Statistical Physics of T-Cell Development and Pathogen Specificity

Andrej Košmrlj,¹ Mehran Kardar,² and Arup K. Chakraborty^{2,3,4,5,6,7}

¹Department of Physics, Harvard University, Cambridge, Massachusetts 02138; email: andrej@physics.harvard.edu

²Departments of Physics, ³Chemical Engineering, ⁴Chemistry, and ⁵Biological Engineering, and ⁶Institute of Medical Engineering and Science, Massachusetts Institute of Technology, Cambridge, Massachusetts 02139; email: kardar@mit.edu, arupc@mit.edu

⁷Ragon Institute of MGH, MIT, and Harvard, Boston, Massachusetts 02129

Annu. Rev. Condens. Matter Phys. 2013. 4:339-60

The Annual Review of Condensed Matter Physics is online at conmatphys.annualreviews.org

This article's doi: 10.1146/annurev-conmatphys-030212-184325

Copyright © 2013 by Annual Reviews. All rights reserved

Keywords

immune system, HIV, extreme value distribution, Hamiltonian minimization

Abstract

In addition to an innate immune system that battles pathogens in a nonspecific fashion, higher organisms, such as humans, possess an adaptive immune system to combat diverse (and evolving) microbial pathogens. Remarkably, the adaptive immune system mounts pathogen-specific responses, which can be recalled upon reinfection with the same pathogen. It is difficult to see how the adaptive immune system can be preprogrammed to respond specifically to a vast and unknown set of pathogens. Although major advances have been made in understanding pertinent molecular and cellular phenomena, the precise principles that govern many aspects of an immune response are largely unknown. We discuss complementary approaches from statistical mechanics and cell biology that can shed light on how key components of the adaptive immune system, T cells, develop to enable pathogen-specific responses against many diverse pathogens. The mechanistic understanding that emerges has implications for how host genetics may influence the development of T cells with differing responses to the human immunodeficiency virus (HIV) infection.

Pathogen: bacterium, virus, or other microorganism that can cause disease

HIV: human immunodeficiency virus

Autoimmune disease:

disease caused by antibodies or lymphocytes produced against substances naturally present in the body

Adaptive immunity:

immunity occurring as a result of prior exposure to an infectious agent or its antigens

Thymus gland: organ where immature T cells are screened against self-peptide MHCs

Peptide: short piece of protein presented to T-cell receptors in a groove of MHC proteins

1. INTRODUCTION¹

The immune system of an organism combats invading pathogens, thereby protecting the host from disease. Jawed vertebrates, such as humans, have an adaptive immune system that enables them to mount pathogen-specific immune responses (1). The opportunistic infections that afflict individuals with compromised adaptive immune systems [e.g., those who have progressed to AIDS after infection with the human immunodeficiency virus (HIV)] highlight the importance of this response for human health. Many other diseases (e.g., multiple sclerosis and type I diabetes) are consequences of the adaptive immune system failing to discriminate between markers of self and nonself. The suffering caused by autoimmune diseases, and the need to combat diverse infectious agents, has motivated a great deal of experimental research aimed at understanding how the adaptive immune system is regulated. These efforts have led to many notable discoveries (2–10), yet a deep understanding of the principles that govern the emergence of an immune or autoimmune response has proven elusive. The inability to rationally design vaccines against many scourges on the planet (such as HIV) highlights the practical consequences of this missing basic science.

An important barrier for the development of mechanistic principles that describe adaptive immunity is that the pertinent processes involve cooperative dynamic events. The many participating components must act collectively for an immune or autoimmune response to emerge. Moreover, these processes span a spectrum of time and length scales that range from interactions between molecules in cells to phenomena that affect the entire organism; feedback loops between processes on different spatiotemporal scales are also important. It is often hard to intuit underlying principles from experimental observations because of the complexity of these hierarchically organized collective processes. The importance of stochastic effects further confounds intuition.

Statistical mechanics provides a conceptual framework and tools (theoretical models and associated computations) that relate microscopic stochastic events to emergent complex behavior. When these insights are coupled closely to biological experiments, underlying physical and chemical mechanisms can be elucidated. In this review, we describe a project that brings together statistical mechanics and cell biology to uncover new concepts in immunology.

The adaptive immune system is not preprogrammed to respond to prescribed pathogens, yet it mounts pathogen-specific responses against diverse microbes and establishes memory of past infections (the basis of vaccination). T lymphocytes (T cells) play an important role in coordinating adaptive immune responses. We explore how the developmental processes in an organ called the thymus gland shape the repertoire of T cells such that adaptive immunity exhibits both remarkable pathogen specificity and the ability to combat myriad pathogens. For the benefit of the uninitiated readers, the following bullet points present a minimal introduction to the features of the immune system relevant for our study (the Appendix provides a slightly expanded description of basic immunology):

- T cells are a type of white blood cell that originate in the bone marrow, mature in the thymus (a gland in front of the heart and behind the sternum), and move to other parts of the body (via blood and lymph vessels) to fight off infections (pathogens).
- Epitopes are protein fragments (peptides) that constitute the molecular signatures of pathogens recognized by T cells. Both self and pathogen proteins are routinely chopped

¹The first two paragraphs of the INTRODUCTION section (with small modifications) are reprinted with kind permission from Reference 11, the *Annual Review of Physical Chemistry*, Volume 61 © 2010, by Annual Reviews, http://www.annualreviews.org.

into pieces within a cell (by protease enzymes and the proteasome). Some fragments [typically 8–15 amino acids long (1)] can bind to another host protein—major histocompatibility complex (MHC)—and these peptide-MHC (pMHC) complexes are displayed on the surface of a cell. Each human inherits several different types of MHC proteins from parents; the different MHC types between individuals are implicated in transplant rejection. T cells inspect pMHC complexes and initiate immune responses if specific foreign peptides (epitopes) are encountered (see Figure 1*a*).

- T-cell receptors (TCRs) are proteins expressed on the surface of T cells, which bind to pMHC complexes. Recognition of the presence of an epitope, and subsequent T-cell response, is determined by the strength of the binding. During synthesis in the bone marrow, immature T cells (thymocytes) acquire distinct TCR sequences through a gene-shuffling process. Each such sequence can potentially bind strongly to a small number of complementary peptides, and a large repertoire of T cells is thus required to ensure proper coverage of the space of potential epitopes.
- Thymic selection: Following synthesis, thymocytes move to the thymus (7, 12–16), where they interact with a variety of self-pMHC molecules [few thousands of different

Major histocompatibility complex (MHC): proteins that present peptides to T-cell receptors

pMHC: peptide-MHC complex

T-cell receptors (TCRs): bind to peptide-MHCs presented on the surface of antigenpresenting cells



Figure 1

T-cell recognition of pathogen signatures. (*a*) Antigen-presenting cells (APCs) engulf pathogens and process their proteins into short peptides, which are bound to major histocompatibility complex (MHC) proteins and presented on the surface. T-cell receptors (TCRs) bind to peptide-MHCs, and sufficiently strong binding enables intracellular signaling and gene transcription, leading to T-cell activation. APCs also present self-peptides derived from self-proteins, but typically T cells are not activated by them. (*b*) TCR recognition of pathogen-derived pMHC molecules is both highly specific and degenerate. It is specific because if a TCR recognizes (*black check mark*) a peptide (*tan rectangle*), most point mutations of the peptide's amino acids (*red*) abrogate recognition (*red X mark*). However, a given TCR can also recognize diverse peptide sequences (*tan, green, magenta*). Panel *a* is adapted from Reference 11, figure 1a.

Thymocyte: immature T cell that goes to the thymus

types (13); see Figure 2]. These self-pMHCs are derived from diverse parts of the host proteome and expressed on the surface of thymic epithelial cells as well as macrophages and dendritic cells. For a thymocyte to exit the thymus and become part of the host's repertoire of T cells, it must pass two tests: (*a*) It must not be negatively selected; i.e., its TCR must not bind to any self-pMHC molecule it encounters with a binding free energy that exceeds a threshold for negative selection. (*b*) It must bind at least one self-pMHC molecule with a binding free energy that exceeds another threshold for positive selection. It is thought that negative selection serves to delete dangerous T cells that may be activated by self-pMHCs and cause autoimmune disease. The positive selection process ensures that the TCRs of selected T cells bind to a host's MHC (this is called MHC restriction), and the MHC can therefore recognize peptides bound to it (e.g., viral peptides).

The focus of this review is to understand how developmental processes in the thymus shape a T-cell repertoire that exhibits both remarkable pathogen specificity, as well as the ability to combat myriad pathogens. This puzzle of specificity/degeneracy is described in Section 2, where we also present a model of the thymic selection process. Computational studies of this model characterize the properties of the selected T-cell repertoire, which in turn elucidate the mechanism



Figure 2

Immature T cells (thymocytes) developed in the thymus. Thymocytes migrate through the thymus and interact with diverse self peptide–major histocompatibility complexes (self-pMHCs) presented on the surface of thymic antigen-presenting cells (APCs). A T cell's receptor (TCR) must bind to at least one of these self-pMHCs moderately to exit the thymus and become a part of the individual's T-cell repertoire (positive selection). A T cell with a TCR that binds to any self-pMHC with an affinity that exceeds a sharply defined threshold dies in the thymus (negative selection). This figure is adapted from Reference 11, figure 3a.

behind their specificity/degeneracy for pathogens. In Section 3, the above model of thymic selection is solved analytically by employing methods from statistical physics, such as extreme value distributions (EVDs) and Hamiltonian minimization. Genetic studies show that people with certain types of MHC are more likely to control HIV infections. We argue that these MHC types may affect thymic selection in a way that influences the statistical properties of the selected T-cell repertoire, and this may provide one contributing factor (of many) for more efficient control of HIV infection (Section 4).

2. SPECIFICITY AND DEGENERACY OF THE T-CELL REPERTOIRE²

TCR recognition of pathogen-derived pMHC molecules is both highly specific and degenerate (**Figure 1***b*). It is specific because if a TCR recognizes a pMHC molecule, most point mutations of the peptide's amino acids abrogate recognition (18, 19). However, a given TCR can also recognize diverse peptides (9, 20–24). This specificity-degeneracy conundrum is made vivid by dividing the world of peptides into classes, with the members of each class having sequences that are closely related. For example, peptides within a class could differ by just one point mutation. A TCR can discriminate quite well between peptides within a class of closely related peptide sequences (as point mutants of the peptides it recognizes are not recognized with high probability). But, at the same time, a given TCR can recognize some other peptides in other classes, which have quite distinct sequences.

This diverse, specific/degenerate (Figure 1b), and largely self-tolerant TCR repertoire is designed during T-cell development in the thymus [Figure 2; (7, 12–16)]. Signaling events, gene transcription programs, and cell migration during T-cell development in the thymus have been extensively studied (7, 12, 13, 15, 16, 25-30). Experiments by Huseby et al. (18, 19) provide important clues about how interactions with self-pMHC complexes in the thymus shape the peptide binding properties of selected TCR amino acid sequences, such that mature T cells exhibit their special properties. These experiments contrasted T cells developed in conventional mice that display a diverse array of self-pMHC complexes (few thousands of types) in the thymus to mice engineered to express only one type of peptide in their thymus. For T cells developing in conventional mice, recognition of antigenic pMHC was found to be sensitive to most point mutations of a recognized foreign peptide's amino acids. In contrast, T cells selected in mice with only one type of peptide in the thymus were much more peptide degenerate, with some T cells being tolerant to several point mutations of recognized foreign peptide amino acids. The thymic selection model (17, 31-33) presented next explains these results and also sheds light on the broader question of how the thymus designs diverse TCR sequences that mediate specific/degenerate pathogen recognition.

2.1. Thymic Selection Model

The key initiating event in T-cell activation is the binding of a TCR to a pMHC complex. The binding interface of a TCR is composed of a more conserved region that is in contact with the MHC molecule and a highly variable (CDR3) region that makes the majority of contacts with the peptide (1). Accordingly, we divide the TCR/pMHC interaction free energy into two parts: a more conserved part represented by a continuous variable and a part that explicitly depends on the variable TCR-peptide contact residues and peptide sequences of amino acids. The former is given

²Some parts of Section 2 are reprinted with kind permission from Reference 17 © 2011, by Springer Science+Business Media.

a value E_c , which may be varied to describe different TCRs and MHCs. The latter is obtained by aligning the TCR and pMHC amino acids that are treated explicitly and adding (in the simplest incarnation of the model) pairwise interactions between corresponding pairs (see, Figure 3*a*). If the amino acid sequences of the TCR and the peptide are represented by strings $\vec{t} \equiv (t_1, t_2, \dots, t_N)$ and $\vec{s} \equiv (s_1, s_2, \dots, s_N)$, respectively, the interaction free energy is

$$E_{\text{int}}\left(E_{c}, \overrightarrow{t}, \overrightarrow{s}\right) = E_{c} + \sum_{i=1}^{N} J(t_{i}, s_{i}).$$

$$1.$$

The contribution from the *i*th amino acid of the TCR (t_i) and the corresponding peptide residue (s_i) is indicated by the matrix element $J(t_i, s_i)$. For numerical purposes we use the Miyazawa-Jernigan amino acid interaction matrix (34) developed in the context of protein folding, but as emphasized later, the qualitative results do not depend on the form of J. The length of the variable TCR-peptide region is taken to be $N \sim 5$. This is the typical number of peptide amino acids in contact with the TCR; the remaining peptide residues are important for binding to the MHC groove and/or are buried within the groove.

Earlier versions of such string models of TCR-pMHC interactions were used to study thymic selection (28, 29, 35), but they did not employ an explicit treatment of amino acids (e.g., a formal string of numbers, bits, etc., was used). These studies provided estimates for certain properties of the selected TCR repertoire (for example, the number of selected TCRs activated by a foreign peptide or a foreign MHC—as in an organ transplant) that are consistent with experimental estimates. They also showed that negative selection in the thymus increases TCR specificity for foreign peptides but did not suggest any mechanistic explanation. Other string models of TCR-pMHC interactions (36–38), with analogies to spin-glass models, were used also to study vaccination strategies for viral diseases and cancers (for a review, see Reference 39).

To model thymic selection, we first construct a set $S \equiv \{\vec{s}\}$ of M peptides of length N representing the self-peptides encountered in the thymus. Each self-peptide is generated as a sequence of N amino acids, each randomly and independently picked with frequencies corresponding to the human proteome (31, 40) [using the mouse proteome does not change the qualitative results (31)]. We next generate many candidate sequences for the peptide contact residues of TCRs, \vec{t} , also randomly with the same amino acid frequencies. To mimic thymic selection, TCR sequences that bind to any of the M self-pMHC too strongly ($E_{int} < E_n$, with more negative free energies corresponding to stronger binding) are deleted (negative selection). However, a TCR must also bind sufficiently strongly ($E_{int} < E_p$) to at least one self-pMHC to receive survival signals and emerge from the thymus (positive selection). Recent experiments show that the difference between the thresholds for positive and negative selection is relatively small [a few $k_BT(27)$]. The threshold for negative selection (E_n) is quite sharp, whereas the thresholds at E_n and E_p does not change the qualitative behavior of the selected T-cell repertoire (17, 31).

To completely specify the interaction free energy between a TCR and pMHC, we need to discuss the value of E_c . Selected TCRs are expected to bind moderately to MHCs, because binding too strongly to MHC (large $|E_c|$) would result in negative selection with any peptide, and binding too weakly (small $|E_c|$) results in TCRs not being positively selected. Each human can have up to 12 different MHC types. A TCR that binds strongly to more than one MHC type is likely to be eliminated during negative selection. Therefore, only TCRs binding to a particular MHC type are considered. This is consistent with the fact that there are no firm reports of a TCR restricted by more than one MHC type within a single human. Variations in E_c for selected TCRs are expected to be small. A rough estimate on the bounds is obtained from the condition that the average



Figure 3

Effects of thymic selection on the characteristics of T-cell receptors (TCRs) selected against M types of self-peptides. (a) Schematic representation of the interface between TCR and peptide major histocompatibility complexes (pMHCs). The region of the TCR contacting the peptide is highly variable and is modeled by a string of amino acids of length $N \sim 5$. The peptide is also treated similarly. (b) Amino acid composition of selected TCRs. TCRs selected against many types of self-peptides in the thymus have peptide contact residues that are enriched in amino acids that interact weakly with other amino acids. (c) Probability density distribution of E_c values (strength of TCR binding to MHC) of TCRs selected against M types of self-peptides. TCRs selected against many types of self-peptides that are recognized by at least one of the selected TCRs. TCRs selected against many types of self-peptides that are enriched with strongly interacting amino acids. Amino acids on the abscissa in panels b and d are ordered according to their largest interaction strength with other amino acids in the interaction matrix, J. This figure is adapted from Reference 17, figures 1 and 4.

interaction free energy between TCR and pMHC for selected TCRs should be between the thresholds for positive and negative selection; i.e.,

$$E_n < E_c + N\overline{J} < E_p, \qquad 2.$$

where \overline{J} is the average value of interaction between amino acids. The two bounds $(E_{c,\max} = E_p - N\overline{J} \text{ and } E_{c,\min} = E_n - N\overline{J})$ ensure that average interactions enable a TCR to survive both positive and negative selection. Because it is enough that a TCR sequence is positively selected by any one of many self-peptides, and not negatively selected by all M self-peptides, the precise bounds for E_c are different, but one expects that the range of E_c values is still small; viz., $E_{c,\max} - E_{c,\min} \propto E_p - E_n$. TCRs whose interactions with MHCs are too weak are unlikely to be properly positioned on MHCs, and hence will be unable to interact with the peptide. Thus, one cannot tune E_c to very low values to escape negative selection. We thus assign to every TCR sequence a random value of E_c chosen uniformly from the interval ($E_{c,\min}, E_{c,\max}$) and then proceed with computing the consequences of the selection process.

2.2. Abundance of Weak Interactions in T-Cell Receptors Selected Against Many Self-Peptides

First, we summarize the results of computational analyses regarding how thymic selection shapes TCR sequences and TCR interactions with MHC. The peptide contact residues of TCR sequences selected against many self-peptides in mouse and humans $[M \sim 10^3 (13)]$ are statistically enriched with weakly interacting amino acids [Figure 3b; (17, 31, 32)], and TCRs with weaker binding to MHC (within the allowed range) are more likely to get selected against many self-peptides, TCR sequences with peptide contact residues containing strongly interacting amino acids (e.g., hydrophobic amino acids or those with flexible side chains), or TCRs that bind strongly to MHC, are more likely to have strong binding with at least one encountered self-pMHC and thus be negatively selected. These qualitative results are robust and are not restricted to particular forms of the interaction potential *J*, or the sharpness of the thresholds for positive and negative selection [see Section 3; (17, 31, 32)]. Using different interaction potentials only changes the identities of the amino acids that interact weakly or strongly, or the criterion used to define interaction strength.

The analysis of available crystal structures of TCR-pMHCs supports the conclusion that the peptide contact residues of selected TCRs are enriched in weakly interacting amino acids (31): Amino acid frequencies of peptide contacting residues on TCRs in these crystal structures were determined and compared to amino acid frequencies in the human proteome [assumed to be the relevant frequencies for TCRs before thymic selection (31)]. Measured amino acid frequencies in the TCRs' peptide contact residues were found to be smaller than in the human proteome for the most strongly interacting amino acids (IVYWREL; 42) and larger for the weakly interacting amino acids (QSNTAG; 42).

2.3. Selection Against Many Self-Peptides Leads to Pathogen-Specific T Cells

Does the selected T-cell repertoire lead to specific recognition of a pathogenic peptide? To study the specificity of mature T cells in peptide recognition, selected TCR sequences were challenged with a collection of many randomly generated pathogenic peptides whose amino acid frequencies correspond to *Listeria monocytogenes* (31, 43), a pathogen that infects humans and is cleared by a T-cell response. TCR recognition of pathogenic peptides occurs if TCR-pMHC binding is sufficiently strong ($E_{int} < E_r$), where the recognition threshold in mouse experiments is such that $E_r \sim E_n$ (44). For each TCR that recognizes a particular pathogenic peptide sequence, the specificity of recognition was tested as follows: Each site on the peptide was mutated to all other 19 possibilities, and recognition of the mutated sequence by the original TCR was assessed. If more than half the mutations at a particular site abrogated recognition by the same TCR, the site was labeled an "important contact." For each TCR-pMHC pair for which recognition occurred, the number of important contacts was determined, and the resulting histogram is plotted in **Figure 4***a*. The higher the number of important contacts, the more specific is the TCR recognition of pathogenic peptides. Small numbers of important contacts correspond to cross-reactive TCRs that are able to recognize many pathogenic peptide mutants.

In agreement with experiments (18, 19), this model finds that TCRs selected against many different self-peptides are very specific, whereas TCRs selected against only one self-peptide are more cross-reactive (Figure 4*a*). On the basis of the amino acid composition of selected TCRs, we can provide a mechanistic explanation for the specificity/degeneracy of pathogen recognition (Figure 4*b*). Because TCR-peptide contact residues are enriched with weakly interacting amino acids, they can interact sufficiently strongly for recognition to occur with pathogenic peptides that are statistically enriched in amino acids that are the stronger binding complements of TCR-peptide contact residues (Figure 3*d*). Such TCR-peptide pairs rely on many weak to moderate interactions, which sum up to provide sufficient binding strength for recognition. Each interaction contributes a significant percentage of the total binding affinity. If there is a mutation to an amino acid of a recognized peptide, it is likely to weaken the interaction it participates in (as recognized peptides are statistically enriched in amino acids that interact strongly with the TCR's amino



Figure 4

Mechanism for specificity and degeneracy of T-cell receptor (TCR) recognition of antigenic peptides. (*a*) Histogram of the number of important contacts with which T cells recognize pathogenic peptides. Single point mutations of pathogenic peptides at important contacts likely abrogate T-cell recognition. T cells selected against many self-peptides recognize pathogenic peptides via many important contacts and are thus specific. In contrast, T cells selected against few types of self-peptides recognize pathogenic peptides with only a few important contacts and are thus cross-reactive. (*b*) The weakly interacting amino acids (*light blue*) on the TCR bind to strongly interacting amino acids (*pink*, *brown*) on antigenic peptides, resulting in multiple moderate-scale interactions that add up to a total binding free energy that is large enough for recognition. Because antigen recognition is mediated by multiple interactions of moderate value, each contact makes a significant contribution to the total interaction free energy necessary for recognition. Therefore, disrupting any interaction by mutating one of the strongly interacting amino acids on the peptide results (shown as a change from *pink* to *gray* color) in abrogation of recognition. At the same time, TCR recognition of antigenic peptides is degenerate, because there are many combinatorial ways of distributing strongly interacting amino acids (*pink*, *brown*) along the peptide, which results in a sufficiently strong binding with TCR for recognition. This figure is adapted from Reference 11, figure 3d, and Reference 17, figure 4a.

acids). Weakening an interaction that contributes a significant fraction of the binding free energy is likely to abrogate recognition because the recognition threshold is sharply defined (27).

In contrast, TCR sequences selected against only one type of self-peptide have a higher chance of containing strongly interacting amino acids (**Figure 3**). Such TCRs can recognize a lot more pathogenic peptides including those composed of weakly or moderately interacting amino acids. In many cases, mutating such amino acids on the peptide does not prevent recognition of the same TCR because a small number of strong contacts dominate recognition [**Figure 4***a* and experiments (19)]. Unless these specific contacts are disrupted by mutations to the peptide, recognition is not abrogated. Accordingly, TCR recognition of pathogenic peptides is more cross-reactive. When selected against fewer types of self-peptides, TCRs that bind strongly to MHCs can escape (**Figure** *3c*). Thus in this case the escape of TCRs that bind strongly or moderately to more than one MHC type (or MHCs with mutations) might also be possible, leading to more cross-reactivity to MHC types [or substitutions of MHC amino acids (18)].

This mechanism for TCR-pMHC specificity is distinct from Fischer's (45) lock-and-key metaphor. Interactions between the TCR and the MHC dock the TCR over its ligand in essentially the same orientation (46, 47)—this may be analogous to shape complementarity, but it is not peptide specific. The complementary residues of the TCR then scan the peptide to assess if there is a sufficient number of moderate interactions to mediate recognition (Figure 4b). An appropriate metaphor may be that the TCR-peptide contact residues scan a bar code, and if there are a sufficient number of lines of moderate width (moderate TCR-peptide interactions), then recognition is possible. This statistical view of TCR specificity for pathogens may describe the initial step of binding, which may then allow modest conformational adjustments, leading to stronger binding (20). This view is consistent with experiments suggesting a two-stage model for TCR-pMHC binding (48).

The statistical view of TCR-pMHC recognition also makes degeneracy or cross-reactivity to peptides with different sequences the flip side of the coin. Although point mutations can abrogate recognition with high probability, making a number of changes to the peptide sequence such that a sufficient number of moderate interactions is still obtained will allow recognition by the same TCR (**Figure 4b**). This may also explain why two peptides with different sequences and conformations in the MHC groove can be recognized by the same TCR (20).

3. T-CELL DEVELOPMENT AND PATHOGEN RECOGNITION AS EXTREME VALUE PROBLEMS³

3.1. Thymic selection of T cells

Interestingly, the thymic selection model presented in Section 2.1 can be solved exactly in the limit of long peptide sequences $(N \rightarrow \infty)$ (17, 32). A T cell expressing TCR with string \vec{t} is selected in the thymus if its strongest interaction with a set S of M self-pMHCs is between the thresholds for negative and positive selection; i.e.,

$$E_n < \min_{\overrightarrow{s} \in S} \left\{ E_{\text{int}} \left(E_c, \overrightarrow{t}, \overrightarrow{s} \right) \right\} < E_p.$$
3.

Equation 3 casts thymic selection as an extreme value problem (49), enabling us to calculate the probability $P_{sel}(E_c, \vec{t})$ that a TCR sequence \vec{t} is selected in the thymus. Let us indicate by

³Some parts of Section 3 are reprinted with kind permission from Reference 32 © 2009, by the American Physical Society, and from Reference 17 © 2011, by Springer Science+Business Media.

 $g(x|E_c, \vec{t})$ the probability density function (PDF) of the interaction free energy between the TCR \vec{t} and a random peptide. The PDF $\Pi(x|E_c, \vec{t})$ of the strongest (minimum) of the *M*-independent random free energies is then obtained by multiplying *g* with the probability of all remaining (M-1) free energy values being larger; i.e.,

$$\Pi\left(x|E_{c}, \overrightarrow{t}\right) = Mg\left(x|E_{c}, \overrightarrow{t}\right) \left(1 - P\left(E < x|E_{c}, \overrightarrow{t}\right)\right)^{M-1},$$
4

where $P(E < x | E_c, \vec{t})$ is the cumulative probability, and noting the multiplicity *M* of a particular interaction free energy being lowest. The probability that TCR \vec{t} is selected is then obtained by integrating $\Pi(x|E_c, \vec{t})$ over the allowed range, as

$$P_{\rm sel}\Big(E_c, \overrightarrow{t}\Big) = \int_{E_n}^{E_p} \Pi\Big(x|E_c, \overrightarrow{t}\Big) dx.$$
 5.

For $M \gg 1$, this EVD converges to one of three possible forms (49), depending on the tail of the PDF for each entry. Equation 1 indicates that in our case, as each interaction free energy is the sum of *N* contributions, $g(x|E_c, \vec{t})$ should be a Gaussian for large *N*, in which case the relevant EVD is the Gumbel distribution (49).

To obtain an explicit form for $\Pi(x|E_c, \vec{t})$, we model the set of self-peptides as *M* strings in which each amino acid is chosen independently. The probability f_a for selecting amino acid *a* at each site is taken to be the frequency of this amino acid in the self-proteome. For a specific TCR sequence \vec{t} , the average interaction free energy with self-peptides then follows from Equation 1 as

$$E_{\rm av}\left(E_c, \overrightarrow{t}\right) = E_c + \sum_{i=1}^N \mathcal{E}(t_i), \qquad 6$$

with $\mathcal{E}(t_i) = [J(t_i, a)]_a$, where we have denoted the average over self-amino acid frequencies by $[G(a)]_a \equiv \sum_{a=1}^{20} f_a G(a)$. Similarly, the variance of the interaction free energy is

$$V(E_c, \overrightarrow{t}) = \sum_{i=1}^{N} \mathcal{V}(t_i), \qquad 7.$$

where $\mathcal{V}(t_i) = \left[J(t_i, a)^2\right]_a - \left[J(t_i, a)\right]_a^2$.

For very long peptide sequences (large N), we can approximate $g(x|E_c, \vec{t})$ with a Gaussian PDF with the above mean and variance. From standard results for the Gumbel distribution (49), we conclude that in the limit of $M \gg 1$, the peak of the distribution $\Pi(x|E_c, \vec{t})$ drifts to lower values as

$$E_0(E_c, \vec{t}) = E_{\rm av}(E_c, \vec{t}) - \sqrt{2V(E_c, \vec{t})} \ln M, \qquad 8.$$

while its width is reduced to

$$\Sigma_0 \left(E_c, \overrightarrow{t} \right) = \sqrt{\frac{\pi^2 V \left(E_c, \overrightarrow{t} \right)}{12 \ln M}}.$$
9

[Because the PDF $g(x|E_c, \vec{t})$ originates from a bounded set of free energies, it is strictly not Gaussian in the tails. Hence, once the extreme values begin to probe the tail of the distribution, the above results will no longer be valid. Indeed, in the limit when $M \sim \mathcal{O}(20^N)$, the EVD will approach

a delta function centered at the *M*-independent value corresponding to the optimal binding free energy.]

From Equation 8 and the selection condition in Equation 3, we see that as the number of selfpeptides, M, increases, the chance of negative selection does too. To counterbalance this pressure for large M, TCRs are enriched with weakly interacting amino acids in their peptide contact residues [small $\mathcal{E}(t_i)$ values] and with weaker interactions with MHC (small E_c value) (see Figure 3). A similar effect relates to the variance of interactions (avoiding negative selection against many self-peptides picks out TCRs with amino acids that exhibit a smaller variance in their interactions with other amino acids), but this tendency is less pronounced because of the square root. The preference for weak binding is independent of the potential J between contacting amino acids; different potentials merely reorder weak and strong amino acids.

Statistical mechanics suggests an analytic expression for the probability that a TCR sequence, \vec{t} , is selected according to Equation 3 in the limit of large N and M. Remarkably, the results seem to be accurate even for short peptides (17, 32). A proper thermodynamic limit is obtained when $\{E_c, E_p, E_n\} \propto N$, and $\ln M \propto N$. The latter ensures that the peak of the EVD distribution, $E_0(E_c, \vec{t})$, in Equation 8 is proportional to N. The same condition also implies that the width $\Sigma_0(E_c, \vec{t})$, in Equation 9 is sharp and independent of N. The relation $\ln M = \alpha N$ can be justified from the expectation that M should grow proportionately to the proteome size P, while $N \propto \ln P$ to enable encoding of the proteome. (The number of distinct peptide sequences of length N grows as 20^N , thus enabling encoding of proteomes with $P \leq 20^N$.) In this large N limit, the EVD is sufficiently narrow that the value of the optimal free energy can be precisely equated with the peak $E_0(E_c, \vec{t})$, and Equation 3 for the selection condition can be replaced with

$$E_n < E_0 \left(E_c, \overrightarrow{t} \right) < E_p.$$
 10.

The above thymic selection condition can now be interpreted as defining a microcanonical ensemble of sequences \vec{t} , which are accepted if the value of the Hamiltonian $E_0(E_c, \vec{t})$ falls on the interval (E_n, E_p) . In the large N limit, canonical and microcanonical ensembles are equivalent and the probability is given by the Boltzmann weight of this Hamiltonian. More formally, the probability for TCR selection, $P_{sel}(E_c, \vec{t})$, is obtained by using the least biased estimate, i.e., maximizing the entropy

$$S = \sum_{E_{c}, \vec{t}} P_{\text{sel}}(E_{c}, \vec{t}) \ln \left[P_{\text{sel}}(E_{c}, \vec{t}) \right], \qquad 11.$$

subject to a constraint that the average free energy,

$$\langle E_0(E_c, \vec{t}) \rangle = \sum_{E_c, \vec{t}} P_{\text{sel}}(E_c, \vec{t}) E_0(E_c, \vec{t}),$$
 12.

is restricted to the interval (E_n, E_p) . This leads to a probability for TCR selection governed by the Boltzmann-like weight (17, 32)

$$P_{\rm sel}\left(E_c, \overrightarrow{t}\right) \propto \left(\prod_{i=1}^N f_{t_i}\right) \rho(E_c) \exp\left[-\beta E_0\left(E_c, \overrightarrow{t}\right)\right].$$
 13.

Here, $\{f_a\}$ and $\rho(E_c)$ are the prior frequencies of amino acids, and the distribution of E_c values before selection, whereas the effect of thymic selection is captured by the Boltzmann weight, with a Lagrange multiplier β determined by the condition $E_n < \langle E_0(E_c, \vec{t}) \rangle < E_p$. Because the allowed

values of $E_0(E_c, \vec{t})$ are bounded from above and below, the Lagrange multiplier β can be either negative or positive.

A difference compared to the standard microcanonical ensemble is that the average free energy is constrained to an interval, rather than a precise value, necessitating a discussion on the choice of β . The possible values for $E_0(E_c, \vec{t})$ span a range from E_{\min} to E_{\max} , and the corresponding number of states form a bell-shaped curve between these extremes with a maximum at some E_{\min} . If $E_{\min} > E_p$, to maximize entropy we must set $\langle E_0(E_c, \vec{t}) \rangle = E_p$ and choose β accordingly. In this case, $\beta > 0$, positive selection is dominant, and stronger amino acids are selected. If $E_{\min} < E_n$, we must set β such that $\langle E_0(E_c, \vec{t}) \rangle = E_n$, $\beta < 0$, negative selection is dominant, and weaker amino acids are selected. For $E_n < E_{\min} < E_p$, we must set $\beta = 0$, and there is no modification due to thymic selection.

Finally, due to the appearance of $\sum_{i=1}^{N} \mathcal{V}(t_i)$ under the square root term, Equation 8 corresponds to an interacting Hamiltonian in which variables at different sites are apparently not independent. This is, however, not the case as the interaction is easily removed by standard procedures such as Legendre transforms or Hamiltonian minimization (50), as follows: We need to solve a Hamiltonian $\mathcal{H}(U, V)$ that depends on two extensive quantities $U = \sum_{i=1}^{N} \mathcal{E}(t_i)$ and $V = \sum_{i=1}^{N} \mathcal{V}(t_i)$. The corresponding partition function can be decomposed as $Z = \sum_{U,V} \Omega(U, V) e^{-\beta \mathcal{H}(U,V)}$, but can be approximated with its largest term. The same density of states $\Omega(U, V) \equiv e^{S(U,V)/k_B}$ appears, irrespective of the specific form of $\mathcal{H}(U, V)$. In particular, the choice

$$\mathcal{H}_0(U, V) = E_c + U - \gamma V - \frac{\ln M}{2\gamma} = E_c + \sum_{i=1}^N \left[\left(\mathcal{E}(t_i) - \gamma \mathcal{V}(t_i) \right) \right] - \frac{\ln M}{2\gamma}$$
 14

corresponds to a set of noninteracting variables, with

$$P_{\text{sel}}\left(E_{c}, \overrightarrow{t}\right) \propto \rho(E_{c}) \exp[-\beta E_{c}] \prod_{i=1}^{N} \left\{ f_{t_{i}} \exp[-\beta(\mathcal{E}(t_{i}) - \gamma \mathcal{V}(t_{i}))] \right\},$$
 15.

for which thermodynamic quantities (such as entropy) are easily computed. By judicious choice of γ , we can then ensure that the same average free energy appears for $\mathcal{H}_0(E_c, \vec{t})$ and our $E_0(E_c, \vec{t})$. Using Legendre transforms, which is equivalent to minimizing $\mathcal{H}_0(E_c, \vec{t})$ with respect to γ , one finds that the required $E_0(E_c, \vec{t})$ is obtained by setting

$$\gamma(\beta) = \sqrt{\frac{\ln M}{2N\langle \mathcal{V} \rangle}},$$
16

where $\langle \ldots \rangle$ refers to the average with the noninteracting weights in Equation 15.

In practice we determine parameters β and γ as follows: Because the average free energy $\langle E_0(E_c, \vec{t}) \rangle$ is a monotonic function of β , we use a bisection method to find the appropriate β that corresponds to the specified value of the average. To do that, we need to discuss how to evaluate the average free energy for a particular β . First we use a bisection method to find a self-consistent value of γ from Equation 16 and then calculate the average free energy using the Boltzmann weight in Equation 15. We thus find β_p and β_n corresponding to $\langle E_0(E_c, \vec{t}) \rangle = E_p$ and $\langle E_0(E_c, \vec{t}) \rangle = E_n$, respectively. On the basis of earlier discussion, we set $\beta = \beta_p$, when $0 < \beta_p < \beta_n$; $\beta = 0$, when $\beta_p < 0 < \beta_n$; and $\beta = \beta_n$, when $\beta_p < \beta_n < 0$.

Figure *5a* depicts the variation of β as a function of ln (*M*)/*N* and the threshold for negative selection E_n , with $(E_p - E_n)/N = 0.5k_BT$. With the thus obtained parameters β and γ we find the amino acid frequencies of selected TCRs as



Figure 5

Representation of the dependency of the parameters (*a*) β , a measure of amino acid composition of selected T-cell receptors (TCRs), and (*b*) β^* , a measure of amino acid composition of pathogenic peptides recognized by selected TCRs, on the number of selected TCRs (*K*), the number of self-peptides (*M*) against which TCRs were selected, and the threshold for negative selection E_n . In panel *a*, the region between the white lines corresponds to $\beta = 0$, to the right (left) of which negative (positive) selection is dominant, and weak (strong) amino acids are selected. The blue dashed lines in panels *a* and *b* indicate the relevant parameter values for thymic selection in mouse. In panel *b*, the solid white line separates regions with $\beta^* > 0$ (only foreign peptides with strongly interacting amino acids are recognized) and $\beta^* = 0$ (every foreign peptide is recognized). The region below the black dashed line in panel *b* corresponds to $\beta = 0$ (every TCR is selected). In panel *b*, the threshold for negative selection (E_n) is fixed. This figure is adapted from Reference 17, figures 2 and 4.

$$f_a^{(\text{sel})} = \frac{f_a \exp[-\beta(\mathcal{E}(a) - \gamma \mathcal{V}(a))]}{\sum_{b=1}^{20} f_b \exp[-\beta(\mathcal{E}(b) - \gamma \mathcal{V}(b))]},$$
17.

and the distribution of selected TCRs' interactions with MHCs as

$$\rho^{(\text{sel})}(E_c) = \frac{\rho(E_c) \exp[-\beta E_c]}{\int_{E_{c_1 \min}}^{E_{c_1 \max}} \rho(E) \exp[-\beta E] dE}.$$
18.

The above analytic expressions agree very well with numerical results from computer simulations of short peptides (N = 5) presented in the previous section (17, 32).

3.2. Nature of Foreign Peptides Recognized by T Cells

After T cells complete thymic selection, a set \mathcal{T} of TCRs, K in number, is available to respond to pathogens. A T cell recognizes infected cells when its TCR binds sufficiently strongly ($E_{int} < E_n$) to a foreign pMHC. This means that a foreign peptide of sequence \vec{s} is recognized by some TCRs if its strongest interaction with the set of TCRs exceeds the threshold for recognition; i.e.,

$$\min_{\overrightarrow{t} \in \mathcal{T}} \left\{ E_{\text{int}} \left(E_c, \overrightarrow{t}, \overrightarrow{s} \right) \right\} < E_n,$$
19.

where the minimization is over the set of K TCRs (each with given E_c and \vec{t}) selected in the thymus.

Equation 19 casts recognition of foreign peptides as another extreme value problem. If we model the set T as K strings in which each amino acid is chosen independently with frequencies $f_a^{(sel)}$ (i.e., ignoring correlations among different positions on one string and also between strings), then in the limit of large $K \gg 1$, the EVD is sharply peaked around (17)

$$E_0^*(\overrightarrow{s}) = \langle E_c \rangle + \sum_{i=1}^N \mathcal{E}^*(s_i) - \sqrt{(2\ln K) \left[\langle E_c^2 \rangle_c + \sum_{i=1}^N \mathcal{V}^*(s_i) \right]}, \qquad 20.$$

and its width is

$$\Sigma_0^*(\vec{s}) = \sqrt{\frac{\pi^2 \left[\langle E_c^2 \rangle_c + \sum_{i=1}^N \mathcal{V}^*(s_i) \right]}{12 \ln K}}.$$
 21.

As $\ln K \propto N \rightarrow \infty$, the distribution becomes very narrow, and the condition for recognition of foreign peptides becomes

$$E_0^*(\vec{s}) < E_n. \tag{22}$$

The mean $\mathcal{E}^*(s_i)$ and the variance $\mathcal{V}^*(s_i)$ of the amino acid interaction free energies are obtained as in the previous section after replacing f_a with $f_a^{(sel)}$. The mean $\langle E_c \rangle$ and the variance $\langle E_c^2 \rangle_c =$ $\langle E_c^2 \rangle - \langle E_c \rangle^2$ of selected TCR interactions with MHCs are obtained using $\langle X \rangle = \int_{E_c, \min}^{E_c, \max} X \rho^{(sel)}(E_c) \exp[-\beta E_c] dE_c$, with $\rho^{(sel)}(E_c)$ given in Equation 18.

Repeating the reasoning of the previous section, the probability for a sequence \vec{s} to be recognized is governed by the Boltzmann weight $P_{\text{rec}}(\vec{s}) \propto \left(\prod_{i=1}^{N} \tilde{f}_{s_i}\right) \exp[-\beta^* E_0^*(\vec{s})]$, where $\{\tilde{f}_a\}$ are prior frequencies of amino acids in the pathogen proteome, whereas the effect of TCR

HLA: human MHC proteins

recognition is captured by the parameter β^* . As before, we introduce a new Hamiltonian $H_0^*(\vec{s}) = \langle E_c \rangle - \gamma^* \langle E_c^2 \rangle_c + \sum_{i=1}^N [\mathcal{E}^*(s_i) - \gamma^* \mathcal{V}^*(s_i)] - \ln K/(2\gamma^*)$, and to ensure the same average free energies, $\langle E_0^*(\vec{s}) \rangle = \langle H_0^*(\vec{s}) \rangle$, we set $\gamma^*(\beta^*) = \sqrt{\ln K/(2\langle E_c^2 \rangle_c + 2N\langle \mathcal{V}^* \rangle)}$. Finally, β^* is determined by constraining $\langle E_0^*(\vec{s}) \rangle < E_n$, while maximizing entropy. If $\beta^* > 0$, only foreign peptides with stronger amino acids are recognized. If $\beta^* = 0$, recognized peptides are not enriched or attenuated in strongly interacting amino acids. Unlike the parameter β for thymic selection of T-cell receptors, β^* cannot be negative as there is no lower free energy bound for recognition in Equation 22. The amino acid frequencies of recognized foreign peptides are then

$$\tilde{f}_{a}^{(\text{rec})} = \frac{\tilde{f}_{a} \exp[-\beta^{*}(\mathcal{E}^{*}(a) - \gamma^{*}\mathcal{V}^{*}(a))]}{\sum_{b=1}^{20} \tilde{f}_{b} \exp[-\beta^{*}(\mathcal{E}^{*}(b) - \gamma^{*}\mathcal{V}^{*}(b))]}.$$
23.

Figure 5*b* depicts variation of β^* as a function of the number of selected TCRs (*K*), the number of self-peptides (*M*) against which TCRs were selected, and the threshold for negative selection E_n with $(E_p - E_n)/N = 0.5k_BT$. For selected TCRs to recognize many foreign peptides (i.e., small value of β^*), we must have $K \gg M$ (i.e., a lot more selected TCRs than self-peptides presented in the thymus). This is consistent with biological values of $K \sim 10^9$ T cells (1) and $M \sim 10^3$ self-peptides (13) in humans.

Equation 23 does not agree as well with the numerical results of computer simulations for short peptides (N = 5), as the corresponding ones for the selected TCR sequences presented before. The reason for the discrepancies is likely in the incorrect assumption that the selected TCR sequences are uncorrelated for small N = 5 (17). However, the qualitative behavior of the parameter β^* as other model parameters are varied is expected to remain valid (Figure 5b).

4. AN ASPECT OF THE ROLE OF HOST GENETICS IN CONTROL OF THE HUMAN IMMUNODEFICIENCY VIRUS THAT MAY BE RELATED TO THYMIC DEVELOPMENT⁴

Each individual inherits a particular set of MHC molecules (up to six types of each MHC class I and class II protein) from their parents. Insights in to how thymic development shapes the T-cell repertoire suggest a previously unknown aspect of how these differences in host genetics can influence the ability of humans to combat infectious diseases (such as HIV). HIV is a highly mutable and rapidly replicating virus that infects human T cells (among other cell types). HIV infection initially leads to acute high-level viremia (the measurable presence of virus in the bloodstream), which is subsequently reduced to lower levels by the immune system. Without therapy, most patients experience a subsequent increase in viral load and ultimately the development of AIDS. AIDS is associated with the occurrence of opportunistic infections because of the degradation of the immune system (T cells). Viremia levels and time to disease vary widely, and the differences correlate with the expression of different MHC class I molecules (as reviewed in Reference 51). Rare individuals (elite controllers) maintain very low levels of HIV without therapy, thereby making disease progression and transmission unlikely. Certain MHC types appear more in elite controllers, with the highest association observed for the so-called HLA-B57 (52, 53). Although many complex factors may be at play, this fact suggests the involvement of T cells in viral control given that T cells activated by MHC-bound viral peptides play an important role during various phases of disease (54-59). T cells in people with different MHC genes could influence viral

⁴Some parts of Section 4 are reprinted with kind permission from Reference 33 © 2010, by Nature Publishing Group.

control in diverse ways. For example, it is known that MHC molecules associated with control present peptides derived from the HIV proteome that are vulnerable to mutations (51), especially because of collective effects of multiple simultaneous deleterious mutations (60). Thus, T cells in people with these MHC molecules are thought to target more vulnerable regions of HIV, thereby hindering mutational escape from the host immune pressure.

A puzzling finding is that the MHC molecules most associated with enhanced control of HIV, HLA-B57, and HLA-B27, are also associated with increased proclivity for certain autoimmune disorders. Indeed, HLA-B57 has been associated with autoimmune psoriasis (61) and hypersensitivity reactions (62), and HLA-B27 with ankylosing spondylitis (63). The understanding of the role of thymic development in shaping the T-cell repertoire that has emerged from experimental and theoretical studies (vide supra) may explain the mechanistic origins of these observations.

Bioinformatics algorithms (64) based on experimental data predict whether a particular peptide will bind to a given MHC molecule (33). Using these algorithms, the fraction of peptides derived from the human proteome (65) that bind to various MHC molecules were computed. Of the roughly 10⁷ unique peptide sequences, only 70,000 are predicted to bind to HLA-B57, while 130,000 bind to a typical HLA-B molecule, and 180,000 bind to HLA-B7 (an MHC type that is associated with faster progression to AIDS) (33).

The intrinsic differences in self-peptide binding among MHC molecules can be important during development of immature T cells in the thymus. As fewer self-peptides are able to bind to HLA-B57 molecules, a smaller diversity of self-pMHCs are encountered by HLA-B57-restricted T cells in the thymus. Thus, as described earlier, HLA-B57-restricted T cells are likely to be more cross-reactive to point mutants of targeted viral peptides than T cells restricted by MHC types that present a greater diversity of self-peptides (Figure 4*a*). This finding is supported by experiments measuring the cross-reactivity of T cells from people with diverse MHCs for HIV peptides (66–68).

A model of host-HIV dynamics showed that a repertoire of T cells more cross-reactive to point mutants of targeted epitopes results in better control of HIV infection (33). This is because such T cells can exert immune pressure on the infecting strain and mutants that rapidly emerge to escape the immune pressure more effectively. Thus, it was predicted that HIV-infected individuals with MHC types that bind fewer self-peptides are more likely to control viral loads to low values. Supporting these predictions, in a large cohort of HLA-type individuals, experiments showed that the relative ability of HLA-B MHC types to control HIV infection correlates with their peptidebinding characteristics that affect thymic development [Figure 6; (33)]. Furthermore, there is also evidence that the immune response in individuals with the HLA-B27 gene who control HIV exhibits a greater proportion of cross-reactive T cells than that of HLA-B27 positive individuals who do not control HIV (69). Even though we do not fully understand why individuals with the HLA-B27 gene exhibit different proportions of activated cross-reactive T cells upon HIV infection. its effects on the control of HIV support our conclusions. Undoubtedly, many complex factors influence the relationship between MHC type and disease outcome. The effect of the factor related to differential thymic selection should be greatest for MHC molecules that bind relatively few (for example, HLA-B57) or many (for example, HLA-B7, -B35, -B8) self-peptides.

Superior control of viral load due to the greater precursor frequency and cross-reactivity of Tcell repertoires restricted by MHC molecules that bind to few self-peptides (for example, HLA-B57) should also confer protection against diseases caused by other fast-mutating viruses. Indeed, HLA-B57 is protective against hepatitis C virus [HCV (70)], another highly mutable viral disease in which T cells are important. Also, HLA-B8, which binds a greater diversity of self-peptides (33), is associated with faster disease progression in HCV (71) and HIV (66). Thus, the correlation between the diversity of peptides presented in the thymus during T-cell development and control or progression of disease may be general.



Figure 6

HLA-B alleles associated with greater ability to control HIV correlate with smaller self-peptide binding propensities. A large group of HIV-infected people were divided into a controller cohort (low levels of HIV RNA) and a progressor cohort (high levels of HIV RNA). The odds ratio is defined as $\frac{p_w/p_{wo}}{c_w/c_{wo}}$, where p_w and p_{wo} (c_w and c_{wo}) are the numbers of individuals in the progressor cohort (the controller cohort) with and without this HLA, respectively. People with HLA alleles associated with an odds ratio value greater or less than one are more likely to be progressors or controllers, respectively. The fraction of peptides derived from the human proteome that bind to a given HLA allele was determined with predictive bioinformatics algorithms (33). The error bars represent the 95% confidence intervals for odds ratios. The gray dashed line corresponds to equal odds for an allele being associated with progressors and controllers. Included are only those HLA-B alleles that were statistically significantly associated with HIV control or progression. This figure is adapted from Reference 33, figure 3.

The results we summarize above also point to a mechanistic explanation for the previously unexplained associations between HLA alleles that confer protection against HIV and autoimmune diseases. T cells restricted by MHC types that bind to few self-peptides are subject to less stringent negative selection in the thymus and should therefore be more prone to recognizing selfpeptides. This may explain the enhanced proclivity for autoimmune disorders in people with MHC genes that are also associated with superior control of HIV infections.

5. FUTURE PROSPECTS

In this review we address how simple statistical mechanical models can be used to shed light on certain aspects of the immune response. However, due to the highly complex characteristics of the adaptive immune system, many basic questions remain unresolved. The richness and intricacy of the problem invites a multitude of approaches from the physical and life sciences to uncover new principles. Below we discuss some additional questions pertinent to development and actions of the T-cell repertoire, where models similar to the ones presented here could lead to new insights.

Thymic selection attempts to remove dangerous T cells that could cause autoimmune disease. But thymic selection is not perfect and some autoreactive T cells may escape, possibly due to the fact that not all self-peptide types are expressed in the thymus and that immature T cells spend a finite time in the thymus. The immune system thus has other protective mechanisms, e.g., certain regulatory T cells suppress immune responses directed against self-tissues. Are T cells that do not encounter certain self-peptides in the thymus just as reactive to these peptides as those derived from pathogens? If so, how is autoimmunity suppressed but not reactivity to pathogens? Most of the time, the immune system is very efficient at preventing autoimmune diseases, but what leads to a higher frequency of escape of autoreactive T cells that target cells of the nervous system and the pancreas in the case of autoimmune diseases such as multiple sclerosis and type I diabetes? Why do the escaping autoreactive T cells attack only particular tissues? By adapting the thymic selection model to include the variability in expression levels of different types of self-peptides in the thymus, one could potentially get insights into the last two questions. The escape probability of autoreactive T cells from the thymus can also be studied as diffusion in a random field of immobile traps (72).

Autoimmune diseases are correlated to a combination of genetic and environmental factors. People who express certain genes have a higher propensity for certain autoimmune diseases, but not everyone with these genes develops disease; e.g., most people with the inflammatory disease ankylosing spondylitis express HLA-B27 (a type of MHC), but most people expressing HLA-B27 do not develop ankylosing spondylitis. Certain genes and viral infections are known to increase the risk of triggering multiple sclerosis. Insights into these puzzles could emerge from stochastic dynamical models of host-pathogen interactions coupled with models of T-cell development.

DISCLOSURE STATEMENT

The authors are not aware of any affiliations, memberships, funding, or financial holdings that might be perceived as affecting the objectivity of this review.

ACKNOWLEDGMENTS

Our work in this area has greatly benefited from fruitful discussions and collaborations with Eric S. Huseby, Eugene I. Shakhnovich, Bruce D. Walker, Herman N. Eisen, and Elizabeth L. Read. This work was supported by the Ragon Institute (A.K.C. and A.K.), National Institutes of Health (NIH) Grant No. 1-PO1-AI071195-01 (A.K.C. and M.K.), NSF Grant No. DMR-12-06323 (M.K.), and an NIH Directors Pioneer award (to A.K.C.).

A. APPENDIX: BASIC IMMUNOLOGY⁵

Higher organisms are constantly exposed to infectious microbial pathogens, yet rarely develop disease. This is because the variety of cells that comprise the innate immune system are efficient in controlling pathogenic microorganisms. The components of the innate immune system respond to common features of diverse microorganisms but are not specific for individual pathogens. Some bacteria and many viruses can evade or overcome the innate mechanisms of host defense. The adaptive immune system mounts pathogen-specific responses against such invading microorganisms. Adaptive immunity also establishes memory of past infections, thereby conferring the ability to mount rapid immune responses to pathogens encountered previously. This immunological memory is the basis for vaccination.

inherent, nonspecific form of immunity to a specific pathogen

⁵Some parts of this Appendix are reprinted with kind permission from Reference 11, the Annual Review of Physical Chemistry, Volume 61 © 2010, by Annual Reviews, http://www.annualreviews.org.

Cellular immunity:

adaptive immunity in which the role of T lymphocytes is predominant

Humoral immunity:

form of immunity mediated by circulating antibodies

Antigen: foreign substance that induces an immune response in the body

Antigen-presenting cells (APCs): express molecular signatures (peptides) of ingested antigens on their surface

A.1. The Two Arms of the Adaptive Immune System

The adaptive immune system has two arms, called cellular immunity and humoral immunity. T lymphocytes (T cells) and B lymphocytes (B cells) are the key regulators of cellular and humoral immunity, respectively. T cells and B cells express immunoglobulin proteins on their surfaces, which are called T-cell receptors (TCRs) and B-cell receptors (BCRs), respectively. The genes encoding these receptors are inherited as gene segments that stochastically recombine during the synthesis of T cells and B cells in the bone marrow. Each gene assembled in a given lymphocyte is thus likely to be distinct, enabling the generation of a great diversity of T cells and B cells expressing different receptors. Different lymphocytes can potentially respond to specific pathogens as distinct receptors can potentially recognize (i.e., bind sufficiently strongly to) molecular signatures of specific foreign invaders. Although the adaptive immune system can mount pathogen-specific responses against varied microbes, the number of possible pathogens suggests that one T cell (or B cell) for every possible pathogen is an unlikely scenario.

The diverse lymphocytes bearing different TCRs and BCRs generated in the bone marrow do not all become part of an organism's army of T cells and B cells. Rather, T cells and B cells undergo development processes that allow only a small fraction of the generated cells to become part of an organism's repertoire of lymphocytes. T cells develop in the thymus (the T stands for thymus); B cells develop in the bone marrow (the B stands for bone marrow) and also, upon activation, in lymphoid organs.

A.2. The T-Cell Recognition Process

Cells of the innate immune system (e.g., dendritic cells, macrophages) engulf pathogens (also called antigens) present in different parts of an organism's body. These cells are called antigenpresenting cells (APCs) because they express molecular signatures of the ingested antigens on their surface. Extracellular fluid from tissues, which contains pathogens or APCs harboring pathogens, drains into lymphoid organs (e.g., lymph nodes, spleen) via the lymphatic vessels. In lymphoid organs, lymphocytes can interact with pathogen-bearing APCs and pathogens and recognize them as foreign.

If a lymphocyte recognizes pathogens in a lymph node, a series of intracellular biochemical reactions occurs (called signaling) that results in gene transcription programs that cause the lymphocyte to become activated; i.e., it begins to proliferate and acquire the ability to carry out functions that can mediate an immune response. Activated lymphocytes thus generated, bearing receptors specific for the infecting pathogen, then leave the lymph node and enter the blood via lymphatic vessels. The lymphatic vessels enable lymphocytes to circulate among the blood, lymphoid organs, and tissues. When activated lymphocytes encounter the same pathogen's molecular markers in the blood or tissues (see below), they can carry out effector functions to eliminate the infection.

The BCRs and TCRs expressed on B cells and T cells can bind to molecular markers called ligands. B cells protect against pathogens in blood or extracellular spaces. The ligands of the BCR include proteins, fragments of proteins, and molecules on the surface of viruses or bacteria. T cells evolved to combat intracellular pathogens. Proteins synthesized by intracellular pathogens are cut up into short peptide fragments [typically 8–15 amino acids long (1)] by enzymes in cells harboring the pathogen. These peptide fragments may bind to the host's MHC proteins. There are two kinds of MHC proteins, called class I and class II. Typically, a human will have up to six types of MHC class I proteins and up to six types of MHC class II proteins. Pathogen-derived peptides bound to

MHC proteins are ultimately expressed on the surface of APCs (encountered by T cells in lymph nodes prior to activation) and infected cells (encountered by T cells patrolling blood and tissues) (see Figure 1*a*).

T-cell recognition of a particular pathogen-derived pMHC implies that its TCR binds to it sufficiently strongly, leading to productive intracellular signaling and activation. The T-cell signaling network does not respond progressively to increasing the stimulus (e.g., TCR-pMHC binding strength); rather, it only responds strongly above a threshold stimulus level (27, 41). In the lymph nodes, T cells activated by peptides presented by MHC class II proteins proliferate and differentiate into many cell types called T helper cells, as they help activate B cells and perform other important functions. T cells activated by peptides presented by MHC class I molecules are called cytotoxic T lymphocytes (CTLs). When activated CTLs encounter cells in tissues that express the pMHC molecules that originally activated them, they can kill these cells by secreting various chemicals.

LITERATURE CITED

- 1. Janeway C, Travers P, Walport M, Shlomchik M. 2004. Immunobiology. New York: Garland Sci. 6th ed.
- 2. Allison JP, McIntyre BW, Bloch D. 1982. J. Immunol. 129:2293-300
- 3. Chan AC, Iwashima M, Turck CW, Weiss A. 1992. Cell 71:649-62
- 4. Dialynas DP, Wilde DB, Marrack P, Pierres A, Wall KA, et al. 1983. Immunol. Rev. 74:29-56
- 5. Gallegos AM, Bevan MJ. 2006. Immunol. Rev. 209:290-96
- 6. Hedrick SM, Cohen DI, Nielsen EA, Davis MM. 1984. Nature 308:149-53
- 7. Hogquist KA, Baldwin TA, Jameson SC. 2005. Nat. Rev. Immunol. 5:772-82
- 8. Irving AB, Weiss A. 1991. Cell 64:891-901
- 9. Unanue ER. 1984. Annu. Rev. Immunol. 2:395-428
- 10. Yanagi Y, Yoshikai Y, Leggett K, Clark SP, Aleksander I, Mak TW. 1984. Nature 308:145-49
- 11. Chakraborty AK, Košmrlj A. 2010. Annu. Rev. Phys. Chem. 61:283-303
- 12. Jameson SC, Hogquist KA, Bevan MJ. 1995. Annu. Rev. Immunol. 13:93-126
- 13. Siggs OM, Makaroff LE, Liston A. 2006. Curr. Opin. Immunol. 18:175-83
- 14. Starr TK, Jameson SC, Hogquist KA. 2003. Annu. Rev. Immunol. 21:139-76
- 15. von Boehmer H, Aifantis I, Gounari F, Azogui O, Haughn L, et al. 2003. Immunol. Rev. 191:62-78
- 16. Werlen G, Hausmann B, Naeher D, Palmer E. 2003. Science 299:1859-63
- 17. Košmrlj A, Kardar M, Chakraborty AK. 2012. J. Stat. Phys. 149:203-19
- 18. Huseby ES, White J, Crawford F, Vass T, Becker D, et al. 2005. Cell 122:247-60
- 19. Huseby ES, Crawford F, White J, Marrack P, Kappler JW. 2006. Nat. Immunol. 7:1191-99
- 20. Eisen HN, Chakraborty AK. 2010. Proc. Natl. Acad. Sci. USA 107:22373-80
- 21. Hemmer B, Vergelli M, Gran B, Ling N, Conlon P, et al. 1998. J. Immunol. 160:3631-36
- 22. Kersh GJ, Allen PM. 1996. Nature 380:495-98
- Misko IS, Cross SM, Khanna R, Elliott SL, Schmidt C, et al. 1999. Proc. Natl. Acad. Sci. USA 96: 2279–84
- 24. Sloan-Lancaster J, Allen PM. 1996. Annu. Rev. Immunol. 14:1-27
- 25. Borghans JAM, Noest AJ, de Boer RJ. 2003. Eur. J. Immunol. 33:3353-58
- 26. Bousso P, Bhakta NR, Lewis RS, Robey E. 2002. Science 296:1876-80
- 27. Daniels MA, Teixeiro E, Gill J, Hausmann B, Roubaty D, et al. 2006. Nature 444:724-29
- 28. Detours V, Mehr R, Perelson AS. 1999. J. Theor. Biol. 200:389-403
- 29. Detours V, Perelson AS. 1999. Proc. Natl. Acad. Sci. USA 96:5153-58
- 30. Scherer A, Noest A, de Boer RJ. 2004. Proc. Biol. Sci. 271:609-16
- Košmrlj A, Jha AK, Huseby ES, Kardar M, Chakraborty AK. 2008. Proc. Natl. Acad. Sci. USA 105: 16671–76
- 32. Košmrlj A, Chakraborty AK, Kardar M, Shakhnovich EI. 2009. Phys. Rev. Lett. 103:068103

- 33. Košmrlj A, Read EL, Qi Y, Allen TM, Altfeld M, et al. 2010. Nature 465:350-54
- 34. Miyazawa S, Jernigan RL. 1996. J. Mol. Biol. 256:623-44
- 35. Chao DL, Davenport MP, Forrest S, Perelson AS. 2005. Eur. J. Immunol. 35:3452-59
- 36. Park JM, Deem MW. 2004. Physica A 341:455-70
- 37. Yang M, Park JM, Deem MW. 2006. Lect. Notes Phys. 704:541-62
- 38. Zhou H, Deem MW. 2006. Vaccine 24:2451-59
- 39. Deem MW, Hejazi P. 2010. Annu. Rev. Chem. Biomol. Eng. 1:247-76
- 40. Flicek P, Aken BL, Beal K, Ballester B, Caccamo M, et al. 2008. Nucleic Acids Res. 36:D707-14
- Prasad A, Zikherman J, Das J, Roose JP, Weiss A, Chakraborty AK. 2009. Proc. Natl. Acad. Sci. USA 106: 528–33
- 42. Zeldovich KB, Berezovsky IN, Shakhnovich EI. 2007. PLoS Comput. Biol. 3:62-72
- 43. Moszer I, Glaser P, Danchin A. 1995. Microbiol. UK 141:261-68
- 44. Naeher D, Daniels MA, Hausmann B, Guillaume P, Luescher I, Palmer E. 2007. J. Exp. Med. 204:2553-59
- 45. Fischer E. 1894. Ber. Dtsch. Chem. Ges. 27:2985-93
- 46. Garboczi DN, Ghosh P, Utz U, Fan QR, Biddison WE, Wiley DC. 1996. Nature 384:134-41
- 47. Garcia KC, Degano M, Staneld RL, Brunmark A, Jackson MR, et al. 1996. Science 274:209-19
- 48. Wu LC, Tuot DS, Lyons DS, Garcia KC, Davis MM. 2002. Nature 418:552-56
- Leadbetter MR, Lindgren G, Rootzen H. 1983. Extremes and Related Properties of Random Sequences and Processes. New York/Heidelberg/Berlin: Springer-Verlag
- 50. Kardar M. 1983. Phys. Rev. Lett. 51:523-26
- 51. Deeks SG, Walker BD. 2007. Immunity 27:406-16
- Migueles SA, Sabbaghian MS, Shupert WL, Bettinotti MP, Marincola FM, et al. 2000. Proc. Natl. Acad. Sci. USA 97:2709–14
- 53. Miura T, Brumme CJ, Brockman MA, Brumme ZL, Pereyra F, et al. 2009. J. Virol. 83:3407-19
- 54. Jin X, Bauer DE, Tuttleton SE, Lewin S, Gettie A, et al. 1999. J. Exp. Med. 189:991-98
- 55. Schmitz JE, Kuroda MJ, Santra S, Sasseville VG, Simon MA, et al. 1999. Science 283:857-60
- 56. Borrow P, Lewicki H, Hahn BH, Shaw GM, Oldstone MB. 1994. J. Virol. 68:6103-10
- 57. Koup RA, Safrit JT, Cao Y, Andrews CA, McLeod G, et al. 1994. J. Virol. 68:4650-55
- 58. Allen TM, Altfeld M, Geer SC, Kalife ET, Moore C, et al. 2005. J. Virol. 79:13239-49
- 59. Allen TM, Yu XG, Kalife ET, Reyor LL, Lichterfeld M, et al. 2005. J. Virol. 79:12952-60
- Dahirel V, Shekhar K, Pereyra F, Miura T, Artyomov M, et al. 2011. Proc. Natl. Acad. Sci. USA 108: 11530–35
- 61. Bhalerao J, Bowcock AM. 1998. Hum. Mol. Genet. 7:1537-45
- 62. Chessman D, Kostenko L, Lethborg T, Purcell AW, Williamson NA, et al. 2008. Immunity 28:822-32
- 63. Bowness P. 2002. Rheumatology 41:857-68
- 64. Peters B, Sidney J, Bourne P, Bui HH, Buus S, et al. 2005. PLoS Biol. 3:e91
- 65. Hubbard TJ, Aken BL, Ayling S, Ballester B, Beal K, et al. 2009. Nucleic Acids Res. 37:D690-97
- 66. Turnbull EL, Lopes AR, Jones NA, Cornforth D, Newton P, et al. 2006. J. Immunol. 176:6130-46
- 67. Gillespie GM, Kaul R, Dong T, Yang HB, Rostron T, et al. 2002. AIDS 16:961-72
- 68. Yu XG, Lichterfeld M, Chetty S, Williams KL, Mui SK, et al. 2007. J. Virol. 81:1619-31
- 69. Chen H, Ndhlovu ZM, Liu D, Porter LC, Fang JW, et al. 2012. Nat. Immunol. 13:691-700
- 70. Thio CL, Gao X, Goedert JJ, Vlahov D, Nelson KE, et al. 2002. J. Virol. 76:4792-97
- 71. McKiernan SM, Hagan R, Curry M, McDonald GS, Kelly A, et al. 2004. Hepatology 40:108-14
- 72. Košmrlj A. 2011. J. Stat. Phys. 142:1277-86

Annual Review of Condensed Matter Physics

Volume 4, 2013

Contents

Why I Haven't Retired Theodore H. Geballe 1
Quantum Control over Single Spins in Diamond V.V. Dobrovitski, G.D. Fuchs, A.L. Falk, C. Santori, and D.D. Awschalom
Prospects for Spin-Based Quantum Computing in Quantum Dots Christoph Kloeffel and Daniel Loss
Quantum Interfaces Between Atomic and Solid-State SystemsNikos Daniilidis and Hartmut Häffner83
Search for Majorana Fermions in Superconductors C.W.J. Beenakker
Strong Correlations from Hund's Coupling Antoine Georges, Luca de' Medici, and Jernej Mravlje 137
Bridging Lattice-Scale Physics and Continuum Field Theory withQuantum Monte Carlo Simulations<i>Ribhu K. Kaul, Roger G. Melko, and Anders W. Sandvik</i>
Colloidal Particles: Crystals, Glasses, and Gels Peter J. Lu (陸 述 義) and David A. Weitz 217
Fluctuations, Linear Response, and Currents in Out-of-Equilibrium Systems S. Ciliberto, R. Gomez-Solano, and A. Petrosyan
Glass Transition Thermodynamics and Kinetics Frank H. Stillinger and Pablo G. Debenedetti
Statistical Mechanics of Modularity and Horizontal Gene Transfer Michael W. Deem

Physics of Cardiac Arrhythmogenesis	
Alain Karma	13
Statistical Physics of T-Cell Development and Pathogen Specificity	
Andrej Košmrlj, Mehran Kardar, and Arup K. Chakraborty33	39

Errata

An online log of corrections to *Annual Review of Condensed Matter Physics* articles may be found at http://conmatphys.annualreviews.org/errata.shtml