

1 **Structure-based modeling of SARS-CoV-2 peptide/HLA-A02 antigens**

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3 Santrupti Nerli¹ and Nikolaos G. Sgourakis²

4 Email: nsgourak@ucsc.edu

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6 ¹Department of Biomolecular Engineering, University of California Santa Cruz, Santa Cruz, CA
7 95064, USA.

8 ²Department of Chemistry and Biochemistry, University of California Santa Cruz, Santa Cruz, CA
9 95064, USA.

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11 **ABSTRACT**

12 As a first step toward the development of diagnostic and therapeutic tools to fight the Coronavirus
13 disease (COVID-19), it is important to characterize CD8⁺ T cell epitopes in the SARS-CoV-2
14 peptidome that can trigger adaptive immune responses. Here, we use RosettaMHC, a comparative
15 modeling approach which leverages existing high-resolution X-ray structures from peptide/MHC
16 complexes available in the Protein Data Bank, to derive physically realistic 3D models for high-
17 affinity SARS-CoV-2 epitopes. We outline an application of our method to model 439 9mer and
18 279 10mer predicted epitopes displayed by the common allele HLA-A*02:01, and we make our
19 models publicly available through an online database (<https://rosettamhc.chemistry.ucsc.edu>). As
20 more detailed studies on antigen-specific T cell recognition become available, RosettaMHC
21 models of antigens from different strains and HLA alleles can be used as a basis to understand the
22 link between peptide/HLA complex structure and surface chemistry with immunogenicity, in the
23 context of SARS-CoV-2 infection.

24

25 An ongoing pandemic caused by the novel SARS coronavirus (SARS-CoV-2) has become the
26 focus of extensive efforts to develop vaccines and antiviral therapies (1). Immune modulatory
27 interferons, which promote a widespread antiviral reaction in infected cells, and inhibition of pro-
28 inflammatory cytokine function through anti-IL-6/IL-6R antibodies, have been proposed as
29 possible COVID-19 therapies (2, 3). However, stimulating a targeted T cell response against
30 specific viral antigens is hampered by a lack of detailed knowledge of the immunodominant
31 epitopes displayed by common Human Leukocyte Antigen (HLA) alleles across individuals
32 (public epitopes). The molecules of the class I major histocompatibility complex (MHC-I, or HLA
33 in humans) display on the cell surface a diverse pool of 8 to 15 amino acid peptides derived from
34 the endogenous processing of proteins expressed inside the cell (4). This MHC-I restriction of
35 peptide antigens provides jawed vertebrates with an essential mechanism for adaptive immunity:
36 surveillance of the displayed peptide/MHC-I (pMHC-I) molecules by CD8⁺ cytotoxic T-
37 lymphocytes allows detection of aberrant protein expression patterns, which signify viral infection
38 and can trigger an adaptive immune response (5). A recent study has shown important changes in
39 T cell compartments during the acute phase of SARS-CoV-2 infection (6), suggesting that the
40 ability to quantify antigen-specific T cells would provide new avenues for understanding the

41 expansion and contraction of the TCR repertoire in different disease cohorts and clinical settings.
42 Given the reduction in breadth and functionality of the naïve T cell repertoire during aging (7),
43 identifying a minimal set of viral antigens that can elicit a protective response will enable the
44 design of diagnostic tools to monitor critical gaps in the T cell repertoire of high-risk cohorts,
45 which can be addressed using peptide or epitope string DNA vaccines (8).
46 Human MHC-I molecules are extremely polymorphic, with thousands of known alleles in the
47 classical HLA-A, -B and -C loci. Specific amino acid polymorphisms along the peptide-binding
48 groove (termed A-F pockets) define a repertoire of 10^4 - 10^6 peptide antigens that can be recognized
49 by each HLA allotype (9, 10). Several machine-learning methods have been developed to predict
50 the likelihood that a target peptide will bind to a given allele (reviewed in (11)). Generally these
51 methods make use of available data sets in the Immune Epitope Database (12) to train artificial
52 neural networks that predict peptide processing, binding and display, and their performance varies
53 depending on peptide length and HLA allele representation in the database. Structure-based
54 approaches have also been proposed to model the bound peptide conformation *de novo* (reviewed
55 in (13)). These approaches utilize various algorithms to optimize the backbone and side chain
56 degrees of freedom of the peptide/MHC structure according to an all-atom scoring function,
57 derived from physical principles (14–16), that can be further enhanced using modified scoring
58 terms (17) or mean field theory (18). While these methods do not rely on large training data sets,
59 their performance is affected by bottlenecks in sampling of different backbone conformations, and
60 any possible structural adaptations of the HLA peptide-binding groove.
61 Predicting the bound peptide conformation whose N- and C- termini are anchored within a fixed-
62 length groove is a tractable modeling problem that can be addressed using standard comparative
63 modeling approaches (19). In previous work focusing on the HLA-B*15:01 and HLA-A*01:01
64 alleles in the context of neuroblastoma neoantigens, we have found that a combined backbone and
65 side chain optimization approach can yield accurate pMHC-I models for a pool of target peptides,
66 provided that a reliable template of the same allele and peptide length can be identified in the
67 database (20). In this approach (RosettaMHC), a local optimization of the backbone degrees of
68 freedom is sufficient to capture minor (within 0.5 Å heavy atom RMSD) changes of the target
69 peptide backbone relative to the conformation of the peptide in the template, used as a starting
70 point. For HLA-A*02:01, the most common HLA allele among disease-relevant population
71 cohorts (21), there is a large number of high-resolution X-ray structures available in the PDB (22),
72 suggesting that a similar principle can be applied to produce models of candidate epitopes directly
73 from the proteome of a pathogen of interest. Here, we apply RosettaMHC to all HLA-A*02:01
74 epitopes predicted directly from the ~30 kbp SARS-CoV-2 genome, and make our models publicly
75 available through an online database. The computed binding energies of our models can be used
76 as an additional validation layer to select high-affinity epitopes from large peptide sets. As detailed
77 epitope mapping data from high-throughput tetramer staining (23–25) and T cell functional
78 screens (26) become available, the models presented here can provide a toehold for understanding
79 links between pMHC-I antigen structure and immunogenicity, with actionable value for the
80 development of peptide vaccines to combat the disease.

81 **Materials and Methods**

82 *Identification of SARS-CoV-2 peptide epitopes*

83 The SARS-CoV-2 protein sequences (https://www.ncbi.nlm.nih.gov/nucore/NC_045512.2) were
84 obtained from NCBI and used to generate all possible peptides of lengths 9 and 10 (9,621 9mer
85 and 9,611 10mer peptides). We used NetMHCpan-4.0 (27) to derive binding scores to HLA-
86 A*02:01, and retained only peptides classified as strong or weak binders (selected using the default
87 percentile rank cut-off values). The binding classification was performed using eluted ligand
88 likelihood predictions. While in this study we use NetMHCpan-4.0 predictions as inputs to select
89 candidate epitopes for structure modeling, our workflow is fully compatible with any alternative
90 epitope prediction method.

91 92 *Selection of PDB templates*

93 To model SARS-CoV-2 / HLA-A*02:01 antigens, we identified 3D structures from the PDB that
94 can be used as templates for comparative modeling. First, we selected all HLA-A02 X-ray
95 structures that are below 3.5 Å resolution and retained only those that have 100% identity to the
96 HLA-A*02:01 heavy chain sequence (residues 1-180). We found 241 template structures bound
97 to epitopes of lengths from 8 to 15 residues (of which 170 are 9mers and 61 are 10mers). For each
98 SARS-CoV-2 target peptide, we selected a set of candidate templates of the same length by
99 matching the target peptide anchor positions (P2 and P9/P10) to each peptide in the template
100 structures. Then, we used the BLOSUM62 (28) substitution matrix to score all remaining positions
101 in the pairwise alignment of the target/template sequences, and the PDB template with the top
102 alignment score was selected for modeling. For target peptides where we found no templates which
103 match both peptide anchors, we scored all positions in the pairwise alignment and selected the top
104 scoring template for modeling.

105 106 *RosettaMHC modeling framework and database*

107 RosettaMHC (manuscript in preparation) is a comparative modeling protocol developed using
108 PyRosetta (29) to model pMHC-I complexes. The program accepts as input a list of peptide
109 sequences, an HLA allele definition and a template PDB file (selected as described in the previous
110 step). To minimize "noise" in the simulation from parts of the MHC-I fold that do not contribute
111 to peptide binding, only the α_1 and α_2 domains are considered in all steps. For each peptide, a full
112 alignment between the target and template peptide/MHC sequences is performed using clustal
113 omega (30). The alignment is used as input to Rosetta's threading protocol (*partial_thread.<ext>*).
114 From the threaded model, all residues in the MHC-I groove that are within a heavy-atom distance
115 of 3.5 Å from the peptide are subjected to 10 independent all-atom refinement simulations using
116 the FastRelax method (31) and a custom movemap file. Binding energies are extracted from the
117 refined structures using interface analyzer protocol (*InterfaceAnalyzer.<ext>*). The top three
118 models are selected based on the binding energies, and used to compute an average energy for
119 each peptide in the input list. RosettaMHC models of SARS-CoV-2/HLA-A*02:01 epitopes are

120 made available through an online database (see data availability). The website that hosts our
121 database was constructed using the Django web framework.

122

123

124 **Results and Discussion**

125

126 *Template identification for structure modeling using RosettaMHC*

127 Our full workflow for template identification and structure modeling is outlined in Figure 1a, with
128 a flowchart shown in Figure 1b. To identify all possible regular peptide binders to HLA-A*02:01
129 that are expressed by SARS-CoV-2, we used a recently annotated version of all open reading
130 frames (ORFs) in the viral genome from NCBI (32), made available through the UCSC genome
131 browser (33). We used 9- and 10- residue sliding windows to scan all protein sequences, since
132 these are the optimum peptide lengths for binding to the HLA-A*02:01 groove (34). While spliced
133 peptide epitopes (35) are not considered in the current study, this functionality can be added to our
134 method at a later stage. Using NetMHCpan-4.0 (27), we identified all 439 9mer and 279 10mer
135 epitopes that are predicted to yield positive (classified as both weak and strong) binders. To further
136 validate this set and derive plausible 3D models of the peptide/HLA-A*02:01 complexes, we used
137 a structure-guided approach, RosettaMHC, which aims to derive a physically realistic fitness score
138 for each peptide in the HLA-A*02:01 binding groove using an annotated database of high-
139 resolution structures and Rosetta's all-atom energy function (36). RosettaMHC leverages a
140 database of 241 HLA-A*02:01 X-ray structures encompassing a range of bound peptides, to find
141 the closest match to each target epitope predicted from the SARS-CoV-2 proteome. To identify
142 the best template for structure modeling, we use sequence matching criteria which first consider
143 the peptide anchors (positions P2 and P9/P10 for 9mer/10mer epitopes), followed by a sequence
144 similarity metric calculated from the full alignment between the template and target peptide
145 sequences. The template assignment statistics for the four different classes of SARS-CoV-2
146 epitopes in our set are shown in Figure 2a. We find that we can cover the entire set of 718 predicted
147 binders using a subset of 114 HLA-A*02:01 templates in our annotated database of PDB-derived
148 structures (Figure 2b). Each target peptide sequence is then threaded onto the backbone of its best
149 identified template, followed by all-atom refinement of the side chain and backbone degrees of
150 freedom using Rosetta's Ref2015 energy function (36), and binding energy calculation.

151

152 *RosettaMHC models recapitulate features of high-resolution X-ray structures*

153 The sequence logos derived from 9mer and 10mer peptides with good structural complementarity
154 to the HLA-A*02:01 groove according to Rosetta's binding energy (see below) adhere to the
155 canonical motif, with a preference for hydrophobic, methyl-bearing side chains at the peptide
156 anchor residues P2 and P9 (Figure 3a). The anchor residue preferences are recapitulated in
157 representative 9mer and 10mer models of the two top binders in our set as ranked by Rosetta's
158 energy (Figure 3c and 3d), corresponding to epitopes TMADLVYAL and FLFVAAIFYL derived
159 from the RNA polymerase and nsp3 proteins, respectively, which are both encoded by *orf1ab* in

160 the viral genome (NCBI Reference YP_009724389.1). In accordance with features seen in high-
161 resolution structures of HLA-A*02:01-restricted epitopes, the peptides adopt an extended, bulged
162 backbone conformation. The free N-terminus of both peptides is stabilized by a network of polar
163 contacts with Tyr 7, Tyr 159, Tyr 171 and Glu 63 in the A- and B- pockets of the HLA-A*02:01
164 groove. The Met (9mer) or Leu (10mer) side chain of P2 is buried in a B-pocket hydrophobic cleft
165 formed by Met 45 and Val 67. Equivalently, the C-terminus is coordinated through polar contacts
166 with Asp 77 and Lys 145 from opposite sides of the groove, with the Leu P9/P10 anchor nestled
167 in the F-pocket defined by the side chains of Leu 81, Tyr 116, Tyr 123 and Trp 147. Residues P3-
168 P8 form a series of backbone and side chain contacts with pockets C, D and E, while most
169 backbone amide and carbonyl groups form hydrogen bonds with the side chains of residues lining
170 the MHC-I groove. These high-resolution structural features are consistent across low-energy
171 models of unrelated target peptides in our input set, suggesting that, when provided with a large
172 set of input templates, a combined threading and side chain optimization protocol can derive
173 physically realistic models.

174

175 *Selection of high-affinity peptide epitopes using a structure-based score*

176 To evaluate the accuracy of our models and fitness of each peptide within the HLA-A*02:01
177 binding groove, we computed Rosetta all-atom binding energies across all complexes modeled for
178 different peptide sets. High binding energies can be used as an additional metric to filter low-
179 affinity peptides in the NetMHCpan-4.0 predictions, with the caveat that high energies can be also
180 due to incomplete optimization of the Rosetta energy function as a result of significant deviations
181 between the target and template backbone conformations, not captured by our protocol. We
182 performed 10 independent calculations for each peptide, and the 3 lower-energy models were
183 selected as the final ensemble and used to compute an average binding energy. The results for all
184 9mer peptides are summarized in Figures 3e, f, while additional results for 10mers are provided
185 through our web-interface and outlined in Supplemental Table 1. As a positive reference, we used
186 the binding energies of the idealized and relaxed PDB templates, which are at a local minimum of
187 the Rosetta scoring function. As a reference set for sub-optimal binders, we modeled decoy
188 structures of poly alanine (polyA) peptide sequences (predicted by NetMHCpan-4.0 to be a top
189 9th percentile binder for HLA-A*02:01), threaded onto the same PDB templates.

190

191 We observe a significant, negative (-26 kcal/mol) energy gap between the average binding energies
192 for PDB templates and poly alanine models. The binding energies for all modeled 9mers from the
193 SARS-CoV-2 genome fall between the average energies of the optimal PDB templates and sub-
194 optimal polyA binders, and show a bimodal distribution with significant overlap with the refined
195 PDB template energies (Figure 1e). Comparison of the distributions between epitopes that are
196 classified as strong versus weak binders by NetMHCpan-4.0 shows a moderate bias towards lower
197 binding energies for the strong binders and a larger spread in energies for weak binders, likely due
198 to suboptimal residues at the P2 and P9 anchor positions (Figure 3f). As an intended positive set,
199 we also modeled 28 9mer peptides that are homologous to peptides in the SARS viral genome and

200 have been previously reported to bind HLA-A*02:01 in the IEDB and ViPR (12, 37, 38) databases
201 (Supplemental Table 2). Inspection of Rosetta binding energies derived from models in this set
202 shows a similar distribution to the epitopes classified by NetMHCpan-4.0 as strong binders, with
203 the energies of 19/28 peptides falling well within the distribution of the refined PDB templates
204 (red dots in Figure 3e).

205
206 Based on these observations, we further classified all epitopes in the original set provided by
207 NetMHCpan-4.0 as strong or weak binders according to the Rosetta binding energy. Peptides with
208 binding energies that fall well within the PDB template distribution (green curve and red dots in
209 Figure 3e) are classified as strong binders. We obtained 154 9mer and 72 10mer strong binders
210 which show optimal complementarity within the HLA-A*02:01 peptide-binding groove according
211 to our modeling simulations. These results suggest that the high-resolution features seen in our
212 models (Figure 3c, d) yield optimal binding energies for a significant fraction of the epitopes
213 predicted by NetMHCpan-4.0 (45/33% of strong binders and 30/25% of weak binders for
214 9mers/10mers, respectively), which are comparable to locally refined PDB structures. The average
215 binding energies for all peptides are provided in our web-interface and in Supplemental Table 1.

216

217 *Surface features of peptide/HLA-A*02:01 models for T cell recognition*

218 Visualization of our models through an interactive online interface provides direct information on
219 SARS-CoV-2 peptide residues that are bulging out of the MHC-I groove, and are therefore
220 accessible to interactions with complementarity-determining regions (CDRs) of T cell receptors
221 (TCRs). Given that $\alpha\beta$ TCRs generally employ a diagonal binding mode to engage pMHC-I
222 antigens where the CDR3 α and CDR3 β TCR loops form direct contacts with key peptide residues
223 (39, 40), knowledge of the surface features for different epitopes adds an extra layer of information
224 to interpret sequence variability between different viral strains. For other important antigens with
225 known structures in the PDB, such features can be derived from an annotated database connecting
226 pMHC-I/TCR co-crystal structures with biophysical binding data (41), and were recently
227 employed in an artificial neural network approach to predict the immunogenicity of different HLA-
228 A*02:01 bound peptides in the context of tumor neoantigen display (42). A separate study has
229 shown that the electrostatic compatibility between self vs foreign HLA surfaces can be used to
230 determine antibody alloimmune responses (43). Given that antibodies and TCRs use a common
231 fold and similar principles to engage pMHC-I molecules (40), it is likely that surface electrostatic
232 features play an important role in recognition of peptide/HLA surfaces by their cognate TCRs in
233 the context of SARS-CoV-2 infection.

234 Electrostatic surface potentials calculated using a numerical solution to the Poisson-Boltzmann
235 Equation (44) for our modeled peptide/HLA-A*02:01 complexes allow us to compare important
236 features for TCR recognition between different high-affinity epitopes (Figure 4). We observe a
237 moderate electropositive character of the HLA-A*02:01 α_1 helix, and a moderate negative
238 potential on the α_2 helix, which is consistent between complexes with different bound peptides.
239 However, due to substantial sequence variability in surface-exposed residues at the P2-P8

240 positions, we observe a range of electrostatic features ranging from negative (epitope
241 TMADLVYAL), to neutral (NLIDSYFVV) or positively charged (KLWAQCVQL). Further
242 classification and ranking of the top binders in our set on the basis of their molecular surface
243 features would enable the selection of the most diverse panel of peptides for high-throughput
244 pMHC tetramer library generation (23-25). Tetramer screening of T cells from COVID-19
245 patients, recovered individuals and healthy donors can be used to identify critical gaps in the T cell
246 repertoire of high-risk groups, and to design epitope DNA strings for vaccine development.

247

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255

256 **Code and Data availability**

257 An online web-interface for visualization and download of all models is available at:
258 <https://rosettamhc.chemistry.ucsc.edu>. The RosettaMHC source code is available at
259 <https://github.com/snerligit/mhc-pep-threader>. Rosetta binding energies for all 718 HLA-
260 A*02:01-restricted peptides in our set are provided in Supplemental Table 1.

261

262 **Disclosures**

263 The authors have no financial conflicts of interest.

264

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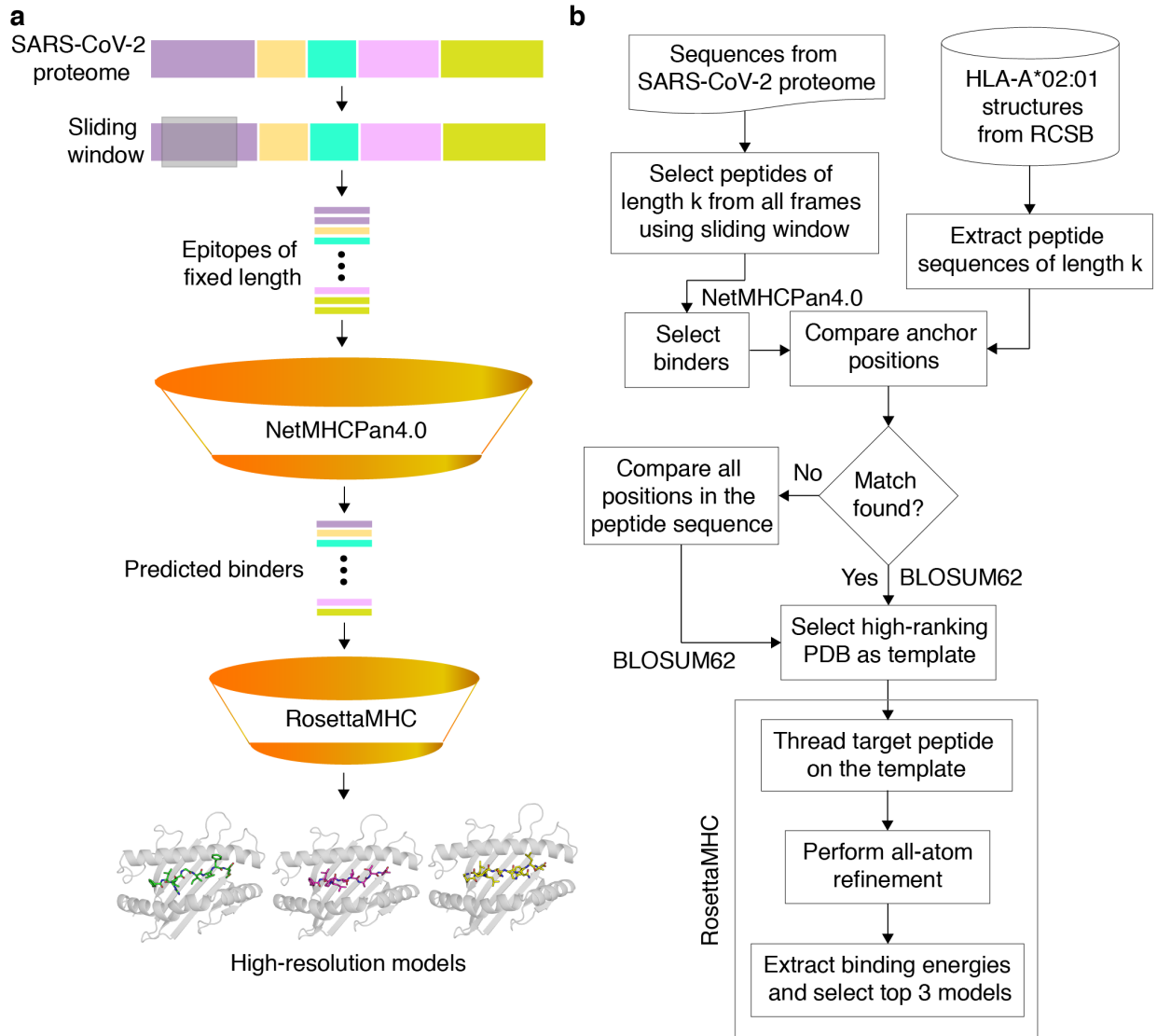
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- 402

403 **FIGURES**

404



405

406 **FIGURE 1. Structure-guided modeling of T cell epitopes in the SARS-CoV-2 proteome**

407 **(a)** General workflow of our pipeline for structure-guided epitope ranking. **(b)** Protein sequences

408 from the annotated SARS-CoV-2 proteome are used to generate peptide epitopes with a sliding

409 window covering all frames of a fixed length (9,621 9mer and 9,611 10mer possible peptides).

410 Candidate peptides are first filtered by NetMHCpan-4.0 (27) to identify all predicted strong and

411 weak binders (439 9mer and 279 10mer epitopes). For rapid template matching and structure

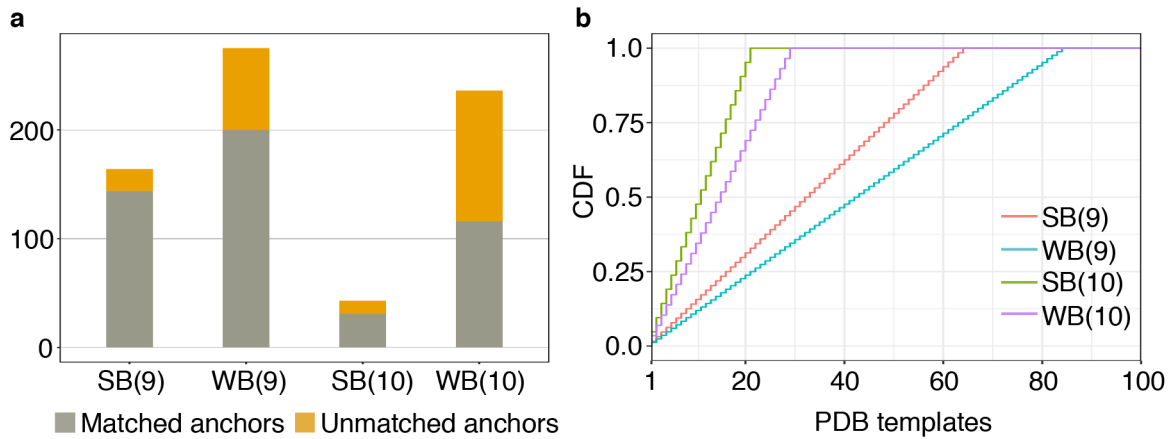
412 modeling, we use a local database of 241 HLA-A*02:01 X-ray structures with resolution below

413 3.5 Å from the Protein Data Bank (22). Each candidate peptide is scanned against all peptide

414 sequences of the same length in the database, and the top-scoring template is used to guide the

415 RosettaMHC comparative modeling protocol and to compute a binding energy.

416



417

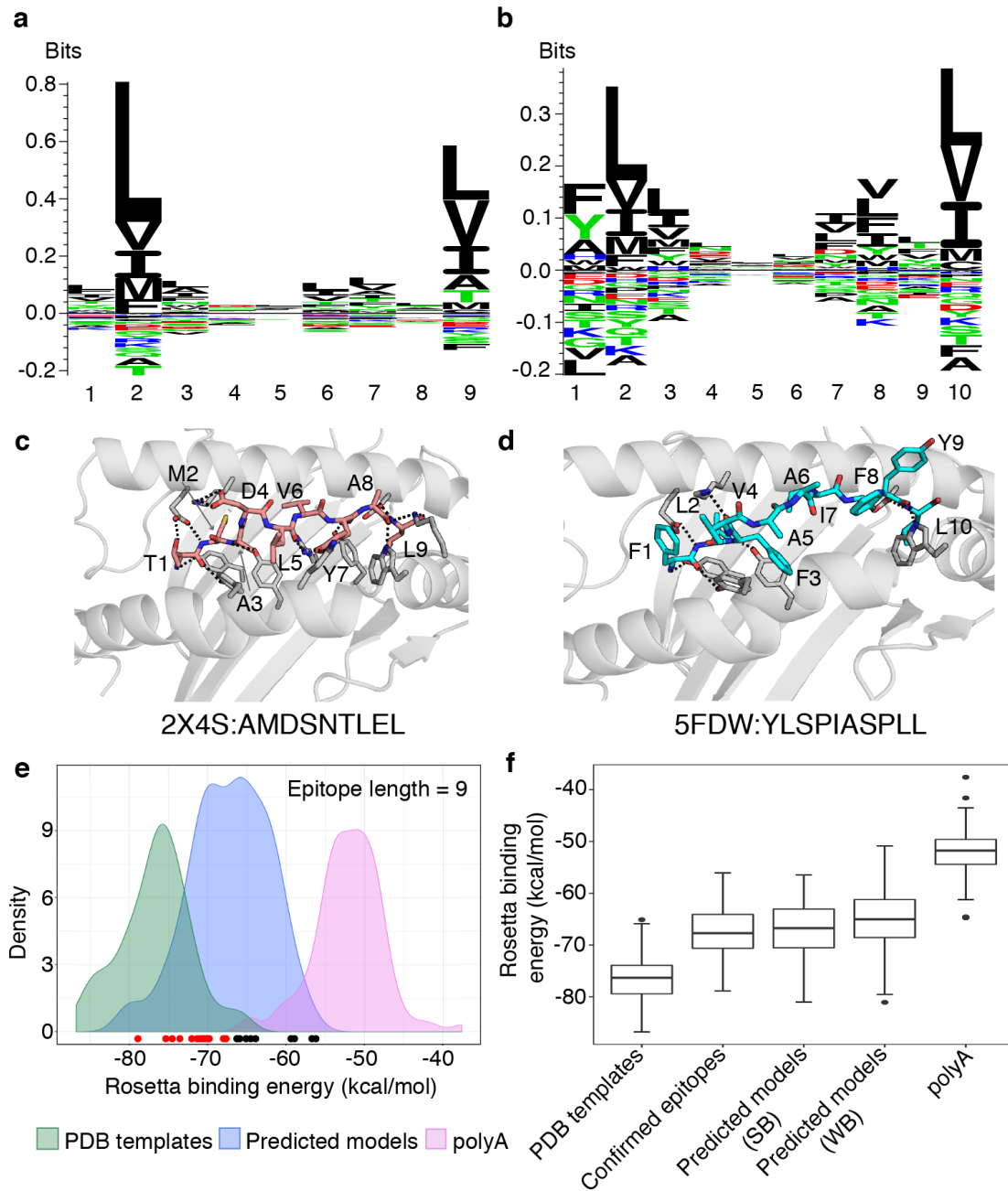
418 **FIGURE 2. Coverage of predicted HLA-A02 epitopes by structural templates in the PDB**

419 (a) Peptide anchor matching statistics of all predicted SARS-CoV-2 strong (SB) and weak binders

420 (WB) of lengths 9 and 10 to a database of 241 high-resolution HLA-A*02:01 X-ray structures (b)

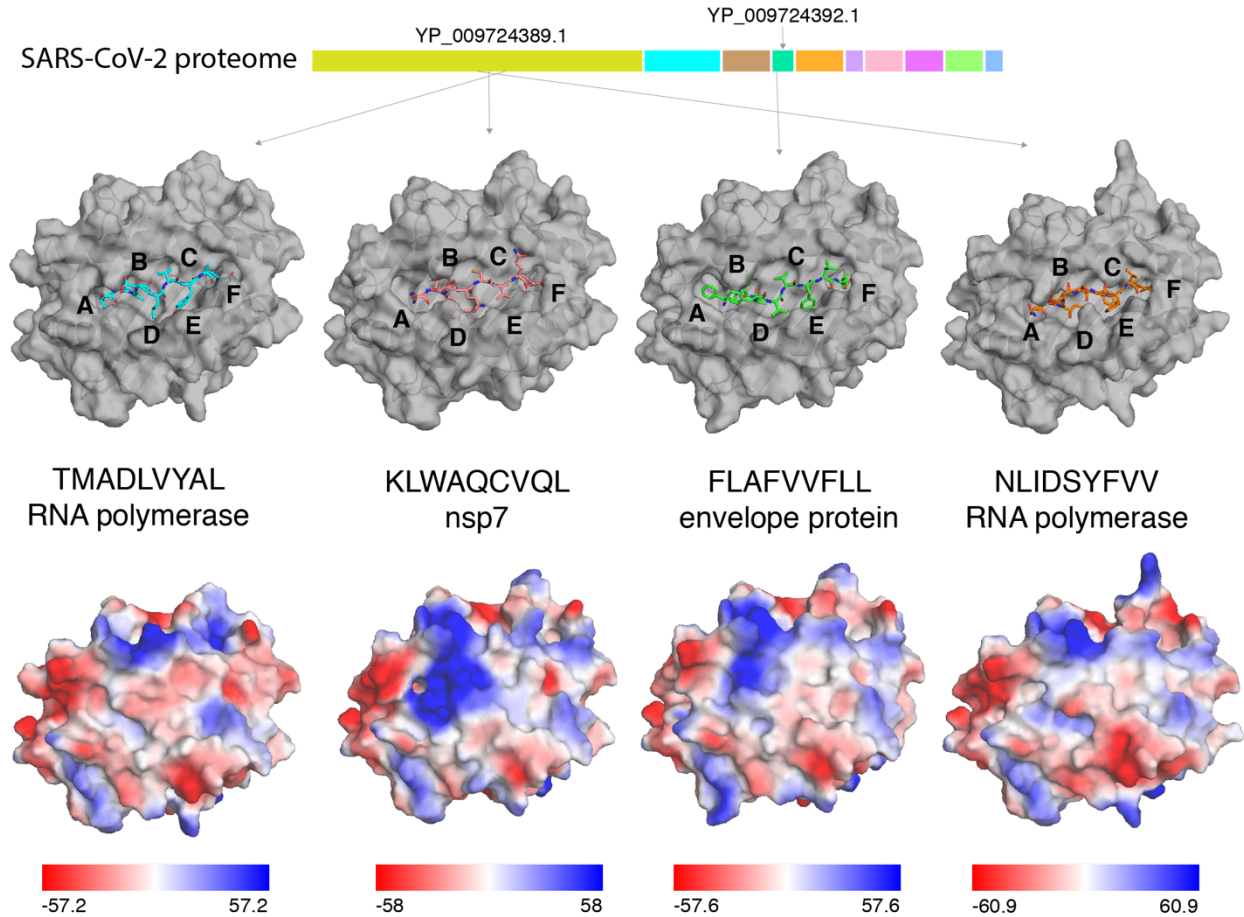
421 Plot showing cumulative distribution (CDF) of strong and weak binder peptides of lengths 9 and

422 10, as a function of the total number of matching templates from the Protein Data Bank (22).



423
 424 **FIGURE 3. Summary of RosettaMHC modeling results for SARS-CoV-2 peptide epitopes**
 425 Sequence logos from the n top ranking epitopes in the SARS-CoV-2 genome, predicted by
 426 NetMHCpan-4.0 (27) and further refined using RosettaMHC binding simulations are shown for:
 427 (a) 9mers ($n=154$) and (b) 10mers ($n=72$). The top 9mer and 10mer epitopes in our refined set are
 428 shown: (c) TMADLVYAL, from RNA polymerase and (d) FLFVAAIFYL, from nsp3. Dotted
 429 lines indicate polar contacts between peptide and heavy chain residues, with peptide residues
 430 labelled. The template PDB IDs and original peptides used for modeling the target peptides are
 431 indicated below each model. (e) Density plots showing distribution of average Rosetta binding
 432 energies (kcal/mol) for all epitopes of length 9. Distributions reflect 93 PDB templates (green),

433 164 strong binder epitopes (according to NetMHCpan-4.0 (27)) (blue), and 93 poly alanine
434 peptides modeled using the same PDB templates and used as a reference set for sub-optimal
435 binders (polyA; pink). The binding energies of models generated for 28 confirmed SARS T cell
436 epitopes from the IEDB and ViPR (37, 38) are indicated by circles at the bottom of the plot. Red
437 circles (19/28) indicate epitopes that lie within the distribution of refined PDB templates and black
438 circles (9/28) indicate epitopes that fall within the distribution of polyA (sub-optimal binders). **(f)**
439 Box plots showing distribution of average binding energies for 93 PDB templates, 93 poly alanine
440 peptides, 28 confirmed epitopes (37, 38) and RosettaMHC models for 164 strong (SB) and 275
441 weak (WB) binder 9mer epitopes predicted from the SARS-CoV-2 proteome using NetMHCpan-
442 4.0 (27).
443



444
445 **FIGURE 4. Variability in TCR recognition features of HLA-A02 with different high-affinity**
446 **peptides.** Molecular surfaces of SARS-CoV-2/HLA-A*02:01 RosettaMHC models are shown for
447 four top-scoring epitopes (ranked by Rosetta binding energy from left to right) captured in the A,
448 B, C, D, E and F pockets of the MHC-I groove (top panel). The origins of the peptide epitopes in
449 the ~30 kbp SARS-CoV-2 genome are noted. Electrostatic surfaces computed for the same models
450 are shown in the bottom panel. Solvent-accessible surface representation with electrostatic
451 potential in the indicated ranges (down to -60 kcal/(mol $\cdot e$) in red and up to $+61$ kcal/(mol $\cdot e$) in
452 blue) were calculated using the APBS solver (45) in Pymol (46). All calculations were performed
453 at 150 mM ionic strength, 298.15 Kelvin, pH 7.2, protein dielectric 2.0, and solvent dielectric
454 78.54. Electrostatic potentials are given in units of kT/e. A 1.4 Å solvent (probe) radius and 10.0
455 points/Å² density was used to calculate molecular surfaces.
456