The ABC of Major Histocompatibility Complexes and T Cell Receptors in Health and Disease

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Abstract

A seminal discovery of major histocompatibility complex (MHC) restriction in T cell recognition by Peter Doherty and Rolf Zinkernagel has led to 45 years of exciting research on the mechanisms governing peptide MHC (pMHC) recognition by T cell receptors (TCRs) and their importance in health and disease. T cells provide a significant level of protection against viral, bacterial, and parasitic infections, as well as tumors, hence, the generation of protective T cell responses is a primary goal for cell-mediated vaccines and immunotherapies. Understanding the mechanisms underlying generation of optimal high-avidity effector T cell responses, memory development, maintenance, and recall is of major importance for the rational design of preventative and therapeutic vaccines/immunotherapies. In this review, we summarize the lessons learned over the last four decades and outline our current understanding of the basis and consequences of pMHC/TCR interactions on T cell development and function, and TCR diversity and composition, driving better clinical outcomes and prevention of viral escape. We also discuss the current models of T cell memory formation and determinants of immunodominant T cell responses in animal models and humans. As TCR composition and diversity can affect both the protective capacity of T cells and protection against viral escape, defining the spectrum of TCR selection has implications for improving the functional efficacy of effector T cell responsiveness and memory formation.

Keywords: MHC, TCR, disease

Introduction

Adaptive immunity forms the basis of successful long-lasting protective immune responses, with humoral immunity being mediated by antibodies produced by B cells, while T cells drive cellular immunity (7). Cytotoxic CD8+ T cells can recognize virus-infected or cancerous cells, induce cell lysis, and produce antiviral cytokines. Helper CD4+ T cells predominantly provide critical help to B cells and CD8+ T cells (16). What distinguishes T cells and B cells from innate immunity is the antigen specificity and generation of long-lasting immunological memory. While antibodies bind to three-dimensional epitopes on proteins, T cell receptors (TCRs) to epitopes formed by small foreign peptide fragments presented by a self-encoded glycoprotein called major histocompatibility complexes (MHCs) (16).

The requirement of the TCR to engage with both peptide and MHC concurrently is called MHC restriction and was first described by Zinkernagel and Doherty in 1974 (147). Their demonstration that T cells require interactions with both self-encoded antigen-presenting molecules and foreign peptides has been instrumental for our understanding of effective T cell immunity against pathogens and cancers, as well as the mechanisms underlying successful transplantation and development of autoimmune diseases. This key discovery was recognized by the Nobel Prize of Medicine and Physiology in 1996.

In this review, we summarize the lessons learned over the last 45 years since the seminal discovery of MHC-restriction by Doherty and Zinkernagel, and outline our current understanding of the basis and consequences of peptide MHC (pMHC)/TCR interactions on T cell development and function.
**T Cell Development and Priming**

T cells originate from bone marrow-derived hematopoietic progenitors, then traffic to the thymus, where double-negative immature CD4+CD8- T cells undergo TCR gene rearrangement, and after successfully doing so, become double-positive (DP) CD4+CD8+ immature T cells. As DP thymocytes undergo positive and negative selection, they commit to the CD4+ or CD8+ T cell lineage (67). The processes of thymic selection generate a diverse TCR repertoire of naive CD4+ or CD8+ restricted by the individual's MHC allotypes, but do not react to self-peptides. These mature naive T cells then circulate between secondary lymphoid organs (SLOs: spleen and lymph nodes), surveying for foreign antigens.

Encounter with a cognate antigen on professional antigen-presenting cells (APCs) in lymph nodes leads to T cell priming, a process requiring three key signals. The first is provided by the TCR-pMHC interaction and activation of the signaling pathways downstream of the TCR complex (40). The second signal is mediated by co-stimulatory molecules like B7 (CD80/CD86) on APCs and their interaction with their respective ligands, like CD28, on T cells (142). The third signal relies on cytokines like interferons (IFNs) or interleukin (IL)-12 released predominantly by APCs (30,34). The events of T cell priming result in epigenetic and transcriptional changes that underlie the well-orchestrated pathways of T cell proliferation and differentiation (51,60), subsequently leading to generation of large pools of antigen-specific effector and memory T cell subsets.

**T Cell Memory Differentiation**

Following antigen clearance, the effector T cell pool contracts by 90–95%, with the remaining 5–10% of antigen-specific T cells forming a population of long-lasting memory T cells capable of responding more rapidly and with greater numbers to antigenic re-encounter. A hallmark of memory T cells is their longevity, with human smallpox-specific CD8+ T cells being detectable even at 75 years postvaccination (48). Elegant human studies by Ahmed's group using in vivo deuterium labeling following vaccination showed that Yellow Fever Virus-specific CD8+ T cells, generated within the first 2 weeks following vaccination, were detectable for as long as 750 days later, estimated to divide once every 485 days (1). Similarly, influenza-specific CD8+ T cells can be detected up to 13 years after an individual's last recorded natural influenza A virus (IAV) infection in humans (133) or for a life-span of a laboratory mouse (65,130). The maintenance of memory CD8+ T cells depends on survival signals provided by cytokines like IL-15 and IL-7, but not by antigen [reviewed by Raebel et al. (99)].

Antigen-specific memory CD8+ T cells are greatly heterogeneous, with four main memory subsets being identified, namely T cell stem cell memory (TSCM, conventionally defined as CD45RA+CD27-CCR7-CD62L+IL-7Rz+/CD95-), T cell central memory (TCM, conventionally defined as CD45RA-CCR7+CD62L-CD95-), T cell effector memory (TEM, conventionally defined as CD45RA-CCR7-CD62L+CD95-), and tissue-resident memory (TRM, conventionally defined as CD45RA-CD69-CD103+/+ present in tissues) (Fig. 1A). These subsets differ with regard to their proliferation, cytokine production, and tissue circulation potential [reviewed in refs. (37,55)]. For instance, TSCM and TCM CD8+ T cells have higher proliferative potential, produce high levels of IL-2, and recirculate between the blood and SLOs. Conversely, effector memory (TEM) CD8+ T cells have a limited proliferation potential, produce mainly IFNγ, and recirculate between blood and peripheral tissue. The fourth subset of memory, TRM, is retained within peripheral tissues where they can provide rapid and potent protection after antigen re-encounter [reviewed by Gebhardt et al. (41)].

Different models exist on how CD8+ T cell memory is formed in relationship to the effector subset (Fig. 1B). The linear model (also called “decreasing potential” model) proposes that T cells progressively go through the memory and effector phases (SCM → CM → EM → EFF) in a process that decreases memory potential and increases effector differentiation (Fig. 1B[1]) (23,51). According to this model, the progression through the different stages is influenced by TCR signal strength/duration and/or the extent of antigenic stimulation on the T cell. The opposing model is called the circular model (or the “on-off-on” model) (Fig. 1B[2]) (60). This model proposed that following antigen encounter, CD8+ T cells differentiate into effector cells and, upon contraction of the response, the same effector cells de-differentiate into memory T cells of different subsets, which then can be recalled and re-differentiated into effector cells after re-encountering the same antigen. Some studies propose an alternative model, whereby the fate of a naive CD8+ T cell is determined as early as the first cell division, with the asymmetric distribution of key transcriptional and/or epigenetic regulators between two daughter cells, whereby one displays increased memory potential, while the other has a greater effector capacity (Fig. 1B[3]) (23,60,61).

These different models are supported, and refuted, by different lines of evidence in various models of infection (23,51,60). Thus, the formation of immunological T cell memory is incredibly complex, with knowledge gaps remaining to be addressed. For instance, although molecular cues and signaling pathways that drive TEM formation have been characterized, the exact origins of how TCM CD8+ T cells are developed are far from clear, that is, whether they are derived from effector cells or memory cells, or both. With the advent of new high-throughput single-cell technologies, novel insights are to be gained into the developmental pathways of memory CD8+ T cells. Given that life-long immune memory is the foundation of effective vaccination, such understanding of memory establishment and recall at different anatomical sites is of utmost importance.

**CD8+ T Cell Effector Function**

Differentiated effector CD8+ T cells traffic to the site of infection, where they recognize their cognate pMHC on target cells by their TCR and subsequently engage in their effector function. Preloaded cytotoxic granules are released following TCR engagement and deliver their contents in the immunological synapse between the T cells and the target cell (Fig. 2A). The granules predominantly release granzymes (A, B, H, K, and M in humans), perforin and granulysin. Perforin forms a pore in the membranes of the target cells for granzyme delivery to the cytosol of infected or cancerous cells (17,138). Granzymes, which are serine proteases, then induce cell-mediated apoptosis of the virus-
The most studied granzyme B causes a rapid induction of caspase-dependent apoptosis through a number of mechanisms, including the cleavage of caspases or anti-apoptotic and proapoptotic factors (17). Granzymes A and K induce caspase-independent apoptosis through mitochondrial pathways, although the molecular details are yet to be fully elucidated (17,138). Granzyme M-induced cell death although ill defined, is caspase and mitochondria independent (17,138). Additional mechanisms of apoptosis induction occur through the engagement of the death receptor Fas or TRAILR on target cells with the Fas ligand or TRAIL, respectively, from/on cytotoxic CD8+ T cells (Fig. 2A). Signaling through these death receptors results in the activation of caspases and subsequent apoptosis. Regardless of
the exact molecular mechanism, the effects of different granzymes and death receptor signaling converge to the aim at hand: death of the target cell. The importance of these pathways can be demonstrated in the context of IAV infection in mice. Following IAV infection, CD8\(^+\) T cells acquire a cytotoxic profile that increases with time and cell division\((57)\), and is maintained up to at least 1 year after infection\((56)\). Importantly, following IAV infection, mice lacking TRAIL\((21)\), Fas, or perforin\((123)\) in CD8\(^+\) T cells show increased morbidity and delayed viral clearance. Interestingly, mice lacking granzymes A and B show normal viral clearance, which may imply a redundancy in the mechanisms of cytotoxicity\((58)\).

TCR engagement also results in the release of antiviral cytokines like IFN\(_c\) and TNF (Fig. 2B). IFN\(_c\) is a pleiotropic cytokine that can enhance the recruitment and activation of immune cells like macrophages and NK cells\((110)\) and the cytolytic capacity of T cells\((11)\), and induce an antiviral state through the expression of antiviral IFN-stimulated genes\((12,107)\). These include genes important for T cell-mediated immunity, like MHC-I and the genes to form the immunoproteasome\((107)\). Proinflammatory cytokines like RANTES and MIP1\(_{\beta}\) are also released by effector CD8\(^+\) T cells, which attract innate immune cells as well as T cells, and overall may aid to viral clearance (Fig. 2B)\((116)\).

Overall, while the defining feature of CD8\(^+\) T cells is the lysis of the target cells, the response of the effector T cell is multifaceted and involves both the delivery of cytotoxic molecules as well as the release of antiviral/inflammatory cytokines and chemokines. Importantly, evidence suggests that polyfunctionality (producing multiple cytokines and cytotoxic molecules) as well as the antigen sensitivity of CD8\(^+\) T cells are important correlates of protection in infectious diseases\((112)\). In HIV-1 elite controllers, an association was found between highly polyfunctional CD8\(^+\) T cells\((5+\) functions) and lower viral load\((10)\). Polyfunctional CD8\(^+\) T cells produced more IFN\(_c\) and were superior effectors than monofunctional CD8\(^+\) T cells\((94,112)\). In

**FIG. 2.** CD8\(^+\) T cell effector function. (A) Cytotoxicity can be induced through a delivery of granzymes and perforin. Perforin forms pores in target cell membranes, allowing delivery of granzymes to the cytosol, where granzymes induce apoptosis through activation of caspases and the mitochondrial pathway. Binding of Fas and TRAIL on the CD8\(^+\) T cell to their receptors FasL and TRAILR on the target cells also induced caspase activation. (B) Recognition of pMHC-I by the TCR results in cytotoxicity of the target cell as well as the release of antiviral cytokine and chemokines, which act on target cells and other immune cells. pMHC, peptide major histocompatibility complex.
addition, HIV-specific human leukocyte antigen (HLA)-
B27-restricted CD8\(^{+}\) T cells, which correlate with disease
protection, are of higher polyfunctionality and functional
avidity (antigen sensitivity) (4). An association between
antigen sensitivity, polyfunctionality, and the ability of
CD8\(^{+}\) T cell to suppress HIV-1 has also been demonstrated
(5). Overall, the quality of the response is an important
factor in CD8\(^{+}\) T cell-mediated protection and needs to be
considered in the design of effective CD8\(^{+}\) T cell-based
vaccines.

**CD8\(^{+}\) T Cell Immunodominance**

While the quality of the CD8\(^{+}\) T cell response can be a
strong correlate of protection (4,5,82,112), the relative
magnitude of different T cell specificities within a response
can still be important in determining immune protection.
From the thousands of potential peptides, which can be
generated from the processing of viral proteins during an
infection, the CD8\(^{+}\) T cell response is generally focused to a
relatively small subset of epitopes. This can be exemplified
by influenza B-derived peptides binding to HLA-A*02:01
(as detected by mass spectrometry), while only three of
them (BHA\(_{543–551}\), BNS\(_{1266–274}\), and BNS\(_{1264–274}\)) being
consistently immunogenic across donors (69). Within this
immunogenic subset, CD8\(^{+}\) T cells against different specific-
ities vary in magnitude and accordingly fall in a repro-
ducible immunodominance hierarchy (124).

A large number of factors dictate these hierarchies, includ-
ing, but not limited to, those relating to antigen presentation
(e.g., antigen abundance and processing, the affinity of peptide
for MHC-I, the stability of the pMHC complex, and the pre-
sentation pathway) and those relating to the responding T
cells [e.g., naïve precursor frequency, extent of their recruit-
ment and expansion, TCR affinity, and competition between
T cells of different specificities, termed immunodomination
(124)]. These factors contribute to immunodominance in a
complex and often context-specific manner. It is important to
note that subdominant epitopes of lower magnitude can
still be highly functional and provide protection against vi-
rnal disease in both humans (38,49) and mouse models
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**CD4\(^{+}\) T Cell Functions**

CD4\(^{+}\) T cells can exert different helper and effector
functions during an immune response [reviewed in refs.
(75,108,120)]. This can be broadly divided into the fol-
lowing: (i) recruitment of immune cells in SLOs or the site
of infection, (ii) provision of help for expansion or effector
function, or (iii) direct antiviral functions (Fig. 3B, C). For
example, in the draining lymph nodes, interactions between
CD4\(^{+}\) T cells and APCs through CD40-CD40L binding in-
duce signaling in the dendritic cell (DC) that results in DC
“licensing” or maturation. This results in the release of
chemokines like CCL3 and CCL4 by the licensed DC to
attract CD8\(^{+}\) T cells at the site of priming (26), and uppre-
gulation of molecules involved in CD8\(^{+}\) T cell priming, like
IL-12, by the licensed DCs (143). IL-12 can, in turn, in-
crease expression of the IL-2R on CD8\(^{+}\) T cells (143).
CD4\(^{+}\) T cells may also provide a source of IL-2 important
for the expansion of CD8\(^{+}\) T cells (Fig. 3B) (143,145). CD4\(^{+}\)
T cells are also found in the infected tissues where they can
display numerous effector roles (Fig. 3C). For instance, they
can act on epithelial cells through the release of antiviral
cytokines like IFN\(_{\gamma}\) to induce an antiviral state (75,120).
In addition, CD4\(^{+}\) T cells can promote conversion of CD8\(^{+}\) T
cells to T\(_{RM}\) by the release of transforming growth factor \(\beta\)
(75,76). It has also become evident that CD4\(^{+}\) T cells can
exert direct cytotoxic function against virally infected cells
mediated by perforin and FAS (20,22). It is important to
note that many of these mechanisms may not be universally
applicable, as studies with different viruses may provide
different results [discussed in refs. (75,120)]. Finally, CD4\(^{+}\)
T cell help is of course not limited to that provided to CD8\(^{+}\)T
cells. In fact, the most studied function of CD4\(^{+}\) T cells is
provision of help to B cells [recently review by Koutsakos
et al. (70)]. Overall, CD4\(^{+}\) T cells are a critical component
of both cellular and humoral immunity through both helper
and non-helper functions.

**Human Leukocyte Antigens: Genetics, Structure,
and Diversity**

The basis of both CD8\(^{+}\) and CD4\(^{+}\) T cell priming,
differentiation, and function is the cognate interaction be-
 tween the TCR and pMHC. As demonstrated by Zinkernagel
and Doherty in 1974 (147), this activation of T cells depends
on simultaneous recognition of both foreign peptide fragments
and self-MHC molecules, a phenomenon called MHC re-
striction. CD8\(^{+}\) T cells are restricted by MHC-I and CD4\(^{+}\) T
cells are restricted by MHC-II. MHC glycoproteins are di-
vided into three classes: MHC-I, MHC-II, and MHC-III.
The MHC-III class comprises of many proteins of the
complement system, cytokines, and heat shock proteins. In
humans, the MHC molecules are termed HLA and are
encoded on chromosome 6 of the human genome (Fig. 4A). HLA Class I (HLA-I) and HLA Class II (HLA-II), which correspond to MHC-I and MHC-II, respectively, are further divided into classical and nonclassical genes (102). Classical HLA-I genes include HLA-A, HLA-B, and HLA-C, while nonclassical alleles include HLA-E, HLA-F, and HLA-G. Classical HLA-II genes include HLA-DR, HLA-DP, and HLA-DQ, while nonclassical alleles include HLA-DO and HLA-DM (Fig. 4B) (102). Human CD8$^+$ T cells recognize peptides presented by classical HLA-A and HLA-B, and to a lesser extent HLA-C (13), although rare populations of HLA-E-restricted CD8$^+$ T cells have also been described (59). Human CD4$^+$ T cells recognize peptides presented by HLA-DR, HLA-DQ, and HLA-DP.

HLA molecules are surface membrane-bound glycoproteins. HLA-I comprises of a heavy chain made of three extracellular domains ($\alpha_1$, $\alpha_2$, and $\alpha_3$) noncovalently attached to a $\beta_2$ microglobulin molecule (Fig. 4C) (102). HLA-II is a heterodimer consisting of an $\alpha$ and a $\beta$ chain (Fig. 4D). The extracellular domains of HLA form an antigen-binding cleft, consisting of two $\alpha$-helices surrounding an antiparallel $\beta$-sheet floor. This generates a platform that can cradle a short amino acid (aa) fragment, termed peptide. These peptides are bound to the floor of the binding groove through the interactions of specific amino acids (known as anchor residues) with specific peptide-binding pockets. Due to the closed structure of the HLA-I binding cleft, small peptides of 8–10 aa are typically presented, although longer peptides of up to 14 aa can be accommodated (Fig. 4C). HLA-II presents longer peptides of >11 aa, with N- and C-terminal “overhang” regions (Fig. 4D) (102).

The pockets and the peptide binding groove of the MHC molecules are highly polymorphic. Variation in those peptide-anchoring residues of the HLA means that different HLA allotypes (the HLA protein encoded by an HLA allele)
will present a different spectrum of peptides. Despite this diversity, HLA alleles can be divided into superfamilies (or supertypes) that share peptide binding specificity, based on shared binding motifs within those peptides. For instance, members of the HLA-A2 superfamily bind to peptides with aliphatic hydrophobic residues (e.g., leucine and isoleucine) in position 2 and at the C-terminus (113). Furthermore, each individual may express up to two different HLA-A, HLA-B, and HLA-C alleles, for a total of six HLA-I alleles. The frequency of alleles and HLA superfamilies varies across ethnic groups. For example, the HLA-A*02:01 allele can be found between 1% and 54.5% of different populations around the world (121). It is worth noting that HLA allotypes can differ by >30 aa or by a handful of amino acids (44). However, even small amino acid changes can have a substantial impact on the peptide repertoire and its recognition by TCRs. This is best exemplified by the HLA-B*35:01 and HLA-B*35:08 allotypes, which only differ by one amino acid (position 156, leucine or arginine, respectively) in the α2 MHC-I helix. This single amino acid alters the conformation of the α2 MHC-I helix, in a manner that allows the prototypical SB27 TCR to recognize a 13-mer peptide from the EBV BZLF1 protein, in the context of HLA-B*35:08, but not HLA-B*35:01 (126). Thus, an

FIG. 4. MHC genes and structure. (A) The HLA locus encoded on Chr 6 of the human genome and is divided into Class I, Class II, and Class III genes. (B) Classification of HLA-I and HLA-II genes in classical and nonclassical groups. (C) Structure of MHC-I protein. A heavy chain made of three extracellular domains (α1, α2, and α3) is noncovalently attached to a β2-microglobulin molecule. The α1 and α2 domains form the peptide-binding cleft, where small peptides can bind. The peptides interact through two anchor residues with the binding pockets of the MHC-I molecule. (D) Structure of MHC-II protein. MHC-II is a heterodimer made of an α and β chain, each of which has two subdomains. The α1 and β1 subdomains form the peptide binding cleft, where peptides can bind. Peptides that bind MHC II are longer than MHC-I peptides and also have anchor residues that interact with binding pockets in the peptide-binding cleft of MHC-II. HLA, human leukocyte antigen.
appreciation of how MHC polymorphism may affect antigen presentation and recognition is crucial for our understanding of the basic principles that may underlie interindividual variation and aberrant T cell responses.

**Antigen Presentation Pathways**

MHC-I proteins present peptides that predominantly originate from within the cell. During homeostasis, cellular proteins are degraded by the proteasome and small peptides are generated. The proteasome is a multimeric protein complex that degrades proteins into small peptide fragments (15,101). During a viral infection, the effects of IFNs induce the formation of an alternative proteasome complex, termed the immunoproteasome, which enhances the generation of peptides for MHC-I presentation. Thus, as viral proteins are synthesized, they are targeted for degradation by the immunoproteasome, either as fully folded proteins or as defective ribosomal products (15,101). This results in the generation of small peptide fragments that originate from the infecting virus, which can be further trimmed by cytoplasmic aminopeptidases. These peptide fragments are then transported into the endoplasmic reticulum (ER) through a protein called transporter associated with antigen processing (TAP), which spans the ER membranes (15,101). After translocation by TAP, any peptide that may be too long for MHC-I binding can be further trimmed by an ER-resident aminopeptidase (ERAP1). In the ER, empty MHC-I molecules are associated with the peptide loading complex (PLC), which includes chaperone proteins like Tapasin and Calnexin (15,101). The PLC holds the empty MHC-I in a peptide-receptive conformation that promotes the binding of peptides as they are translocated by TAP, which is also part of the PLC. Peptide binding stabilizes the MHC-I protein so that it is released from the ER quality control chaperones and is transported through the Golgi apparatus to the cell surface (Fig. 5) (15,101). This allows CD8\(^+\) T cells to survey the intracellular proteome of a cell for signs of infection or malignancy. This pathway also operates in professional APCs, enabling the priming of CD8\(^+\) T cells, and requires infection of the APC. In specialized subsets of dendritic cells, however, alternative pathways exist (15,101). This process, called cross-presentation, involves the uptake of extracellular antigens, its retrotranslocation from phagosomes into the cytoplasm, and subsequent degradation by the proteasome and MHC-I loading in the ER, although other pathways of delivery to MHC exist (15,101).

A different pathway operates for the generation of MHC-II-presented peptides. These peptides originate from extracellular sources or from proteins degraded by the endocytic pathway (15,101). Proteins in the endolysosomal compartments can be cleaved by resident proteases into peptide fragments, which can then be delivered to a late endosomal compartment called the MIIC (MHC-II compartment). As MHC-II is synthesized in the ER, it is associated with an invariant chain that promotes its trafficking to the MIIC. The proteases of the MIIC cleave the invariant chain to a small peptide fragment called the class II-associated invariant chain peptide, which can then be exchanged for high-affinity peptides with the assistance of a nonclassical MHC-II protein called HLA-DM in humans. The p/MHC-II complex will then traffic to the cell surface for CD4\(^+\) T cell recognition (Fig. 5) (15,101).

**TCRs: Structure, Recombination, and Selection**

Recognition of pMHC complexes by \(\alpha\beta\) T cells is mediated by the TCR. The TCR consists of an \(\alpha\) and a \(\beta\) chain (TRA and TRB). Each chain consists of several gene segments: TRA variable (TRAV), TRA joining (TRAJ), and TRA constant (TRAC) for the \(\alpha\) chain (human chromosome 14), and TRBV, TRB diverse (TRBD), TRBJ, and TRBC for the \(\beta\) chain (human chromosome 7), which join during somatic recombination in the thymus. Each chain contains three variable loops called complementarity-determining regions (CDRs). While CDR1 and CDR2 are germline encoded in the V gene segments, the CDR3 loops are generated from the junction of V(D)J segments and thus display a greater level of diversity (83,102) (Fig. 6A).

Diversity in TCR sequences can arise from the following: (i) the permutations of different V(D)J segments for the \(\alpha\) (47 TRAV and 61 TRAJ genes) and \(\beta\) (54 TRBV and 14 TRBJ and 2 TRBD genes) chains, (ii) the imprecise joining of gene segments during recombination by recombination activation genes, (iii) the random addition of nontemplate nucleotides at the junction sites by terminal deoxynucleotidyl, and (iv) the combination of \(\alpha\) and \(\beta\) chains. This collectively results in a theoretical repertoire of \(10^{15}–10^{20}\) potential \(\alpha\beta\) TCRs that can be generated (35,83). However, after thymic selection, \(\sim 2 \times 10^7\) distinct TCRs are maintained per individual in humans (6). An additional source of diversity can be provided by polymorphisms in TCR genes, which can have significant effects in the TCR repertoire. For instance, TCRs with the TRBV9*01 allele, but not the TRBV9*02 allele, recognize the HLA-B*35:01-restricted EBV peptide (HPVG), due to a single amino acid change between the two alleles (Gln55His). While the Gln55 in TRBV9*01 easily binds the P7-Asp of the peptide, the electropositive His of the TRBV9*02 allele cannot accommodate the P7-Asp (46).

The process of thymic selection considerably reduces the available TCRs/\(\alpha\beta\) repertoire by many orders of magnitude. The positive and negative selection of immature thymocytes are based on the strength of their interaction with self-pMHC expressed in the thymic medulla (67). A weak interaction rescues developing thymocytes from death by neglect and promotes their development into either CD4\(^+\) or CD8\(^+\) T cells. Conversely, a strong interaction for self-pMHC results in clonal deletion and thus removal of autoreactive T cells (67).

Overall, through the vast diversity enabled by TCR genetics, the immune system can recognize the extremely large number of potential peptide antigens, which can be derived from the plethora of host and viral proteins. Concomitantly, the intricate processes of thymic selection ensure that self-derived antigens are not targeted by T cells during an immune response.

**TCR Recognition of pMHC: Geometry and Restriction**

The six CDR loops of the TCR are critical for recognition of the pMHC complex. Overall, the TCR docks over the pMHC complex with a conserved geometry. The TCR binds
in a diagonal manner on top of the peptide binding groove with an angle of \(\sim 45^\circ\), although this can range from \(20^\circ\) to \(70^\circ\) (102). Two notable exceptions described in the last few years represent two TCRs, which dock over their cognate pMHC with a reverse geometry (9,45). The conventional docking mode positions the TCR \(\alpha\) chain over the \(\alpha_2\) MHC-I helix and the TCR \(\beta\) over the \(\alpha_1\) helix. In addition, the CDR3\(\alpha\) is positioned over the N-terminus of the peptide and the CDR3\(\beta\) over the C-terminus (Fig. 6B). Consequently, the germline-encoded CDR1 and CDR2 loops interact with the MHC and the hypervariable CDR3 engages the peptide. However, it is pertinent to note that generally, the CDR1/2 loops can account for \(\sim 40\%\) of the buried surface area (a measure of the size of the interface between two interacting molecules) between the TCR and the peptide, and the CDR3 loops account for \(\sim 35\%\) of the contacts with the MHC-I (24).

The requirement of T cells to simultaneously bind the self-MHC as well as peptides is intriguing, and its functional and structural bases have been the focus of intense research [reviewed by La Gruta et al. (71)]. Briefly, two proposed theories explain what drives MHC restriction of TCRs. First, the germline-encoded theory proposes that TCRs are hard-wired, through evolutionary selection of germline-encoded motifs, to recognize MHC molecules. This model is mostly supported by studies identifying conserved TCR-MHC interactions, like the conserved Tyr48 in CDR2\(\beta\) (111) and the intrinsic MHC reactivity of many (\(\sim 30\%\)) TCRs before thymic selection (79,81,146). The second model, called selection theory, proposes that the ability of TCRs to recognize MHC is imposed by selection during T cell development in the thymus, through the requirement of CD4 or CD8 co-receptor binding for efficient signaling through the TCR. A third, unifying model proposes that (i) a predisposition of TCRs to bind MHC ensures that enough premature T cells in the thymus will develop into mature T cells in the periphery, despite the enormous diversity of potential MHC allotypes within a species, and (ii) the process of positive

**FIG. 5.** Antigen presentation pathways. Viral proteins produced during viral replication or antigens taken up by phagocytosis and retrotranslocated into the cytosol are degraded by the proteasome, releasing small peptide fragments. These are transported into the ER by TAP. Peptide length can be further optimized by an ER-resident aminopeptidase (ERAP1), before the peptide loading complex that promotes peptide binding to MHC-I. After peptide binding, MHC-I traffics to the cell surface through the Golgi. MHC-II in the ER is associated with an invariant chain that promotes its trafficking to a late endosomal compartment called MIIC. Proteins in endolysosomes can be broken into peptide fragments by resident proteases and delivered to the MIIC. There proteases cleave the invariant chain to a small peptide fragment (CLIP), which is exchanged for high-affinity peptides. pMHC-II complexes then traffic to the cell surface. ER, endoplasmic reticulum; TAP, transporter associated with antigen processing.
selection then endows the mature T cell pool with reactivity toward the specific MHC allotypes expressed within the individual. As more structural information becomes available and with the advent of system immunology that enables high-throughput analysis of TCR repertoires, a deeper understanding of MHC restriction is to be gained.

**FIG. 6.** TCR genetics, structure, and MHC-I recognition. (A) TCR comprises an α and a β chain, generated through the recombination of V(D)J genes. Each chain contains three CDRs. CDR1 and 2 are germline encoded, but V-segment-specific, while CDR3 is hypervariable as a result of combinatorial and junctional diversity. (B) The TCR docks over the pMHC in a conserved mode that positions the TCRα chain over the MHC α2 domain and the N-terminus of the peptide, while the TCRβ sits over the MHC α1 domain and the C-terminus of the peptide. CDR, complementarity-determining region.

**TCR Repertoire Diversity and Sharing**
Although the theoretical/potential diversity of the human peripheral TCR repertoire is estimated to be around $10^{12}$, the actual TCR diversity of a naive individual is $\sim 2.5 \times 10^7$. This striking difference in numbers means that the likelihood of the same TCRαβ being found in two individuals would be
extremely low (83,90) and indeed inbred mice only share 20–25% of their naive TCR repertoire (19). Some TCR clonotypes may be shared across unrelated individuals. Yet, many T cell responses are characterized by one or a few TCR\(\alpha\beta\) clonotypes that are shared across large numbers of genetically unrelated individuals, which is extraordinarily specific, given the vast diversity and estimated number of potential TCRs generated between individuals (64,83,128). This raises the question of how the same TCRs can be found across individuals, given the numerical paradox described above.

Two key factors that may contribute to this phenomenon are TCR genetics and the structure of the pMHC-I/TCR interface (83). With regard to genetics, many public TCRs, that is, TCRs shared across different individuals, can be generated by convergent recombination (135–137), a process by which sequences converge to produce the same TCR amino acid sequence. Convergence may result from recombining TCR segments converging to the same nucleotide sequence or different nucleotide sequences converging to the same amino acid sequence due to codon redundancy. This, in combination with germline likeliness (low number of N-insertions), is thought to contribute to the generation of public TCRs by increasing the number of naive precursors of a specific TCR clonotype. This is supported by studies by Venturi et al. in the murine IAV model (137), Simian immunodeficiency virus (SIV) infection in macaques (136), and human Cytomegalovirus (CMV) and Epstein Barr virus (EBV) responses (135), revealing that the extent of TCR sharing correlated with a low number of N-insertions and a large number of nucleotide sequences encoding for the same amino acid sequence.

The structure underlying the TCR-pMHC-I interaction is another critical factor that can account for the presence of public TCRs. This is supported by structural studies of multiple public TCRs (44,102,119), which highlight that biases in V/J gene usage and CDR3 motifs (which define public TCRs) have structural foundations, whereby residues encoded within the biases are critical for pMHC engagement. This can be exemplified by the public HLA-A*02-restricted M158-specific TCRs, which are strongly biased for TRBV19 and a non-germline-encoded arginine from CDR3\(\beta\) loop. Based on the prototypical A2/M158-specific JM22 TCR structure (119), the TRBV19 gene segment dominates contacts with the MHC through its CDR1\(\beta\) and CDR2\(\beta\) loops, enabling the TCR to adopt the correct orientation over the peptide, so that the conserved arginine from CDR3\(\beta\) can form the “peg-in-notch” created by the peptide and the MHC α2-helix (119).

Overall, the public TCR clonotypes can be generated more efficiently, which result in a numeric advantage in the naive repertoire. These TCRs then need to optimally interact with their cognate epitope and outcompete other TCRs. Indeed, both criteria need to be met, as more efficiently generated TCRs still need to expand and clonally dominate the response. Conversely, TCRs capable of optimal pMHC engagement and clonal dominance need to be efficiently generated by recombination and preselected for in the thymus.

**TCR Repertoire Diversity and Selection During Immune Responses**

The majority of evidence on generation and persistence of antigen-specific TCRs comes from easily manipulated mouse models of viral infection or peptide immunization. The development of a tetramer-magnetic enrichment approach (86) has facilitated a dissection of naive antigen-specific TCR repertoires and the understanding of how particular clonotypes are recruited into an immune response. Using this method, the naive precursor frequencies of CD4\(^+\) T cells directed against peptides restricted by I-A\(^b\) were estimated between ~20 and 200 per mouse (~190 for 2W1S:1-A\(^b\), ~20 for FlkC:I-A\(^b\), and ~16 for OVA:1-A\(^b\) specificities), when spleen and lymph node were analyzed (86). The naive population size correlated with the magnitude of the primary immunodominant CD4\(^+\) T cell responses. The TCRβ diversity of the naive antigen-specific CD4\(^+\) T cells, as measured by V\(\beta\) antibody staining and fluorescence-activated cell sorting, was highly reflective of the TCR repertoire on d8 after peptide + lipopolysaccharide immunization.

Adapting the enrichment method to CD8\(^+\) T cells, the numbers of naive tetramer-specific CD8\(^+\) T cells per mouse range from 10s (~30 naive precursors for immunodominant influenza-specific D\(^b\)NP366 (72)) to 100s (~600 for murine CMV D\(^b\)M15 (91)) and 1,000s in one extreme example (~3,000 for malaria D\(^b\)GAP50) (131). Although the size of the naive CD8\(^+\) T cell pools can predict the size of the immunodominant effector CD8\(^+\) T cell populations (91), it is CD8\(^+\) T cell quality, rather than merely quantity, which plays a key role in determining immune response magnitude (33,72). Using tetramer enrichment, single-cell sorting, and reverse transcription-polymerase chain reaction (RT-PCR), the composition and diversity of naive and effector TCR repertoires were compared across four different CD8\(^+\) specificities (D\(^b\)NP366, D\(^b\)PA224, D\(^b\)PB1-F262, K\(^b\)NS2, D\(^b\)CD8\(^+\)) in B6 mice (72,122). While the main TCR characteristics (βJ region and CDR3 length) were highly comparable between the naive and effector TCR repertoires, selective expansion of TCR clonotypes and differential clonotype distribution was evident. This most likely results from both the intrinsic and extrinsic (antigen driven) characteristics of the naive TCRs (122). Notably, the subdominant T cell specificities (D\(^b\)PB1-F262, D\(^b\)CD8\(^+\), K\(^b\)NS2, D\(^b\)CD8\(^+\)) demonstrated a failure to fully recruit and expand the naive precursor pool, which correlated with a lower prevalence of high-avidity T cells compared to immunodominant T cell specificities (D\(^b\)NP366, D\(^b\)CD8\(^+\), D\(^b\)PA224, D\(^b\)CD8\(^+\)) (33,72).

The notion of biased TCR recruitment is further supported by the use of a sophisticated transgenic TCR\(\alpha\)β barcoding approach to track T cells during an immune response. Although the majority of the naive TCR repertoire was recruited into the immune response (134), the different T cell families expanded at different rates (42). While the dominant transgenic OT-I TCR clones expanded 400-fold more than an average naive clone, subdominant clones, corresponding to 50% of OT-I families, had fewer than 200 daughter cells. An inverse correlation between CD62L expression and the clonal size suggests that despite a single TCR specificity, naive clones differ markedly in their clonal expansion and differentiation.

Using a human-adapted tetramer enrichment, the naive T cell precursor frequencies for HLA-A*0201-restricted NY-ESO-1\(_{1157–165}\), WT1\(_{126–134}\), HIV Gag p177–85, HCV Core\(_{132–140}\), and CMV pp65 antigens in cancer-free and seronegative individuals have been estimated between 0.6 × 10\(^{-6}\) and 5.3 × 10\(^{-6}\) CD8\(^+\) T cells (2.88). In comparison,
frequencies of epitope-specific memory CD8+ T cells are in the range of 1 antigen-specific cells in 10^5 CD8+ T cells (1\times10^{-5}) (2,69,88). TCR\(\delta\)\(\beta\) signatures within naïve human antigen-specific T cells are still to be elucidated. Overall, these studies shed light on how the naïve dominant and subdominant TCR clonotypes are mobilized into an immune response, important for designing improved vaccines and immunotherapies that aim to promote the magnitude and breadth of the T cell response.

The exact mechanism underlying preferential selection of specific TCR clones over the others is unclear; however, published evidence suggests that the avidity for pMHC is a key determinant of TCR selection and clonotype representation in the immune repertoire. As high-avidity CD8+ T responses can kill virus-infected cells \(~1\,000\) more efficiently than low-avidity T cell pools (3), high-avidity TCR clonotypes are clearly important for more favorable clinical outcomes. An approach combining single-cell RT-PCR and affinity measurements directly linked pMHC-II binding affinities and clonal diversity of responding CD4+ T cells (77). Although naïve CD4+ T cells specific for the pigeon coctreme c (PCC) antigen displayed a wide array of TCR affinities, TCR clonotypes with low-affinity TCRs were lost early after antigen exposure. Thus, it appears that the affinity threshold for pMHC class II is required for the transition of PCC-specific CD4+ T cells from naive to fully expanded immune effectors. This strategy enables the best-fit TCRs to have a selective advantage over less-fit clonotypes in the naïve T cell repertoire (77).

Although only TCRs with the “adequate” threshold pMHC affinity get recruited into the immune response, TCR clonotypes within any given response display a wide range of pMHC avidities, all undoubtedly above the recruitment threshold level. Clonotypic dissection of high pMHC avidity T cell pools performed by single-cell RT-PCR analysis combined with tetramer dilution (63) and tetramer dissociation (85) probed the link between pMHCI avidity and specific effector TCR clonotypes. In comparison to the total pool of antigen-specific CD8+ T cells (stained with tetramers at saturating levels and no anti-H2D\(^{b}\) antibody), a subset of immunodominant D\(^{b}\)PA224+V\(^{\beta}\)7+CD8+ TCRs was preferentially selected under conditions of limited tetramer availability. This resulted in particular D\(^{b}\)PA224+TCR\(\beta\) clonotypes being overrepresented in the “high-avidity” population. Conversely, prominent public D\(^{b}\)NP366+V\(^{\beta}\)8.3+CD8+ T cell clonotypes were present at equivalent prevalence in immune T cells when sorted from the total antigen-specific CD8+ T cells. As D\(^{b}\)PA224+V\(^{\beta}\)7+CD8+ T cells are of much higher pMHCI avidity than D\(^{b}\)NP366+V\(^{\beta}\)8.3+CD8+ pools (73), there appears to be a pattern of differential TCR\(\beta\) clonotype distribution within high and low pMHCI avidity populations. Thus, a higher degree of clonal TCR\(\beta\) diversity within a particular epitope leads to more scope for pMHCI avidity segregation between the clonotypes. In contrast, the finding of the same public D\(^{b}\)NP366+V\(^{\beta}\)8.3+CD8+ clonotypes being present in both high- and low-avidity subsets suggests that other intrinsic factors, apart from pMHCI–TCR interactions, promote the overall effectiveness for T cell specificities with lower overall pMHCI-I avidity.

Active repertoire selection and clonal dominance in immune repertoires are based on TCR avidity for pMHCI. This was supported by a subsequent human study showing that the emergence of dominant TCR clonotypes during infection correlates with higher TCR avidity for pMHCI tetramers in individuals with chronic CMV or EBV infection (96). High pMHCI TCRs are more preferably driven toward extensive expansion and terminal differentiation (95).

While high pMHCI avidity allows rapid recruitment of the “best-fit” clonotypes into an immune response, lower pMHCI avidity clonotypes can use a number of intrinsic mechanisms to enhance overall avidity and functional sensitivity, enabling their competitive recruitment and maintenance within an immune response. These mechanisms can be divided into (i) co-receptors that contribute to the TCR-pMHCI interaction (through binding/stabilization or synapse formation, e.g., CD8\(\beta\)) and (ii) co-receptors that provide the necessary signals for entering the effector or memory pathways (CD27 and CD28 co-receptors or cytokine receptors, including IL-2R and IL-7R).

Differential recruitment into the immune response of barcoded transgenic OT-I cells with a single TCR specificity, as discussed above, provides clear evidence that preferential selection of TCR clonotypes does not solely depend on a TCR\(\alpha\)\(\beta\) signature (42). Similarly, the distribution of public TCR\(\beta\) clonotypes among both low and high pMHCI avidity subsets within the influenza-specific D\(^{b}\)NP366+CD8+ T cell response (63,85) suggests that intrinsic factors other than CDR3\(\beta\) sequence contribute to T cell avidity. The main intrinsic contributor to pMHCI avidity is the dependence of specific T cells on a CD8\(\beta\) co-receptor for stable pMHCI binding. Experimentally, this is assessed with either CD8\(\beta\)-null 227 tetramers, which prevent CD8\(\beta\)-co-receptor binding and thereby detect only high avidity CD8+ T cells (96), or by using a blocking anti-CD8\(\beta\) mAb in combination with peptide stimulation in an ex vivo ICS (127). It is well established that TCR clonotypes with lower pMHCI avidities have higher dependence on CD8\(\beta\) for recruitment into the immune response (85,95,96). Thus, the CD8\(\beta\) co-receptor contributes greatly to the overall avidity of TCRs and serves to promote clonotypic diversity within an immune response.

Second, TCR\(\alpha\)\(\beta\) heterodimer expression levels and adjacent membrane organization also play an important role in determining T cell fitness during the immune response. The efficiency of TCR\(\alpha\)\(\beta\) surface expression depends highly on the intrinsic qualities of the clonally expressed TCR-CD3 complex and strongly correlates with functional activity of T cells (50). Early in the response, the affinity of CD8+ T cells can also be increased by optimization of the TCR-mediated signaling machinery, specifically Lck expression (114). Higher Lck protein levels lead to markedly increased IFN-\(\gamma\) production (functional avidity), thus allowing “affinity maturation” without any genetic modification in actual TCR\(\alpha\)\(\beta\) signatures. The cholesterol organization of a T cell membrane also contributes to pMHCI avidity and the capacity to respond to a low antigen dose. Membrane TCR reorganization upon activation allows antigen-specific TCRs to bind multimeric pMHCI complexes with increased avidity. This enhances responsiveness mediated through increased cross-linking of TCRs (36). Furthermore, the avidity of CD8+ T cells can be generated by co-localization of TCRs together with CD8\(\alpha\)\(\alpha\) and CD8\(\alpha\)\(\beta\) in the lipid rafts (27). Such optimal TCR-CD8 organization requires relatively fewer TCR engagements when compared to low-avidity TCRs. Finally, the expression (and/or levels) of co-stimulatory
molecules like CD27 and CD28 as well as cytokine receptors like IL-2Rα and IL-7Rα can promote more efficient recruitment of T cells into the effector differentiation pathway (62,92).

Thus, while TCRαβ sequence appears to be the major contributor to pMHC avidity in diverse virus-specific T cell populations, other intrinsic mechanisms promoting CD8β and TCR–pMHC interactions play an important role in avidity determination, especially in the absence of TCR diversity.

TCR Repertoire Diversity Across Distinct Stages of CD8+ T Cell Differentiation

The frequency of memory T cells is <1% at birth and greatly increases with age. In humans, memory CD4+ and CD8+ T cells constitute ~50% of all the circulating peripheral blood T cells in a healthy adult (89). Established memory T cell populations provide protection against recurrent infections, leading to milder clinical outcomes and rapid recovery of the host. For example, even in the absence of neutralizing antibodies to the newly emerged influenza viruses, preexisting memory CD8+ and CD4+ T cell pools promote recovery from experimental H3N2, natural H1N1-2009, or avian H7N9 influenza (118,140,144).

The frequency of epitope-specific memory T cells is up to 200–1,000 times (31) greater than the naïve precursor pool (72), allowing rapid responses and enhanced pathogen clearance upon recall. In general, the TCR clonotypic composition of memory T cell pools reflects that of the naïve and early or late effector T cell sets. In the B6 mouse model of influenza infection, longitudinal analyses of the same animals sampled repeatedly (using blood) have allowed dissection of individual clonotypes from acute primary effectors through to early and long-term memory using single-cell sorting, RT-PCR, and CDR3β sequencing (64,125). The prominent TCRβ clonotypes found at the acute effector phase for two immunodominant responses, DضpPA223-Vβ7-CD8+ and DضpNP336-Vβ8.3-CD8+ populations, persisted into both short- and long-term memory, and could be subsequently expanded after the secondary challenge (32,64,125). Furthermore, both TCRβ diversity and clonotypic composition within the antigen-specific memory pools are equivalent to those found at other time points after infection, including early antigen exposure. Such clonotypic stability of the long-term memory T cell populations is supported by earlier data from LCMV infection or CW3 immunization (14,78,117). In contrast, however, TCR narrowing in the memory pool has been reported for Listeria monocytogenes-specific CD8+ T cells and PCC-specific CD4+ T cells (25,80). In this study, the preferential selection of high-avidity TCR clones into the memory pool has been linked to TCR affinity/avidity maturation. Such differential levels of TCR clonotype persistence into memory might be related to a certain pMHC−TCR avidity threshold required for survival into the long-term memory.

Memory T cells are highly heterogeneous with distinct lymph node homing properties (CD62L and CCR7), anatomical localizations and functions (106). An obvious question emerges as to how different TCR clonotypes contribute to the specific memory subsets and their functions. Experiments utilizing clonotypic analysis of TCR signatures in humans (8) and mice (18,66) showed that common TCRs are distributed across the main memory T cell subsets, central T cell memory (Tcm) and effector T cell memory (Tem) populations. However, strikingly, the Tcm sets contain additional TCRβ clonotypes (diversity increased by ~30%) not found within the Tem populations. TCR diversity within both the Tcm and Tem subsets was consistently stable, starting from the early acute phase of infection (d8) through to d28 early and >d500 long-term memory (66). The main conclusion here is that different T cell memory subsets contain stable TCRβ signatures from the early days after antigen exposure, with the memory Tcm pool preserving clonal diversity and preventing “overdominance” by a few greatly expanded TCR clones. Together, these findings support the idea that any antigen-specific T cell (with the sufficient pMHCi avidity) that has not achieved a terminally differentiated stage can become a memory T cell. As the establishment of immunological memory is critical for a rational design of any cell-mediated vaccine, understanding the capacity of the optimal and suboptimal TCR clonotypes to transition between functionally different naive/effector/memory subsets is important.

The Importance of TCR Repertoire Diversity on Disease Outcome

TCR clonotype selection during an antigen-driven T cell response can influence biological outcome, with diversity and composition of the peripheral TCR repertoire having major consequences for subsequent immune responses in both animal models and human disease (83). Nikolich-Zugich and colleagues presented the first elegant evidence for the link between TCR repertoire diversity and immune protection (82). Although the herpes virus (HSV)-derived SL8 (gag35–50) peptide is presented equally well by both the wt H2Kb and the mutant H2-Kbm8 MHC class I glycoproteins, mice expressing the wt H2Kb allele are much more susceptible to HSV infection. Dissection of the SL8-specific TCR repertoire established that SL8-H2Kbm8 selects more diverse and higher-avidity TCRs (82). Thus, TCR repertoire diversity, pMHCi avidity, and protection are interrelated and influence resistance profiles.

A subsequent link between TCR repertoire diversity, avidity, and protection from viral escape was established in the rhesus macaque SIV model (96,97). SIV-specific CD8+ T cell responses are directed against two prominent epitopes characterized by different TCR repertoires. The Tat28–35 (TL8)-Mamu-A*01-specific CD8+ T cells are characterized by expression of TCRs with conserved CDR3β sequences, while the Gag181–189 (CM9)-Mamu-A*01-specific clonotypes display diverse CDR3β sequences. CD8+ T cell responses to the SIV-TL8 peptide with a restricted TCR repertoire were associated with the rapid emergence viral escape mutants at p5 of the TL8 peptide, while the diverse TCR repertoire of SIV CM9-specific CD8+ T cells led to a high level of conservation within the wt CM9 peptide sequence. Although mutations may occur within the CM9 peptide, the CM9-Mamu-A*01-specific TCRs, especially the public clonotypes (95), have the breadth to recognize escape variants, thereby preventing the outgrowth of mutant viruses. Thus, the diverse CM9-specific TCR repertoire allows the preferential selection of high-avidity TCR clonotypes, recognizing the newly emerging viral variants (96).
As with persistent infections, TCR diversity protects from viral escape in acute, readily resolved infections (129). In B6 mice, the diverse and high pMHC avidity DPα225-specific TCR repertoire protects against viral escape during influenza infection. Conversely, sequential emergence of escape variants occurs preferentially within the DPNP366 + epitope, which is recognized by CD8 + T cells expressing a restricted TCRβ repertoire (64). The selection of CD8 + T cell escape mutants is driven by selective pressure from DPNP366 + CD8 + T cells as the variants revert to the wt sequence in the absence of immune pressure in MHC-mismatched mice (129). Preemptive priming against the influenza virus mutants, however, can expand T cells expressing additional distinct TCRs that are capable of mounting robust recall responses against the viral variants (129).

Overall, the data from animal models strongly suggest that the extent of TCR diversity, and the resultant selection of TCR clonotypes with high avidity for pMHC, generates broadly protective optimal responses capable of recognizing both the wt virus and the newly emerging mutants.

Studies point to the importance of TCR selection for viral control and disease-associated morbidity in HIV, hepatitis C virus (HCV), CMV, and influenza. The most solid evidence comes from HIV infection, where HLA association with the disease outcome is well documented. Given the key role of MHC molecules in TCR selection during thymocyte ontogeny, differences in MHC haplotype are likely to shape the available TCR repertoire. While some HLA alleles, HLA-B*27, B*57, and B*58, have been identified as “protective” against HIV-1, others, including HLA-B*35 and HLA-B*53, are considered susceptibility factors for more rapid disease progression. This implies the importance of CD8 + T cells (and specific TCRs involved) in the control of HIV infection. However, as not all HLA-B*2705 individuals control HIV infection, the composition of the specific TCR clonotypes in a selected HLA-B*2705 population drives HIV control. Indeed, a comparative study of five elite controllers and five progressors found that differential control of HIV-1 is mediated through specific TCR clonotypes directed toward the immunodominant Gag epitope, KK10 (28).

Superior control of HIV replication by “protective” TCR clonotypes was linked to the cross-reactive capacity to recognize the wt and the “escape” L6M variant, combined with their optimal capacity to rapidly deliver lytic granules (granzymes/perforin) to the pMHC-I-TCR immunological synapse and kill virally infected targets. In controllers, such effective TCRs are consistently of high prevalence (immunodominant), while the progressors possess only functionally ineffective clonotypes (immunodominant and subdominant). Interestingly, the CDR3 sequences of KK10-specific CD8 + T cells in the HIV controllers are close to germline and have been shown previously to efficiently recognize emerging viral variants, and thus control HIV infection (68).

An independent study also showed that cross-reactive KK10-specific TCR clonotypes, as shown by double-tetramer staining and similar docking sites within ternary pMHC-TCR complexes, can be rapidly generated and recruited to control both and the wt virus and the newly emerged L6M mutant (74). Similarly, the antiviral efficacy and cross-reactivity of the TW10 epitope (Gag 240–246) were superior in HLA-B57-expressing controllers compared to progressors (28). Clearly, such associations between MHC haplotype, TCR repertoire, and disease control are of key interest as we seek to understand and manipulate T cell immunity through vaccination.

Similar to HIV, protective HLA alleles (A*0201, A*0301, B*5701, B*1801, and B*0801) are associated with universal cross-strain CD8 + T cell immunity against acute influenza A virus infections, including the newly emerged H7N9 subtype (98). In contrast, individuals expressing A*2402, A*6801, and B*1501 show poor influenza-specific CD8 + T cell response profiles (52,98,132). The molecular mechanism underlying the universal HLA-mediated protection is mainly unknown. To date, TCR repertoire for influenza-specific CD8 + T cells has been reported across different HLAAs (29,43,89,93,109,128,141). The CD8 + T cells specific for A2-M158 are exclusively restricted to TRBV19 usage and express similar CDR3β sequences. A dominant public TCRαβ clone (TRAV27 GGSQGNL; TRBV19 SIRSSYEQ) can be found at high frequency within M158 CD8 + T cell responses in all HLA-A*0201 individuals tested. This public A2-M158 TCR can also recognize the naturally occurring variants of the M158 peptide (128), which contributes, at least in part, to the high conservation rate of the M158 epitope within the influenza viruses in human circulation. This further suggests that the HLA-A2 + population (up to 50% population coverage) has universal T cell immunity to any M158 variant. In contrast, the immunodominant influenza-specific NP418–426 peptide presented by the HLA-B7 superfamily (including HLA-B*3501, B*3503, B*0702) has sequentially mutated at four different TCR contact positions, generating over 20 different peptide sequences over the past 90 years (47). However, depending on mutations occurring at the most solvent-exposed residues at p4 and p5, B7-NP418-specific CD8 + T cell responses show two distinct profiles of cross-reactivity toward either seasonal (1947–2005) or pandemic (1918, 2009) influenza-derived NP418 peptides. TCR clonotypes within the B7 NP418 CD8 + T cell populations differentially recognize distinct peptides from either the pandemic or seasonal influenza viruses (128). Thus, a preemptive influenza vaccine that elicits TCR pools, which recognize distinct antigenic variants, can provide universal immunity against any future influenza strain or subtype.

In CMV infection, the diversity of TCRαβ repertoire (rather than the magnitude of the CD8 + T cell pool) against the HLA-A*0201-restricted pp65495–503 (NLV) epitope is associated with viral control (139). TCRαβ diversity, analyzed by unbiased single-cell multiplex RT-PCR, inversely correlated with the circulating antibody levels, a hallmark of viral load in patients. Further modification of this technique to utilize IFN-γ capture in clinical settings found a novel TCR signature (TRAV3TRAJ31-TRBV12-TRBJ1-1) within the HLA-A*0201-restricted CMV-NLV + CD8 + response capable of recognizing alloreactive peptide/s in the context of HLA-B27 alleles (B*2705, B*2707, and B*2709) (87). This clearly indicates the importance of any cross-reactive viral-specific TCRs in HLA-mismatched solid organ transplants. Cross-reactive CMV-specific CD8 + TCR clonotypes capable of recognizing distinct viral variants evolve over time, as shown for the immunodominant HLA-B8-restricted CMV IE-1 epitope (115). This is in contrast to CD8 + T cells directed toward a single genetic variant with oligoclonal private TCR repertoires.
Although there is much less understanding of the involvement of TCRs in controlling the HCV disease, comparative analysis of two immunodominant A*0101- and B*0801-restricted epitopes showed that TCR repertoire diversity might be associated with HCV persistence or clearance in humans (84). Such diversity generated by switching of TRBV and TRBJ genes around a conserved CDR3β loop in HCV-specific TCRs suggests a potential mechanism to control antigenic variations in HCV. Recent studies have also focused on TCR clonotype usage in cancer and autoimmunity and found common features between virus- and cancer-recognizing TCRs, including TCR bias and CDR3β length (83).

Concluding Remarks

Studies in the last four decades have made significant progress in our understanding of T cell-mediated immunity and dissecting the complexities of T cell populations, and have generated critical questions that need to be addressed if we are to harness T cells by vaccination and immunotherapies. Published evidence from both animals and humans shows the importance of TCR repertoire in the disease outcome, with a diverse TCR repertoire providing a greater range of clonotypes with scope for the preferential selection of high-avidity TCRs into the immune response. Of a particular advantage for disease control are diverse TCR repertoires with high pMHC avidity TCR clonotypes capable of promoting superior CD8+ T cell function and recognizing viral escape mutants, as well as the public prominent TCR clonotypes. However, the exact mechanisms underlying recruitment, persistence, and recall of dominant TCR clonotypes across the human life span, from early childhood to the aged years, are far from being clear and need to be taken into consideration when developing T cell-based vaccination strategies to provide effective and optimal protection across the life span of an individual. Similarly, the immunodominance of T cells directed at numerous epitopes across different HLAs within the same individual should be addressed, and so should identification of novel T cell epitopes across different ethnicities so the T cell-mediated vaccines can reach a substantial global coverage. Finally, while we now understand how TCRs and pMHCs interact at the molecular level, the very basis of MHC restriction remains enigmatic and highly debatable (71). All this knowledge gained and all the arising questions, which span every aspect of T cell biology and have been pioneered by work by Peter Doherty and colleagues, are only possible because of the 1974 groundbreaking discovery of MHC restriction by Doherty and Zinkernagel.

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