How nonuniform contact profiles of T cell receptors modulate thymic selection outcomes

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T cell receptors (TCRs) bind foreign or self-peptides attached to major histocompatibility complex (MHC) molecules, and the strength of this interaction determines T cell activation. Optimizing the ability of T cells to recognize a diversity of foreign peptides yet be tolerant of self-peptides is crucial for the adaptive immune system to properly function. This is achieved by selection of T cells in the thymus, where immature T cells expressing unique, stochastically generated TCRs interact with a large number of self-peptide-MHC; if a TCR does not bind strongly enough to any self-peptide-MHC, or too strongly with at least one self-peptide-MHC, the T cell dies. Past theoretical work cast thymic selection as an extreme value problem and characterized the statistical enrichment or depletion of amino acids in the postselection TCR repertoire, showing how T cells are selected to be able to specifically recognize peptides derived from diverse pathogens yet have limited self-reactivity. Here, we investigate how the diversity of the postselection TCR repertoire is modified when TCRs make nonuniform contacts with peptide-MHC. Specifically, we were motivated by recent experiments showing that amino acids at certain positions of a TCR sequence have large effects on thymic selection outcomes, and crystal structure data that reveal a nonuniform contact profile between a TCR and its peptide-MHC ligand. Using a representative TCR contact profile as an illustration, we show via simulations that the statistical enrichment or depletion of amino acids now varies by position according to the contact profile, and, importantly, it depends on the implementation of nonuniform contacts during thymic selection. We explain these nontrivial results analytically. Our study has implications for understanding the selection forces that shape the functionality of the postselection TCR repertoire.

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I. INTRODUCTION

T cell receptors (TCRs) bind peptides loaded onto major histocompatibility complex (MHC) molecules (abbreviated as peptide-MHC) on the surface of antigen-presenting cells (APCs), and the strength of this interaction determines T cell activation [1,2]. Such peptides are derived from either the host itself (self-antigens), or, potentially, pathogens infecting the host (foreign antigens). Thus, optimizing the ability of T cells to recognize diverse foreign antigens with high specificity, yet be self-tolerant, is essential for the proper functioning of the adaptive immune system. Immature T cells (or thymocytes) express a distinct TCR on their surface assembled through a stochastic process of gene rearrangement, generating a highly diverse repertoire [3] that is acted upon by selection in the thymus. There thymocytes are screened against a large number of self-peptide-MHC; those that do not productively bind to self-peptide-MHC die (this is called positive selection), and those that bind too strongly are also eliminated (this is called negative selection). T cells that survive thymic selection are exported to the body’s periphery where they participate in the adaptive immune response. In this way, thymic selection shapes the postselection TCR repertoire to potentially recognize diverse foreign antigens yet limit self-reactivity [4]. While detailed statistics of the postselection repertoire are now available [5–7], how thymic selection achieves this outcome is less well understood.

Mathematical models have shed light on various aspects of thymic selection (for a review, see Ref. [8]); in particular, models that represented TCR-peptide-MHC interactions as pairwise interactions between digit strings [9,10] were useful for studying how TCR cross-reactivity can result. In Refs. [11,12], some of the authors considered a more explicit representation of inter-amino acid interaction strengths (the Miyazawa-Jernigan matrix [13]) and characterized the statistical enrichment or depletion of amino acids in the postselection TCR repertoire, computationally [11] and analytically [12], as a function of parameters such as the number of self-peptide-MHC encountered during selection. These studies provided insight into how T cells are selected to be specific for unknown foreign peptides and yet are self-tolerant.

In the past few years, more detailed information about thymic selection outcomes have emerged. Advances in high-throughput sequencing have allowed researchers to quantify the statistics of postselection TCR sequences in detail, revealing positional differences in the enrichment of amino acids [5,6]. Furthermore, recent experiments found that preselection thymocytes that were activated by self-peptide-MHC (and hence would fail negative selection) were enriched in hydrophobic amino acids at positions 6 and 7 of the TCRβ-chain complementarity-determining region (CDR3β), while thymocytes passing both positive and negative selection were enriched in amino acids with moderate hydrophobicity at these positions [7]. These results agree with theoretical
predictions made in previous work [11,12] but additionally find varying levels of enrichment at different CDR3β positions. Furthermore, a large number of crystal structures of TCR-peptide-MHC complexes have been analyzed that reveal a nonuniform contact interface between TCRs and their peptide-MHC ligands (quantified, for example, by atom-atom contact profiles; see Fig. 1) and show that positions 6 and 7 of the CDR3β sequence make the strongest contacts on average. Taken together, these findings show that certain TCR positions are more important than others for influencing thymic selection outcomes, and that this information can be captured by nonuniform contact profiles such as those in Fig. 1.

The paper is organized as follows: in Sec. II we develop a mathematical model of thymic selection that incorporates nonuniform contact profiles. In Sec. III we consider two possible interpretations of nonuniform contacts and study their effects on thymic selection outcomes. The stark differences in levels of enrichment between the two interpretations are explained analytically. In Sec. IV we discuss further work and conclude.

II. MODEL DESCRIPTION

Immature T cells (or thymocytes) undergo positive and negative selection in the thymus before maturation. Each thymocyte expresses a distinct TCR on its surface, generated stochastically through V(D)J gene recombination and insertions and deletions of nucleotides (whose probabilities have been inferred from high-throughput sequencing data [3]), creating a diverse preselection repertoire. In the thymic cortex, thymocytes are presented with self-peptide-MHC by thymic antigen-presenting cells (APCs). Thymocytes that do not bind strongly enough to any self-peptide-MHC die of insufficient survival signals; this is called positive selection. Thymocytes that survive positive selection migrate to the thymic medulla, where they are further screened against self-peptide-MHC, and those that bind too strongly with at least one self-peptide-MHC receive apoptotic signals and are eliminated; this is called negative selection.

We cast this process in a mathematical model as follows: following Ref. [11], TCR sequences, \( t = (t_1, \ldots, t_N) \), of length \( N \) are generated by sampling amino acids \( t_i, i = 1, \ldots, N \), independently from a distribution \( p_{\text{pre}}(t_i) \), that is taken to be the amino acid distribution of the human proteome [14]. While this is highly simplified compared to how the actual preselection repertoire is generated [3], it does not affect our
FIG. 2. Mathematical model of positive and negative selection in the thymus. (a) String representation of TCR-peptide-MHC binding. A TCR sequence of length \( N \) (here \( N = 5 \)) interacts with a peptide sequence, also of length \( N \), bound to a MHC molecule. Each colored square represents a different amino acid. TCR and peptide amino acids interact in a pairwise fashion [see Eq. (1)]. (b) Schematic of possible thymic selection outcomes. Top: Thymocyte failing negative selection. Middle: Thymocyte failing positive selection. Bottom: Thymocyte surviving thymic selection, and their sequences were used to construct the enrichment curve, where all \[\beta\] values less than 1 imply enrichment (depletion) in the postselection repertoire. Blue dotted line: Results of numerical simulations of the model for \( 5 \times 10^6 \) independently generated TCR sequences, each encountering \( M = 10^3 \) independent self-peptide sequences. 20.6% of TCRs survived thymic selection, and their sequences were used to construct the enrichment curve, where all \( i = 1, \ldots, N \) have been averaged together. The statistical uncertainty is smaller than the symbol sizes, as confirmed by simulations with \( 10^6 \) trials. Red solid line: Prediction of Eq. (8) with \( \beta^* \) obtained from Eqs. (9)–(11). Theory gave \( \beta^* = -0.38 \), implying selection for weakly interacting amino acids. Parameter values (same as Ref. [12]): all \( f(c_i) = 1 \). Henceforth, we will specialize to one MHC type (i.e., \( E_c \) a constant) because its diversity is much lower than that of self-peptides (a human for example expresses six different MHC class I molecules), and we are not going to focus on how TCR cross-reactivity to other MHC molecules can arise.

Positive and negative selection are carried out as follows: if the strongest (minimum) interaction energy between a TCR and \( M \) independent self-peptide-MHC is weaker (greater) than a positive selection threshold \( E_{p} \), or if it is stronger (less) than a negative selection threshold \( E_{n} \), the thymocyte dies. These hard constraints are consistent with experiments that found relatively small differences in TCR-ligand affinity at the negative selection threshold [16] (although they found that the positive selection threshold is softer). Thymocytes that interact with \( M \) self-peptide-MHC with strongest (minimum) binding energy within \([E_n, E_p]\) survive thymic selection and mature into naive T cells [see Fig. 2(b)].

A. Parameter values

The CDR3 loops of a TCR typically make the greatest contacts with peptide (as opposed to MHC) [17]. Here, the TCR amino acid string \( t \) represents the CDR3 \( \beta \) sequence, which is typically of length 10–18 [6]. During thymic selection, a thymocyte typically interacts with \( 10^3 \)–\( 10^4 \) self-peptide-MHC. Realistic values of the difference between positive and negative selection thresholds, and the T cell activation free energy with self-peptide alone (without MHC), are given by \( E_p - E_n = 2.5k_BT \) and \( E_n - E_c = -21k_BT \), respectively [11,12].

later results, which concern the action of thymic selection on this distribution. Unlike Ref. [11], here a TCR has the capacity to make nonuniform contacts, which we capture by specifying \( c = (c_1, \ldots, c_N) \). We describe values and interpretations of \( c \) in the next section.

During thymic selection, a TCR interacts with \( M \) independent self-peptide sequences, \( s = (s_1, \ldots, s_N) \), of length \( N \) bound to MHC, that are also randomly generated according to \( p_{\text{pre}} \). Because these are mostly linear peptides, they do not have an associated \( c \). Following Ref. [11], we model the binding strength between a TCR and self-peptide as pairwise interactions between TCR amino acid \( t_i \) and self-peptide amino acid \( s_i \), for \( i = 1, \ldots, N \) [see Fig. 2(a)]. Thus, the overall binding energy, \( E(t,s,c) \), is

\[
E(t,s,c) = E_c + \sum_{i=1}^{N} f(c_i)J(t_i,s_i),
\]

where \( E_c \) captures interactions between TCR and MHC, \( J(t,s) \) is an interaction potential that in principle accounts for biochemical and other properties of inter-amino acid interactions, and \( f(c_i) \) accounts for nonuniform contact at position \( i \). Following Ref. [11], we use the Miyazawa-Jernigan (MJ) matrix for \( J(t,s) \) [13], whose structure largely arises from hydrophobic forces [7,15]; hydrophobic amino acids are strongly interacting (more negative \( J(t,s) \) values), while hydrophilic amino acids are weakly interacting (less negative \( J(t,s) \) values). (Note that the results that follow do not qualitatively depend on the potential, but Ref. [7] noted the importance of hydrophobicity for the strength of TCR-peptide-MHC interactions, implying that use of the MJ matrix is reasonable.) Reference [11] implicitly assumed that all \( f(c_i) = 1 \). Henceforth, we will specialize to one MHC type (i.e., \( E_c \) a constant) because its diversity is much lower than that of self-peptides (a human for example expresses six different MHC class I molecules), and we are not going to focus on how TCR cross-reactivity to other MHC molecules can arise.

Positive and negative selection are carried out as follows: if the strongest (minimum) interaction energy between a TCR and \( M \) independent self-peptide-MHC is weaker (greater) than a positive selection threshold \( E_p \), or if it is stronger (less) than a negative selection threshold \( E_n \), the thymocyte dies. These hard constraints are consistent with experiments that found relatively small differences in TCR-ligand affinity at the negative selection threshold [16] (although they found that the positive selection threshold is softer). Thymocytes that interact with \( M \) self-peptide-MHC with strongest (minimum) binding energy within \([E_n, E_p]\) survive thymic selection and mature into naive T cells [see Fig. 2(b)].
B. Model without nonuniform contacts

References [11,12] considered a model with all \( f(c_i) = 1 \), \( N = 5 \), \( M = 10^3 \), and \( E_p - E_n \) and \( E_n - E_p \) given above. Numerical simulations of this model resulted in a distribution of TCR sequences surviving selection that was statistically different from the preselection one, in that the former was enriched in weakly interacting amino acids and depleted in strongly interacting ones [see Fig. 2(c)]. This result was consistent with later experiments [7].

C. Theory of the postselection TCR repertoire distribution

Reference [12] developed an analytical theory for the postselection TCR repertoire distribution, valid in the limit of \( N, M \to \infty \). Here we extend this theory to include nonuniform contact profiles. A self-contained derivation is in the Appendix; we state below the essential results.

A TCR with sequence \( t = (t_1, \ldots, t_N) \) and contact profile \( c = (c_1, \ldots, c_N) \) experiences a distribution of binding energy selections during thymic selection, with mean

\[
\mu(t,c) = E_c + \sum_{i=1}^{N} \mu(t_i, c_i),
\]

and variance

\[
v(t,c) = \sum_{i=1}^{N} v(t_i, c_i),
\]

where there are no cross-correlation terms in Eq. (3) because sites are independent. In Ref. [12] [where all \( f(c_i) = 1 \)], \( \mu(t_i, c_i) \) and \( v(t_i, c_i) \) are given by

\[
\mu(t_i) = [J(t_i, a)]_a,
\]

and

\[
v(t_i) = [J(t_i, a)]_a^2 - [J(t_i, a)]_a^2,
\]

where \( [J(t_i, a)]_a \) and \( [J(t_i, a)]_a^2 \) are the first and second moments of interaction strength experienced by amino acid \( t \) when interacting with self-peptide sequences,

\[
[J(t,a)]_a \equiv \sum_{a=1}^{20} J(t,a)p_{perc}(a),
\]

where \( a \) runs over the 20 possible amino acids \( t \) encounters. In this paper, the forms of \( \mu(t_i, c_i) \) and \( v(t_i, c_i) \) depend on the interpretation of \( c \) and will be specified in the next section.

For large \( N \) and \( M \), the minimum binding energy selection by a TCR during selection tends to the Gumbel distribution (see the Appendix), whose peak is at

\[
\rho_G(t,c) = \mu(t,c) - 2\sqrt{v(t,c)}\alpha,
\]

where \( \alpha = \sqrt{\log M} \), and whose variance is \( v_G(t,c) = \pi^2 v(t,c)/(12 \log M) \). In the limit of \( N, M \to \infty \) (keeping \( N \propto \log M \)), this distribution concentrates around its peak, which lies somewhere between \( E_n \) and \( E_p \). In this limit, the postselection TCR repertoire distribution becomes (see the Appendix)

\[
p_{\text{post}}(t_1|c_1) = \frac{1}{Z_{\beta,\gamma,c_1}} e^{-\beta \rho_G(t_1,c_1) - \gamma v_G(t_1,c_1)} p_{\text{perc}}(t_1),
\]

where \( Z_{\beta,\gamma,c_1} \) ensures normalization,

\[
\gamma = \frac{\alpha}{\sqrt{2 \sum_{i=1}^{N} [v(t_i,c_i)]_{\beta,\gamma,c_i}}},
\]

and

\[
\langle v(t_i,c_i) \rangle_{\beta,\gamma,c_i} = \sum_{i=1}^{20} v(t_i,c_i)p_{\text{post}}(t_i|c_i).
\]

The value of \( \beta \) in Eq. (8) is chosen such that \( \langle \rho_G(t,c) \rangle \) [from Eq. (7)] lies within \( [E_n, E_p] \). \( p_{\text{post}}(t_1|c_1) \) depends on \( c_1 \) explicitly, and on the full contact profile implicitly through \( \gamma \). Also in this limit, \( \langle \rho_G(t,c) \rangle \) becomes

\[
\langle \rho_G(t, c) \rangle_{\beta,\gamma,c} = \sum_{i=1}^{N} (\mu(t_i,c_i) - \gamma v(t_i,c_i))_{\beta,\gamma,c_i} - \frac{\alpha^2}{2\gamma}.
\]

In practice, for each value of \( \beta \), one iterates between Eqs. (9) and (10) until a self-consistent value of \( \gamma \) is obtained. Values of \( \beta \) corresponding to \( \langle \rho_G(t,c) \rangle_{\beta,\gamma,c} = E_p \) and \( \langle \rho_G(t,c) \rangle_{\beta,\gamma,c} = E_n \) are found, and the one closer to zero is taken to be \( \beta^* \) (or \( \beta^* = 0 \) if they straddle 0). Intuitively, \( \beta^* \) parametrizes the degree to which weakly or strongly interacting amino acids are enriched in the postselection TCR repertoire; positive \( \beta^* \) implies selecting for strongly interacting amino acids, and negative \( \beta^* \) implies selecting for weakly interacting ones.

III. INTERPRETATIONS OF NONUNIFORM TCR CONTACT PROFILES

How is the picture of Fig. 2(c) modified when nonuniform contact profiles are taken into account? We were initially inspired by crystal structures of TCR-peptide-MHC complexes, which show a variation in the number of contacts made between TCR and peptide-MHC along the TCR sequence for every peptide-MHC on average [see Fig. 4(a)], which is consistent with experimental findings [7] and suggests that nonuniform contact profiles are useful in capturing positional differences that influence thymic selection outcomes. Figure 4(a) was obtained by rescaling the total number of contacts measured from 53 TCR-peptide-MHC structures such that the maximum is 1. We will make use of this representative, “average” contact profile as \( c \) in the following.

A. Deterministic screening interpretation

Motivated by crystal structure analyses, we first considered a model where every interaction between TCR amino acid \( t_i \) and self-peptide amino acid \( s_j \) was weakened by a factor \( c_i,j \), \( 0 \leq c_i \leq 1 \) [see Fig. 3(a)]. This is represented by setting \( f(c_i) = c_i \) in Eq. (1). In principle, there is no reason to exclude values of \( c_i > 1 \) (depending on how interactions are modified by number and type of contacts), but here we considered only reduced interactions, for example, due to screening by intervening water molecules.

We performed numerical simulations of our model using the deterministic interpretation of the contact profile of Fig. 4(a),...
with TCRs of length \( N = 15 \) and keeping the other parameter values the same as in Ref. [12] and Fig. 2(c) (i.e., \( M = 10^3 \), \( E_p - E_n = 2.5 \, k_B T \), and \( E_n - E_c = -21 \, k_B T \)). Statistics of TCR amino acids in the postselection repertoire are shown in Figs. 4(b)–4(c) (dotted lines). Enrichment curves for each site are plotted in a different color. Now, the degree of enrichment depends on position; sites corresponding to the largest \( c_i \) (sites 6 and 7) experience the greatest degree of enrichment, while sites making no contact (sites 1–3, 14, and 15) have enrichment values close to 1. Furthermore, sites making contacts are now enriched in strongly interacting amino acids, which is the opposite of Fig. 2(c).

To understand this result, we repeated the theoretical analysis of the previous section. The mean and variance of interaction energies for amino acid \( t_i \) at position \( i \) are now modified to

\[
\mu_A(t_i, c_i) = c_i [J(t_i, a)]_a, \tag{12}
\]

and

\[
v_A(t_i, c_i) = c_i^2 [J(t_i, a)^2]_a - c_i^2 [J(t_i, a)]_a^2, \tag{13}
\]

by replacing \( J(t_i, a) \rightarrow c_i J(t_i, a) \) in Eqs. (4) and (5). Using these expressions in Eqs. (9)–(11), we found \( \beta^* = 0.79 > 0 \), which indeed implies enrichment of strongly interacting amino acids. Plots of Eq. (8) for each position (solid lines in Figs. 4(b) and 4(e)) also resemble enrichment curves obtained from simulations.

Upon reflection, this change is not surprising. The contact profile of Fig. 4(a) has an “effective length” of \( \sum_{i=1}^{15} c_i \approx 3.2 < 5 \), implying that the mean binding energy experienced by a TCR during selection is roughly 64% of that experienced in the model with \( N = 5 \) and without nonuniform contacts, which produced Fig. 2(c). (Indeed, the same analytical computation for \( N = 3 \) and without nonuniform contacts also gives \( \beta^* > 0 \).) Thus, it is plausible that here TCR sequences need to be enriched in more strongly interacting amino acids in order to have their strongest binding energy during selection fall between \( E_n \) and \( E_p \). Indeed, the analytical theory makes this intuition concrete.

Note that the analytical curves systematically underpredict the degree of enrichment, i.e., the value of \( \beta^* \) obtained from theory was slightly too small. For finite \( M \), the extreme value distribution in fact has a finite width, and so one might underestimate \( \beta^* \) by matching the peak \( \langle \rho(t, c) \rangle_{y, c} \) to \( E_p \). Indeed, when we include the next-order correction to \( \rho(\mathbf{c}, t) \), which is positive [see after Eq. (A2) in the Appendix], the analytical curves match those from simulations much better (not shown).

The results of the deterministic interpretation of the nonuniform contact profile of Fig. 4(a), however, contradict the experiments of Ref. [7], which find that TCRs with strongly interacting amino acids at sites 6 and 7 should fail negative selection. While we have attempted to use a realistic contact profile, relaxing some of the assumptions we have made, such as \( c_i < 1 \) or the functional form of the interaction potential being \( f(c_i) \propto c_i \), may lead to results that better agree with experiments. Additionally, the hypothesis of deterministic screening need not be correct, and we were led to another interpretation of nonuniform contacts, which we describe in the following section.

**B. Stochastic binding interpretation**

In fact, crystal structures are merely static pictures of TCR-peptide-MHC binding, whereas the events leading to T cell activation following encounters with peptide-MHC are much more dynamic [17]. Indeed, it is known that the CDR3 loop of TCRs is relatively flexible, with a range of conformations that can bind different ligands [18]. There also exist crystal structures of the same TCR bound to different peptide-MHC that have different parts of the TCR binding the different ligands [19].

These facts motivate another interpretation of nonuniform contacts: for every encounter during thymic selection, TCR amino acid \( t_i \) binds to self-peptide amino acid \( s_j \) with probability \( c_i \), \( 0 \leq c_i \leq 1 \) [see Fig. 3(b)]. This is represented by setting \( f(c_i) = X_i \), where \( X_i \) is a Bernoulli random variable with parameter \( c_i \). This stochastic binding interpretation gives the same average interaction energy as the deterministic one:

\[
\mu_B(t_i, c_i) = c_i [J(t_i, a)]_a, \tag{14}
\]

while its variance is instead

\[
v_B(t_i, c_i) = c_i^2 [J(t_i, a)^2]_a - c_i^2 [J(t_i, a)]_a^2, \tag{15}
\]

which has one factor of \( c_i \) multiplying \( [J(t_i, a)^2]_a \) [as opposed to two in Eq. (13)].

We performed numerical simulations of this model, and obtained a postselection repertoire shown in Figs. 4(d) and 4(e) (dotted lines). Again, the degree of enrichment is greater for sites making greater contacts. However, unlike the deterministic interpretation, here sites making contacts are enriched in weakly interacting amino acids. Repeating the previous analytical computations using Eqs. (14) and (15) also give \( \beta^* = -0.38 < 0 \). Thus, the argument that we made previously was incomplete; even though the “effective length” here is roughly 3.2 as well because the mean interaction energies are equal [Eqs. (12) versus (14)], their variances are different, and it is the variance that nontrivially modifies thymic selection outcomes. Because of the larger variance here, it is plausible that weakly interacting amino acids are sufficient for having
FIG. 4. Thymic selection with nonuniform contact profiles. (a) Total number of atom-atom contacts for 53 TCR-peptide-MHC crystal structures, rescaled such that the maximum is 1 (contact data taken from Ref. [7]; see Protein Data Bank IDs therein). (b–e) Numerical simulations (dotted lines) and analytical computations (solid lines) for deterministic (b and c) and stochastic (d and e) interpretations of nonuniform contacts. Each colored line represents the degree of enrichment of amino acids at a position along TCR sequences surviving thymic selection. For ease of viewing, sites 1–7 are plotted in panels (b) and (d) and sites 8–15 in panels (c) and (e). To create the dotted lines, $5 \times 10^6$ simulations were performed for each model with independently drawn TCR and self-peptide sequences. The deterministic model had 2.29% of TCRs surviving thymic selection, while the stochastic model had 13.4% of TCRs surviving; these sequences were used to construct the enrichment curves shown. Solid lines are plots of Eq. (8) with $\beta^*$ obtained from Eqs. (9)–(11). Strongly interacting amino acids were enriched ($\beta^* > 0$) for the deterministic model, but depleted ($\beta^* < 0$) for the stochastic model. Parameter values: $N = 15$, $M = 10^3$, $E_p - E_c = 2.5$, $E_n - E_c = -21$, $\gamma^* = 1.05$, $\beta^* = -21$ (in units of $k_B T$). Note: In panels (c) and (e), the curves for sites 8 and 9 almost coincide as $c_8$ and $c_9$ are almost equal.

Another way to think of this is in terms of extreme values in lengths. During each interaction, a subset of positions of a TCR interact with self-peptide amino acids. In the negative selection-dominated regime, TCRs are more likely to be eliminated because of one interaction that is too strong, rather than all $M$ interactions being too weak. Performing the analytical computation for $N \geq 7$ and without nonuniform contacts gives $\beta^* = -\infty$, implying that no TCR that makes seven or more contacts during selection should survive. Also, the analytical theory for $N = 6$ and without nonuniform contacts gives $\beta^* = -\infty$. The analytical theory makes this intuition concrete.
Nonuniform contact profiles are a step towards modeling the complicated, structure-dependent nature of TCR-peptide-MHC interactions. However, a more informed method should be developed to infer such a profile from crystal structures. For example, we made use of measurements of the number of peptide-MHC atoms a distance of 4Å away from TCR amino acids, but characteristic distances should depend on the kind of interaction (hydrogen bonding, van der Waals, etc.).

We limited computations in this paper to one contact profile to illustrate its effect on thymic selection outcomes. Separately, it would be interesting to characterize the statistics of contact profiles from crystal structures. This task, however, is limited by the relatively small number of crystal structures known, as opposed to the large quantities of high-throughput sequencing data available. We note that an additional step needs to be taken to connect the model in this paper to statistics of aligned TCR sequences that appear in, e.g., Refs. [5,6] (which do not focus on residues making contact with peptide-MHC but rather the entire aligned CDR3 region, and hence do not find enrichment at TCR positions corresponding to recent experiments [7]): the model should be run separately for different contact profiles, and the enrichment curves should be averaged together according to the passing rates for the different contact profiles. Note that this is a possible mechanism for obtaining enrichment of an amino acid at one position in a sequence alignment, and depletion of the same amino acid at another, because these positions might feature in different contact profiles that have different $\beta^\ast$. This also implies that the results from an average contact profile are, in general, different from running the model for separate contact profiles and averaging the results together, because the former does not account for different passing rates. Thus, it would be interesting to attempt the inverse problem of inferring differential contacts and binding tendencies at different positions of a TCR sequence from positional differences in the postselection TCR repertoire, but this is complicated by the underdeterminacy of the problem.

Inferring overall patterns determining a TCR’s specificity to peptide-MHC from knowledge of TCR-peptide-MHC crystal structures is challenging because there is no canonical way by which a TCR interacts with peptide-MHC [17,20,21]; the same TCR may bind different peptide-MHC in very different ways [19]. Predicting TCR sequences that recognize a given set of peptide-MHC by inferring “sequence motifs” of TCRs has been achieved very recently [22,23]. We believe that analyzing thymic selection outcomes has implications for antigenic specificity, because surviving thymic selection involves interactions with a large number of self-peptide-MHC, and thus features relevant for thymic selection outcomes are also relevant for antigenic specificity. The interpretations of nonuniform contacts we studied here have distinct and measurable effects on the postselection repertoire, and thus they probably play a role in antigenic specificity as well, which perhaps gives a mechanistic basis for the sequence motifs discovered in the recent studies [22,23].

Recently, a paper that also modified the thymic selection model of Refs. [11,12] to include positional differences in TCR-peptide-MHC interactions appeared [24]. In essence, for a given TCR, they drew the values of $f(c_i)J(t_i,s_i)$ in Eq. (1), $i = 1, \ldots, N$, independently from a Gaussian distribution. Thus, they reduced thymic selection to an extreme value
problem with a random energy model, in which “TCRs” and “amino acids” lose their meaning; this contradicts studies that find predictive features determining specificity that are based on TCR amino acid sequences \[7,22,23\]. However, their model is simpler to analyze and may be a useful null model, and it would be interesting to compare it with a statistical ensemble of contact profiles. The authors also commented that the model of Refs. \[11,12\] fell short in that very few self-peptides (i.e., the most strongly interacting ones) perform the job of negative selection of Refs. \[11,12\] fell short in that very few self-peptides (i.e., the most strongly interacting ones) perform the job of negative selection effectively as the full panel of self-peptides. While this is true (which follows directly from specifying an inter-amino acid interaction potential), it is not known how large this fraction is for the real thymic selection process. Experimentally, this could be tested by engineering the thymus to contain peptides consisting of only strongly interacting amino acids \[4\]. Also, it is possible that these peptides are somehow found rarely or not at all in the thymus, because self-peptides are chopp-up versions of actual proteins. Large values of \(M\) are likely still required to randomly generate such special peptides. We note that the model we study here moves away from the limitations they raised, as different TCRs with special peptides. We note that the model we study here moves away from the limitations they raised, as different TCRs with special peptides. We note that the model we study here moves away from the limitations they raised, as different TCRs with special peptides. We note that the model we study here moves away from the limitations they raised, as different TCRs with special peptides. We note that the model we study here moves away from the limitations they raised, as different TCRs with special peptides. We note that the model we study here moves away from the limitations they raised, as different TCRs with special peptides.
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