

1 Transmembrane helices are an over-presented and
2 evolutionarily conserved source of major
3 histocompatibility complex class I and II epitopes

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Abstract

Cytolytic T cell responses are predicted to be biased towards membrane proteins. The peptide-binding grooves of most alleles of histocompatibility complex class I (MHC-I) are relatively hydrophobic, therefore peptide fragments derived from human transmembrane helices (TMHs) are predicted to be presented more often as would be expected based on their abundance in the proteome. However, the physiological reason of why membrane proteins might be over-presented is unclear. In this study, we show that the predicted over-presentation of TMH-derived peptides is general, as it is predicted for bacteria and viruses and for both MHC-I and MHC-II, and confirmed by re-analysis of epitope databases. Moreover, we show that TMHs are evolutionarily more conserved, because single nucleotide polymorphisms (SNPs) are present relatively less frequently in TMH-coding chromosomal regions compared to regions coding for extracellular and cytoplasmic protein regions. Thus, our findings suggest that both cytolytic and helper T cells are more tuned to respond to membrane proteins, because these are evolutionary more conserved. We speculate that TMHs are less prone to mutations that enable pathogens to evade T cell responses.

Keywords: antigen presentation, membrane proteins, adaptive immunity, transmembrane domain, epitopes, MHC-I, MHC-II, evolutionary conservation

29 **Abbreviations**

Abbreviation	Full
ER	Endoplasmatic reticulum
ERAD	ER-associated degradation
HLA	Human leukocyte antigen
IEDB	Immune Epitope Database
LB	lipid body
MAP	Membrane-associated protein
MHC	Major histocompatibility complex
MVB	Multivesicular body
PLC	Peptide-loading complex
SNP	Single nucleotide polymorphism
TMH	Transmembrane helix
TMP	Transmembrane protein

1 Introduction

Our immune system fights diseases and infections from pathogens, such as fungi, bacteria or viruses. An important part of the acquired immune response, that develops specialized and more specific recognition of pathogens than the innate immune response, are T cells which recognize peptides, called epitopes, derived from antigenic proteins presented on Major Histocompatibility Complexes (MHC) class I and II on the cell surface.

The MHC proteins are heterodimeric complexes encoded by the HLA (Human Leukocyte Antigens) genes. In humans, the peptide binding groove of MHC-I is made by only the alpha subunit. There are three classical alleles of MHC-I, hallmarked by a highly polymorphic alpha chain called HLA-A, HLA-B and HLA-C, that all present epitopes to cytolytic T cells. For MHC-II, both the alpha and the beta chains contribute to the peptide binding groove. There are three classical alleles of MHC-II as well, called HLA-DR, HLA-DQ and HLA-DP, that all present epitopes to helper T cells. Each MHC complex can present a subset of all possible peptides. For example, HLA-A and HLA-B have no overlap in which epitopes they bind [1]. Moreover, the HLA genes of humans are highly polymorphic, with hundreds to thousands of different alleles, and each different allele presents a different subset of peptides [2].

Humans express a limited set of MHC alleles and therefore an individual's immune system detects only a fraction of all possible peptide fragments. However, at the population level, the coverage of pathogenic peptides that are detected is very high, because of the highly polymorphic MHC genes. It is therefore believed that MHC polymorphism improves immunity at the population level, as mutations in a protein that disrupt a particular MHC presentation at the individual level, so-called escape mutations, will not affect MHC presentation for all alleles present in the population [3].

57 Many studies are aimed at identifying the repertoire of epitopes that are
58 presented in any of the different alleles to determine which epitopes will result
59 in an immune response, as this will for instance aid the design of vaccines.
60 These studies have led to the development of prediction algorithms that allow
61 for very reliable *in silico* predictions of the peptide binding affinities [4, 5, 6]. For
62 example, S. Tang et al. [6] found that, of the 432 peptides that were predicted
63 to bind to an MHC allele, 86% were experimentally confirmed to do so.

64 Using these prediction algorithms, we recently showed that peptides derived
65 from transmembrane helices (TMHs) are likely to be more frequently presented
66 by MHC-I than expected based on their abundance [7], which is in line with
67 a previous study by Istrail et al [8], demonstrating that N-terminal signal se-
68 quences are likely to be presented within major histocompatibility complexes,
69 due their hydrophobic nature. Moreover, we showed that some well-known im-
70 munodominant peptides stem from TMHs. This over-presentation is attributed
71 to the fact that the peptide-binding groove of most MHC-I alleles is relatively
72 hydrophobic, and therefore hydrophobic TMH-derived peptides have a higher
73 affinity to bind than their soluble hydrophilic counterparts.

74 TMHs are hydrophobic as they need to span the hydrophobic lipid bilayer
75 of cellular membranes. They consist of an alpha helix of, on average, 23 amino
76 acids in length. TMHs can also be predicted with high accuracy from a pro-
77 tein sequence by bioinformatics approaches [9, 10, 11, 12, 13, 14]. For example,
78 a study by Jones [12] found that, from 184 transmembrane proteins (TMPs)
79 with known topology, 80% of the TMH predictions of these proteins matched
80 the experimental findings. TMHs are common structures in the proteins of hu-
81 mans and microbes. Different TMH prediction tools estimate that 15-39% of all
82 proteins in the human proteome contain at least one TMH [15]. However, the
83 physiological reason why peptides derived from TMHs would be presented more

84 often than peptides stemming from soluble (i.e., extracellular or cytoplasmic)
85 protein regions is unknown. In this study, we hypothesized that the presen-
86 tation of TMH residues is evolutionarily preferred, since TMHs are less prone
87 to undergo escape mutations. One reason to expect such a reduced variability
88 (and hence evolutionary conservation) in TMHs, is that these are restricted in
89 their variability by the functional requirement to span a lipid bilayer. This lim-
90 its many of the amino acids present in TMHs to have hydrophobic side chains
91 [16, 17]. Therefore, we speculated that the TMHs of pathogens might have a
92 lower chance to develop escape mutations, as that will result in a dysfunctional
93 TMH and render the protein inactive.

94 This study had two objectives. First, we aimed to generalize our findings
95 by predicting the antigenic presentation from different kingdoms of life in both
96 MHC-I and -II. From these *in silico* predictions, we conclude that TMH-derived
97 epitopes from a human, viral and bacterial proteome are likely to be presented
98 more often than expected by chance for most alleles of MHC-I and II. We con-
99 firmed the presentation of TMH-derived peptides by re-analysis of peptides from
100 The Immune Epitope Database (IEDB) [18]. Second, we tested our hypothe-
101 sis that TMHs are more evolutionary conserved than soluble protein regions.
102 Our analysis of human single nucleotide polymorphisms (SNPs) showed that
103 random point mutations are indeed less likely to occur within TMHs. These
104 findings strengthen the emerging notion that TMHs are important for the T
105 cell-mediated adaptive immune system, and hence are of importance in vaccine
106 development.

107 2 Methods

108 2.1 Predicting TMH epitopes

109 To predict how frequently epitopes overlapping with TMHs are presented, a sim-
110 ilar analysis strategy was applied as described in [7] for several alleles of both
111 MHC-I and MHC-II, and for a human, viral and bacterial proteome. To sum-
112 marize, for each proteome, all possible 9-mers (for MHC-I) or 14-mers (MHC-II)
113 were derived. For each of these peptides, we determined if it overlapped with a
114 predicted TMH and if it was predicted to bind to the most frequent alleles of
115 each MHC allele.

116 For MHC-I, 9-mers were used, as this is the length most frequently presented
117 in MHC-I and was used in our earlier study [7]. For MHC-II, 14-mers were used,
118 as this is the most frequently occurring epitope length [19]. A human (UniProt
119 ID UP000005640.9606), viral (SARS-CoV-2, UniProt ID UP000464024) and
120 bacterial (*Mycobacterium tuberculosis*, UniProt ID UP000001584) reference pro-
121 teome was used. TMHMM [9] was used to predict the topology of the proteins
122 within these proteomes. To predict the affinity of an epitope to a certain HLA
123 allele, **EpitopePrediction** [7] for MHC-I and **MHCnuggets** [20] for MHC-II was
124 used. Both MHC-I and MHC-II alleles were selected to have a high prevalence
125 in the population, where the alleles of MHC-I are the alleles representing the
126 13 supertypes with over 99.6% coverage of the population’s MHC-I repertoire
127 as defined by [1] [21], and the 21 MHC-II alleles, have a phenotypic frequency
128 of 14% or more in the human population [22].

129 We define a protein to be a binder if, for a certain MHC allele, any of its
130 9-mer or 14-mer peptides have an IC50 value in the lowest 2% of all peptides
131 within a *proteome* (see supplementary Tables S1 and S2 for values), this differs
132 from our previous study where we defined a binder as having an IC50 in the
133 lowest 2% of the peptides within a *protein*. This revised definition precludes

134 bias of proteins that give rise to no or only very few MHC epitopes. To verify
135 that the slight change in method yields similar results, a side by side comparison
136 is shown in the supplementary materials, Figures S1A and S1B.

137 **2.2 TMH epitopes obtained from experimental data**

138 To obtain experimental confirmation that peptides stemming from TMHs are
139 presented by MHC-I and MHC-II, we mined the IEDB [18] for confirmed human
140 MHC-ligands. We queried the IEDB for all linear epitopes obtained from MHC
141 ligand assays in healthy humans, carrying the MHC alleles as used in this study.
142 From these epitopes, we kept those that were present exactly once in the human
143 reference proteome with UniProt ID UP000005640.9606. We concluded that
144 the epitope overlapped with a TMH if at least 1 amino acid was overlapping
145 with a TMH, as predicted with TMHMM [9]. The full analysis can be found at
146 https://github.com/richelbilderbeek/bbbq_article_issue_157.

147 **2.2.1 Evolutionary conservation of TMHs**

148 To determine the evolutionary conservation of TMHs, we first collected human
149 single nucleotide polymorphisms (SNPs) resulting in a single amino acid substi-
150 tution and determined if this occurred within a predicted TMH or not.

151 As a data source, multiple NCBI (<https://www.ncbi.nlm.nih.gov/>) databases
152 were used: the *dbSNP* [23] database, which contains 650 million cataloged non-
153 redundant human variations (called *RefSNPs*, https://www.ncbi.nlm.nih.gov/snp/docs/RefSNP_about/), and the databases *gene* (for gene names [24])
154 and *protein* (for proteins sequences [25]).

156 The first query was a call to the *gene* database for the term 'membrane
157 protein' (in all fields) for the organism *Homo sapiens*. This resulted in 1,077
158 gene IDs (on December 2020). The next query was a call to the *gene* database

159 to obtain the gene names from the gene IDs. Per gene name, the *dbSNP* NCBI
160 database was queried for variations associated with the gene name. As the
161 NCBI API constrains its users to three calls per second (to assure fair use), we
162 had to limit the extent of our analysis.

163 The number of SNPs was limited to the first 250 variations per gene, resulting
164 in $\approx 61k$ variations. Only variations that result in a SNP for a single amino acid
165 substitution were analyzed, resulting in $\approx 38k$ SNPs. The exact amounts can be
166 found in the supplementary materials, Tables S3 and S4.

167 SNPs were picked based on ID number, which is linked to their discovery
168 date. To verify that these ID numbers are unrelated to SNP positions, the
169 relative positions of all analyzed SNPs in a protein were determined. This
170 analysis showed no positional bias of the SNPs, as shown in supplementary
171 figure S2.

172 Per SNP, the *protein* NCBI database was queried for the protein sequence.
173 For each protein sequence, the protein topology was determined using **Pureseq**TM.
174 Using these predicted protein topologies, the SNPs were scored to be located
175 within or outside TMHs.

176 3 Results

177 3.1 TMH-derived peptides are predicted to be over-presented 178 in MHC-I

179 Figure 1A shows the predicted presentation of TMH-derived peptides in MHC-
180 I, for a human, viral and bacterial proteome. Per MHC-I allele, it shows the
181 percentage of binders that overlap with a TMH with at least one residue. The
182 horizontal line shows the expected percentage of TMH-derived epitopes that
183 would be presented, if TMH-derived epitopes would be presented just as likely

as epitopes derived from soluble regions, when assuming equal incidence of soluble and TMH-derived epitope presentation. For 11 out of 13 MHC-I alleles, TMH-derived epitopes are predicted to be presented more often than the null expectation, for a human and bacterial proteome. For the viral proteome, 12 out of 13 MHC-I alleles present TMH-derived epitopes more often than expected by chance. The extent of the over-presentation between the different alleles is similar for the probed proteomes, which strengthens our previous conclusion [7] that the hydrophobicity of the MHC-binding groove is the main factor responsible for the predicted over-presentation of TMH-derived peptides.

3.2 TMH-derived peptides are predicted to be over-presented in MHC-II

We next wondered if the over-representation of TMH-derived peptides would also be present for MHC-II. Figure 1A shows the percentages of MHC-II epitopes predicted to be overlapping with TMHs for our human, viral and bacterial proteomes. We found that TMH-derived peptides are over-presented in all of the 21 MHC-II alleles, for a human, bacterial and viral proteome, except for HLA-DRB3*0101 in *M. tuberculosis*. See supplementary Table S5 for the exact TMH and epitope counts.

3.3 The over-presentation of TMH-derived peptides is caused by the hydrophobicity of the MHC peptide binding groove

For MHC-I, we previously showed that the over-presentation of TMH-derived peptides is caused by the hydrophobicity of the peptide binding grooves [7]. Figures 1B and 1C show the extent of over-presentation of TMH-derived epitopes as a function of the hydrophobicity preference score for the different human MHC

alleles. An assumed linear correlation explains 88% of the variability in MHC-I. For MHC-II, 62% of the variability is explained by hydrophobicity. This indicates that TMH-derived peptides are over-presented, because the peptide binding grooves of most MHC-I and -II alleles are relatively hydrophobic.

3.4 Experimental validation of presentation of TMH-derived peptides

The Immune Epitope Database (IEDB) from the National Institutes of Health contains millions of linear epitope sequences obtained by MHC ligand assays. For the MHC alleles used in this study, we obtained 54,303 and 2,484 linear epitope sequences for the MHC-I and MHC-II alleles from human origin respectively. There are relatively few epitopes for MHC-II, as MHC-II has many more different alleles than MHC-I, whereas we selected only the human epitopes found for the 21 MHC-II alleles used in this study.

Figure 2A and S3 show there are similar levels of over-presentation of TMH-derived epitopes between (1) the percentage of TMH-derived epitopes that is reported in the IEDB database versus (2) the percentage of TMH-derived epitopes that is predicted to be presented in MHC-I alleles. For MHC-II alleles, there were too few epitopes per MHC allele to result in an informative figure.

In figure 2B we grouped all the epitopes presented by MHC-I and MHC-II alleles by the percentage of TMH-derived epitopes, which are 22% and 10%, respectively.

These findings robustly confirm that epitopes derived from human TMHs are presented in both MHC-I and MHC-II, and support that they are over-presented. See the supplementary Table S6 for the exact values.

We also mined the IEDB database for epitopes for any type of T cell response from the specified alleles. From the total reports, 36% and 7% concerned TMH-

235 derived epitopes in MHC class I and II, respectively (see Figure S4).

236 This data confirms that not only TMH derived epitopes are presented on
237 MHC, but this also elicits T-cell mediated immune responses.

238 **3.5 Human TMHs are evolutionarily conserved**

239 We addressed the question whether there is an evolutionary advantage in pre-
240 senting TMHs. We determined the conservation of TMHs by comparing the
241 occurrences of SNPs located in TMHs or soluble protein regions for the genes
242 coding for membrane proteins. We obtained 911 unique gene names associated
243 with the phrase 'membrane protein', which are genes coding for both membrane-
244 associated proteins (MAPs, which have no TMH) and transmembrane proteins
245 (TMPs, which have at least one TMH). These genes are linked to 4,780 pro-
246 tein isoforms, of which 2,553 are predicted to be TMPs and 2,237 proteins are
247 predicted to be MAPs. We obtained 37,630 unique variations, of which 9,621
248 are SNPs that resulted in a straightforward amino acids substitution, of which
249 6,062 were located in predicted TMPs. See supplementary Tables S3 and S4 for
250 the detailed numbers and distributions of SNPs.

251 Per protein, we calculated two percentages: (1) the percentage of a protein
252 sequence length bearing TMHs, and (2) the percentage of SNPs located within
253 these predicted TMHs. Each percentage pair was plotted in figure 3A. The
254 proportion of SNPs found in TMHs varied from none (i.e., all SNPs were in
255 soluble regions) to all (i.e., all SNPs were in TMHs). To determine if SNPs
256 were randomly distributed over the protein, we performed a linear regression
257 analysis, and added a 95% confidence interval on this regression. This linear fit
258 nearly goes through the origin and has a slope below the line of equality, which
259 shows that less SNPs are found in TMHs than expected by chance.

260 We determined the probability to find the observed amount of SNPs in TMHs

261 by chance, i.e., when assuming SNPs occur just as likely in soluble domains as
 262 in TMHs. We used a binomial Poisson distribution, where the number of trials
 263 (n) equals the number of SNPs, which is 21,208. The probability of success
 264 for the i th TMP (p_i), is the percentage of residues within a TMH per TMP.
 265 These percentages are shown as a histogram in figure 3B. The expected number
 266 of SNPs expected to be found in TMHs by chance equals $\sum p \approx 4,141$. As
 267 we observed 3,803 SNPs in TMHs, we calculated the probability of having that
 268 amount or less successes. We used the type I error cut-off value of $\alpha = 2.5\%$. The
 269 chance to find, within TMHs, this amount or less SNPs equals $6.8208 \cdot 10^{-11}$. We
 270 determined the relevance of this finding, by calculating how much less SNPs are
 271 found in TMHs, when compared to soluble regions, which is the ratio between
 272 the number of SNPs found in TMHs versus the number of SNPs as expected
 273 by chance. In effect, per 1000 SNPs found in soluble protein domains, one finds
 274 918 SNPs in TMHs, as depicted (as percentages) in figure 3C.

275 We split this analysis for TMPs containing only a single TMH (so-called
 276 single-membrane spanners) and TMPs containing multiple TMHs (multi-membrane
 277 spanners). We hypothesized that single-membrane spanners are less conserved
 278 than multi-membrane spanners, because multi-membrane spanners might have
 279 protein-protein interactions between their TMHs, for example to accommodate
 280 active sites, and thus might have additional structural constraints. From the
 281 split data, we did the same analysis as for the total TMPs. Figure 4A shows the
 282 percentages of TMHs for individual proteins as a function of the percentage of
 283 SNPs located in TMHs. For both single- and multi-spanners, a linear regression
 284 shows that less SNPs are found in TMHs, than expected by chance.

285 We also determined the probability to find the observed amount of SNPs by
 286 chance in single- and multi-spanners. For single-spanners, we found 452 SNPs
 287 in TMH, where ≈ 462 were expected by chance. The chance to observe this or a

288 lower number by chance is 0.319. As this chance was higher than our $\alpha = 0.025$,
289 we consider this no significant effect. For the multi-spanners, we found 3,351
290 SNPs in TMH, where $\approx 3,678$ were expected by chance. The chance to observe
291 this or a lower number by chance is $8.315841 \cdot 10^{-12}$, which means this number
292 is significantly less as explained by variation. The TMHs of multi-spanners are
293 thus significantly more conserved than soluble protein regions, whereas this is
294 not the case for single-spanners.

295 Also, for single- and multi-spanners, we determined the relevance of this
296 finding by calculating where and how much less SNPs are found in TMHs when
297 compared to soluble regions, as depicted in Figure 4B and 4B. In effect, per
298 1,000 SNPs found in soluble protein domains, one finds 978 SNPs in TMHs of
299 single-spanners and 911 SNPs in TMHs of multi-spanners.

300 4 Discussion

301 Epitope prediction is important to understand the immune system function and
302 for the design of vaccines. In this study, we provide evidence that epitopes
303 derived from TMHs are a major source of MHC epitopes. Our bioinformat-
304 ics predictions indicate that the TMH-derived epitope repertoire is larger than
305 expected by chance for both MHC-I and MHC-II, regardless of the organism.
306 Moreover, reanalysis of MHC-ligands from the IEDB database confirmed the
307 presentation of TMH-derived epitopes. Therefore, it seems likely that TMH-
308 derived epitopes would also result in enhanced T cell responses, although the
309 conservation of TMHs might promote the deletion of T cells responsive to TMH-
310 derived epitopes by central tolerance mechanisms. Finally, our SNP analysis
311 shows that TMHs are evolutionary more conserved than solvent-exposed pro-
312 tein regions.

313 4.1 Mechanism of MHC presentation of TMH-derived epi- 314 topes

315 Although our data show that TMH-derived epitopes are presented in all clas-
316 sical MHC-I and MHC-II alleles, the molecular mechanisms of how integral
317 membrane proteins are processed for MHC presentation are largely unknown
318 [7]. Most prominently, the fundamental principles of how TMHs are extracted
319 from their hydrophobic lipid environments into the aqueous vacuolar lumen,
320 leading to subsequent proteolytic processing are unresolved.

321 A first possibility is that the extraction of TMPs from the membrane is
322 mediated by the ER-associated degradation (ERAD) machinery. For MHC class
323 I (MHC-I) antigen presentation of soluble proteins, the loading of the epitope
324 primarily occurs at the endoplasmatic reticulum (ER). The chaperones tapasin
325 (TAPBP), ERp57 (PDIA3), and calreticulin (CALR) [26] first assemble and
326 stabilize the heavy and light chains of MHC-I. Later, this complex binds to the
327 transporter associated with antigen processing (TAP) leading to the formation of
328 the so-called peptide-loading complex (PLC). The PLC drives import of peptides
329 into the ER and mediates their subsequent loading into the peptide-binding
330 groove of MHC-I [27]. Membrane proteins first will have to be extracted from
331 the membrane before they become amenable to this MHC-I loading by the
332 PLC. In the ER, this process can be orchestrated by the ERAD machinery,
333 consisting of several chaperones that recognize TMPs, ubiquitinate them, and
334 extract them from the ER membrane into the cytosol (retrotranslocation) for
335 proteasomal degradation [28, 29]. Similar to the peptides generated from soluble
336 proteins, the TMP-derived peptides might then be re-imported by TAP into the
337 ER for MHC-I loading. This ERAD-driven antigen retrotranslocation might be
338 facilitated by lipid bodies (LBs) [30], since LBs can serve as cytosolic sites for
339 ubiquitination of ER-derived cargo [31].

340 A second possibility is that TMPs are proteolytically processed by intramem-
341 brane proteases that cleave TMHs while they are still membrane embedded.
342 Supporting this hypothesis is the well-established notion that peptides gener-
343 ated by signal peptide peptidases (SPPs), an important class of intramembrane
344 proteases that cleave TMH-like signal sequences, are presented on a specialized
345 class of MHC-I called HLA-E [32]. The loading of peptides generated by SPP
346 onto MHC-I does not depend on the proteasome and TAP, possibly because
347 the peptides are directly released into the lumen of the ER [32]. However, this
348 mechanism cannot explain how most membrane proteins can be processed for
349 antigen presentation, because SPPs only cleave TMH-like signal sequences at
350 their C-termini, and N-terminal domains will hence not be removed. Neverthe-
351 less, the presentation of peptides with a high hydrophobicity index was shown
352 to be independent of TAP as well [33], suggesting that the TMH peptides might
353 perhaps be released directly in the ER lumen by other intramembrane proteases.

354 A third possibility is that peptide processing and MHC-loading occur in
355 multivesicular bodies (MVBs) [32]. TMPs can be routed from the plasma mem-
356 brane and other organelles by vesicular trafficking to endosomes. Eventually,
357 these TMPs can be sorted by the endosomal sorting complexes required for
358 transport (ESCRT) pathway into luminal invaginations that pinch off from the
359 limiting membrane and form intraluminal vesicles. This thus results in MVBs
360 where the membrane proteins destined for degradation are located in intralumi-
361 nal vesicles. Upon the fusion of MVBs with lysosomes, the entire intraluminal
362 vesicles including the TMPs are degraded [34]. Via this mechanism, TMPs
363 might well be processed for antigen presentation, particularly since the loading
364 of MHC-II molecules is well understood to occur in MVBs [35, 36, 37]. However,
365 such processing of membrane proteins in MVBs for antigen presentation poses
366 a problem, because complexes of HLA-DR with its antigen-loading chaperon

HLA-DM were only observed on intraluminal vesicles, but not on the limiting membranes of MVBs [37], indicating that epitope loading of MHC-II also occurs at intraluminal vesicles. This observation hence raises the question how the intraluminal vesicles carrying the TMPs destined for antigen presentation can be selectively degraded, while the intraluminal vesicles carrying the MHC-II remain intact. A second problem is that phagosomes carrying internalized microbes lack intraluminal vesicles, and it is hence unclear how TMPs from these microbes would be routed to MVBs for MHC-II loading [37].

Alternatively to the enzymatic degradation of lipids in MVBs by lipases [38, 39], they might be oxidatively degraded by reactions with radical oxygen species produced by the NADPH oxidase NOX2 [40]. This oxidation can result in a destabilization and disruption of membranes [40] and might thereby lead to the extraction of TMPs. Due to the hydrophobic nature of TMHs, however, the extracted proteins will likely aggregate and it is unclear how these aggregates would be processed further for MHC loading.

4.2 Evolutionary conservation of TMHs

In general, one might expect that evolutionary selection shapes an immune system where surveillance is directed towards protein regions essential for the survival, proliferation and/or virulence or pathogenic microbes, as these will be most conserved. In SARS-CoV-2, for example, there is preliminary evidence that the strongest selection pressure is directed upon residues that change its virulence [41]. These regions, however, may only account for a small part of a pathogen's proteome. Additionally, the structure and function of these essential regions might differ widely between different pathogenic proteins. Because of this scarcity and variance in targets, one can imagine that it will be mostly unfeasible to provide innate immune responses against such rare essential protein

393 regions, as suggested in a study on influenza [42], where it was found that the
394 selection pressure exerted by the immune system was either weak or absent.

395 Evolutionary selection of pathogens by a host’s immune system, however, is
396 more likely to occur for protein patterns that are general, over patterns that are
397 rare. While essential catalytic sites in a pathogenic proteome might be relatively
398 rare, TMHs are common and thus might be a more feasible target for evolution
399 to respond to. Indeed, we have found the signature of evolution when both
400 factors, that is, TMHs and catalytic sites are likely to co-occur, which is in TMPs
401 that span the membrane at least twice. In contrast to single-spanners, where
402 we found no significant evolutionary conservation, the TMHs of multi-spanners
403 are more evolutionarily conserved than soluble protein regions. Likely, the TMHs
404 in many multi-spanners need to interact with each other for correct protein
405 structure and function and they might hence be more structurally constrained
406 compared to the TMHs of single-spanners. Thus, we speculate that the human
407 immune system is more attentive towards TMHs in multi-spanners, as these are
408 evolutionarily more conserved.

409 There have been more efforts to assess the conservation of TMHs, using
410 different methodologies. One such example is a study by Stevens and Arkin [43],
411 in which aligned protein sequence data was used. Also this study found that
412 TMHs are evolutionarily more conserved, as the mean amino acid substitution
413 rate in TMHs is about ten percent lower, which is a similar value as we found.
414 Another example is a study by Oberai, et al. [44] that estimated the conservation
415 scores for TMHs and soluble regions based on alignments of evolutionary related
416 proteins, and also found that TMHs are more conserved, with a conservation
417 score that was 17% higher in TMHs. Note that the last study also found that
418 mutations in human TMHs are likelier to cause a disease, in line with our
419 conclusion that TMHs are more conserved.

420 Together, from this study, two important conclusions can be drawn. First,
421 the MHC over-presentation of TMHs is likely a general feature and predicted to
422 occur for most alleles of both MHC-I and -II and for humans as well as bacterial
423 and viral pathogens. Second, TMHs are genuinely more evolutionary conserved
424 than soluble protein motifs, at least in the human proteome.

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437 6 Data Accessibility

438 The data presented in the study are deposited in the Zenodo repository, acces-
439 sion number 10.5281/zenodo.5809139. Additionally, all code, intermediate and
440 final results are archived at [https://github.com/richelbilderbeek/bbbq_](https://github.com/richelbilderbeek/bbbq_article)
441 [article](https://github.com/richelbilderbeek/bbbq_article).

442 7 Authors' contributions

443 RJCB and FB conceived the idea for this research. MVB helped with the
444 proteome analysis of *M. tuberculosis*. RJCB wrote the code. RJCB, MB, GvdB
445 and FB wrote the article.

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662 8 Figures

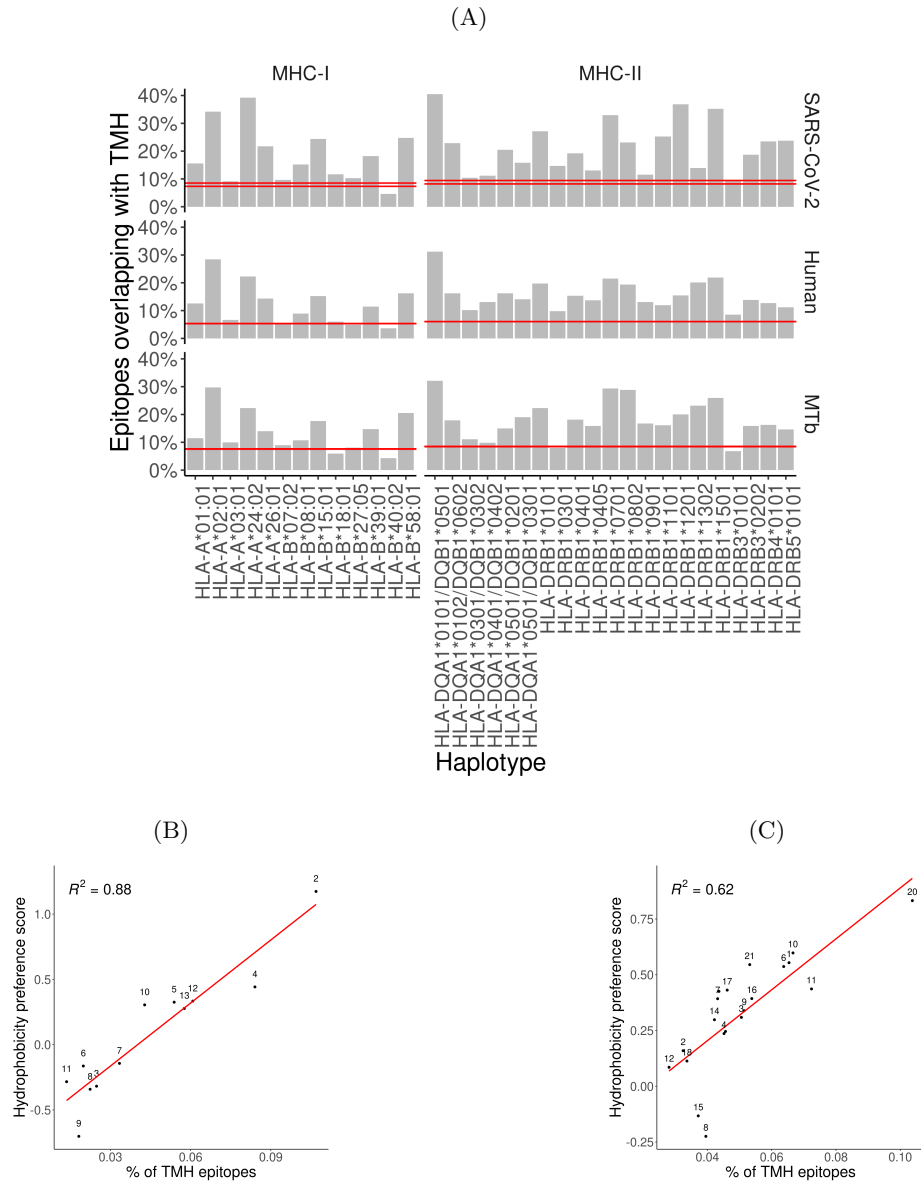


Figure 1: Over-presentation of TMH-derived epitopes on most MHC-I and -II alleles (A) The percentage of epitopes for MHC-I and -II alleles that are predicted to overlap with TMHs for the proteomes of SARS-CoV-2 (top row), human (middle row) and *M. tuberculosis* (MtB; bottom row). The pair of horizontal red lines in each plot indicate the lower and upper bound of the 99% confidence interval. See supplementary Tables S5 and S7 for the exact TMH and epitope counts. **(B-C)** Correlation between the percentages of predicted TMH-derived epitopes and the hydrophobicity score of all predicted epitopes for human MHC-I **(B)** and MHC-II alleles **(C)**. Diagonal red line: linear regression analysis. Labels are shorthand for the HLA alleles, see the supplementary Table S8 for the names.

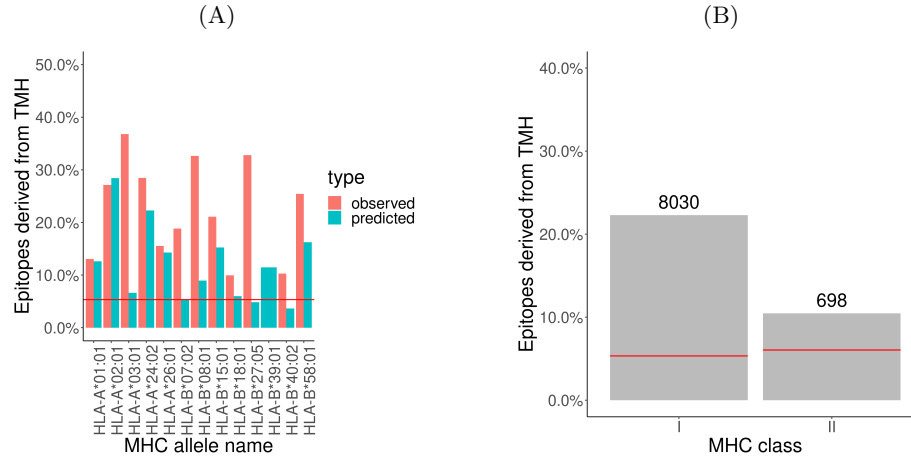


Figure 2: **Analysis of epitope database shows that TMH derived epitopes are over-presented.** The percentage of epitopes for MHC-I and -II alleles that overlap with TMHs that are presented. The pair of horizontal red lines in each plot indicate the lower and upper bound of the 99% confidence interval. Note that only one line is visible as this interval is relatively narrow. Alleles are listed in Table S8. **(A)** Observed and predicted percentage of TMH-derived epitopes for MHC-I alleles. **(B)** MHC ligands from IEDB corresponding to TMH-derived epitopes. The numbers above the bars denotes the number of TMH derived epitopes obtained.

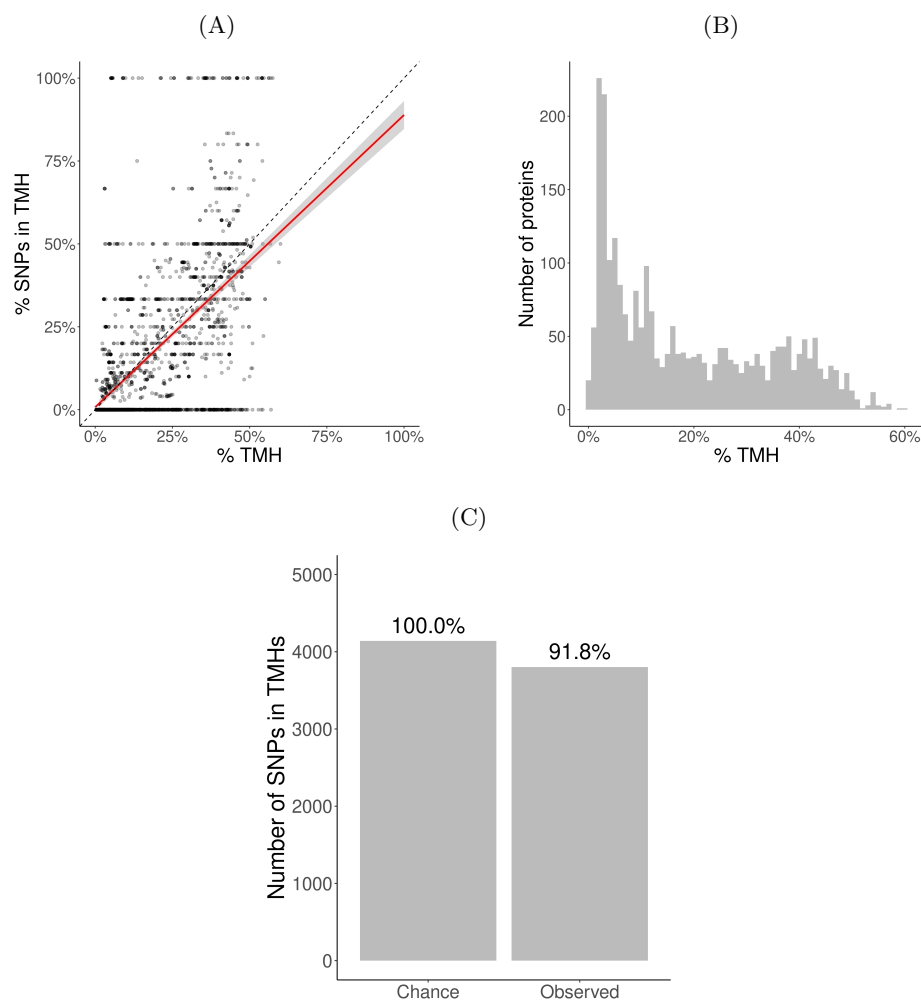


Figure 3: **Evolutionary conservation of human TMHs.** (A) Percentage of SNPs found in TMHs. Each point shows for one protein the predicted percentage of amino acids that are part of a TMH (x -axis) and the observed occurrence of SNPs being located within a TMH (y -axis). The dashed diagonal line shows the line of equality (i.e., equal conservation of TMHs and soluble protein regions). The diagonal red line indicates a linear fit, the gray area its 95% confidence interval. (B) Distribution of the percentages of TMH in the TMPs used in this study. (C) The number of SNPs in TMHs as expected by chance (left bar) and found in the dbSNP database (right bar). Percentages show the relative conservation of SNPs in TMHs found relative to stochastic chance.

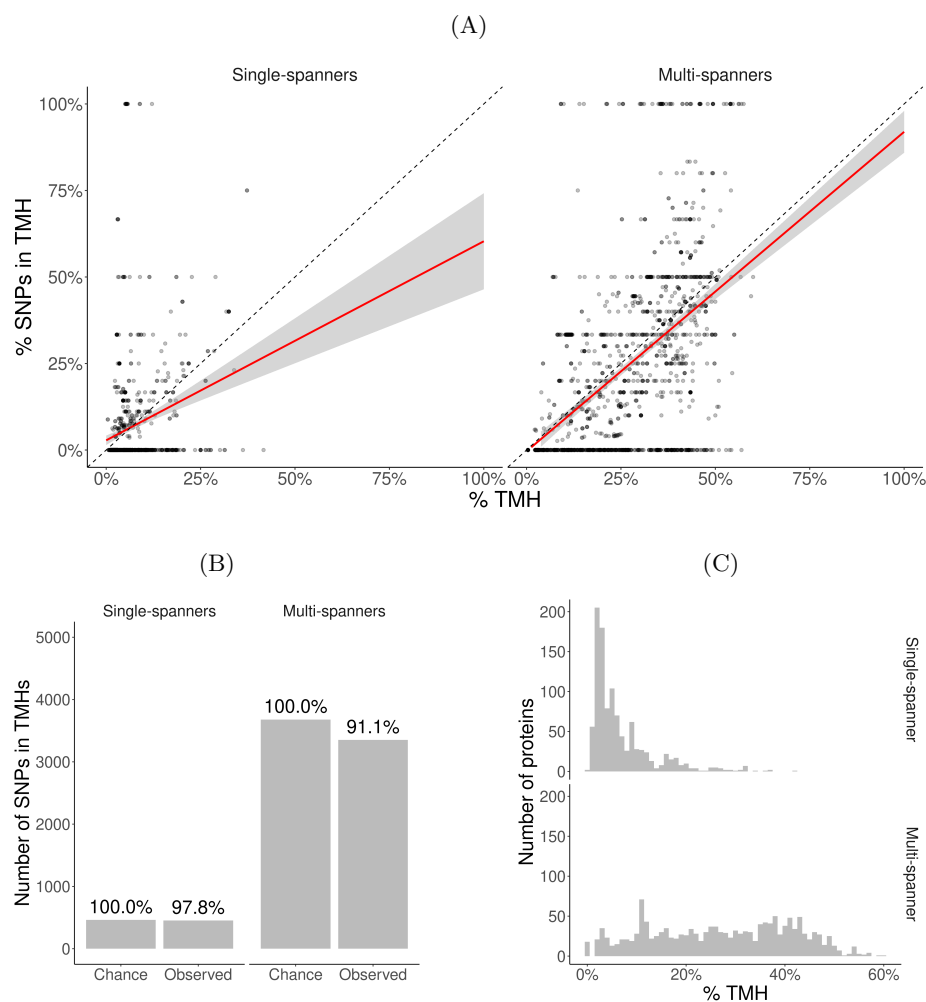


Figure 4: Membrane proteins with multiple TMHs are evolutionary more conserved than proteins with only a single TMH. (A) Percentage of SNPs found in TMPs predicted to have only a single (left) or multiple (right) TMHs. Each point shows for one protein the predicted percentage of amino acids that are part of a TMH (x -axis) and the observed occurrence of SNPs being located within a TMH (y -axis). The dashed diagonal lines show the line of equality (i.e., equal conservation of TMHs and soluble protein regions). The diagonal red lines indicate a linear fit, the gray areas their 95% confidence intervals. (B) The number of SNPs in TMHs as expected by chance and observed in the dbSNP database, for TMPs with one TMH (single-spanners) and multiple TMHs (multi-spanners). Percentages show the relative conservation of SNPs in TMHs found relative to the stochastic chances. (C) Distribution of the proportion of amino acids residing in the plasma membrane.

663 A Supplementary materials

664 A.1 Differences with Bianchi et al., 2017

665 A part of this study does the same analysis as Bianchi et al., 2017. mainly
666 concern the use of different software and a different definition of what an MHC
667 binder is.

668 The earlier study defined a peptide an MHC binder if *within the protein* in
669 which it was found, is was among the peptides with the 2% lowest IC50 val-
670 ues. This can be seen at [https://github.com/richelbilderbeek/bianchi_](https://github.com/richelbilderbeek/bianchi_et_al_2017/blob/master/predict-binders.R)
671 [et_al_2017/blob/master/predict-binders.R](https://github.com/richelbilderbeek/bianchi_et_al_2017/blob/master/predict-binders.R), where the binders are written
672 to file.

673 However, in this study, an MHC binder is defined as a peptide within a
674 *proteome* in which it is found, that is among the peptides with the 2% lowest
675 IC50 values. Subsection A.2 shows the IC50 values for a binder per MHC allele.

676 Our previous study used the TMHMM web server to predict TMHs. The
677 desktop version of TMHMM, however, gives an error message on the 25 seleno-
678 proteins found in the human reference proteome. For the sake of reproducible
679 research, we used the desktop version (as we can call it from scripts) and, due
680 to this, we removed the selenoproteins from this analysis.

681 To verify if the previous and the current method give rise to notable differ-
682 ence, we show a side-by-side comparison in figures S1A and S1B. The figures
683 that MHC molecules that over-present or under-present TMH-derived epitopes,
684 do so in both studies. The extent to which TMH-derived epitopes are presented,
685 however, is more extreme in our current setup.

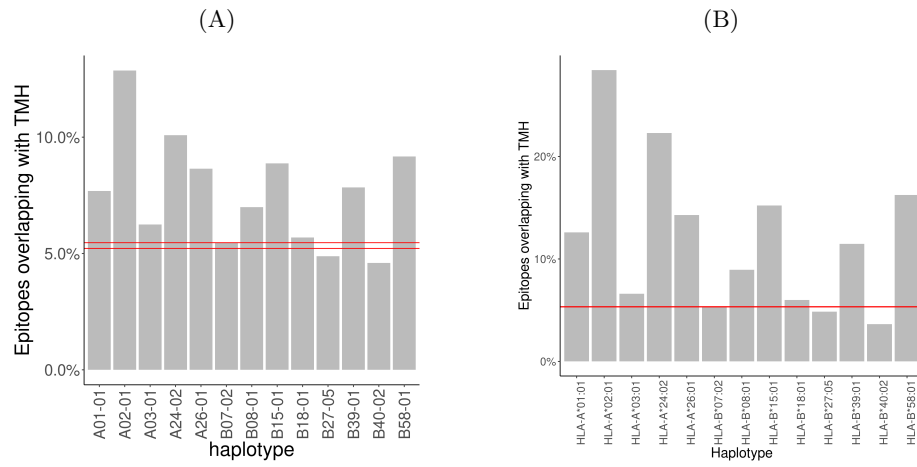


Figure S1: **(A)** Results for [7]. Dashed lines denotes the coincidence interval. **(B)** Results for this study. Dashed line denotes the percentage as expected by chance.

Table S1: IC50 values (in nM) per haplotype below which a peptide is considered a binder. percentage used: 2

haplotype	covid	human	myco
HLA-A*01:01	1470.5912	2545.9537	2812.1714
HLA-A*02:01	118.9596	218.7274	186.7565
HLA-A*03:01	537.0144	804.7455	1544.1073
HLA-A*24:02	984.8147	1590.0623	1971.8258
HLA-A*26:01	1095.2591	1771.6924	1526.1101
HLA-B*07:02	1215.7734	705.6514	435.5361
HLA-B*08:01	886.5661	883.0951	1023.2213
HLA-B*18:01	921.4157	1063.2215	1319.0445
HLA-B*27:05	1186.0963	689.8815	475.6130
HLA-B*39:01	437.3506	484.3843	399.3873
HLA-B*40:02	585.6308	541.2392	600.1688
HLA-B*58:01	435.4693	591.0526	538.9063
HLA-B*15:01	281.9129	440.6541	482.8369

A.2 IC50 values of binders per MHC allele

Per target proteome (i.e. human, SARS-CoV-2, *M tuberculosis*), we collected all 9-mers (for MHC-I) and 14-mers (for MHC-II), after removing the selenoproteins and proteins that are shorter than the epitope length. From these epitopes, per MHC allele, we predicted the IC50 (in nM) using `epitope-prediction` (for MHC-I) and `MHCnuggets` (for MHC-II). Here, we show the IC50 value per MHC allele that is used to determine if a peptide binds to the allele's MHC for MHC-I (see supplementary Table S1) and MHC-II (see supplementary Table S2).

Table S2: IC50 values (in nM) per haplotype below which a peptide is considered a binder. percentage used: 2

haplotype	covid	human	myco
HLA-DRB1*0101	7.3896	9.72	9.9600
HLA-DRB1*0301	121.8420	198.40	164.4900
HLA-DRB1*0401	59.8780	74.92	84.3112
HLA-DRB1*0405	46.2324	51.88	66.7100
HLA-DRB1*0701	17.7464	22.40	28.1700
HLA-DRB1*0802	99.7592	137.16	67.9900
HLA-DRB1*0901	42.3464	53.52	41.5400
HLA-DRB1*1101	35.9988	39.01	48.9200
HLA-DRB1*1201	194.4408	248.72	289.7300
HLA-DRB1*1302	21.1084	40.59	35.4100
HLA-DRB1*1501	32.6196	40.69	46.6700
HLA-DRB3*0101	175.2984	298.94	218.7300
HLA-DRB3*0202	176.8168	291.95	405.8724
HLA-DRB4*0101	47.6384	51.04	62.7800
HLA-DRB5*0101	32.8872	43.52	60.2312
HLA-DQA1*0501/DQB1*0201	193.1108	209.89	174.2124
HLA-DQA1*0501/DQB1*0301	51.2028	43.47	20.3200
HLA-DQA1*0301/DQB1*0302	361.8180	365.96	296.4712
HLA-DQA1*0401/DQB1*0402	214.1932	242.68	199.8912
HLA-DQA1*0101/DQB1*0501	550.4488	674.95	930.9612
HLA-DQA1*0102/DQB1*0602	157.4480	174.82	114.3512

Table S3: Amounts. raw = all variations, including DNA variations. all_proteins = all proteins. map = membrane associated protein. tmp = transmembrane protein. in_tmh = in transmembrane helix of TMP. in_sol = in soluble region of TMP.

what	raw	all_proteins	map	tmp	in_tmh	in_sol
Number of variations	60931	37831	16623	21208	3803	17405
Number of unique variations	60544	37630	16606	21024	3789	17235
Number of unique SNPs	NA	9621	4219	6026	1140	4936
Number of unique gene names	953	911	457	605	325	590
Number of unique protein names	5163	4780	2227	2553	1280	2467
Percentage TMH	NA	10	0	19	26	18

Table S4: Amounts. single_in_tmh = in transmembrane helix of single-spanner. single_in_sol = in soluble region of single-spanner. multi_in_tmh = in transmembrane helix of multi-spanner. multi_in_sol = in soluble region of multi-spanner.

what	single_in_tmh	single_in_sol	multi_in_tmh	multi_in_sol
Number of variations	452	7734	3351	9671
Number of unique variations	451	7733	3338	9502
Number of unique SNPs	160	2393	994	2762
Number of unique gene names	96	282	243	344
Number of unique protein names	304	1032	976	1435
Percentage TMH	11	5	35	26

A.3 Counts

See supplementary Tables S3 and S4 for an overview of all amounts. Note that, for the analyses using the SARS-CoV-2 virus proteome, we labeled this by its disease (covid) to prevent typos. In supplementary Table S3 there are multiple instances where the amounts are expected to add up, yet don't, as one SNP can work on multiple isoforms. For example, there are 9,621 unique SNPs found in all proteins, of which 4,219 around found in MAPs and 6,026 in TMPs. Apparently, 624 SNPs work on a set of isoforms that contains both MAPs and TMPs.

704 **A.4 Relative positions**

705 See Supplementary Figure S2 for the distribution of the relative position of the
706 SNPs.

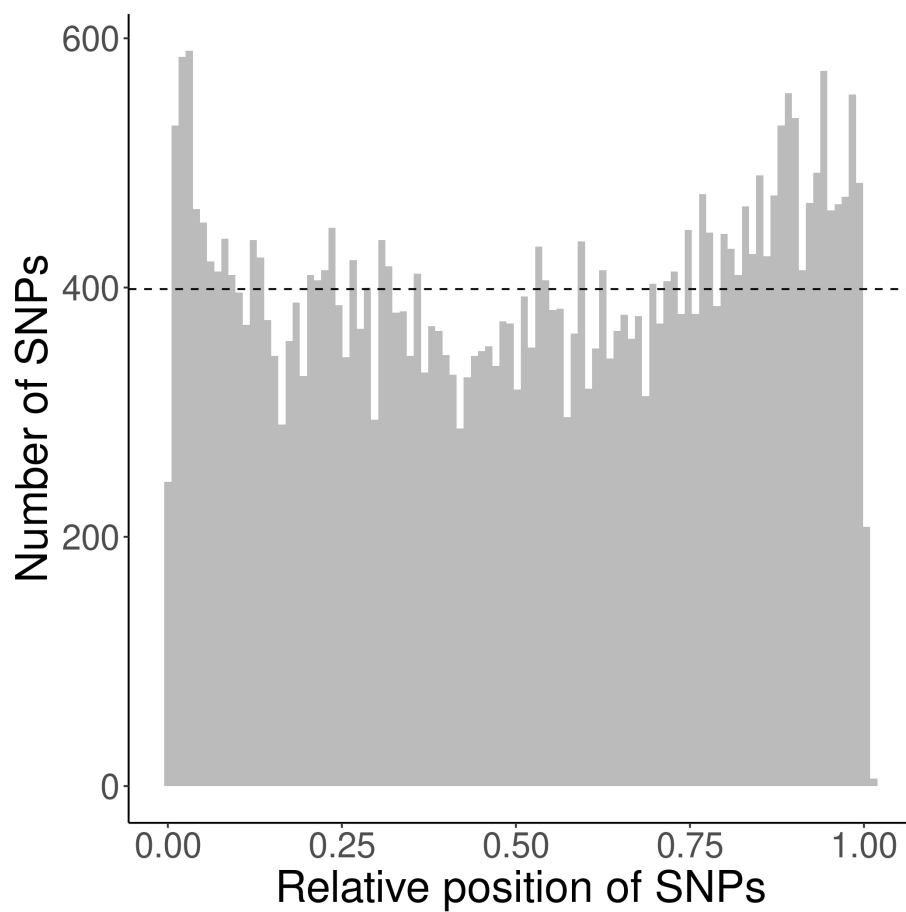


Figure S2: Distribution of the relative position of the SNPs used, where a relative position of zero denotes the first amino acid at the N-terminus, where a relative position of one indicates the last residue at the C-terminus.

Table S5: Percentage of MHC-II 14-mers overlapping with TMH. Values in brackets show the number of binders that have at least one residue overlapping with a TMH (first value) as well as the number of binders (second value). percentage used: 2

haplotype	covid	human	myco
HLA-DQA1*0101/DQB1*0501	40.433 (112/277)	31.214 (69752/223464)	32.158 (8187/25459)
HLA-DQA1*0102/DQB1*0602	22.910 (74/323)	16.167 (35753/221147)	17.950 (4608/25671)
HLA-DQA1*0301/DQB1*0302	10.381 (30/289)	10.179 (22623/222248)	11.144 (2842/25502)
HLA-DQA1*0401/DQB1*0402	11.111 (32/288)	13.135 (29319/223219)	9.890 (2524/25522)
HLA-DQA1*0501/DQB1*0201	20.430 (57/279)	16.240 (36186/222820)	14.999 (3823/25489)
HLA-DQA1*0501/DQB1*0301	15.808 (46/291)	14.106 (31046/220089)	18.969 (4878/25715)
HLA-DRB1*0101	27.119 (80/295)	19.774 (43968/222349)	22.293 (5692/25533)
HLA-DRB1*0301	14.676 (43/293)	9.801 (21831/222752)	7.956 (2025/25451)
HLA-DRB1*0401	19.231 (55/286)	15.325 (34011/221930)	18.113 (4641/25623)
HLA-DRB1*0405	12.996 (36/277)	13.684 (30380/222012)	15.837 (4036/25484)
HLA-DRB1*0701	32.877 (96/292)	21.512 (47856/222465)	29.304 (7471/25495)
HLA-DRB1*0802	23.132 (65/281)	19.339 (42859/221623)	28.805 (7358/25544)
HLA-DRB1*0901	11.565 (34/294)	13.111 (29043/221520)	16.798 (4301/25605)
HLA-DRB1*1101	25.197 (64/254)	11.924 (26582/222928)	16.103 (4101/25467)
HLA-DRB1*1201	36.897 (107/290)	15.482 (34596/223464)	20.018 (5098/25467)
HLA-DRB1*1302	13.962 (37/265)	20.121 (44798/222646)	23.141 (5935/25647)
HLA-DRB1*1501	35.206 (94/267)	21.836 (48671/222893)	25.891 (6584/25430)
HLA-DRB3*0101	9.158 (25/273)	8.496 (18884/222274)	6.819 (1740/25517)
HLA-DRB3*0202	18.657 (50/268)	13.832 (30687/221859)	15.843 (4059/25620)
HLA-DRB4*0101	23.529 (68/289)	12.749 (28376/222568)	16.221 (4131/25467)
HLA-DRB5*0101	23.776 (68/286)	11.235 (24993/222464)	14.648 (3732/25478)

A.5 Presentation of TMH-derived epitopes

See supplementary Table S5 for the percentage of MHC-II 14-mers overlapping with TMH.

710 **A.6 The percentage of TMH-derived epitopes from IEDB**

711 **epitopes**

712 We display the over-presentation of epitopes taken from the IEDB database, for
 713 two assays: an MHC ligand assay (Figure 2A) and a T cell assay (see figure S4),
 714 as a bar plot. Supplementary Table S6 below shows the exact numbers.

MHC class	Dataset	n
I	iedb_mhc_ligand	22.28% (1789/8030)
I	iedb_t_cell	35.91% (93/259)
II	iedb_mhc_ligand	10.46% (73/698)
II	iedb_t_cell	6.66% (42/631)

Table S6: Percentage of epitopes derived from a TMH for epitopes taken from the IEDB, for two different types of assays: an MHC ligand assay, as well as a T cell assay. The values between brackets show the the number of epitopes that were predicted to overlapping with a TMH per all epitopes that could be uniquely mapped to the representative human reference proteome.

715 A.7 Correlation of epitope presentation

716 In the main text of this research, we use two sources of epitopes to determine
 717 if TMH-derived epitopes are presented. The first source of epitopes are all the
 718 9-mers (for MHC-I) (and 14-mers for MHC-II) derived from a human reference
 719 proteome, where this over-presentation is displayed in figure 1A. The second
 720 source of epitopes are those that are present in the IEDB that are obtained
 721 from MHC ligand assays, as displayed in figure 2A.

722 Here we correlate between the over-presentation of TMH-derived epitopes
 723 between these two sources of data. Figure S3 shows per MHC allele the per-
 724 centage of TMH-derived epitopes, with a linear trendline.

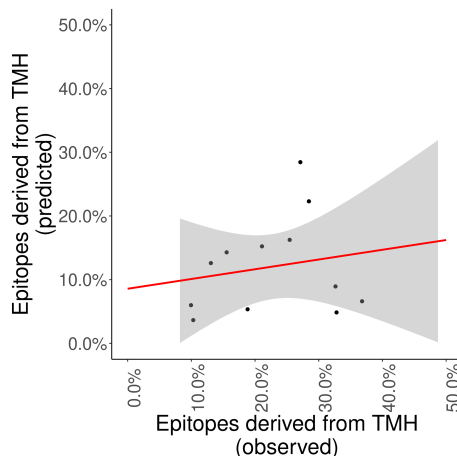


Figure S3: **TMH-derived epitopes are over-presented when using predicted as well as experimental data** For the MHC class I alleles, the over-presentation of TMH-derived epitopes is correlated between IEDB MHC ligand epitopes (horizontal axis) and the 9-mers derived from a human reference proteome (vertical axis). Alleles are listed in Table S8). The trendline shows the linear correlation between these percentages, where the gray area is the 95% confidence interval.

725 **A.8 Presentation of TMH-derived epitopes result in T cell**
 726 **responses**

727 Figure S4 shows the percentage of TMH-derived epitopes of the reported epi-
 728 topes from human origin for which T-cell responses were established. The data
 729 was obtained from the IEDB and includes only the MHC alleles used in this
 730 study. As there are many (especially class II) MHC alleles, only a small per-
 731 centage of the full IEDB data could be used.

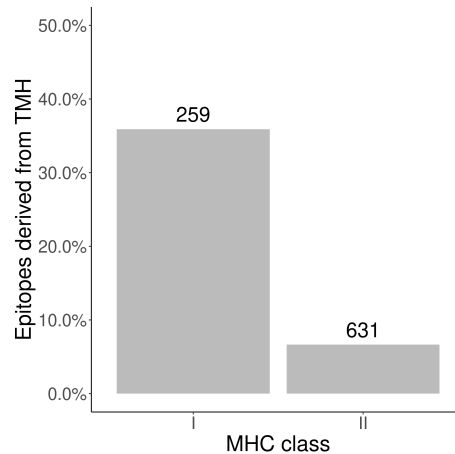


Figure S4: **TMH-derived epitopes evoke T-cell responses** The numbers above the bars denotes the number of epitopes found in the IEDB for the MHC alleles used in this study.

Table S7: Percentage of MHC-I 9-mers overlapping with TMH. Values in brackets show the number of binders that have at least one residue overlapping with a TMH (first value) as well as the number of binders (second value). percentage used: 2

haplotype	covid	human	myco
HLA-A*01:01	15.603 (44/282)	12.600 (28377/225209)	11.424 (2947/25797)
HLA-A*02:01	34.155 (97/284)	28.441 (63994/225003)	29.749 (7646/25702)
HLA-A*03:01	9.122 (27/296)	6.606 (14851/224796)	9.972 (2565/25721)
HLA-A*24:02	39.223 (111/283)	22.297 (50313/225648)	22.346 (5752/25741)
HLA-A*26:01	21.739 (65/299)	14.287 (32232/225598)	13.950 (3598/25793)
HLA-B*07:02	9.712 (27/278)	5.347 (11893/222429)	8.899 (2291/25744)
HLA-B*08:01	15.248 (43/282)	8.935 (19981/223616)	10.714 (2750/25667)
HLA-B*15:01	24.324 (72/296)	15.228 (34498/226542)	17.600 (4547/25835)
HLA-B*18:01	11.724 (34/290)	5.993 (13409/223745)	5.960 (1536/25773)
HLA-B*27:05	10.227 (27/264)	4.854 (10882/224178)	8.031 (2063/25688)
HLA-B*39:01	18.182 (50/275)	11.468 (25621/223419)	14.682 (3787/25793)
HLA-B*40:02	4.594 (13/283)	3.647 (8147/223408)	4.264 (1097/25729)
HLA-B*58:01	24.731 (69/279)	16.245 (36409/224119)	20.558 (5292/25742)

A.9 Presentation of TMH-derived epitopes

See supplementary Table S7 for the percentage of MHC-I 9-mers overlapping with TMH.

Supplementary Table S8 shows the shorthand notation for the HLA alleles.

Supplementary Tables S7 and S5 show the exact number of binders, binders that overlap with TMHs and the percentage of binders that overlap with TMHs, as visualized by figure 1A.

index	haplotype_name
1	HLA-A*01:01
2	HLA-A*02:01
3	HLA-A*03:01
4	HLA-A*24:02
5	HLA-A*26:01
6	HLA-B*07:02
7	HLA-B*08:01
8	HLA-B*18:01
9	HLA-B*27:05
10	HLA-B*39:01
11	HLA-B*40:02
12	HLA-B*58:01
13	HLA-B*15:01
1	HLA-DRB1*0101
2	HLA-DRB1*0301
3	HLA-DRB1*0401
4	HLA-DRB1*0405
5	HLA-DRB1*0701
6	HLA-DRB1*0802
7	HLA-DRB1*0901
8	HLA-DRB1*1101
9	HLA-DRB1*1201
10	HLA-DRB1*1302
11	HLA-DRB1*1501
12	HLA-DRB3*0101
13	HLA-DRB3*0202
14	HLA-DRB4*0101
15	HLA-DRB5*0101
16	HLA-DQA1*0501/DQB1*0201
17	HLA-DQA1*0501/DQB1*0301
18	HLA-DQA1*0301/DQB1*0302
19	HLA-DQA1*0401/DQB1*0402
20	HLA-DQA1*0101/DQB1*0501
21	HLA-DQA1*0102/DQB1*0602

Table S8: Abbreviations of the haplotype names

Goal	Tool	Reference
Predict topology	TMHMM	[9]
Predict topology	PureseqTM	[14]
Predict epitopes MHC-I	epitope-prediction	[7]
Predict epitopes MHC-II	NetMHCIIpan	[45, 46]
Call TMHMM from R	tmhmm	[47]
Call PureseqTM from R	pureseqtmr	[48]
Call NetMHCIIpan from R	netmhc2pan	[49]
Work with IEDB	iedbr	[50]
Work with rentrez	sprentrez	[51]
Combine all	bbbq	[52]

Table S9: Overview of all software used in this research.

739 A.10 Prediction software used

740 For this research, we needed software to predict protein topology, as well as the
741 MHC-I and MHC-II binding affinities of epitopes. We selected our software, by
742 searching the scientific literature to identify the most recent free and open source
743 (FOSS) prediction software. This was done by searching for papers that (1) cite
744 older prediction software, and (2) present a novel method to make predictions.
745 As a starting point, per type of prediction software, a review paper was used
746 ([53] for protein topology, [54] for MHC-I binding affinities and [55] for MHC-II
747 binding affinities).

748 There are multiple computational tools developed to predict which parts of
749 a protein forms a TMH. In 2001, multiple of such prediction tools have been
750 compared [53], of which TMHMM [9] turned out to be the most accurate, as
751 is used in the previous study [7]. However, TMHMM has a restrictive software
752 license and is nearly two decades old. Therefore, PureseqTM [14], was also used
753 in this study, which has been more recently developed and has a free software
754 license.

755 For MHC-I, there are multiple computational tools developed to predict epi-
756 topes. According to [54], at that time, NetMHCcons [56] gave the best predic-

757 tions. We used the same tool as used in our earlier study, **epitope-prediction**
758 [7],

759 Also for MHC-II, there are multiple computational tools developed to pre-
760 dict epitopes, such as using a trained neural network [55] or a Gibbs sam-
761 pling approach [57]. According to [54], in 2011, from a set of multiple tools,
762 NetMHCIIpan [45, 46] made the most accurate predictions. The most recent
763 FOSS tool available now appears to be MHCnuggets [20], which can do both
764 MHC-I and MHC-II predictions. As we already use **epitope-prediction** [7]
765 for MHC-I predictions, we use MHCnuggets only for MHC-II predictions.

766 To retrieve the data from the NCBI databases the **rentrez** R package [58]
767 was used that calls the NCBI database’s API. The NCBI database provides a
768 stable user experience for all users, by limiting its API to 3 calls per second
769 per user. Additionally, the API splits the result of a bigger query into multiple
770 pages, each of which needs one API call. The **sprentrez** package [51] provides
771 for bigger queries of multiple (and delayed) API calls.

772 To retrieve the data from the IEDB databases [18], the **iedbr** R package [50]
773 was written, to calls the IEDB database’s API. Similar to the NCBI database,
774 the IEDB has a limit to 1 call per second per user and allows a query results to
775 return 10k results maximally. The **iedbr** package [50] allows for bigger queries.

776 A.11 Prediction software written

777 The R programming language is used for the complete experiment, including the
778 analysis. The complete experiment is bundled in the 'bbbq' R package, which
779 is dependent on 'tmhmm', 'pureseqtmr', 'epitope-prediction' and 'mhc-nuggets-r'
780 as described below.

781 The R package 'tmhmm' was developed to do the similar topology predic-
782 tions as our earlier study (that used 'TMHMM'), yet in an automated way.
783 'TMHMM' has a restrictive software license [9] and allows a user to download a
784 pre-compiled executable after confirmation that he/she is in academia. The R
785 package respects this restriction and allows the user to install and use TMHMM
786 from within R, as done in this study. 'tmhmm' has been submitted to and is
787 accepted by the Comprehensive R Archive Network (CRAN).

788 To be able to call, from R, the TMH prediction software 'PureseqTM' [14],
789 which is written in C, the package 'pureseqtmr' has been developed. 'purese-
790 qtmr' allows to install 'PureseqTM' and use most of its features. 'pureseqtmr'
791 has been submitted to and is accepted by CRAN.

792 MHCnuggets is a free and open-source Python package to predict epitope
793 affinity for many MHC-I and MHC-II variants [20]. The R package 'mhc-
794 nuggets-r' allows one to install and use MHCnuggets from within R. Also 'mhc-
795 nuggets-r' has been submitted to and is accepted by CRAN.

796 To reproduce the full experiment presented in this paper, the functions
797 needed are bundled in the 'bbbq' R package. This package is too specific to
798 be submitted to CRAN.

Table S10: Percentage of spots and spots that overlap with a TMH

target	mhc_class	n_spots	n_spots_tmh	f_tmh
covid	1	14207	1124	7.91
covid	2	14137	1245	8.81
human	1	11220940	598391	5.33
human	2	11118448	672273	6.05
myco	1	1299707	98613	7.59
myco	2	1279742	108419	8.47

799 A.12 Prediction of percentage of epitopes overlapping with 800 a TMH

801 Supplementary Table S10 shows an overview of the findings, where a target
802 specifies the source of the proteome, where `covid` denotes SARS-CoV-2 and
803 `myco` denotes *Mycobacterium tuberculosis*. `mhc_class` denotes the MHC class,
804 `n_spots` the number of possible 9-mers (for MHC-I) or 14-mers (for MHC-II)
805 possible. `n_spots_tmh` the number of epitopes that overlapped with a TMH
806 that were binders. `f_tmh` the percentage of peptides that had at least 1 residue
807 overlapping with a TMH.

808 **A.13 Minor methods**

809 These are details that are removed from the 'Methods' section.

810 PureseqTM does not predict the topology of proteins that have less than
811 three amino acids. The TRDD1 ('T cell receptor delta diversity 1') protein,
812 however, is two amino acids long. The R package `pureseqtmr`, however, predicts
813 that mono- and di-peptides are cytosolic.

814 **A.14 Minor discussion**

815 These are details that are removed from the 'Discussion' section.

816 In this experiment we predicted epitopes that overlap with TMHs from a
817 human, bacterial and viral proteome, would these proteins be expressed in a
818 human host. Bacteria, however have different cell membranes and cell walls,
819 hence different structural requirements for a TMH. Both topology prediction
820 tools were trained to recognize human TMHs, thus we cannot be sure that
821 the transmembrane regions predicted in bacterial proteins are actually part of a
822 TMH. For the purpose of this study, we assume the error in topology predictions
823 to be unbiased way towards topology. In other words: that a bacterial TMH is
824 incorrectly predicted to be absent just as often as it is incorrectly predicted to
825 be present elsewhere.

826 Regarding the evolutionary conservation of TMHs using SNPs, again, it is
827 estimated that approximately ten percent of SNPs is a false positive that result
828 from the methods to determine a SNP. One example is that sequence variations
829 are incorrectly detected due to highly similar duplicated sequences [59]. We
830 assume that these duplications occur as often in TMHs as in regions around
831 these, hence we expect this not to affect our results.

832 In our evolutionary experiment, we removed variations that were synony-
833 mous mutations (i.e. resulted in the same amino acid, from a different genetic

code) from our analysis. There is evidence, however, that these synonymous mutations do have an effect and may even be evolutionary selected for [60]. As the possible effect of synonymous mutations is ignored by our topology prediction software, we do so as well.

838 A.15 Relative presentation of TMH-derived epitopes

839 To compare the over-presentation of TMH-derived epitopes between the differ-
 840 ent proteomes, we normalized this percentages in such a way that 1.0 is the
 841 percentage of TMH-derived epitopes that would be expected by chance. Fig-
 842 ure S5 and S6 show these normalized values for the MHC-I and MHC-II alleles
 843 respectively.

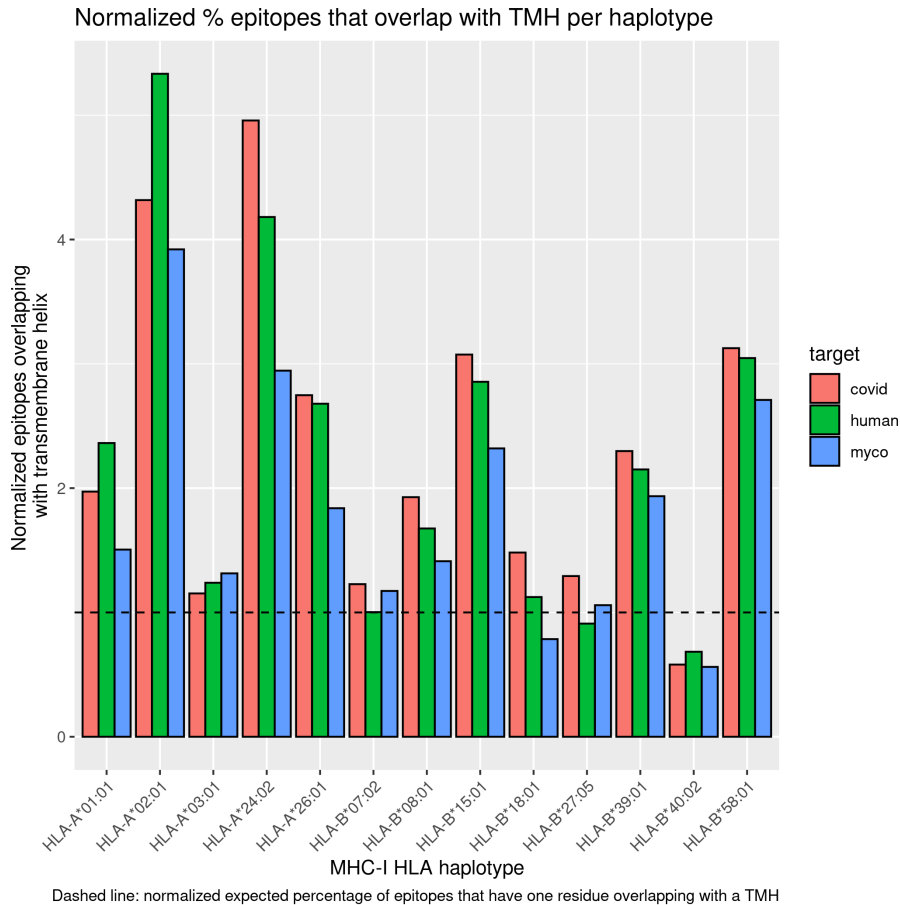


Figure S5: Normalized proportion of MHC-I epitopes overlapping with TMHs for human, viral and bacterial proteomes. Legend: covid = SARS-CoV-2, human = *Homo sapiens*, myco = *Mycobacterium tuberculosis*

844 To determine the additional over-presentation of TMH-derived epitopes in

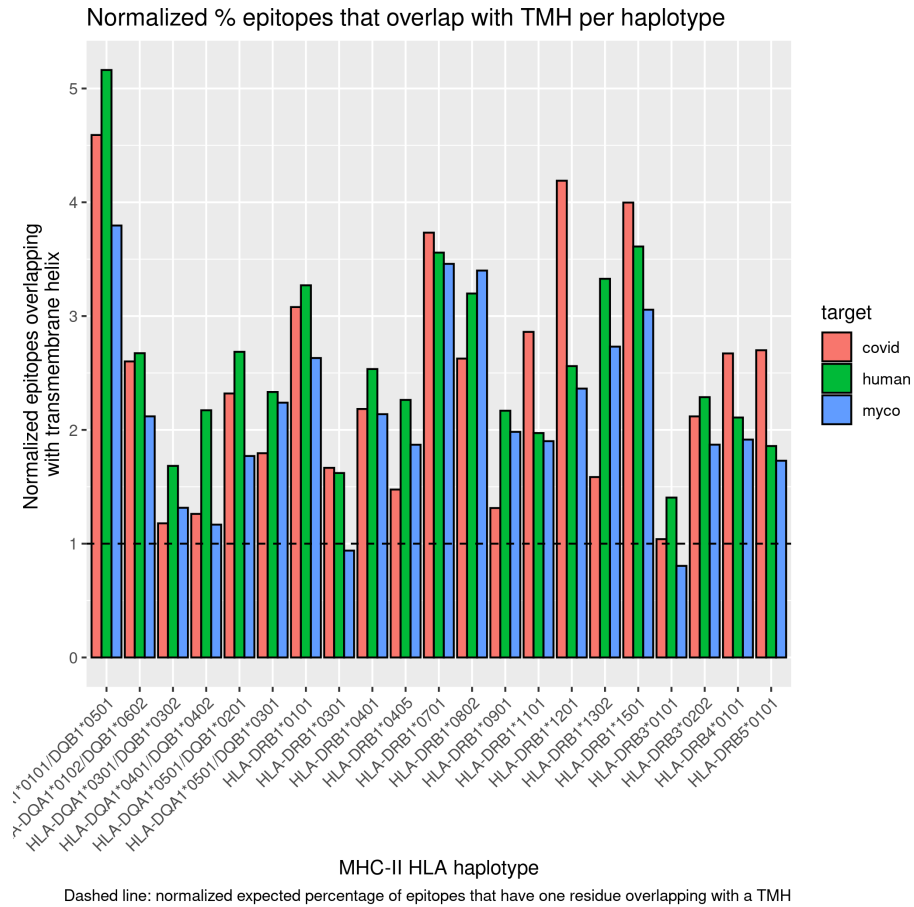


Figure S6: Normalized proportion of MHC-II epitopes overlapping with TMHs for human, viral and bacterial proteomes. Legend: covid = SARS-CoV-2, human = *Homo sapiens*, myco = *Mycobacterium tuberculosis*

845 MHC-II (as compared to MHC-I), we normalized the data to enable a side-
 846 by-side comparison. The percentage of TMH-derived epitopes presented was
 847 normalized to the expected percentage of TMH-derived epitopes, where 1.0
 848 denotes that the percentage of presented TMH-derived epitopes matches the
 849 values as expected by chance. The normalized values per MHC allele are shown
 850 in figure S7. To compare the TMH-derived over-presentation per MHC class,
 851 we grouped the normalized values per allele, and plot the mean and standard
 852 error, as shown in figure S8.

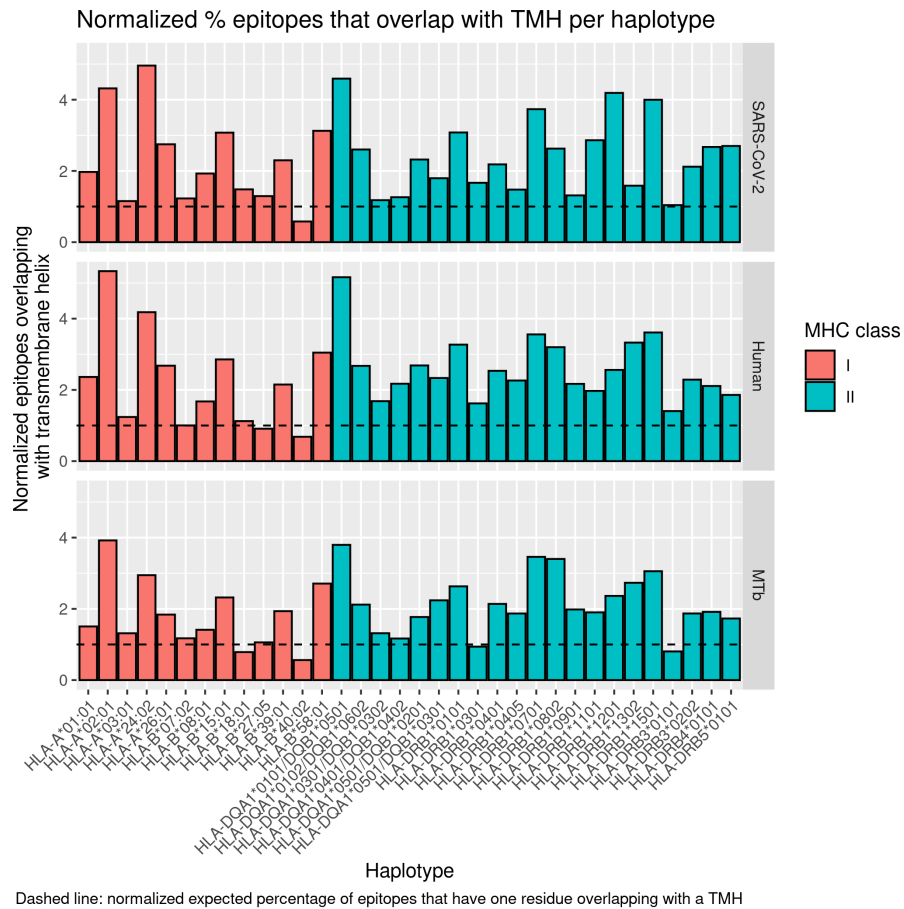


Figure S7: Normalized proportion of MHC-I and MHC-II epitopes overlapping with TMHs, for the different MHC alleles and proteomes

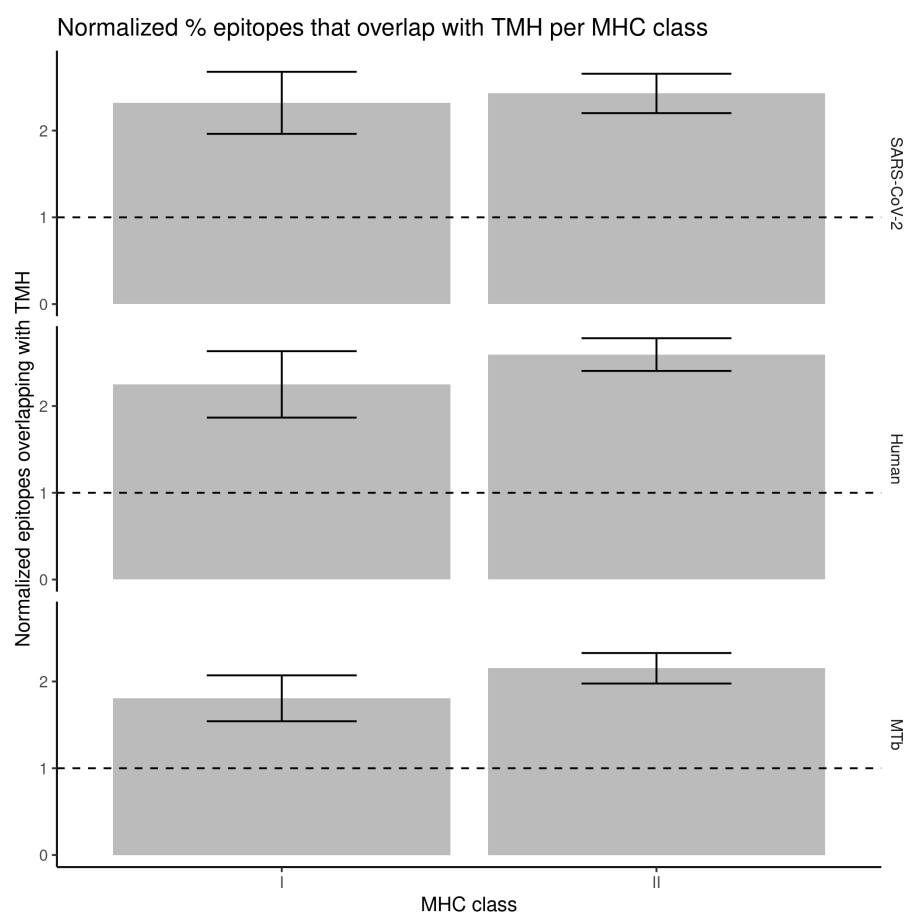


Figure S8: Normalized proportion of MHC-I and MHC-II epitopes overlapping with TMHs, for the different MHC classes and proteomes. Error bars denote the standard error.

853 **A.16 Evolutionary conservation**

854 Figure S9 shows the distribution of the number of SNPs per gene name, at the
 855 date we started the experiment, at December 14th 2020.

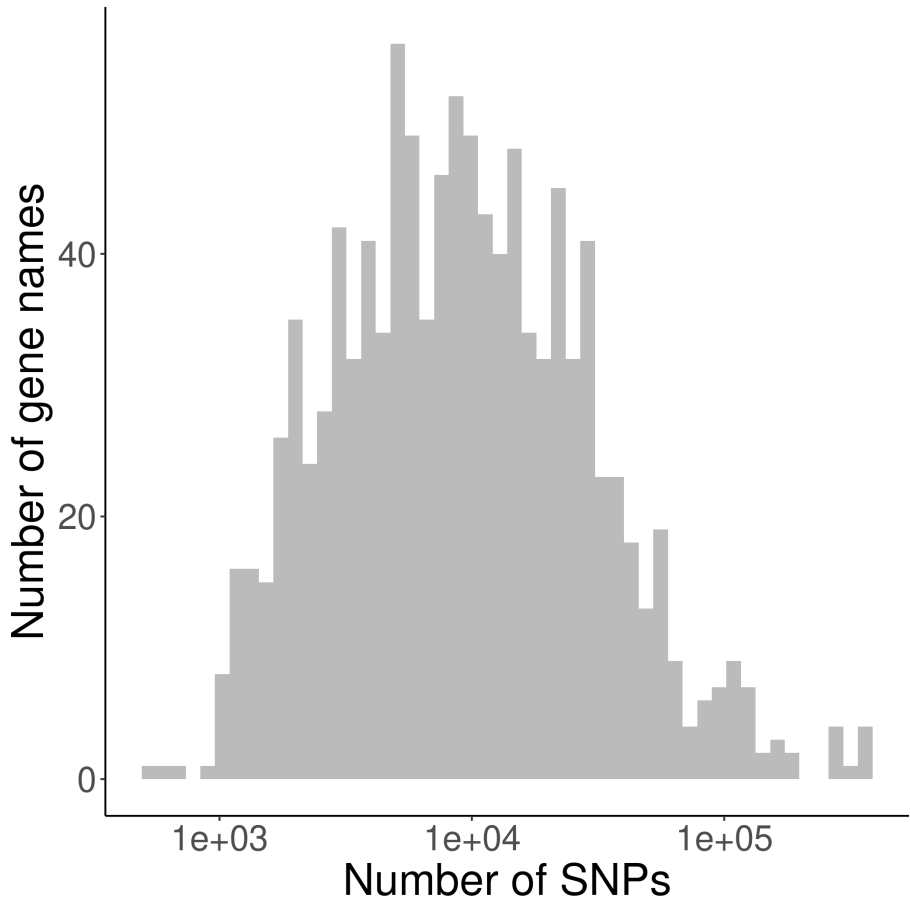


Figure S9: Distribution of the number of SNPs per gene name in the NCBI database.

856 To verify if SNPs were sampled uniformly over proteins, we show the dis-
 857 tribution of the relative position in figure S2. We find no clear evidence of a
 858 bias.

859 Supplementary Table S11 shows the statistics for all SNPs, where supple-

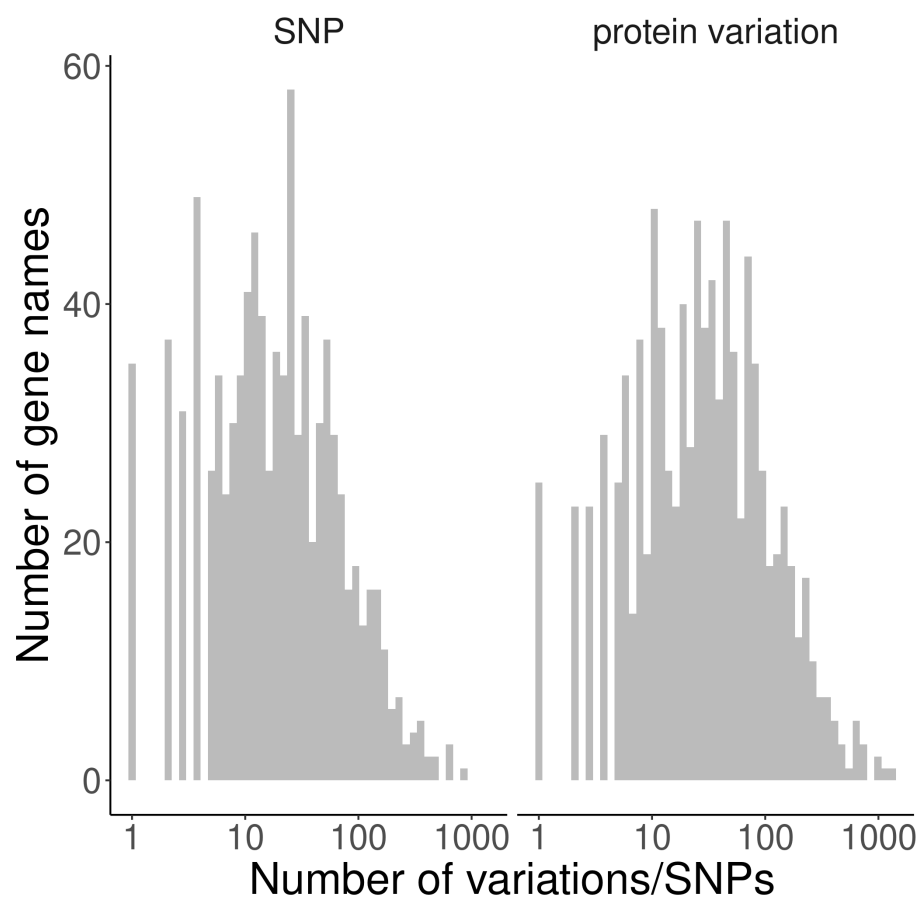


Figure S10: Distribution of the number of protein variations and SNPs per gene name processed.

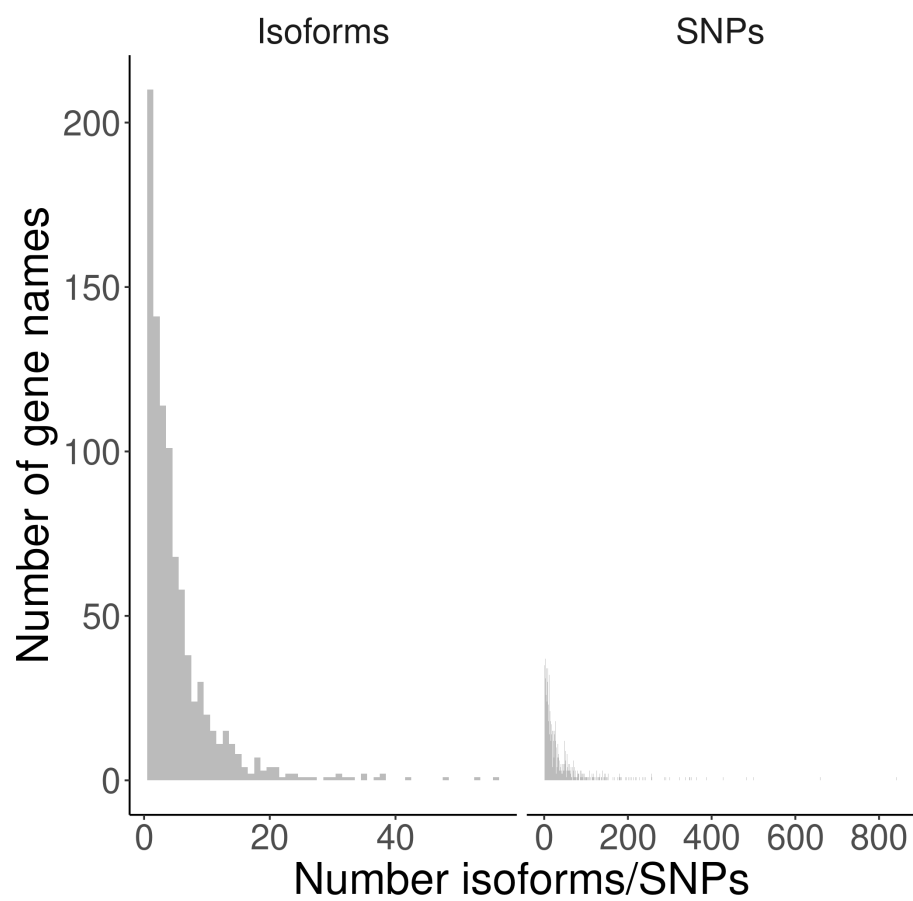


Figure S11: Histogram of the number of proteins found per gene name. Most often, a gene name is associated with one protein.

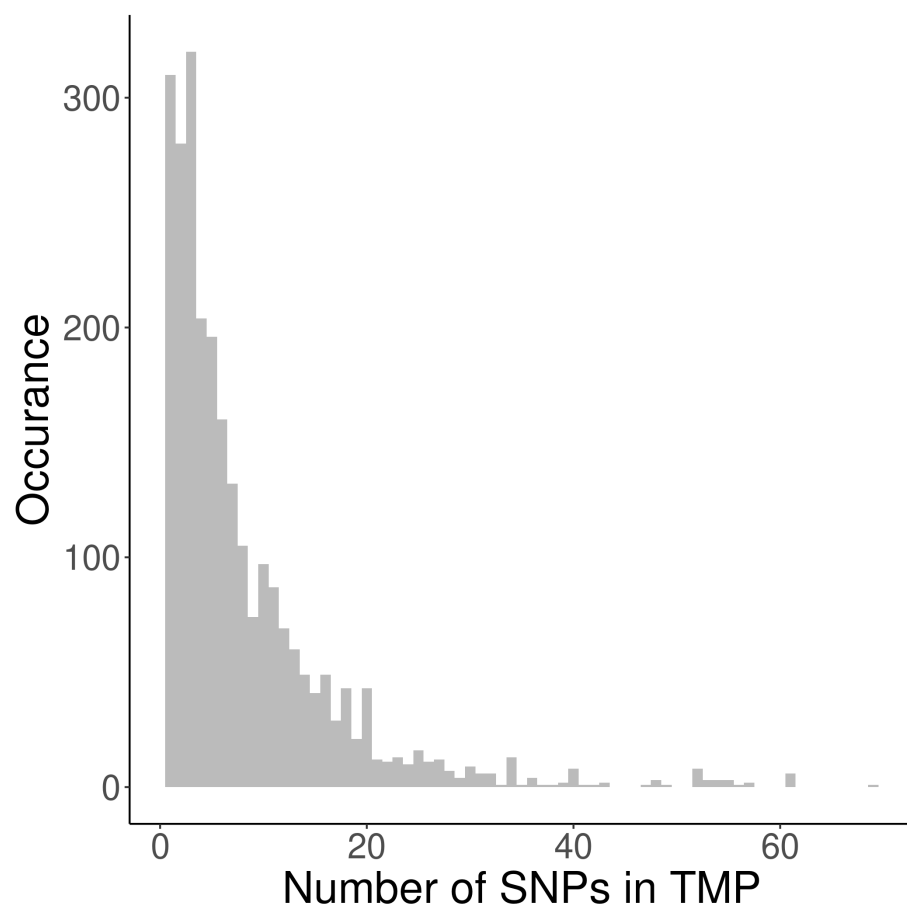


Figure S12: Histogram of the number of SNPs per trans-membrane protein.

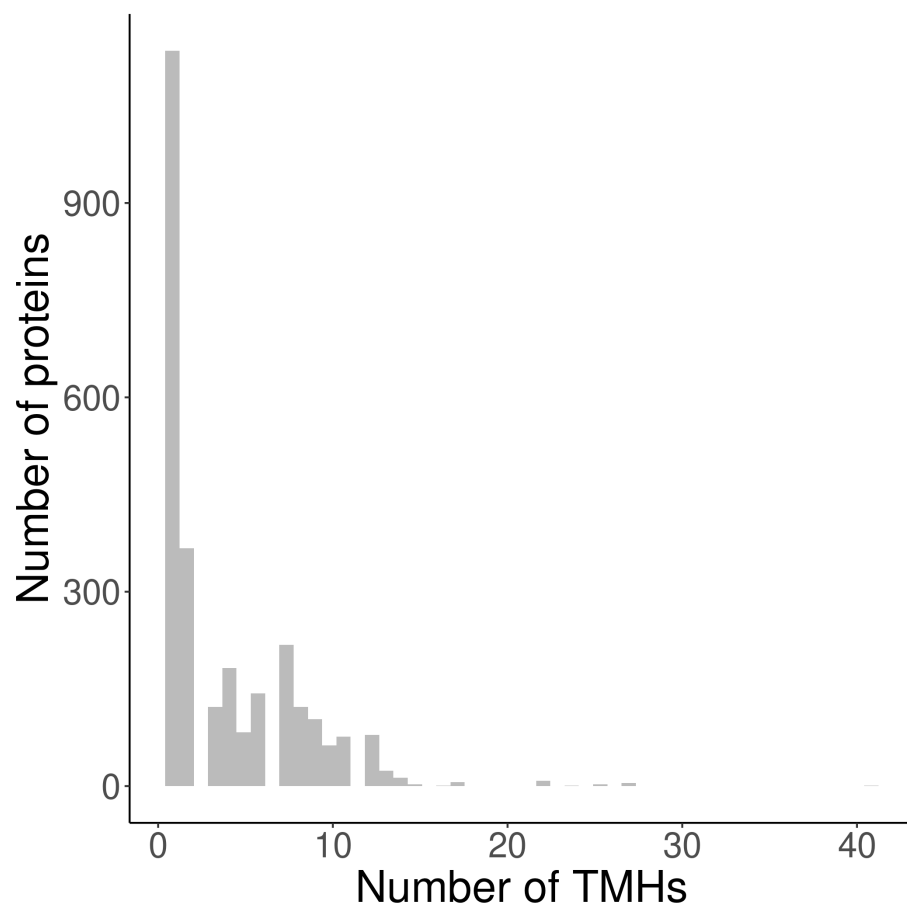


Figure S13: Histogram of the number of TMHs predicted per protein, for the trans-membrane proteins used.

Table S11: Statistics for all TMPs. p = p value. n = number of SNPs. n_{success} = number of SNPs found in TMHs (dashed blue line). $E(n_{\text{success}})$ = expected number of SNPs to be found in TMHs.

parameter	value
p	6.820823e-11
n	21208
n_{success}	3803
$E(n_{\text{success}})$	4140.56

Table S12: Statistics for the single-spanners. p = p value. n = number of SNPs in single-spanners. n_{success} = number of SNPs found in TMHs of single-spanners (dashed blue line). $E(n_{\text{success}})$ = expected number of SNPs to be found in TMHs of single-spanners.

parameter	value
p	0.3189532
n	8186
n_{success}	452
$E(n_{\text{success}})$	462.1535

860 mentary Tables S12 and S13 show the statistics for only single-spanners and
861 multi-spanners respectively.

Table S13: Statistics for the multi-spanners. p = p value. n = number of SNPs in multi-spanners. n_{success} = number of SNPs found in TMHs of multi-spanners (dashed blue line). $E(n_{\text{success}})$ = expected number of SNPs to be found in TMHs of multi-spanners.

parameter	value
p	8.315841e-12
n	13022
n_{success}	3351
$E(n_{\text{success}})$	3678.406

862 **A.17 Presentation of TMH-derived epitopes when two amino** 863 **acids overlap**

864 In our experiment, we define a TMH-derived epitope as a peptide that overlaps
865 with a TMH for at least one amino acid. One could argue that we should use
866 a higher number of overlapping amino acids, so that the epitope has a higher
867 TMH coverage. We chose not to, for two reason: (1) epitopes that overlap with
868 a TMH for 1 AA already, cannot be processed by the proteasome in a known and
869 conventional way as it still requires extraction from the membrane (2) whatever
870 number of overlapping amino acids we use, we expect the pattern to be the same
871 as the chance that an epitope stems from a TMH is equally reduced. However,
872 using only 1 AA gives the most TMH-derived epitopes and hence the highest
873 statistical power.

874 To prove this point, we did exactly the same analysis as shown in Figure
875 1A, yet with defining a TMH-derived epitope as an epitope that overlaps with
876 a TMH for at least 2 AAs, as shown in Figure S14. As these two figures look
877 identical, we also added the counts as numbers, with Table S14 showing the
878 same data as S5, except the former uses 2 AAs overlap. Likewise, Table S15
879 showing the same data as S7, except the former uses 2 AAs overlap.

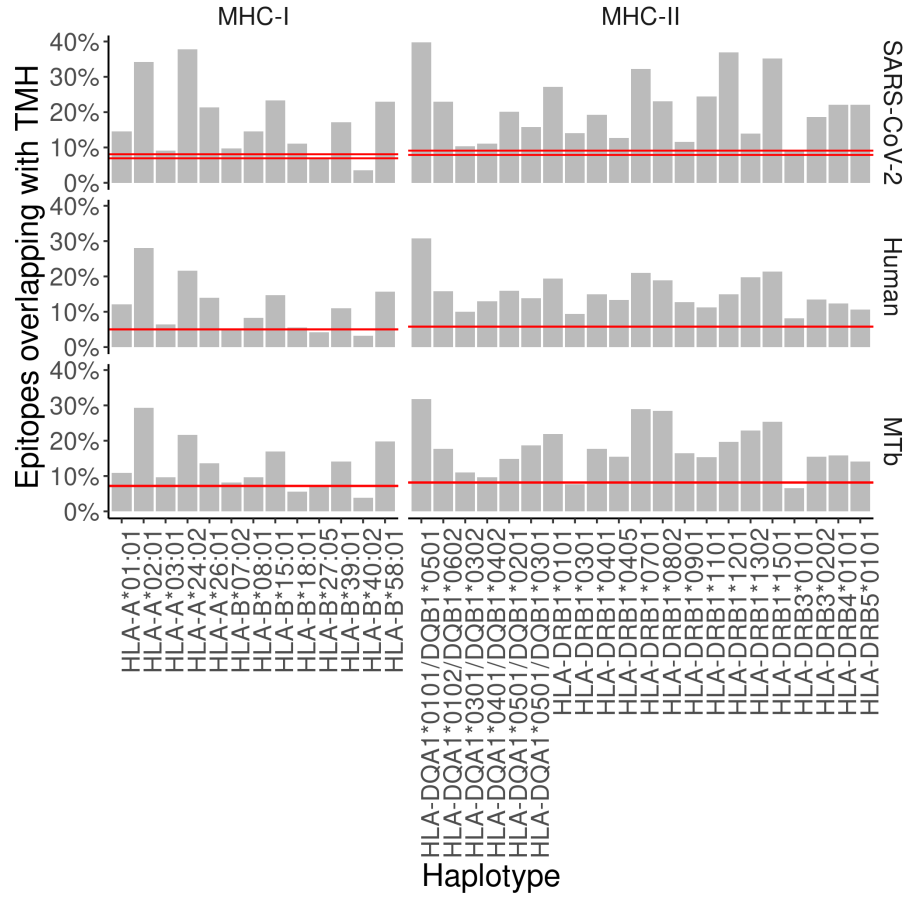


Figure S14: The percentage of epitopes for MHC-I and -II alleles that are predicted to overlap with TMHs (for at least two amino acids) for the proteomes of SARS-CoV-2 (top row), human (middle row) and *M. tuberculosis* (bottom row). The pair of dashed lines in each plot indicate the lower and upper bound of the 99% confidence interval. See supplementary Tables S14 and S15 for the exact TMH and epitope counts.

Table S14: Percentage of MHC-II 14-mers overlapping with TMH. Values in brackets show the number of binders that have at least two residues overlapping with a TMH (first value) as well as the number of binders (second value). percentage used: 2

haplotype	covid	human	myco
HLA-DQA1*0101/DQB1*0501	39.711 (110/277)	30.813 (68855/223464)	31.777 (8090/25459)
HLA-DQA1*0102/DQB1*0602	22.910 (74/323)	15.858 (35070/221147)	17.713 (4547/25671)
HLA-DQA1*0301/DQB1*0302	10.381 (30/289)	9.996 (22217/222248)	10.960 (2795/25502)
HLA-DQA1*0401/DQB1*0402	11.111 (32/288)	12.915 (28829/223219)	9.670 (2468/25522)
HLA-DQA1*0501/DQB1*0201	20.072 (56/279)	15.969 (35582/222820)	14.830 (3780/25489)
HLA-DQA1*0501/DQB1*0301	15.808 (46/291)	13.890 (30570/220089)	18.682 (4804/25715)
HLA-DRB1*0101	27.119 (80/295)	19.401 (43139/222349)	21.944 (5603/25533)
HLA-DRB1*0301	13.993 (41/293)	9.415 (20972/222752)	7.638 (1944/25451)
HLA-DRB1*0401	19.231 (55/286)	14.925 (33122/221930)	17.652 (4523/25623)
HLA-DRB1*0405	12.635 (35/277)	13.298 (29523/222012)	15.469 (3942/25484)
HLA-DRB1*0701	32.192 (94/292)	21.057 (46845/222465)	28.884 (7364/25495)
HLA-DRB1*0802	23.132 (65/281)	18.909 (41907/221623)	28.496 (7279/25544)
HLA-DRB1*0901	11.565 (34/294)	12.730 (28199/221520)	16.505 (4226/25605)
HLA-DRB1*1101	24.409 (62/254)	11.282 (25151/222928)	15.357 (3911/25467)
HLA-DRB1*1201	36.897 (107/290)	14.985 (33487/223464)	19.633 (5000/25467)
HLA-DRB1*1302	13.962 (37/265)	19.774 (44027/222646)	22.903 (5874/25647)
HLA-DRB1*1501	35.206 (94/267)	21.341 (47568/222893)	25.415 (6463/25430)
HLA-DRB3*0101	9.158 (25/273)	8.145 (18105/222274)	6.556 (1673/25517)
HLA-DRB3*0202	18.657 (50/268)	13.445 (29830/221859)	15.457 (3960/25620)
HLA-DRB4*0101	22.145 (64/289)	12.341 (27467/222568)	15.856 (4038/25467)
HLA-DRB5*0101	22.028 (63/286)	10.677 (23753/222464)	14.138 (3602/25478)

Table S15: Percentage of MHC-I 9-mers overlapping with TMH. Values in brackets show the number of binders that have at least two residues overlapping with a TMH (first value) as well as the number of binders (second value). percentage used: 2

haplotype	covid	human	myco
HLA-A*01:01	14.539 (41/282)	12.092 (27232/225209)	10.912 (2815/25797)
HLA-A*02:01	34.155 (97/284)	28.037 (63085/225003)	29.360 (7546/25702)
HLA-A*03:01	9.122 (27/296)	6.388 (14361/224796)	9.673 (2488/25721)
HLA-A*24:02	37.809 (107/283)	21.677 (48913/225648)	21.643 (5571/25741)
HLA-A*26:01	21.405 (64/299)	13.905 (31370/225598)	13.632 (3516/25793)
HLA-B*07:02	9.712 (27/278)	4.880 (10854/222429)	8.184 (2107/25744)
HLA-B*08:01	14.539 (41/282)	8.218 (18376/223616)	9.662 (2480/25667)
HLA-B*15:01	23.311 (69/296)	14.686 (33269/226542)	16.961 (4382/25835)
HLA-B*18:01	11.034 (32/290)	5.603 (12537/223745)	5.560 (1433/25773)
HLA-B*27:05	6.818 (18/264)	4.171 (9350/224178)	7.054 (1812/25688)
HLA-B*39:01	17.091 (47/275)	10.983 (24538/223419)	14.159 (3652/25793)
HLA-B*40:02	3.534 (10/283)	3.251 (7264/223408)	3.852 (991/25729)
HLA-B*58:01	22.939 (64/279)	15.627 (35022/224119)	19.793 (5095/25742)