9,000 MTBC genomes. After reconstructing a phylogenetic tree, the dataset underwent a trimming 758 process to reduce the number of samples while maintaining as much genetic diversity as possible. From 759 these reduced datasets, we reconstructed a tree and an alignment. PAUP mapped each detected 760 polymorphism into the phylogeny. Finally, knowing the annotation of the polymorphisms and the branch in 761 which they appeared allowed us to generate pN/pS trajectories. b. TBportals was used to obtain a dataset 762 enriched for resistant strains. The same approach as described above was applied (except for the trimming 763 step), thereby obtaining pN/pS trajectories for each gene based on the information of this new dataset. We 764 also downloaded drug-susceptibility test (DST) information for each resistant strain. Combining both the 765 genomic and the phenotypic information allowed the generation of computational models linking the 766 observed phenotypes to mutations in specific genes.

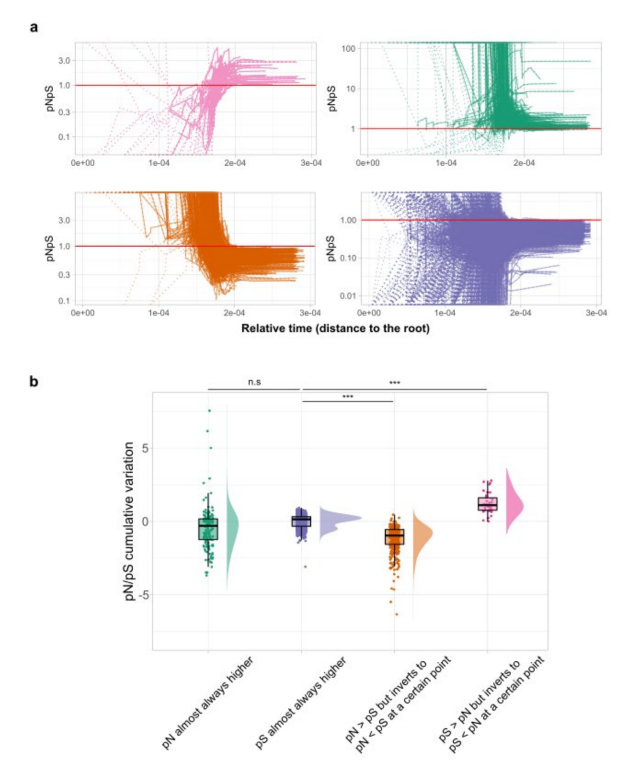
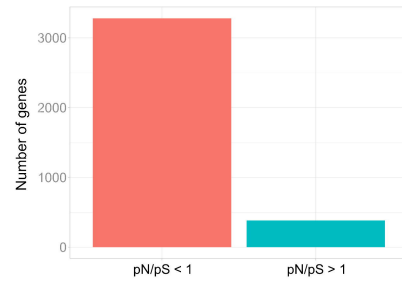
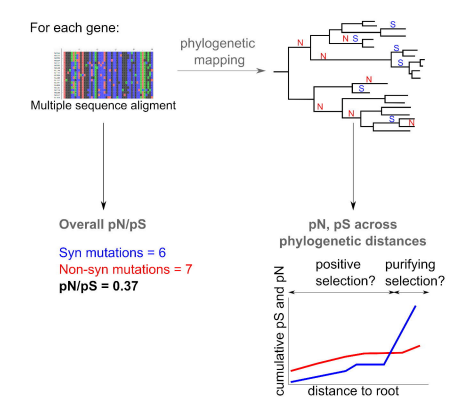


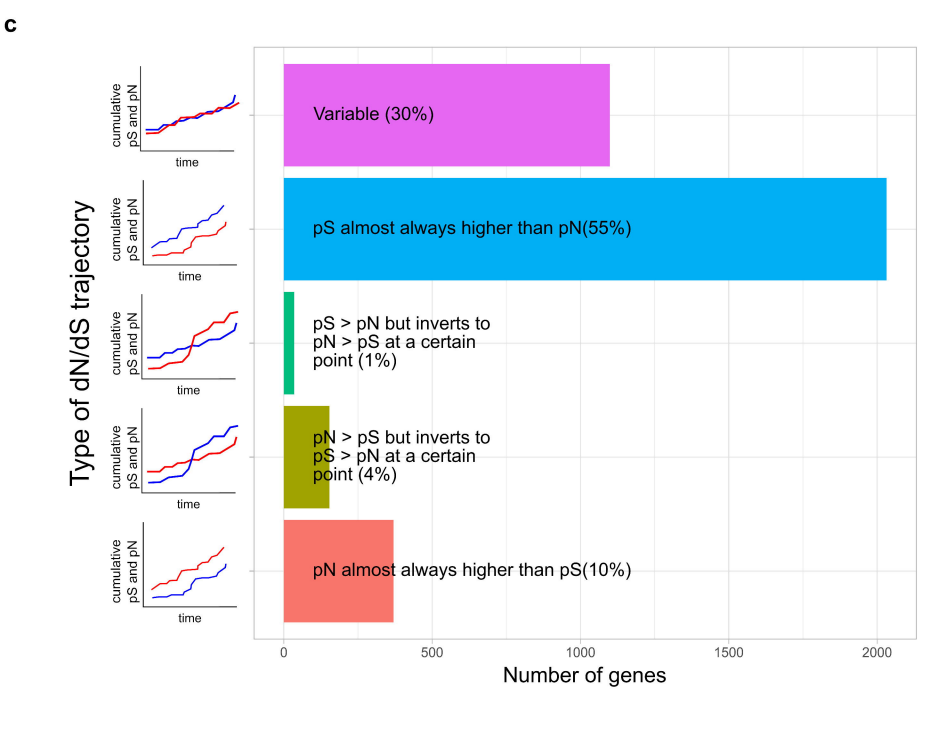
Figure S2. Classification of Genes According to pN/pS Trajectory. a. pN/pS variation across the phylogeny, from root to tips. Each line corresponds to a different gene. Genes were classified as: (i) pS almost always higher than pN (blue); (ii) pN almost always higher than pS (green); (iii) pS > pN but inverts to pN > pS at a certain point (pink); (iv) pN > pS but inverts to pS > pN at a certain point (orange); (v) pN

773 and pS had complex trajectories (not plotted). The red horizontal line marks pN/pS = 1. The first three 774 values of the trajectory (dashed in the plots) were not considered for classification, and the rest of the 775 analysis as they present with high variability due to a small number of mutations. b. Cumulative pN/pS 776 variation distribution for each gene category. Categories reflecting 'stable' trajectories ('pN almost always 777 higher' and 'pS almost always higher') accumulated low variance in pN/pS and displayed no significant 778 differences (Welch t-test, p-value > 0.05). In both cases, the pN/pS cumulative variation is around zero. In contrast, categories with changing trajectories display significant differences (Welch t-test, p-value << 0.01),

- 779
- 780 using 'pS almost always higher category' as the reference category.
- 781 **Table S1.** Samples used in the analyses, including accession numbers and the main phylogenetic 782 lineage
- 783 Table S2. Classification of genes in the five main categories defined in the main results
- 784 Table S3. Classification of the antigens/epitopes studied, including the categories proposed for 785 each of the features and the lineages in which they show differential trajectories.
- 786 **Table S4.** P-values of the computational models generated - Genes marked in yellow display
- 787 significant values, probably due to phylogenetic markers.
- 788 **Table S5.** Homoplastic variants called in the intergenic regions analyzed.
- 789 **Table S6.** 123 intergenic regions that exhibited pl/pS values that are outliers of the genomic pl/pS
- 790 distribution. Observed and expected mutations in the intergenic regions, probability of observing
- 791 SNPs by chance (Poisson distribution), and the pl/pS calculated are shown.
- 792
 Table S7. Genomic regions not considered for analysis.
- 793 Data S1. Plots of all trajectories calculated.

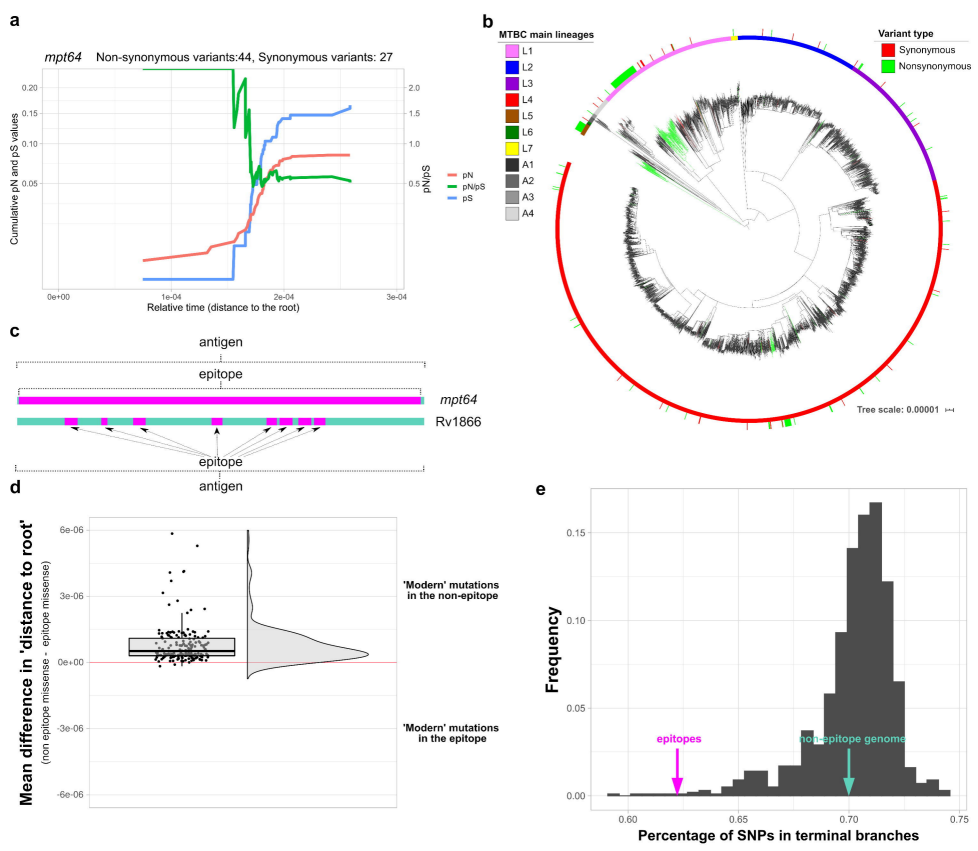


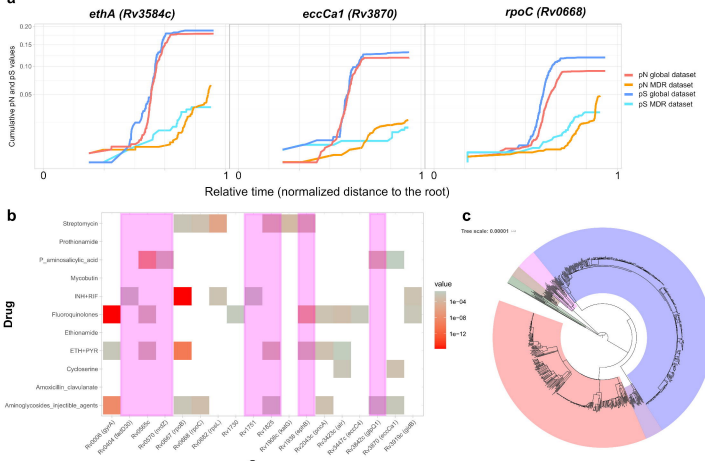




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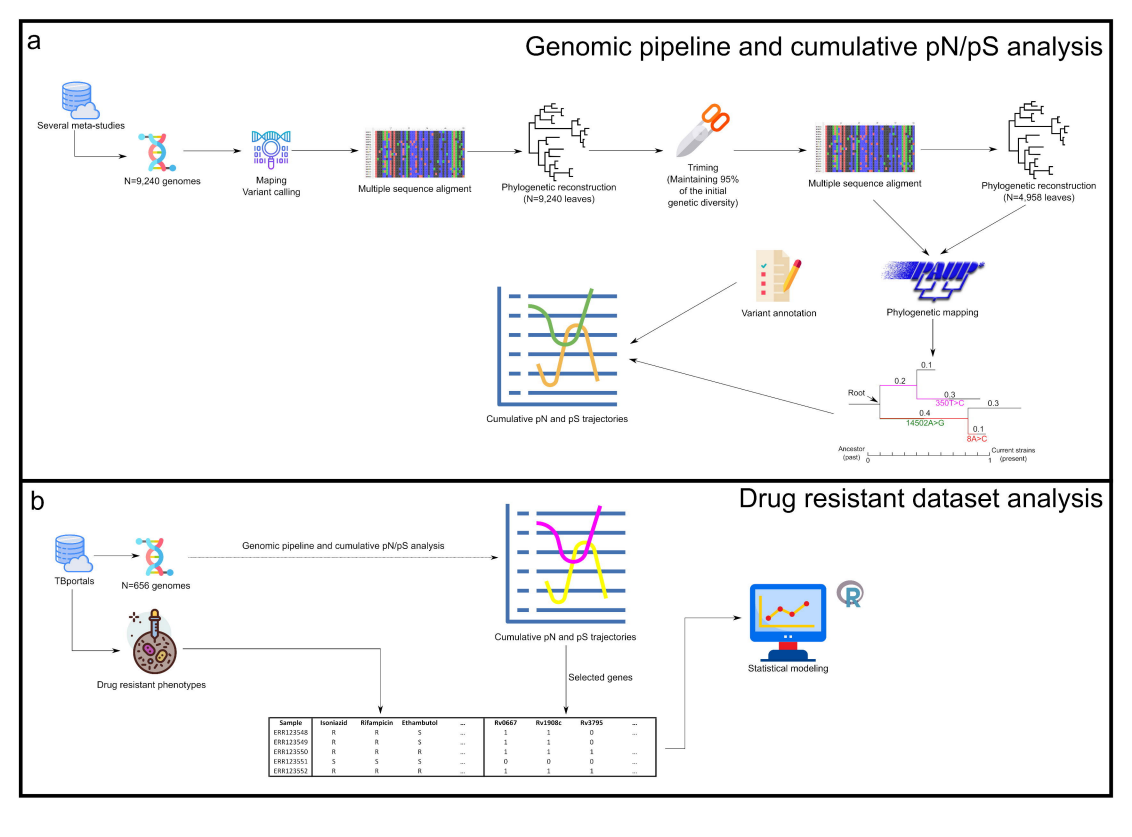
b

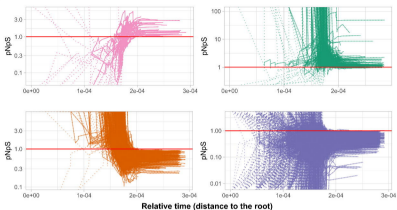


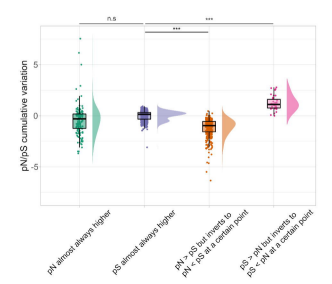


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Gene







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