

1 **Genome sequencing of 196 *Treponema pallidum* strains from six continents reveals**
2 **additional variability in vaccine candidate genes and dominance of Nichols clade strains in**
3 **Madagascar**

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45 **Abstract**

46 In spite of its immutable susceptibility to penicillin, *Treponema pallidum* (*T. pallidum*)
47 subsp. *pallidum* continues to cause millions of cases of syphilis each year worldwide, resulting in
48 significant morbidity and mortality and underscoring the urgency of developing an effective
49 vaccine to curtail the spread of the infection. Several technical challenges, including absence of
50 an *in vitro* culture system until very recently, have hampered efforts to catalog the diversity of
51 strains collected worldwide. Here, we provide near-complete genomes from 196 *T. pallidum*
52 strains – including 191 *T. pallidum* subsp. *pallidum* – sequenced directly from patient samples
53 collected from 8 countries and 6 continents. Maximum likelihood phylogeny revealed that
54 samples from most sites were predominantly SS14 clade. However, 99% (84/85) of the samples
55 from Madagascar formed two of the five distinct Nichols subclades. Although recombination
56 was uncommon in the evolution of modern circulating strains, we found multiple putative
57 recombination events between *T. pallidum* subsp. *pallidum* and subsp. *endemicum*, shaping the
58 genomes of several subclades. Temporal analysis dated the most recent common ancestor of
59 Nichols and SS14 clades to 1717 (95% HPD: 1543-1869), in agreement with other recent
60 studies. Rates of SNP accumulation varied significantly among subclades, particularly among
61 different Nichols subclades, and was associated in the Nichols A subclade with a C394F
62 substitution in TP0380, a ERCC3-like DNA repair helicase. Our data highlight the role played by
63 variation in genes encoding putative surface-exposed outer membrane proteins in defining
64 separate lineages, and provide a critical resource for the design of broadly protective syphilis
65 vaccines targeting surface antigens.

66

67 **Author Summary**

68 Each year, millions of new cases of venereal and congenital syphilis, caused by the
69 bacterium *Treponema pallidum* (*T. pallidum*) subsp. *pallidum*, are diagnosed worldwide,
70 resulting in significant morbidity and mortality. Alongside endemic circulation of syphilis in
71 low-income countries, disease resurgence in high-income nations has underscored the need for a
72 vaccine. Due to prior technological limitations in culturing and sequencing the organism, the
73 extent of the genetic diversity within modern strains of *T. pallidum* subsp. *pallidum* remains
74 poorly understood, hampering development of a broadly protective vaccine. In this study, we
75 obtained 196 near-complete *T. pallidum* genomes directly from clinical swabs from eight
76 countries across six continents. Of these, 191 were identified as *T. pallidum* subsp. *pallidum*,
77 including 90 Nichols clade genomes. Bayesian analysis revealed a high degree of variance in
78 mutation rate among subclades. Interestingly, a Nichols subclade with a particularly high
79 mutation rate harbors a non-synonymous mutation in a putative DNA repair helicase. Coupling
80 sequencing data with protein structure prediction, we identified multiple novel amino acid
81 variants in several proteins previously identified as potential vaccine candidates. Our data help
82 inform current efforts to develop a broadly protective syphilis vaccine.

83

84

85 **Introduction**

86 Syphilis, caused by the spirochete bacterium *Treponema pallidum* subspecies *pallidum*
87 (TPA) remains endemic in low-income countries, where the majority of cases of this infection
88 occurs. A surge in syphilis incidence, however, has been recorded as well in mid- and high-
89 income nations, primarily among men who have sex with men (MSM) and persons living with
90 HIV (PLHIV). The United States saw a 6.5-fold increase in primary and secondary syphilis cases
91 between 2000 and 2019 (1,2), driven in large part by cases among MSM, although cases among
92 heterosexual individuals are now rising rapidly as well. Globally, there were approximately 6
93 million new cases per year among 15-49 year olds in 2016 (3). One million of these cases occur
94 in pregnant women, of which 63% are in sub-Saharan Africa alone (4). Preventing cases among
95 women of childbearing age is a critical worldwide public health initiative, as TPA can cross the
96 placenta and cause spontaneous abortion and stillbirth. Maternal-fetal transmission of syphilis
97 caused approximately 661,000 adverse birth outcomes globally in 2016 alone (5). In the United
98 States, congenital syphilis is also rising, from 9.2 per 100,000 live births in 2013 to 48.5 per
99 100,000 live births in 2019, more than a five-fold increase (2).

100 Given rising infection rates, increasing difficulties in procuring benzathine penicillin G
101 (BPG) for treatment (6), and widespread *T. pallidum* resistance to azithromycin (7–9) which is
102 no longer a viable alternative to BPG, the development of a vaccine against syphilis has become
103 a public health priority. To this end, the syphilis spirochete poses a particular challenge. In
104 contrast to other gram-negative bacteria, TPA has a remarkably low surface density of integral
105 outer membrane proteins (OMPs) (10,11) and uses phase variation (random ON-OFF switching
106 of expression) to further vary its overall surface antigenic profile (12,13). In parallel, this
107 pathogen has evolved a highly efficient gene conversion-based system able to generate millions

108 of variants of the putative surface-exposed loops of the TprK OMP, thus creating an ever-
109 changing target for the host defenses, which fosters immune evasion, pathogen persistence, and
110 re-infection (14–16). Furthermore, TPA cannot be cultured in axenic culture, instead requiring
111 propagation in rabbit testes or, more recently, in co-culture with rabbit epithelial cells (17). This
112 has further hampered efforts to sequence clinical specimens to catalog regions of conservation
113 and diversity, particularly of the OMPs, which is critical for development of an effective vaccine.
114 As of this writing, consensus sequences of only 67 TPA strains have been deposited in INSDC
115 databases, and of these not more than 53 were recovered directly (or following low passage
116 rabbit culture) from clinical specimens. Additional data exist within the Sequence Read Archive
117 (SRA) for up to 600-800 samples annotated as TPA but, without extensive manual curation and
118 reliable assembly pipelines, high-quality data contained within the SRA remain inaccessible to
119 most users. To inform vaccine development efforts, we generated high quality (<1% ambiguous
120 or missing data) near-complete genomes from 196 *T. pallidum* genomes using hybrid capture,
121 enabling direct determination of sequences from clinical specimens without the need for
122 enrichment by culture in rabbits or *in vitro*. These newly available genomes were analyzed to
123 unveil diversity in potential TPA vaccine targets in combination with *in silico* protein folding
124 technology. Our work broadens our understanding of the molecular underpinnings of TPA, and
125 serve as a resource for developing a broadly protective vaccine effective against syphilis.

126

127 **Results**

128 **Nichols clade strains are predominant in Madagascar**

129 As part of ongoing efforts to catalog global *T. pallidum* genomic diversity, we received
130 samples containing *T. pallidum* genomic DNA recovered from primary or secondary lesions. We

131 attempted whole genome sequencing on those with > 100 copies of *tp0574* per 17.5 μ L genomic
132 DNA and obtained 196 high quality genomes consisting of <1% ambiguities using a custom
133 hybridization capture panel to enrich for *T. pallidum* DNA followed by processing through a
134 custom bioinformatic pipeline for consensus genome calling involving both de novo assembly
135 and reference mapping to the SS14 reference genome (NC_021508.1). Summary demographic
136 characteristics for all samples, including 191 TPA, four *T. pallidum* subsp. *endemicum* (TEN),
137 and one *T. pallidum* subsp. *pertenue* (TPE) used in this study are presented in Table 1. Samples
138 were collected between 1998 and 2020 from 8 countries (Peru, Ireland, USA, Papua New
139 Guinea, Madagascar, Italy, Japan, and China) across 6 continents. Median coverage of the
140 reference genome by trimmed, deduplicated reads was 76.8x (range 16.5 - 1293.4), with a
141 minimum of 6 reads required to unambiguously call a base. Median input genomes, as
142 determined by *tp0574* qRT-PCR, for successful genome recovery was 4,319 copies (range 101 –
143 304,484) (Supporting Information 1 – Sample Statistics). The median number of ambiguities in
144 the finished genomes prior to masking was 49 (range 0-10,753).

145

146 **Table 1: Demographic information of samples sequenced in this study**

Country	Year(s) of Collection	Number of Samples	Sex (n)			Stage (n)			Previous Study
			Male	Female	Unknown	Primary	Secondary	Unknown	
Peru	2019	9	5	0	4	5	0	4	n/a
Ireland	2002	11	0	0	11	0	0	11	(18–20)
USA	1998-2002	15	11	3	1	14	1	0	(19,20)
Papua New Guinea	2019	1	0	0	1	0	0	1	n/a
Madagascar	2000-2007	85	0	0	85	10	16	59	(20–22)
Italy	2017	10	10	0	0	10	0	0	n/a
Japan	2019-2020	57	34	22	1	0	0	57	n/a
China	2018	8	0	0	8	0	0	8	n/a
TOTAL	1998-2020	196	60	25	111	39	17	140	

147

148 Following the assembly of the 196 genomes, we combined our strains with an additional
149 55 publicly available consensus genomes, including five TPE, two TEN, 11 TPA laboratory
150 isolates highly passaged in rabbit, and 37 direct clinical specimens/low passage rabbit TPA
151 strains. Due to differences in library preparation expected to affect performance of our assembly
152 pipeline, we chose not to reassemble genomes that did not have an available consensus sequence.
153 All genomes were masked at the intra-rRNA tRNA-Ala and tRNA-Ile and highly repetitive *arp*
154 and *tp0470* genes for which short read Illumina sequencing could not resolve position or relative
155 length. Genomes were further masked at all paralogous *tpr* genes prior to recombination masking
156 by Gubbins (23).

157 The maximum-likelihood phylogenetic tree shown in Figure 1 (and in tabular format in
158 Supporting Information 1 – Sample Metadata) is defined by approximately 130-150 non-
159 recombining SNPs separating any two Nichols and SS14 tips and approximately 1,200-1,450
160 SNPs separating any two TPA and TPE or TEN tips. It recapitulates several features seen in
161 previous phylogenies of *T. pallidum*. Notably, it includes a SS14 Omega node that contains
162 nearly all SS14 clade samples, as well as tight geographic clustering of samples predominantly
163 collected in China and Japan (24) and characterized by uniform azithromycin resistance caused
164 by the A2058G mutation in the 23S rRNA allele (Figure 1A-C; SS14 Omega – East Asia node).
165 None of these samples was resistant via the A2059G allele. We also observed genotypic
166 azithromycin resistance in geographically diverse samples in both the SS14 and Nichols clades,
167 further supporting the hypothesis that this mutation arises spontaneously (24).

168 The most striking feature of our *T. pallidum* phylogeny is the extensive circulation of
169 strains belonging to the Nichols clade in Madagascar. All but one of the 85 Madagascar strains
170 belonged to one of two Nichols subclades, A and B. The former consists of only Madagascar

171 strains except for a single strain from Cuba, and the latter containing only Madagascar samples.
172 Except for an A2059G 23S rRNA mutation (25) observed in one sample, all Madagascar Nichols
173 strains were azithromycin sensitive.

174 In addition to Nichols subclades A and B primarily from Madagascar, three additional
175 distinct subclades were observed, containing samples collected throughout the world. The
176 Nichols C subclade shares a common ancestor with the Nichols B subclade and is uniformly
177 azithromycin resistant, in contrast to most other Nichols clade samples. Both Nichols D (which
178 contains all laboratory strains) and Nichols E subclades are more distantly related to the
179 Madagascar samples in Nichols A and B. The Nichols E subclade includes two previously
180 reported samples from France as well as two newly sequenced samples from Japan and Italy.
181 Interestingly, both the Japanese and Italian patients whose samples are included in this subclade
182 report their sexual orientation as MSM; one French sample, CW59, was collected from an anal
183 smear. Although this is hardly conclusive, the appearance of distinct TPA clades circulating
184 among MSM individuals has recently been documented in Japan (26), suggesting that this
185 phenomenon may be occurring worldwide.

186 In addition to the unexpected number of Nichols clade samples, we were also surprised to
187 observe two samples from Maryland that clustered with the very distant SS14 clade genome,
188 MexicoA, originally collected in 1953 from a male living in Mexico. These strains, MD06 and
189 MD18B, diverge from the MexicoA strain by 33 and 24 non-recombining SNPs, respectively;
190 from SS14 Omega strains by about 45 SNPs; and Nichols clade strains by about 150 SNPs. The
191 MexicoA strain is unique in that it shares signatures of both syphilis and yaws organisms in
192 several virulence factors (27,28). To our knowledge, clinical specimens clustering with the
193 MexicoA strain have not previously been reported. Although the MD18B and MD06 samples

194 were collected in 1998 and 2002, respectively, and little demographic information for the
195 samples exists, this is further evidence that our definitions of subspecies of *T. pallidum* may need
196 periodic revisiting.

197 We also found that four Japanese samples that were clinically diagnosed as syphilis but,
198 based on our whole genome analysis, appear to be part of the TEN subspecies, sharing an
199 ancestor with the canonical TEN genomes IraqB and BosniaA. The observation of TEN samples
200 following a syphilis diagnosis has been previously reported in Japan (29), Cuba (30), and France
201 (presumed to have been contracted in Pakistan) (31). While both the previously discovered and
202 new Japanese TEN samples are resistant to azithromycin via the canonical mutation in the 23S
203 rRNA alleles, neither the Cuban nor French samples are resistant. Furthermore, three of the four
204 samples reported herein were collected from individuals with diverse travel histories (China,
205 Japan, and the Philippines), suggesting that a sexually transmitted TEN outbreak may be even
206 more widespread than previously suspected.

207 As a mechanism to begin cataloging diversity of several genes known to be polyallelic,
208 we also examined the multi-locus sequence type (MLST) types (32) of all TPA samples using
209 whole genome sequence as well as the combined MLST. Figure 1D highlights the six most
210 common MLST at each locus for TPA samples and whether the overall subtype had been
211 previously reported. Complete MLST data can be found in Supporting Information 1. Across all
212 238 TPA samples, we found a total of 15 unique complete *tp0136* sequences, including six not
213 previously reported in the MLST database, which contains 26 alleles. Seventeen unique *tp0548*
214 sequences were found, including seven novel sequences, relative to 58 known alleles. All five
215 observed *tp0705* alleles had been previously reported. In total, excluding the 13 samples that
216 were indeterminate at any of the three loci, we found a total of 40 unique haplotypes, including

217 22 not previously reported in the MLST database. Overall, 88 of 225 *T. pallidum* subsp. *pallidum*
218 samples had a novel overall haplotype, including at least one sample from each country from
219 which samples were obtained, and all 76 (100%) from Madagascar, underscoring the importance
220 of wide geographic sampling to catalog the diversity of TPA strains.

221

222 **Putative recombination shapes modern *T. pallidum* subsp. *pallidum* genomes but remains a
223 rare event**

224 Although *T. pallidum* does not have any known plasmids or infecting phages,
225 recombination has nonetheless been shown to be an important mechanism by which genetic
226 diversity may be generated in this pathogen (27,33,34). In particular, the *tpr* family of paralogs is
227 thought to have arisen through gene duplication (28,35); for this reason, all *tpr* genes have been
228 masked for all analyses in this study. Figure 2A shows the comparison of ML tree topology
229 between genomes that have been recombination masked (left) or unmasked (right) (with tip order
230 included in Supporting Information 2). Although no samples were classified to different
231 subclades when recombinant loci were not masked, the overall tree topology was altered.
232 Notably, the SS14 Omega node had more distinct subclades in the absence of recombination
233 masking, suggesting that much of the diversity within SS14 Omega is due to recombination
234 rather than mutation. Furthermore, the Nichols B subclade of Madagascar samples becomes the
235 outgroup within the Nichols clade in the absence of recombination masking.

236 The method of recombination detection we employed relies on identification of an
237 increased density of SNPs per sliding window throughout the clonal frame rather than
238 identification of a discrete donor for each putative recombination event. Although previous
239 analyses have found more recombination in the Nichols clade than SS14 (33), our use of more

240 than 90 clinical specimens belonging to multiple Nichols subclades, albeit with a geographic
241 bias, likely provides a more complete picture of the evolutionary processes that shaped the
242 Nichols clade. In spite of the number of samples examined, recombination remained a rare event
243 in Tpr-masked genomes. Of the 474 nodes on the ML tree, including 238 tips and 236 internal
244 nodes, only 27 branches with recombination were detected. Sixteen of these were on internal
245 nodes and 11 on extant. Of the extant recombination events, four were detected in the 101
246 Nichols clade samples, and seven of 137 in the SS14 clade samples, suggesting no clade-specific
247 differences in recombination ($p=0.7633$, Fisher's Exact test).

248 Figure 2B highlights the positions of identified recombinant regions in the aligned
249 genomes, with grey blocks corresponding to recombination that occurred during the separation
250 of SS14 and Nichols clades, and colored blocks corresponding to recombination events that
251 occurred during the evolution of individual subclades. The grey and red striped block represents
252 a second recombination event in the SS14 Mexico clade that occurred in the same region as the
253 ancestral recombination. As has been previously reported (33,36), many of the identified
254 recombinant regions correspond to the most diverse genes in *T. pallidum*, such as *tp0136*, *tp0326*
255 (*BamA*), and *tp0515* (*LptD*) (Supporting Information 2). Notably, many of the ORFs identified as
256 recombinant encode proteins that are predicted to be at least partially surface exposed, and
257 therefore the increased SNP density may represent either bona fide recombination or selective
258 pressure of the host immune system on non-recombinant genes.

259 Recombination events specific to each subclade shown in Figure 2B were examined, with
260 representative data in Figure 2C-F for SS14 Mexico, Nichols A, Nichols B, and Nichols E
261 subclades, respectively. Windows of approximately 60 bases of the alignments of putative
262 recombinant regions are shown, and include additional *T. pallidum* species members TPE, TEN,

263 and the *T. pallidum* Fribourg-Blanc treponeme, recently proposed to be reclassified as a TPE
264 strain, due to its genetic similarity to other yaws strains (37), with non-identical nucleotides
265 highlighted. Interestingly, several of the identified recombinant loci, including Block G in SS14
266 Mexico, Block F in Nichols A, and Blocks E and L in Nichols E, have sequences identical to
267 those found in all 6 TEN genomes included in Figure 1. TEN or TPE sequences have been found
268 previously in several Nichols clade samples, suggesting prior recombination (33,36). However,
269 our markedly extended phylogeny of the Nichols clade suggests that recombination between
270 TPA and TEN has independently occurred on multiple occasions. This demonstrates that inter-
271 subspecies recombination continues to play an important role in the diversification of *T. pallidum*
272 subspecies.

273

274 ***T. pallidum* subsp. *pallidum* subclades have different rates of SNP accumulation**

275 The evolutionary history of TPA has been a point of considerable debate in recent years,
276 particularly in light of new evidence that could not exclude the presence of TPA in Northern
277 Europe in the late 15th century, casting doubt on the popular theory that venereal syphilis was
278 introduced to Europe by the returning Columbian expeditions (36). In order to determine the date
279 of the most recent common ancestor (MRCA) of the samples included in our study, we first
280 analyzed the temporal signal present among TPA strains by regressing the root-to-tip distances in
281 the SNP-only maximum-likelihood tree (Figure 3A). The left panel shows this calculation
282 performed on a tree that included 11 highly passaged laboratory strains (eight in Nichols clade
283 and three in SS14 clade), identified by open circles, while the right panel is based on a tree that
284 excluded laboratory strains. Notably, the negative slope seen for the Nichols clade appears to be
285 due to the presence of laboratory strains. This is consistent with accelerated accumulation of

286 SNPs during routine passage of the laboratory strains for decades between collection and
287 sequencing. Therefore, laboratory strains were excluded from further dating analysis.

288 We were curious as to why the Pearson correlation coefficients of the SS14 and Nichols
289 clades (0.200 and 0.023, respectively) were so poor even in the absence of laboratory strains, and
290 hypothesized that this may be due to differences inherent to the polyphyletic structure of both
291 clades. We tested this by plotting the residuals of the regression by subclade and found
292 significant differences between groups (Figure 3B, $p < 2e^{-16}$, ANOVA), suggesting that rates of
293 SNP accumulation may differ across the TPA phylogeny.

294 Therefore, we proceeded to Bayesian ancestral reconstruction and dating of clinical
295 specimens by BEAST 2 (38), using an uncorrelated relaxed clock with a starting rate of 3.6×10^{-4}
296 (24,39) as a prior model to account for differences in rates of mutation in different branches of
297 the tree. Figure 3C shows the dated Bayesian phylogeny, with branches colored to reflect the rate
298 of SNP accumulation. Black nodes have a posterior probability of >95%. Consistent with
299 previous studies (24,36,39), we dated the MRCA of TPA to 1717 (95% HPD 1543-1869), the
300 Nichols clade to 1893 (1839-1940), and the SS14 clade to 1921 (1868-1964), and found that the
301 rates of SNP accumulation on branches with >95% posterior probability ranged between 0.2 and
302 0.73 fixed SNPs/year. The inset figure shows the mean rates of diversification on branches with
303 >95% posterior support for each tip, supporting our hypothesis that different subclades have
304 different rates of mutation ($p < 2e^{-16}$, ANOVA).

305

306 **Host immune pressure drives mutation in the same putative antigens in SS14 and Nichols**
307 **clades**

308 Observed differences in accumulation of SNPs among subclades may represent the
309 effects of sampling bias or bottlenecks or may reflect differences in the underlying biology. To
310 examine the functional differences that define each subclade (including loci identified as
311 recombinant (Figure 2), we used augur (40) to reconstruct the ancestral nodes identified in the
312 recombination-masked ML phylogeny, transferred ORF annotations from the TPA reference
313 genome NC_021508.1, and translated each ORF to detect coding changes. Figure 4A shows all
314 nodes used for these analyses, with subclade tips collapsed for simplicity. All coding changes
315 detected in Node 101 (SS14 clade ancestral) relative to Node 001 (Nichols clade ancestral,
316 considered equivalent to the TPA root node for these calculations) are shown; data for all
317 additional parent-child node pairs are included in Supporting Information 3. Forty-nine of 1002
318 putative ORFs were altered between SS14 and Nichols ancestral nodes, with a total of 134 non-
319 synonymous mutation events. We defined a mutation event as a single amino acid change,
320 insertion/deletion, or frameshift. We did not separately include the effects of putative
321 recombination events because we did not attempt to formally characterize recombination donors,
322 and therefore could not disentangle the effects of recombination from selective pressure driving
323 increased mutation.

324 We next attempted to define functional changes between the SS14 and Nichols clades by
325 examining overrepresentation of altered loci in categories annotated by structural similarity (41).
326 We used the annotation of the Protein Data Bank (PDB) structure of the highest scoring model,
327 with a confidence cutoff of 75%, allowing 798 coding sequences (CDSs) to be assigned to a total
328 of 62 unique PDB categories. We then performed Fisher's exact tests to test for
329 overrepresentation of altered proteins in each category. For SS14 vs Nichols ancestral nodes (101
330 vs 001, Supplementary Figure 1), we only found significant overrepresentation in a single

331 category, “Signaling Protein”, with 3 (*tp0073*, *tp0640*, and *tp0995*) loci out of the 16 in the
332 category altered. However, because these annotations are by structural similarity rather than
333 known function, it is likely that testing for overrepresentation of structural annotations does not
334 fully capture the functional differences between any two clades.

335 Because functional annotation of *T. pallidum* proteins is still hampered by the absence of
336 a reverse genetics system, we chose next to focus on alteration of proteins known or suspected to
337 interact with the host immune system. We included proteins that reacted with pooled sera from
338 individuals with known syphilis infection (42,43) or otherwise known to be surface-exposed
339 (Supporting Information 3 - Antigens) and again performed overrepresentation tests (Figure 4B).
340 Along branches with more than 10 altered proteins, only two nodes (N015, Nichols C, and N005,
341 Nichols D Lab Strains) did not have significant p values ($p < 0.05$) relative to their parent. When
342 examining individual mutation events in nodes with more than four altered proteins, mutation in
343 antigenic proteins represents more than 30% of the amino acid variability in more than half of
344 nodes, and at least 10% in all nodes (Figure 4C). Antigenic proteins are enriched among proteins
345 that become mutated relative to their parent node in multiple subclades, representing separate
346 events (Figure 4D). Furthermore, among antigens that were mutated relative to the parent node in
347 more than one subclade, none was exclusive to either the SS14 or Nichols clade. These data
348 suggest that interaction with the host immune system drives a large proportion of the evolution
349 of both major clades of this pathogen, either via individual SNPs or horizontal gene transfer.

350 However, although antigens are enriched for non-synonymous mutations relative to the
351 rest of the proteome, mutation of non-antigenic proteins may make considerable contributions to
352 *T. pallidum* pathogenicity and immune interaction. When examining proteins whose mutation
353 was unique to a single clade (Figure 4D), we found a C394F mutation in the ERCC3-like DNA

354 repair helicase TP0380 (44) only in the Nichols A subclade, which had a much higher median
355 rate of SNP accumulation than any other subclade (Figure 3C). It is plausible that mutation of
356 this helicase compromises DNA repair and contributes to a more rapid rate of evolution within
357 this clade.

358

359 **Predicted structural changes of putative surface proteins not limited to polymorphic**
360 **residues**

361 For any protein, multiple independent mutation events along several branches of the TPA
362 phylogeny strongly suggest the protein is under selective pressure. Of the six proteins that
363 undergo mutation along four or more of the 14 branches in the phylogeny (Figure 4D and
364 Supporting Information 3 – Heatmap), five (TP0136, TP0326, TP0548, TP0966, and TP0967)
365 are known to be antigenic. TP0326, TP0548, TP0966, and TP0967 are likely outer membrane
366 proteins based on their homology to *N. gonorrhoeae* BamA (TP0326) and *E. coli* FadL (TP0548)
367 and TolC (TP0966, TP0967) (45) and reviewed in (46). TP0136 is a lipoprotein that appears to
368 be localized to the outer membrane, where it functions as a fibronectin- and laminin-binding
369 adhesin (47–49). To date, recombinant TP0136 and TP0326 have been tested as potential
370 vaccine candidates in rabbits, with TP0136 delaying ulceration but not providing full protection
371 upon challenge (47), and TP0326 providing partial protection in some studies (50,51), while not
372 protective in others (52). Although antigens harboring polymorphisms would not traditionally be
373 considered viable vaccine candidates, the paucity of outer membrane proteins in *T. pallidum* (46)
374 demands evaluation of imperfect candidates.

375 Accordingly, for the five most frequently mutated putative outer membrane antigens, we
376 developed models that highlight the positions predicted to undergo the most structural change

377 upon mutation, including those at orthogonal sites. We first performed global alignments of
378 sequence variants for each of the five proteins using hhpred (53–55) (Supplementary Figures 2-6,
379 panel A, and Supporting Information 4). We then generated composite homology models of the
380 SS14 variant using RosettaCM (56) guided by hhpred sequence alignment. Ribbon structures and
381 surface contours with highlighted polymorphic residues of the SS14 variant are shown in
382 Supplementary Figures 2-6, panels B and C, respectively. Then, the SS14 model was used as a
383 template for predicting the structure of variants of other strains. We performed a global
384 superposition of variant structures and computed an average per-atom displacement relative to
385 the reference model (taking sequence changes into account, see Methods). The resulting per-
386 atom deviations were then mapped onto the model of the SS14 variant, with blue representing
387 regions of the lowest displacement from the SS14 model and red the highest (Supplementary
388 Figures 2-6, panel D). This approach allowed detection of structural changes not simply at the
389 site of the polymorphism, but also orthogonal changes due to disruption of hydrogen and other
390 bonds. Furthermore, it allows “tuning” of the structural effect of a mutation on each atom, with
391 the mutation of similar residues (such as leucine to isoleucine) resulting in less displacement of
392 each atom than substitution of dissimilar residues (such as arginine to histidine). N terminal
393 residues comprising predicted secretion sequences are not shown for TP0136 (48) or TP0326
394 (57). Best estimates for Gram negative signal peptides were predicted by SignalP 5.0 (58) for
395 TP0548, TP0966, and TP0967 SS14 variants and excluded from display.

396 In spite of slightly different approaches employed in their generation and our use of the
397 SS14 variant rather than Nichols, our structural models for TP0326, TP0548, TP0966, and
398 TP0967 generally agree with the models recently proposed by Hawley *et al.* (45). TP0326 is a
399 large multidomain component of the β -barrel Assembly Complex (BAM) and includes a C-

400 terminal β -barrel. Consistent with previous studies (34,45,57,59), we found that extracellular
401 loop (ECL)-4 and the serine-rich tract of ECL-7 contribute to much of the between-strain
402 structural diversity (arrows and single arrowheads, respectively, Supplementary Figure 2B-D).
403 We also found that the large ECL-3 (double arrowheads, Supplementary Figure 2B-D) had nine
404 polymorphic residues, rendering the entire exposed surface of the protein variable due to strain-
405 to-strain variation in ECLs, particularly 3, 4, and 7 (Supplementary Figure 2C-D).

406 In contrast to TP0326, the structure and function of which has been studied extensively,
407 less is known about TP0548, a predicted homolog of the *E. coli* fatty acid transporter FadL. We
408 predict the structure to be a 14-stranded β -barrel, with periplasmic C-terminal α -helices,
409 consistent with previous studies (45). Prediction of linear B cell epitopes (BCEs) using BepiPred
410 2.0 (60) revealed that, depending on the as-yet unknown position of the cleavage of the N
411 terminal signal sequence, up to four linear BCEs eight residues or longer are predicted to occur
412 in invariant, extended host-facing loops at the N-terminus and ECL-2 (Supplementary Figure 3A,
413 arrows in Supplementary Figure 3B show the relevant loops), rendering them potentially of use
414 in a vaccine cocktail. Notably, the displacement seen in ECL-2, containing BCEs 3 and 4,
415 (Supplementary Figure 3D, arrow) is most likely due to stochastic differences in predicting the
416 conformation of the flexible loop rather than true structural variation.

417 Both TP0966 and TP0967 are predicted to be orthologs of the *E. coli* efflux pump TolC
418 (45), and are predicted to have a tri-partite structure, with each monomer contributing four β -
419 strands to β -barrel that spans the outer membrane with BCEs predicted within the ECLs (45).
420 Supplementary Figures 4B and 5B highlight a single monomer for TP0966 and TP0967,
421 respectively; both ECLs of TP0966, and ECL1 of TP0967, contain polymorphic residues that
422 disrupt predicted linear B cell epitopes (Supplementary Figures 4A and 5A, Supplementary

423 Figures 4C and 5C, arrows). For both TP0966 and TP0967, the residues with the most
424 displacement that disrupt the extracellular surface are not the polymorphic positions
425 (Supplementary Figures 4D and 5D, arrows). Rather, in TP0966, the polymorphic charged
426 residues in and adjacent to the ECLs may cause changes to electrostatic interactions that
427 influence loop position. In TP0967, the length of the poly-glycine tract alters the position of
428 ECL1. The likely result is disruption of the conformational epitopes formed by the surface loops
429 in TP0966 and TP0967.

430 Finally, we generated a structural model of TP0136, and found it to adopt a 7-bladed
431 beta-propeller fold in its N-terminal domain, followed by a relatively unstructured C-terminal
432 domain (Supplementary Figure 6B). The beta-propeller structure is noteworthy as it is
433 homologous to structures found in several eukaryotic integrins that mediate binding to the
434 extracellular matrix (61), as well as to bacterial lectins (62). Several tracts of serine and lysine
435 repeats are a unique structural feature of TP0136; the beta-propeller fold of TP0136 allows these
436 intrinsically disordered regions to form unstructured loops between beta strands. Unsurprisingly,
437 the surfaces that comprise the β -strands have some polymorphisms (Supplementary Figure 6B
438 and C, boxed region) but they are not predicted to cause extensive structural displacement and
439 disruption of the fold, as shown by primarily blue coloring in the boxed region of Supplementary
440 Figure 6D.

441 Interestingly, the deletion in TP0136 that appears in 4 sequence variants (2, 5, 22, 23, and
442 24, Supplementary Figure 6A, alignment position 161-192) and entirely removes the large
443 flexible loop annotated by an arrow in Supplementary Figure 6B-D is not found in any ancestral
444 node sequences (Figure 4), but arises independently in strains from multiple geographic
445 locations, including Nichols clade strains from Madagascar and the United States (subclades A,

446 B, and D), and SS14 clade strains from Japan, Peru, and Ireland (subclades Omega – East Asia
447 and SS14 Omega`), consistent with this genomic region being a hotspot for recombination
448 (Figure 2).

449

450 **Discussion**

451 In recent years, *T. pallidum* genomics has been significantly advanced by projects aimed
452 at studying the origin and spread of strains responsible for the modern syphilis pandemic
453 (24,26,36,39,63), as well as the emergence of azithromycin resistance (24,26,63). Increasingly
454 the challenge in *T. pallidum* genomics will be attaining complete genomic sequences from
455 undersampled regions, associating genomic sequencing with spirochete biochemical functions,
456 and gaining actionable insights into *T. pallidum* evolution that inform vaccine design.

457 With these goals, we generated 196 near-complete *T. pallidum* genomes from diverse
458 locations, including three countries – Peru, Italy, and Madagascar – with no previous complete
459 genomes publicly available. Peruvian samples (n=9) belonged exclusively to the SS14 Omega`
460 subclade, which contains samples collected worldwide and corresponds to the largest SS14 sub-
461 lineage in a recent analysis of SNPs in TPA strains (63). Eight of the ten Italian strains also
462 belonged to the Omega` subclade.

463 The remaining two Italian strains, collected in Turin and Bologna, were of two distinct
464 Nichols subclades, one of which clustered with three Japanese syphilis strains in Nichols
465 subclade C, and the other clustered with samples of Japanese and French origin, forming the
466 distantly related Nichols subclade E. Notably, none of the Japanese or Italian samples clustered
467 with the Malagasy Nichols samples, which, but for a single Cuban strain in Nichols subclade A,
468 formed two private subclades. Because the samples from Madagascar were collected between

469 2000-2007, it is unknown whether there has been introduction of additional lineages of TPA in
470 the intervening years, or whether the two nearly private subclades are reflective of the currently
471 circulating strains.

472 Sample collection date is also an important consideration to the interpretation of
473 azithromycin resistance data. None of the strains collected in the USA between 1998 and 2002
474 were resistant to azithromycin. However, this was prior to the detection of widespread
475 azithromycin resistance in the United States (8); therefore, the lack of resistance detected in the
476 strains sequenced for the present study should not be considered representative of the current
477 status. Only one of the strains collected from Madagascar between 2000 and 2007 was resistant
478 to azithromycin; no subsequent sampling has been performed, thus, no conclusions about
479 azithromycin resistance in strains currently circulating in Madagascar can be drawn.

480 In our study, as in other recent global *T. pallidum* subsp. *pallidum* genomics initiatives
481 (24,63), samples were collected and sequenced based on availability rather than representing an
482 even distribution based on global burden of disease. The result of this is that, although we gained
483 a broader picture of worldwide diversity, some regions (North America, western Europe, eastern
484 Asia) continue to be overrepresented, while other regions (Africa – particularly Sub-Saharan
485 Africa, which bears the largest share of cases worldwide – and South Asia and South America)
486 are still vastly under-sampled. However, an important takeaway from our study as well as the
487 recent paper from Beale *et al.* (63) is that the general understanding that SS14 represents the vast
488 majority of circulating strains may require revisiting. Although the island nation of Madagascar
489 is unlikely truly representative of the diversity of strains currently circulating in Sub-Saharan
490 Africa, particularly because the samples are 15-20 years old, our finding that 99% of Malagasy
491 strains belong to the Nichols clade, coupled with Beale *et al.*'s discovery of Nichols strains

492 circulating in Zimbabwe and South Africa (63) strongly suggests widespread circulation of
493 Nichols clade TPA in Africa. Clearly, increased sampling must be a priority to enable
494 understanding of syphilis epidemiology in Africa, and to ensure a vaccine covers strains
495 circulating in the regions most hard hit by the modern pandemic.

496 Our temporal analysis generally agreed with previous estimates of mutation rate (39,63)
497 in spite of the fact that we used a relaxed, rather than fixed, clock model to determine whether
498 there were differences in the rate of mutation along different branches of the *T. pallidum* subsp.
499 *pallidum* phylogeny, which could indicate either different selection pressures or underlying
500 biological differences contributing to the phenotype. Indeed, we found significant differences in
501 the rates of mutation among the subclades (Figure 3C). The Nichols A subclade was particularly
502 interesting to us, given its high median rate of mutation along branches within the subclade with
503 high posterior support. Notably, when we examined the non-synonymous mutations that defined
504 the Nichols A subclade relative to its ancestral node, shared by Nichols subclades A, B, and C
505 (Figure 4A/D), we found that one of the non-synonymous mutations found only within the
506 Nichols A subclade was in TP0380, a putative ERCC3-like DNA repair helicase that interacts
507 with DNA replication machinery by yeast two-hybrid analysis (41,44,64). Although the
508 functional significance of the C394F mutation (C1181A in *tp0380*) is unknown, it is tempting to
509 speculate that it may directly affect DNA repair. This hypothesis of a potential mutator
510 phenotype in *T. pallidum* can now be examined *in vitro*, given the recent description of the first
511 genetic transformation in *T. pallidum* (65). If TP0380 mutation is indeed responsible for the
512 elevated rate of mutation seen within the Nichols A subclade, the implications for vaccine design
513 may be significant.

514 By definition, an effective syphilis vaccine needs to protect against most strains
515 circulating where the vaccine is administered. Our work further supports that the majority of
516 non-synonymous mutations that define *T. pallidum* subsp. *pallidum* subclades are in proteins
517 putatively located in the outer membrane, or known to react with serum from syphilis patients
518 (Figure 4) (42,43). These data, along with recent structural modeling of *T. pallidum* outer
519 membrane proteins showing that putative B cell epitopes are primarily found on the protein
520 surface predicted to face the host (45), strongly suggest that immune pressure is the most
521 important driver of mutation in *T. pallidum* subsp. *pallidum*. Indeed, our own structural
522 modeling, which highlights regions with the highest structural displacement due to sequence
523 variability, confirms that the regions with the highest displacement are frequently polyallelic
524 (Supplementary Figures 2-6). Given the paucity of *T. pallidum* outer membrane proteins, and the
525 extensive mutation of predicted epitopes, a multivalent vaccination strategy may engender a
526 polyclonal humoral response capable of neutralizing a wider array of strains, a strategy currently
527 being adopted in our laboratory.

528 Finally, an important caveat to these data is that, due to their extensive recombination and
529 duplication, we excluded arguably the most important *T. pallidum* proteins that interact with the
530 host immune system, the Tpr family (14,28). Although this approach has been used before to
531 ensure an accurate phylogeny free from the confounding effects of recombination (24,63), as
532 well as to prevent mistakes due to improper resolution of their repetitive elements during de
533 novo assembly (39), an understanding of how the *tpr* genes evolve and influence host immunity
534 is critical to developing an efficacious vaccine to *T. pallidum*. Accordingly, we are currently
535 undertaking additional analyses of the Tpr family in these strains, including the hypervariable
536 regions of TprK.

537 The data presented in this study represent a step forward toward developing a successful
538 vaccine against syphilis. Alongside increased sequencing of strains from regions without
539 extensive sampling, particularly Africa and South America, improved biophysical and
540 computational methods are necessary to unequivocally determine which proteins are expressed
541 on the surface of the bacterium during human infection. The new system to genetically engineer
542 *T. pallidum* (65) will undoubtedly aid these studies, as well as allow the development of strains
543 to test vaccine candidates in animal experiments. Finally, a successful vaccine must not only be
544 efficacious against all circulating strains, but must also be sufficiently low cost and robust to
545 ambient temperatures to allow distribution in the developing world, which is currently bearing
546 the burden of the modern pandemic.

547

548 **Methods**

549 **Ethics Statement:** All human samples were collected and deidentified following protocols
550 established at each institution. Samples from Ireland, Madagascar, and USA have been
551 previously published (18–22). IRB protocol numbers for collection of the remaining samples are
552 as follows: China: Nanjing Medical University, 2016□050; Italy: Universities of Turin and
553 Genoa, PR033REG2016, University of Bologna, 2103/2016; Japan: National Institute of
554 Infectious Diseases, 508 and 705; Peru: University of Southern California, HS-21-00353; Papua
555 New Guinea, Lihir Medical Center, Medical Research Advisory Committee of the PNG NDOH
556 No: 17.19. Sequencing of deidentified strains was covered by the University of Washington
557 Institutional Review Board (IRB) protocol number STUDY00000885.

558 **Library Preparation:** Samples were collected and DNA extracted using standard protocols (66).
559 Treponemal burden was assessed by quantitative PCR (qPCR) for *TP47* multiplexed with human

560 β-globin, using primer sequences *TP47-F*: 5'-CAAGTACGAGGGAACATCGAT, *TP47-R*: 5':
561 TGATCGCTGACAAGCTTAGG, *TP47-probe*: 5'-6FAM-
562 CGGAGACTCTGATGGATGCTGCAGTT-NFQMGB. Pre-capture libraries were prepared
563 from up to 100 ng input genomic DNA using the Kapa Hyperplus kit (Roche), using a
564 fragmentation time of 8 minutes and standard-chemistry end repair/A-tailing, then ligated to
565 TruSeq adapters (Illumina). Adapter-ligated samples were cleaned with 0.8x Ampure beads
566 (Beckman Coulter) and amplified with barcoded primers for 14-16 cycles, followed by another
567 0.8x Ampure purification.

568 ***T. pallidum* capture:** Capture of *T. pallidum* genomes was performed according to Integrated
569 DNA Technology's (IDT's) xGen Hybridization Capture protocol. Briefly, pools of 3-4 libraries
570 were created by grouping samples with similar treponemal load for a total of 500 ng DNA, and
571 Human Cot 1 DNA and TruSeq blocking oligos (IDT) added prior to vacuum drying. The
572 hybridization master mix, containing biotinylated probes from a custom IDT oPool tiling across
573 the NC_010741.1 reference genome, was then added overnight (>16 hr) at 65C. The following
574 day, streptavidin beads were added to the capture reaction, followed by extensive washing, 14-16
575 cycles of post-capture amplification, and purification with 0.8x Ampure beads. Pool
576 concentration was determined by Qubit assay (Thermo Fisher) and size verified by Tapestation
577 (Agilent). Libraries were sequenced on a 2x150 paired end run on a HiseqX.

578 **Fastq processing:** Fastqs were processed and genomes assembled using our custom pipeline,
579 available at https://github.com/greninger-lab/Tpallidum_WGS. Paired end reads were adapter-
580 and quality-trimmed by Trimmomatic 0.35 (67), using a 4 base sliding window with average
581 quality of 15 and a minimum length of 20, retaining only paired reads. Trimmed reads were then
582 filtered with bbduk v38.86 (68) in two separate steps. First, reads were filtered very stringently,

583 allowing removal of contaminating non-*T. pallidum* reads, against a reference containing the two
584 rRNA loci, with a 100 bp 5' and 3' flank, from each of five reference *T. pallidum* genomes
585 (NC_021508.1 (*T. pallidum* subsp. *pallidum* strain SS14), NC_016842.1 (*T. pallidum* subsp.
586 *pertenuis* strain SamoaD), NC_016843.1 (*T. pallidum* subsp. *pertenuis* strain Gauthier),
587 NC_021179 (*T. pallidum* strain Fribourg-Blanc treponeme), NZ_CP034918.1 (*T. pallidum*
588 subsp. *pallidum* strain CW65)). We used a kmer size of 31, a Hamming distance of 1, a
589 minimum of 98% of kmers to match reference, and removal of both reads if either does not pass
590 these criteria. Second, unmatched reads from the rRNA filtration step were then filtered against
591 the complete reference genomes that had been masked with N at the rRNA loci, using a kmer
592 size of 31 and a Hamming distance of 2. Matching reads from the two steps were concatenated
593 and used for input for genome mapping and assembly.

594 **Genome assembly:** Filtered reads were mapped to the *T. pallidum* street 14 reference genome,
595 NC_021508.1, using Bowtie2 v2.4.1 (69) with default parameters and converted to bam with
596 samtools v1.6 (70), followed by deduplication by MarkDuplicates in Picard v2.23.3 (71). Prior to
597 *de novo* assembly, rRNA-stripped reads were filtered with bbduk (68) to remove repetitive
598 regions of the genome, including the repeat regions of the *arp* and *TP0470* genes, as well as
599 *tprC*, *tprD*, and the *tprEGF* and *tprIJ* loci, using a pseudo-kmer size of 45 and Hamming
600 distance of 2. *De novo* assembly was performed using Unicycler v0.4.4 (72) using default
601 settings, with rRNA- and repetitive region-stripped paired fastqs as input. Contigs longer than
602 200 bp were then mapped back to NC_021508.1 reference genome using bwa-mem 0.7.17-r1188
603 (73) and a custom R script (74) used to generate a hybrid fasta merging contigs and filling gaps
604 with the reference genome. Deduplicated reads were initially remapped to this hybrid using
605 default Bowtie2 settings, local misalignments corrected with Pilon v1.23.0 (75), and a final

606 Bowtie2 remapping to the Pilon consensus used as input to a custom R script (74) to close gaps
607 and generate a final consensus sequence, with each position called at a threshold of 50% of reads
608 supporting a single base. A minimum of six reads were required to call bases; coverage lower
609 than 6x was left ambiguous by calling “N”. All steps of genome generation were visualized and
610 manually confirmed in Geneious Prime v2020.1.2 (76). Following consensus generation, the
611 tRNA-Ile and tRNA-Ala sites that occur within the rRNA loci were masked to N due to short
612 reads being unable to resolve the order of the sites. Consensus genomes were further masked at
613 the *arp* and *tp0470* repeats and *tprK* variable regions prior to further analysis and deposition in
614 the NCBI genome database.

615 **Phylogeny:** Consensus genomes that had been masked at the *arp* and *tp0470* repeats, intra-rRNA
616 tRNAs, and *tprK* variable regions were further masked to N at all *tpr* genes, which are known to
617 be recombinogenic (35). Masked genomes were aligned with MAFFT v7.271 (77) with a gap
618 open penalty of 2.0 and an offset (gap extension penalty) of 0.123. Aligned genomes were
619 recombination masked using 25 iterations of Gubbins v2.4.1 (23). Recombination masking was
620 performed separately with and without *T. pallidum* subsp. *pertenue* and *T. pallidum* subsp.
621 *endemicum* sequences as appropriate. Recombination masking was mapped back onto whole
622 genome sequences and visualized using maskrc-svg v0.5 (78), and iqtree v2.0.3 (79) used to
623 generate a whole genome maximum likelihood phylogeny using 1000 ultrafast bootstraps and
624 automated selection of the best substitution model. A non-recombination-masked maximum
625 likelihood tree was generated using the same parameters but with the raw MAFFT output.
626 Sequences of *tp0136*, *tp0548*, and *tp0705* were extracted and batch queried using the PubMLST
627 database ((32), accessed 01-22-2021).

628 **Bayesian Dating:** TempEst v1.5.3 (80) was used to calculate root-to-tip distances for the SNP-
629 only maximum likelihood phylogeny calculated with or without *T. pallidum* subsp. *pallidum*
630 laboratory strains, assuming one year uncertainty in strains with collection dates estimated.
631 Regressions of distance vs sample date were performed per clade in R. Bayesian dating was
632 performed in the BEAST2 suite (38) using the recombination masked SNP-only (n=600 sites)
633 alignment of *T. pallidum* subsp. *pallidum*, excluding laboratory strains. Priors included a relaxed
634 clock lognormal model with a starting rate of 3.6×10^{-4} (24,39), constant population size, and a
635 GTR +gamma substitution model. Three separate runs, each with 100,000,000 MCMC cycles
636 were performed and the first 10,000,000 cycles discarded as burn-in. All runs converged and
637 were merged prior to calculation of the maximum clade credibility (MCC) tree.
638 **Ancestral node reconstruction:** Augur v10.1.1 (40) was first used to map all tips without
639 recombination masking onto the whole genome phylogeny generated following recombination
640 masking, ensuring appropriate ancestral relationships unconfounded by recombination, using the
641 “refine” function. The “ancestral” function was next used with default settings to infer ancestral
642 node sequences. Sequences of select nodes were aligned to reference NC_021508.1 using
643 MAFFT as above, and annotations of the reference transferred to ancestral node sequences in
644 Geneious. Pairwise global alignments of protein sequences were performed in R using the
645 Biostrings package (81), and analysis and statistical measurements performed in R using custom
646 scripts.
647 **Antigens:** Antigens were manually curated based on being reactive against *T. pallidum* subsp.
648 *pallidum* positive human sera in either of two previous studies (42,43) or, to control for low
649 expression hampering detection by these in vitro methods, by being selected as likely surface
650 proteins or lipoproteins based on extensive literature searches.

651 **Structural modeling:** Genomes were annotated using Prokka v1.14.6 (82), using the --proteins
652 flag to force annotations to comply with NC_021508.1. Translated coding sequences for vaccine-
653 relevant genes were extracted with a custom R script. Sequences containing ambiguities or
654 truncations likely due to assembly gaps in the genome were manually reviewed in Geneious and
655 excluded from further analysis.

656 Homology modelling in RosettaCM (56) was used to build initial models of the SS14
657 variant of each protein. For all sequences collected as part of this study, hhpred (53–55) was
658 used to identify homologous structures, and only those sequences with fulllength alignments
659 (covering >70% of the target) with high probability (>95% hhpred score) were considered for
660 structural modelling. Given these alignments, 100 independent modelling trajectories were
661 carried out for a reference sequence, guided by the top 1-7 templates for each target. We used
662 the following templates in modelling each target: TP0136 used 4a2l and 5oj5; TP0326 used 4k3b
663 and 5d0o; TP0548 used 6h3i; TP0966 used 1yc9, 3d5k, 4k7r, 4mt0, 4mt4, 5azs, and 6u94;
664 TP0967 used 1yc9, 3d5k, 5azp, 5azs, and 6u94. For targets TP0966 and TP0967, modelling was
665 carried out considering the complete homotrimeric configuration, using the symmetry of the
666 templates as a guide.

667 Following homology modelling, the lowest-energy model was selected and used as a
668 starting point for modelling the mutant sequences. We again used RosettaCM, providing the
669 reference model as the “template” and each mutation as the “target” sequence. For each mutant
670 sequence, three models were predicted and the lowest-energy one was used in analysis of
671 structural deviations.

672 Structural deviation analysis involved comparing the structures of proteins with different
673 sequences, and standard difference metrics (like backbone RMSd) do not properly report

674 differences in sidechain identities. Instead, we used a “per-atom RMSd” metric, where the
675 structures were first superimposed on the reference structure by aligning common backbone
676 atoms. Then, for each atom in the reference structure, the distance was computed not to a
677 corresponding atom, but rather the closest atom of the same chemical identity (e.g., the oxygens
678 of glutamate and aspartate would map to one another). This was then used to calculate the per-
679 atom and per-residue RMS deviations reported in the manuscript. In this part of the analysis, the
680 homotrimeric configuration of targets TP0966 and TP0967 was again used.

681 Bacterial signal peptide predictions were performed with SignalP 5.0 (58) Linear B cell
682 epitopes were predicted using the BepiPred-2.0 (60).

683 **Statistics and Visualization:** Unless otherwise noted, all statistical analysis was performed in R
684 v 4.0.0. Phylogenetic trees and metadata were visualized with the R packages ggtree (83), treeio
685 (84), and ggplot (85), multiple sequence alignments by R package ggmsa (86) and figures
686 generated using cowplot and Adobe Illustrator v24.1.3.

687 **Data Availability:** Paired end reads have been uploaded to the NCBI Sequencing Read Archive,
688 Bioproject PRJNA723099. Consensus genomes have been deposited to NCBI Genome,
689 accession numbers CP073381-CP073576 (Supporting Information 1 - Sample Accessions).

690

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694

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946

947 **Figure Legends:**

948 **Figure 1: Whole genome phylogeny of *T. pallidum* patient isolates.** A) Whole genomes were
949 MAFFT-aligned, recombination-mask, and maximum-likelihood phylogeny determined. Tips
950 are shown as grey triangles and nodes with >0.95 support from 1000 ultrafast bootstraps shown
951 as black circles. B) Subspecies/lineage, subclade, and continent of origin of all samples included
952 in phylogeny. C) Azithromycin sensitivity/resistance as conferred by the 23S rRNA 2058/2059
953 alleles. Data represents alleles at both rRNA loci. D) MLST subtypes, including novel
954 sequences, for *tp0136*, *tp0548*, and *tp0705*, as well as whether the three alleles constitute a
955 known or novel MLST. Top 6 most abundant sequences at each locus are colored, while other
956 less abundant known and novel sequences are grouped and colored in light and medium grey,

957 respectively. Sequences containing N bases are denoted as indeterminate and shown in dark
958 grey. Expanded metadata for all samples is included in Supporting Information 1.

959

960 **Figure 2: Effect of recombination on *T. pallidum* subsp. *pallidum* evolution. A)**

961 Recombination-masked (left) and unmasked (right) phylogenies, with equivalent subclades
962 highlighted. Relative position of each tip is traced between the two panels. B) Putative
963 recombinogenic regions in each clade. Genomic position is relative to the length of the MAFFT
964 alignment. Consensus alignment of all tips is shown on the grey panel, with recombination
965 blocks lettered above. Grey blocks represent recombination that occurred during evolution of the
966 SS14 clade. Red and blue blocks represent recombination events unique to each clade. Mixed
967 grey and colored blocks are regions of ancestral recombination that had a second event unique to
968 that clade. C-F) Two example regions of recombination in SS14 Mexico (C), Nichols A (D),
969 Nichols B (E), and Nichols E (F). Genomic position of the first divergent base in the window
970 shown are shown with NC_021508.1 numbering.

971

972 **Figure 3: SS14 and Nichols subclades have different rates of SNP accumulation. A)** Linear
973 regressions for recombination-masked root-to-tip distances from maximum likelihood phylogeny
974 as a function of year of collection, including (left) or not including (right) highly passaged
975 laboratory strains. B) Residuals from linear regression without laboratory strains were plotted per
976 subclade, $p < 2e^{-16}$, ANOVA. C) Bayesian maximum clade credibility tree showing mean
977 common ancestor heights. Highlighted nodes have a posterior probability of >0.95, and branch
978 colors reveal rate of change (SNPs per genome per year). Ages and 95% highest posterior
979 density are included for nodes of interest including the TPA, SS14, and Nichols ancestral nodes,

980 as well as those of each subclade. Inset: For each tip, mean rates of SNP accumulation along
981 branches with >0.95 posterior probability were plotted per subclade, $p < 2e^{-16}$, ANOVA.

982

983 **Figure 4: Coding mutations in the *T. pallidum* subsp. *pallidum* phylogeny.** A) Whole genome
984 ML phylogeny of TPA, with tips collapsed to the subclade node. Open reading frames of
985 inferred ancestral sequences for each node were annotated based on the SS14 reference sequence
986 NC_021508. Coding mutations, including for putative recombinant genes, for each child node
987 were determined relative to its parent node (complete list in Supporting Information 4). Loci
988 with amino acid differences (n=49 loci, n=134 individual AA mutation events) in the SS14
989 ancestral clade node (N101) are shown relative to the Nichols ancestral node (N001). B)
990 Positions are equivalent to those shown in A. Black square represents the Nichols Ancestral
991 Node (N001). Number of antigens with coding mutations on each child node relative to parent
992 node. Color represents p value of for overrepresentation by Fisher's Exact test of antigens among
993 all mutated proteins per branch; those in grey have a p value > 0.05. C) Percentage of total
994 individual mutation events per branch. Raw numbers of mutation events in antigens per total
995 mutation events are shown for each branch. D) Tile plot showing mutated proteins in the
996 ancestral node for each subclade relative to its parent node, colored by antigen or not. Proteins
997 are arranged by number of subclades bearing mutations. Data is recapitulated in Supporting
998 Information 3

999

1000 **Supplementary Figure 1: Distribution of putative protein functional annotation based on**
1001 **high-confidence Phyre2 models.** Percent of proteins in each category different between the
1002 SS14 ancestral clade node (N101) and the Nichols ancestral node (N001) (orange) were

1003 compared to annotations across the whole genome (purple). Overrepresentation was tested by
1004 Fisher's exact test, $*p < 0.05$.

1005

1006 **Supplementary Figure 2: Multiple sequence alignment and structural modeling of TP0326.**

1007 A) Multiple sequence alignment for all amino acid sequence variants. Polymorphic residues are
1008 highlighted, and positions of extracellular loops 3, 4, and 7 are shown. B) Side (left) and top
1009 (right) cartoon representation of TP0326, with a color gradient between blue at the N-terminus to
1010 red at the C-terminus. C) Side (left) and top (right) space-filling representation of TP0326, with
1011 polymorphic residue positions colored magenta. D) Side (left) and top (right) space-filling
1012 representation of TP0326, with atoms colored by average per atom displacement in all variants
1013 relative to the SS14 reference sequence. Arrows, single arrowheads, and double arrowheads
1014 point to positions of ECLs 4, 7, and 3, respectively.

1015

1016 **Supplementary Figure 3: Multiple sequence alignment and structural modeling of TP0548.**

1017 A) Multiple sequence alignment for all amino acid sequence variants. Polymorphic residues are
1018 highlighted, and positions of relevant predicted B cell epitopes are shown. B) Side (left) and top
1019 (right) cartoon representation of TP0548, with a color gradient between blue at the N-terminus to
1020 red at the C-terminus. Arrows point to the flexible loops that contain predicted linear BCEs. C)
1021 Side (left) and top (right) space-filling representation of TP0548, with polymorphic residue
1022 positions colored magenta. D) Side (left) and top (right) space-filling representation of TP0966,
1023 with atoms colored by average per atom displacement in all variants relative to the SS14
1024 reference sequence. Arrow points to ECL-2, which contains two invariant predicted BCEs.

1025

1026 **Supplementary Figure 4: Multiple sequence alignment and structural modeling of TP0966.**

1027 A) Multiple sequence alignment for all amino acid sequence variants. Polymorphic residues are
1028 highlighted. ECLs 1 and 2 are boxed, and the linear BCEs contained in the SS14 variant (#1) are
1029 marked in red. B) Side (left) and top (right) cartoon representation of TP0966, with a color
1030 gradient between blue at the N-terminus to red at the C-terminus. C) Side (left) and top (right)
1031 space-filling representation of TP0966, with polymorphic residue positions colored magenta.
1032 Arrow points to polymorphic residues in surface loops. D) Side (left) and top (right) space-filling
1033 representation of TP0966, with atoms colored by average per atom displacement in all variants
1034 relative to the SS14 reference sequence. Arrows point to the high displacement, non-
1035 polymorphic residues.

1036

1037 **Supplementary Figure 5: Multiple sequence alignment and structural modeling of TP0967.**

1038 A) Multiple sequence alignment for all amino acid sequence variants. Polymorphic residues are
1039 highlighted. ECL1 is boxed, and the linear BCE contained in the SS14 variant (#4) is marked in
1040 red. B) Side (left) and top (right) cartoon representation of TP0967, with a color gradient
1041 between blue at the N-terminus to red at the C-terminus. C) Side (left) and top (right) space-
1042 filling representation of TP0967, with polymorphic residue positions colored magenta. Arrow
1043 points to polymorphic residues in surface loops. D) Side (left) and top (right) space-filling
1044 representation of TP0967, with atoms colored by average per atom displacement in all variants
1045 relative to the SS14 reference sequence. Arrows point to the high displacement, non-
1046 polymorphic residues.

1047

1048 **Supplementary Figure 6: Multiple sequence alignment and structural modeling of TP0136.**

1049 A) Multiple sequence alignment for all amino acid sequence variants. Polymorphic residues are
1050 highlighted. B) Side (left) and top (right) cartoon representation of TP0136, with a color gradient
1051 between blue at the N-terminus to red at the C-terminus. C) Side (left) and top (right) space-
1052 filling representation of TP0136, with polymorphic residue positions colored magenta. D) Side
1053 (left) and top (right) space-filling representation of TP0136, with atoms colored by average per
1054 atom displacement in all variants relative to the SS14 reference sequence. Boxed areas represent
1055 regions of low displacement in the β -strands. In all panels, arrows point to the large extracellular
1056 loop that is removed in variants found in several subclades.

1057

1058 **Supporting Information 1: Expanded metadata for all samples presented in Figure 1.**

1059 **Sample Statistics:** “Input Genomes” refers to the number of copies of TP47 (*tp0574*) included in
1060 pre-capture library preparation. “Coverage” refers to the average deduplicated read depth at any
1061 position in the initial mapping of reads to the reference sequence NC_021508. “Length” is the
1062 length of the final consensus genome. “Number Ambiguities” and “Percent Ambiguities” refers
1063 to Ns in samples prior to masking. **Sample Accessions:** NCBI Biosample and assembly
1064 accessions per sample. **Sample Metadata:** “Tip Number” refers to sample position on the
1065 phylogenetic tree in Figure 1, with #1 at the bottom of the page. ***tp0136 MLST, tp0548 MLST,***
1066 ***tp0705 MLST:*** *Complete* information on MLST alleles, including how novel sequences relate to
1067 the closest known match.

1068

1069 **Supporting Information 2: Expanded Recombination Data from Figure 2: Tip Order:** Top
1070 to bottom order of tips in recombination masked and unmasked phylogenies. **Recombination**

1071 **Blocks:** Genes included in each recombination block. **Blocks per Clade:** Precise locations of
1072 recombination events detected per clade.

1073

1074 **Supporting Information 3: Expanded data for Figure 4. Summary Data by Locus:**

1075 Information on number of mutations per locus per branch, including functional annotations.

1076 **Antigens:** List of loci included as antigens. CDS name and genomic position information is from

1077 NC_021508.1. **Heatmap Data:** Data included in Figure 4D. “Sum” represents the number of

1078 nodes at which there is a change in that locus. **NXXX vs NYYY:** Detailed information on all

1079 detected mutations per branch, named by parent and child node number.

1080

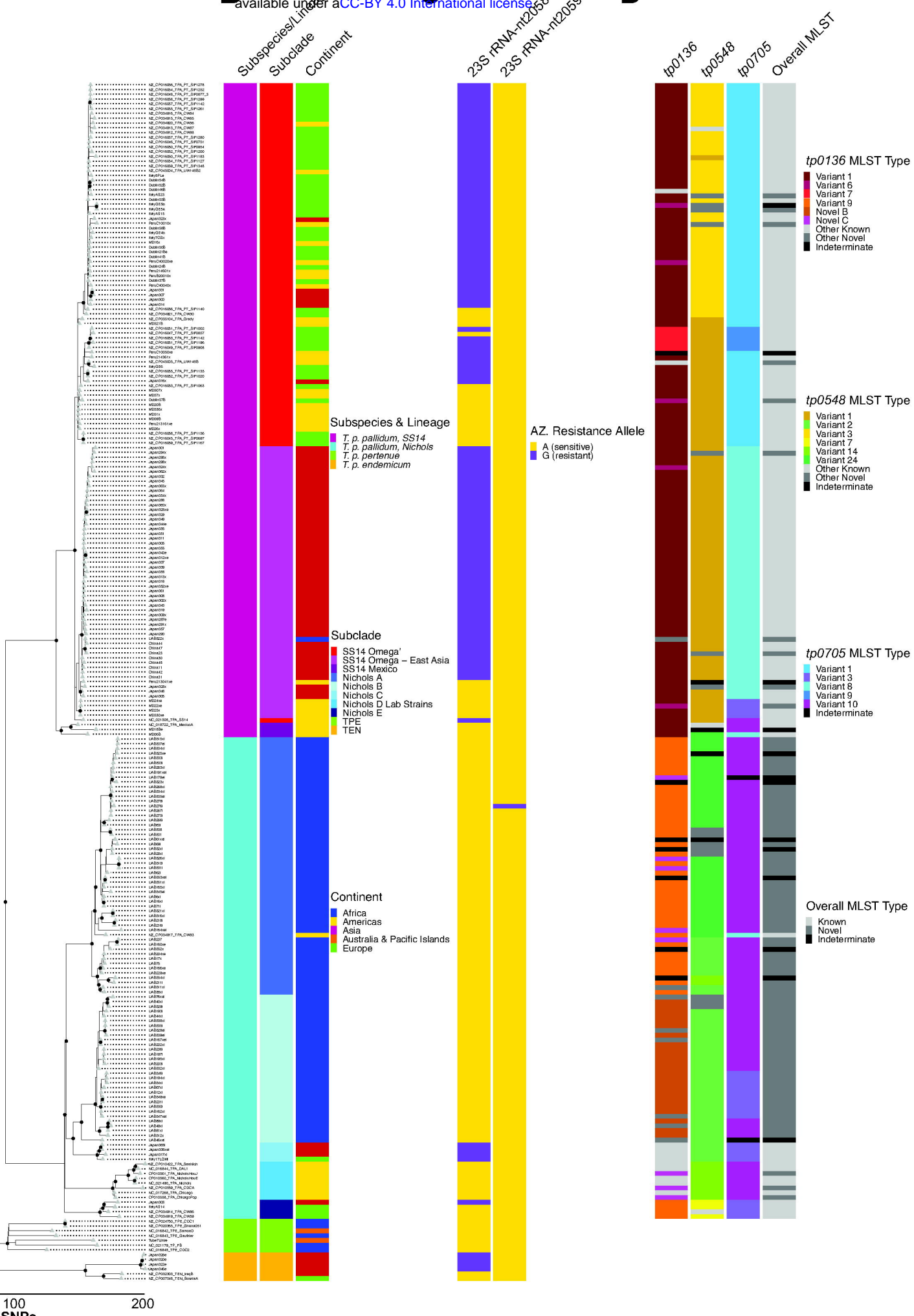
1081 **Supporting Information 4: Expanded data for Supplementary Figures 2-6.** TP0136,

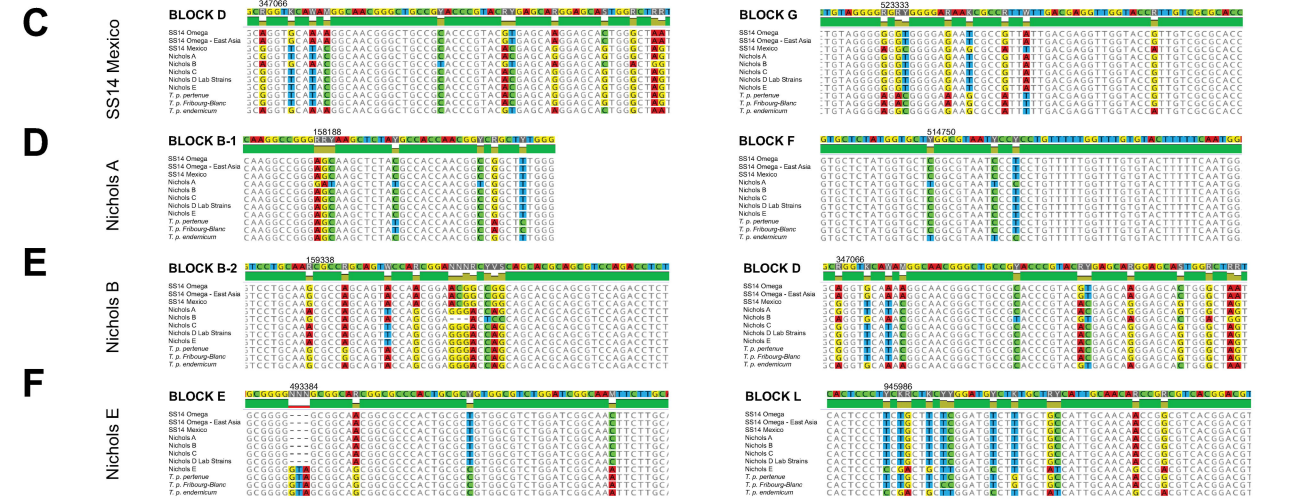
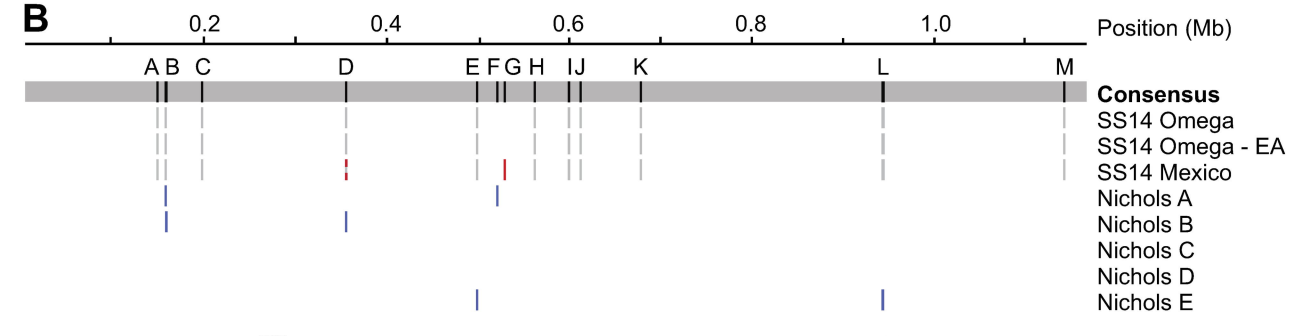
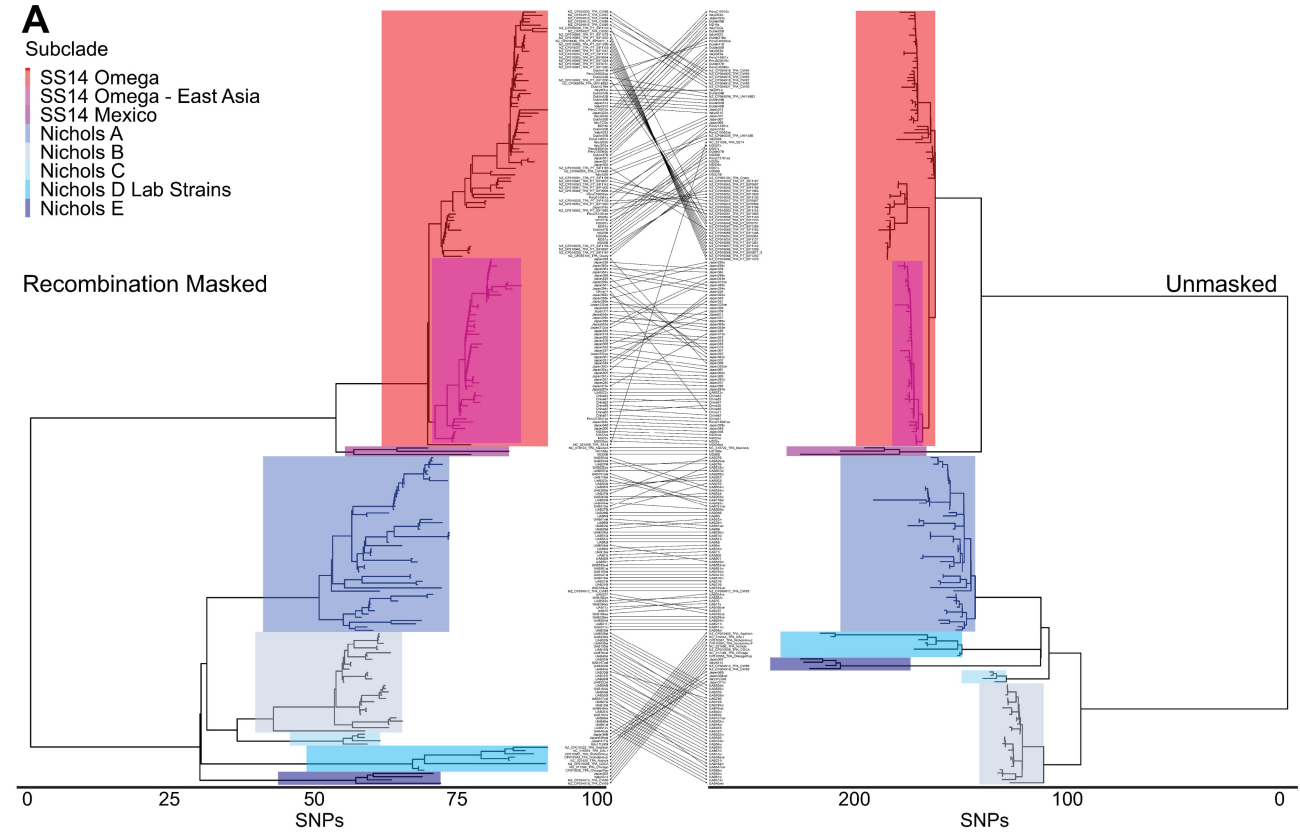
1082 TP0326, TP0548, TP0966, and TP0967 variant amino acid sequence by sample and subclade.

1083

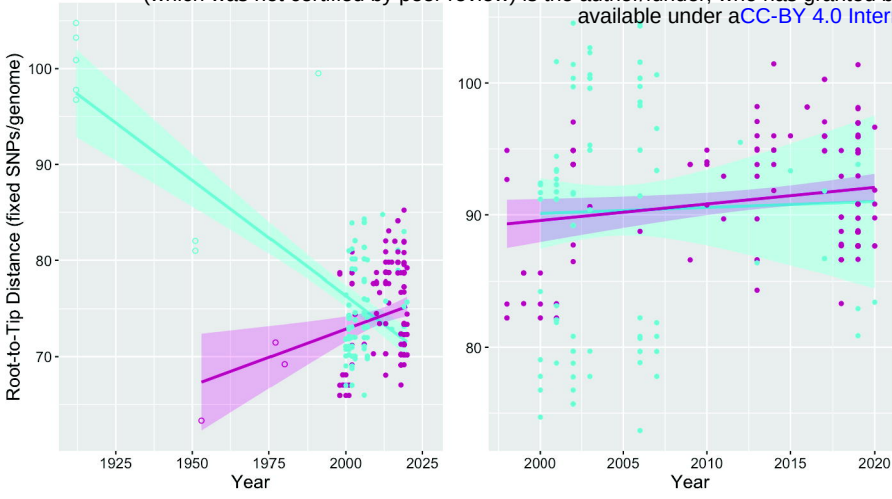
1084

A

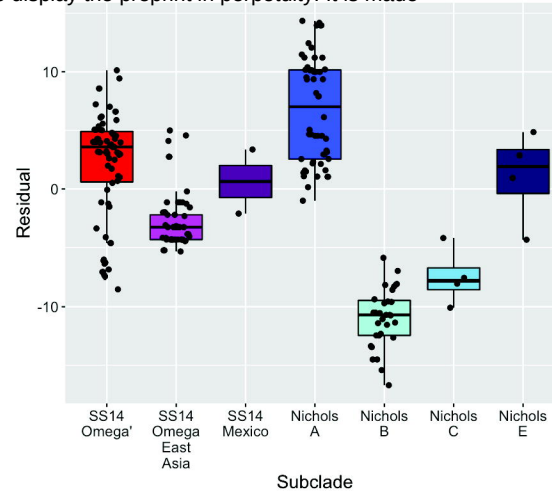




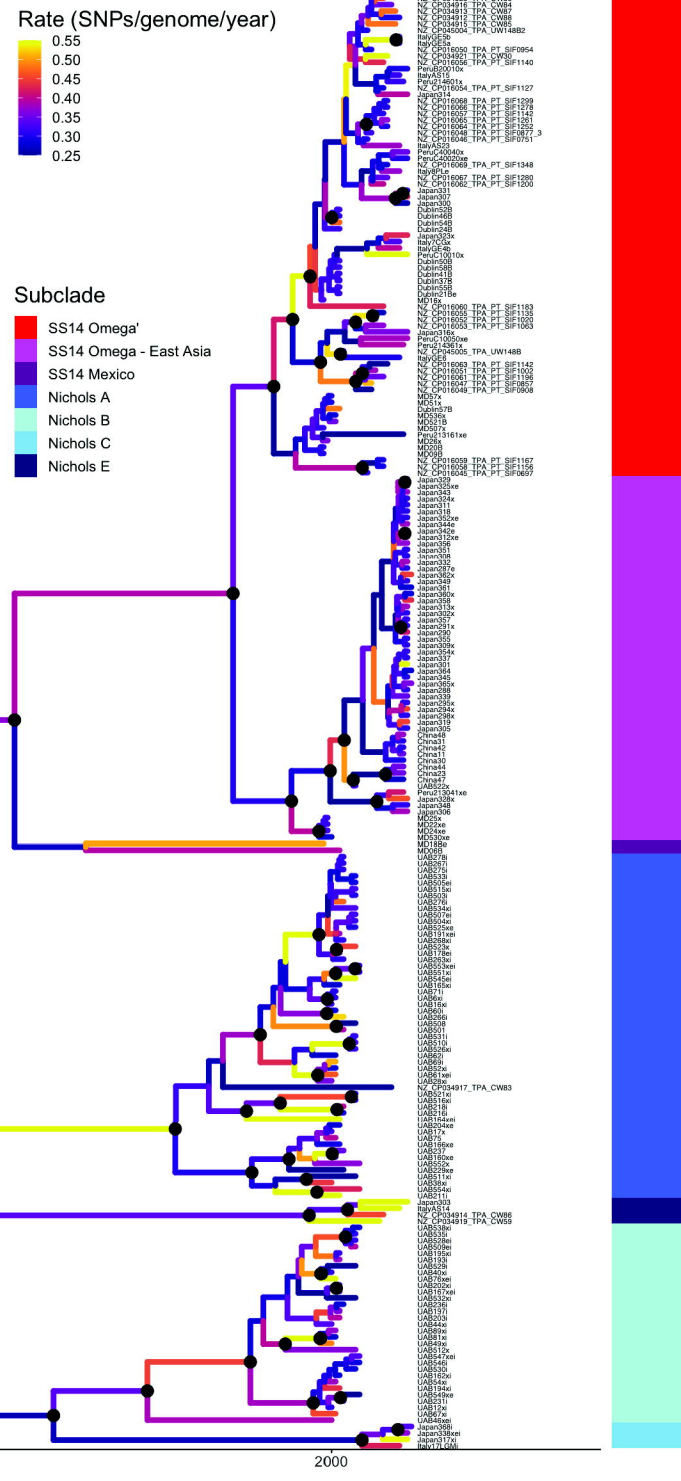
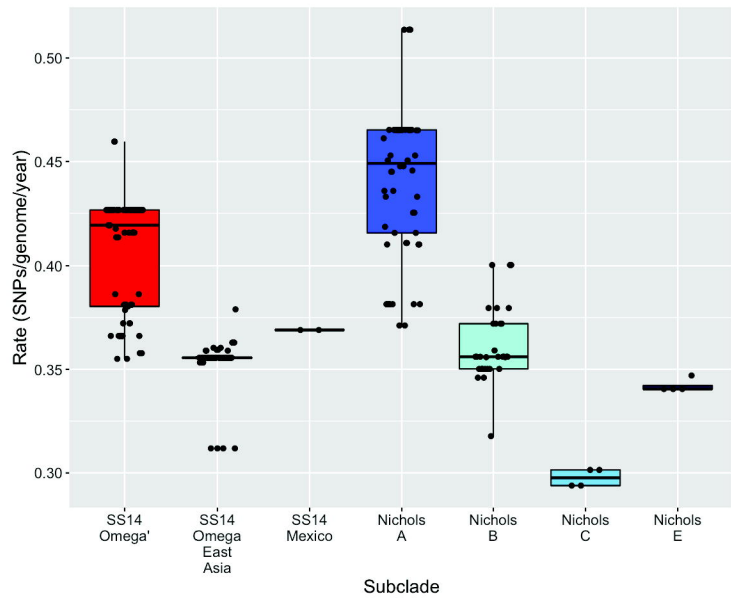
A



B



C



Node Ages (95% HPD) MRCA: 1717 (1543 - 1869)

SS14: 1921 (1868 - 1964)
 SS14 Omega: 1975 (1960 - 1988)
 SS14 Omega': 1985 (1977-1993)
 SS14 Omega - East Asia: 1990 (1981 - 1997)
 SS14 Mexico: 1938 (1892 - 1979)

Nichols: 1893 (1839 - 1940)
 Nichols AE: 1903 (1850 - 1947)
 Nichols BC: 1930 (1891 - 1965)
 Nichols A: 1961 (1942 - 1978)
 Nichols B: 1954 (1926 - 1978)
 Nichols C: 2008 (2001 - 2014)
 Nichols E: 1995 (1976 - 2007)

