Biomechanics of T Cell Dysfunctions in Chronic Diseases

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Understanding the mechanisms behind T cell dysfunctions during chronic diseases is critical in developing effective immunotherapies. As demonstrated by several animal models and human studies, T cell dysfunctions are induced during chronic diseases, spanning from infections to cancer. Although factors governing the onset and the extent of the functional impairment of T cells can differ during infections and cancer, most dysfunctional phenotypes share common phenotypic traits in their immune receptor and biophysical landscape. Through the latest developments in biophysical techniques applied to explore cell membrane and receptor–ligand dynamics, we are able to dissect and gain further insights into the driving mechanisms behind T cell dysfunctions. These insights may prove useful in developing immunotherapies aimed at reinvigorating our immune system to fight off infections and malignancies more effectively. The recent success with checkpoint inhibitors in treating cancer opens new avenues to develop more effective, targeted immunotherapies. Here, we highlight the studies focused on the transformation of the biophysical landscape during infections and cancer, and how T cell biomechanics shaped the immunopathology associated with chronic diseases.

Keywords: T cell dysfunction, chronic diseases, infections, cancer, tumor microenvironment, immune receptor landscape, biophysical landscape, biomechanics

INTRODUCTION

T cells are at the frontline of immune surveillance, acting against pathogens and malignancies to maintain host homeostasis. Upon recognition of antigenic peptides presented on major histocompatibility complex (MHC) or MHC-like molecules (1, 2), T cells become activated and undergo clonal expansion, resulting in the generation of effector cells that help contain the spread of the disease. During clonal expansion, changes can occur at transcriptional, epigenetic and metabolic levels that enhance the effector functions of T cells (3). Effector T cells produce high amounts of cytokines, including interferon (IFNγ) and tumor necrosis factor (TNFα), and cytoplasmic granules containing granzymes and perforin (4). During antigenic clearance, the majority of effector CD8⁺ T cells follow an apoptotic cell death, but 5–10% of cells differentiate into memory T cells (5). Memory T cells are capable of rapidly executing their effector functions upon re-encounter of the same antigen or pathogen (6, 7). They have a unique transcriptional makeup, which allows them to be distinguished from naïve and effector T cells (8). This unique transcriptional profile shapes the functional characteristics of memory T cells as well as their phenotype to maintain the acquired
Dependong on phenotypic and functional features, T cell dysfunctions can be classified into ignorance, tolerance, exhaustion, anergy or senescence (Table 1). These different states of dysfunctions can operate as mechanisms to reduce autoimmunity (ignorance and tolerance), minimise repercussions from inappropriate T cell stimulation (anergy), or simply regulate T cell division (senescence) (9). However, exhaustion brings a unique perspective to the state of T cell dysfunction as it occurs despite physiologically appropriate T cell stimulation. During the transformation of effector T cells into dysfunctional phenotypes, the proliferative capacity along with cytokine production gets reduced (10–12). Moreover, alterations occur at the membrane level transform the immune receptor landscape of T cells (13) and their biophysical properties (14–16). This has ramifications in terms of maintaining optimal T cell responses against pathogens and malignancies. In this review, we highlight recent findings that helped to broaden our understanding on how T cell dysfunctions can reform the immune receptor and biophysical landscape of T cells, and how it can ultimately influence the state of disease progression. We primarily discuss dysfunctional T cell phenotypes in the context of chronic infections and cancer to draw our conclusions.

**MECHANISMS OF T CELL DYSFUNCTION IN CHRONIC DISEASES**

It is conceivable that all T cell dysfunctions stem from alterations occur in the biological process of T cell activation or differentiation. As a result, different dysfunctional T cells share common phenotypic traits, which make the differentiation between T cell dysfunctions subtypes difficult. However, in depth understanding of different molecular mechanisms driving T cell dysfunctions will help to identify signature phenotypic traits to build much clearer and distinguishable profiles for each subtype. Here, we attempt to highlight different mechanisms that drive T cell dysfunctions and the most commonly associated dysfunctional T cells found in chronic diseases.

**T Cell Tolerance and Ignorance**

Complete T cell activation requires three signals; first signal provided by TCR-cognitive pMHC interaction, the second is a costimulatory or coinhibitory receptor activation signal provided by APCs, and the third is provided by extracellular cytokines. Of these signals, the second signal becomes crucial in determining the functional outcome of T cell signaling, which may promote T cell effector functions (costimulation) or dampen excessive immune responses (coinhibition) to maintain immunological tolerance (17). Thus, both T cell activation and tolerance are interconnected to tightly regulate and maintain optimal immune responses against foreign antigens while preventing autoimmunity against self-antigens. Failure to maintain immunological tolerance may result in various types of autoimmune diseases. T cell tolerance is primarily enforced by central and peripheral tolerance. Central tolerance operates by eliminating self-reactive T cell through negative selection in the thymus during early stages of T cell development. These T cells express high-avidity TCRs to self-antigens and are mainly eliminated from the system via clonal deletion or diverted to differentiate into regulatory T cells (Treg) through thymic negative selection. These elimination mechanisms have been reviewed elsewhere (18). Although majority of self-reactive T cells get screened and eliminated though negative selection in the thymus, this process alone is not sufficient to safeguard against autoimmunity. Self-reactive T cells that escape thymic negative selection are eliminated by peripheral tolerance which acts as the second barrier to maintain immunological tolerance. It was shown that peripheral tolerance is most effective in detecting and eliminating mature T cells that express low-avidity self-reactive TCRs, while central tolerance is effective against eliminating thymocytes expressing high-avidity self-reactive TCRs (19). Peripheral tolerance operates with various mechanisms to inactivate self-reactive T cells that escaped central tolerance. These mechanisms include clonal deletion (20, 21), clonal suppression by Tregs (22–24) and induction of functional non-responsiveness via intrinsic cell programming mechanisms (25). It has been suggested that manifestation of a large proportion of autoimmune diseases are linked with the breakdown of peripheral tolerance mechanisms (26–29). In some instances, self-antigens fail to induce negative selection of self-reactive T cells and they become clonally ignorant (30–32). This can be due to low expression of the self-antigen or its physical sequestration at immune-privileged sites like the blood-brain barrier (31). During self-antigen encounter, unlike self-tolerant T cells, self-ignorant T cells remain functional. Most self-ignorant T cells in the periphery are naive, but given the right stimulatory conditions, they can initiate autoimmune responses (33–35).

**T Cell Anergy**

Another important state of T cell dysfunction is anergy. The consensus that describe the mechanism behind T cell anergy is based on T cell antigen-stimulation in the absence of the second signal i.e. costimulation, which drives T cells into a
hyporesponsive state for an extended period of time (36). Upon re-encounter of the same stimuli with optimal costimulation, anergic T cells fail to proliferate and produce cytokines (36). It has been shown that one of the hallmarks of T cell anergy is the reduced interleukin (IL)-2 production (37). T cell anergy has been broadly classified into clonal anergy and in vivo anergy (36). Clonal anergy can be induced in CD4+ T cells when stimulated with a strong first signal (TCR-pMHC interaction) and in the absence of the second signal. Low doses of agonist in the presence of costimulation has also been shown to induce clonal anergy (38). In vivo anergy also known as adaptive tolerance can occur in the thymus or in the periphery and often associates with naïve T cells during self-antigen stimulation in a costimulation deficient or high coinhibitory environment (37, 39). For example, cancer cells and tumor-antigen presenting cells are shown to express high levels of coinhibitory receptor ligands (PD-L1, PD-L2 and many other) with relatively low levels of costimulatory receptor ligands (CD80 and CD86) in the tumor microenvironment to promote T cell anergy (40–43). Despite overlapping functional and phenotypic features, clonal anergy and in vivo anergy are driven by distinct molecular mechanisms. For instance, while both anergic phenotypes display either impaired IL-2 production (clonal anergy) or impairment in all TCR-induced cytokine production (in vivo anergy) as a key dysfunctional feature during antigen stimulation, only clonal anergy can be rescued by the addition of exogenous IL-2 or by diacylglycerol kinase-α (DGK) inhibitor (44, 45). Clonal and in vivo anergy also differ in their signaling defects. TCR-based signaling pathway seems to have impairment in Zap-70 phosphorylation of LAT in the in vivo anergy model (37). The signaling pathway of clonal anergy shown to have defects in MAP-kinases activation and mobilisation of NF-κB to the nucleus (46). Anergic phenotypes are associated with a number of autoimmune diseases including human type-1 diabetes (26), systemic lupus erythematosus (47), autoimmune gastritis (48) and myasthenia gravis (49).

**T Cell Senescence**

Senescence is recognized as a T cell dysfunction that can play paradoxical roles in adaptive immunity. Based on the triggering mechanisms, T cell senescence can be classified into replicative senescence or premature senescence (30). Replicative senescence occurs as a consequence of telomere shortening after several rounds of cell division, which is associated with the natural aging process (51–53). Hence, an increased number of senescent T cells have been found in the elderly population, which increase their susceptibility for malignancies and chronic diseases (54, 55). However, a number of studies have shown that accumulation of senescent T cells is not limited to the aged population, but can be found in younger patients with chronic infections and cancer (56–59). The second form of senescence: premature senescence is independent of telomere shortening, and is induced by external factors including cellular stress, particularly oncogenic stress (60, 61). During tumorigenesis, oncogenes become activated and promote uncontrolled cell division, which is commonly observed in many types of cancers. Adversely, a high proliferation rate in cancer cells can become a genetic and metabolic burden which triggers cellular senescence pathways, causing irreversible cell cycle arrest (62, 63). This demonstrates the paradoxical nature of cellular senescence. Moreover, by inducing DNA damage responses, both Tregs and tumor cells can convert T cells to become senescent (64–66). Transformation of effector to senescent T cells dramatically change the immune receptor landscape of T cells. The marked decline of costimulatory receptor expression (CD27 and CD28) (67, 68) is one of the main biomarkers of senescent T cells alongside higher expression of killer cell lectin-like receptor subfamily G member 1 (KLRG-1), Tim-3, CD57 and CD45RA (69–71).

**T Cell Exhaustion**

Since its first characterisation in lymphocytic choriomeningitis virus (LCMV) infection model in mice (72), T cell exhaustion has been the topic of much debate and is implicated in a number of chronic infections (primarily caused by viruses) and cancer (73–76) (Figure 1). Over the past few years, the topic of T cell exhaustion has become more relevant as we attempt to uncover the molecular mechanisms behind chronic T cell dysfunctions and develop effective immunotherapies to manage these conditions.

Although driving forces of T cell exhaustion may differ base on different pathological settings, most, if not all proposed mechanisms of T cell exhaustion centres around the three-signal model of T cell activation. Persistent antigen-stimulation, effects of native-regulatory cytokines and immune-suppressive influence of immunoregulatory cells like Tregs are known to promote exhaustion in effector T cells (79). Among these exhaustion inducible factors, persistent antigen-stimulation has been observed across several chronic infection and cancer models in humans and mice (80–82). Accordingly, the dose and the duration of antigen exposure can contribute to the degree of T cell exhaustion. This has been reviewed in later sections of the review.

A key difference has been identified in T cell differentiation during acute and chronic phases of a disease. In the acute phase, T cell mediated antigen clearance comprise of T cell expansion, contraction, and generation of memory T cells. This pattern diverges from the classic differentiation pathway during chronic infections and cancer as a consequence of persistent, higher prevalence of antigens. In this environment, T cells undergo persistent antigen stimulation, which progressively impair their effector functions and drive them to exhaustion. This functional impairment in exhausted T cells, however, does not describe a complete loss of effector functions. As reported by numerous accounts, exhausted T cells still retain some degree of effector functionality having control over the spread of the disease (83–85). Moreover, exhausted T cells share characteristics with memory T cells, which shows their capacity to survive long-term and respond to rechallenge of the antigen (86–89). An important differentiation arises from exhausted T cells having memory characteristics with those do not. Exhausted T cells with memory characteristics have shown to express transcription factor TCF1 (86, 87, 89–91). This subpopulation is responsible for maintaining immune responses during chronic diseases (86,
FIGURE 1 | Immune receptor landscape during T cell exhaustion. Exhaustion can be induced by chronic infections (in this instance viral infections) or cancer. Factors that influence the onset and the extent of T cell exhaustion differ in these two exhaustion models. During a chronic infection, pathogen clearance becomes inefficient, leading to persistent inflammation and chronic antigen stimulation of T cells which results in clonal deletion or exhaustion. In cancer, the immunosuppressive tumor microenvironment plays a crucial role in shaping the outcome of T cell exhaustion. Tumor microenvironment comprised of stroma containing a fibroblast network and a number of immune cells including regulatory T cells (Tregs) and tumor-associated macrophages (TAMs) which together promote tumorigenesis (77). Tumor microenvironment can induce stromal cells to secrete growth factors to promote angiogenesis (i.e. grow new blood vessels that feed the tumor) (78). Overall, T cell exhaustion in both chronic infections and cancer known to have several overlapping functional and phenotypic characteristics. The most common feature is sustained upregulation of inhibitory receptors during the course of the disease.
87, 92) and remains to be critical for the success of anit-PD-1 blockade therapy (87, 93, 94), while TCF1-exhausted T cells fail to provide such responses. In numerous studies TCF3 exhausted T cells were described as "stem-like" T cells (87, 89) or "progenitor exhausted" cells (93) and more recently these terms have been unified under "precursor exhausted T cells" (T_{PEX}) in contrast to TCF terminally exhausted effector T cells (T_{EX}) (95). Several other transcription factors have been identified to be coexpressed with TCF. Most notably, the T-box transcription factors T-bet and eomesodermin homologue (EOMES) which are known to regulate immune responses during acute and chronic infections (96–98). Their role in maintaining exhaustion phenotypes in both T_{PEX} and T_{EX} needs further investigation. B lymphocyte-induced maturation protein 1 (BLIMP1) is another critical transcription factor required in lymphocyte subset differentiation (99, 100). BLIMP1 suppresses multiple genes linked to T cell memory regulation, therefore, found to be partially expressed in T_{PEX} but not in T_{EX} (87, 91). Other transcription factors relevant in maintaining T cell exhaustion phenotype have been reviewed elsewhere (89, 101–103).

Cytokines play a major role in shaping the outcomes of T cell activation. During the acute phase of infection, proinflammatory cytokines (TNF-\(\alpha\), IL-1\(\beta\), and IL-8) promote the development of effector T cells to fight off the infection. When the pathogen persists through to the chronic phase, negative regulatory cytokines like IL-10 and TGF-\(\beta\) target different pathways to suppress T cell activation and induce T cell exhaustion. IL-10 is produced by multiple different cells in the tumor microenvironment (TME) including Tregs, tumor-associated macrophages (TAM) and cancer cells (104–106). By inducing PD-L1 expression in dendritic cells, IL-10 promotes T cell exhaustion (107). Downregulation of MHCs, intercellular adhesion molecule 1 (ICAM-1) and costimulatory ligands (CD80 and CD86) on APCs promoted by IL-10 also contribute to immunosuppression (108). Importantly, previous studies have shown that IL-10 plays a role as an anti-inflammatory cytokine by demonstrating its capacity to inhibit or downregulate the production of proinflammatory cytokines (109). It is well known that the surface lattice formed by galectin-glycoprotein can influence membrane remodelling to suppress T cell mediated immune responses (110, 111). Recently, IL-10 was shown to be involved in an immune loop regulatory N-glycan branching, which heightened galectin-3 binding, thereby decreasing T cell antigen sensitivity (112). Although, IL-10 has been broadly characterized as an immune suppressor, at higher concentrations IL-10 and PEGylated IL-10 (pegilodecakin) has shown to have properties that enhance cytotoxicity and proliferative capacity of tumor-specific CD8\(^+\) T cells (113–115). The mechanisms underlie this paradoxical nature of IL-10 remains to be investigated. TGF-\(\beta\) is a pleiotropic cytokine and is produced in large amounts in the TME. Generally, TGF-\(\beta\) restrict tumor growth in early stages by inducing the Smad signaling pathway (116). However, in late stages of the cancer, TGF-\(\beta\) has been linked to tumor progression by modulating immune responses most likely through a Smad-independent signaling pathway (117).

Given the paradoxical role of TGF-\(\beta\) in cancer, makes it one of the most complex factors to be studied in the TME. Understanding how and when TGF-\(\beta\) switch from tumor suppressor to tumor promoter is being actively investigated. Immunoregulatory cells including Tregs (CD4 and CD8), myeloid-derived suppressor cells (MDSC) and NK cells are shown to induce T cell exhaustion in effector T cells during chronic infections (118–122). Tregs at the site of the infection or at the TME secrete negative regulatory cytokines, IL-10 and TGF-\(\beta\), to promote immune suppression (122), thereby limiting anti-pathogen or anti-tumor activity of effector T cells. The exact mechanism of how Treg induce immune suppression to drive effector T cells into exhaustion is still unclear. However, reinivigoration studies of exhausted CD8\(^+\) T cells associated with chronic LCMV by blocking PD-1 signaling pathway and depleting Tregs simultaneously, suggest a role for Tregs in T cell exhaustion (123).

In summary, although persistent antigen stimulation remains as a key driving force for T cell exhaustion, numerous other factors differentially contribute to the development of exhaustion. In the next section we attempt to highlight two main models of T cell exhaustion found in chronic diseases.

**T CELL EXHAUSTION MODELS**

From studies reporting common phenotypic characteristics of exhausted T cells, there is an emerging profile that describes T cell exhaustion as a distinct state of cell differentiation. Accordingly, exhaustion phenotypes have been identified in both chronic viral infections and in cancer (Figure 1). Although these exhaustion models share common functional features, they differ substantially in some respects.

**T Cell Exhaustion in Chronic Infections**

The persistent overload of pathogens during chronic infections leads to persistent antigenic stimulation of T cells. This drive T cells into clonal deletion or exhaustion, both of which lead to reduced pathogen clearance. This is more commonly reported in infections associated with viruses, though T cell exhaustion has also been identified in bacterial and parasitic infections (124, 125). Here, our focus will be on T cell exhaustion during chronic viral infections. During chronic infections, reduced proliferative capacity and low interleukin-2 (IL-2) production (126) are known to be some of the earliest signs of loss of T cell effector functions. At the intermediate state, TNF-\(\alpha\) and IFN-\(\gamma\) production are reduced (73). The low cytotoxicity in CD8\(^+\) T cells is also observed at this stage. Loss of these functional properties occur partially or in severe exhaustion, completely. Finally, these exhausted virus-specific T cells are deleted from the system (72, 74). Hence, a stage-by-stage descent into exhaustion has been observed (10). The level of T cell exhaustion primarily depends on the amount and the strength of antigen stimulation (127). Although the "strength" of stimulus is difficult to define, prolonged exposure to a persistent viral load is an important determinant in the process of exhaustion. For example, higher
antigen load with prolong exposure results in severe exhaustion phenotypes seen in LCMV, untreated HBV, and HIV chronic infections (79, 128–130). The role of helper CD4+ T cells are also important for promoting effector CD8+ T cell functions, thus their low availability has been linked to T cell exhaustion (131, 132). Hence, high viral load and low availability of helper CD4+ T cells generally correlates with severe exhaustion phenotype (75). Overall, a number of factors including the viral load, location of viral replication and the immunosuppressive environment, contribute to the level of effector function impairment in T cells during chronic infections.

### Sustained Upregulation of Inhibitory Receptors

In addition to the gradual loss of effector functions, another classic feature of exhausted T cell is the sustained upregulation of inhibitory receptors (Figure 1). These surface expressed inhibitory receptors include programmed cell death protein 1 (PD-1) (133), cytotoxic T lymphocyte associated antigen 4 (CTLA-4), lymphocyte-activation gene 3 (LAG-3), T cell immunoglobulin and mucin-domain containing protein 3 (TIM-3), B and T lymphocyte attenuator (BTLA) and many others (13). In non-pathological settings, the transient expression of inhibitory receptors along with their co-stimulatory counterparts (CD28 and ICOS) serve a key role in maintaining the immunological tolerance. This is readily observed in acute infections where inhibitory receptors contribute to restrain immunopathology after pathogen clearance has been achieved. In fact, upregulation of inhibitory receptors are commonly observed during T cell activation (133–135), although steady-state expression levels may vary depending on the state of cell differentiation (136–138). As pathogen clearance progresses, inhibitory receptors are downregulated and maintained at low levels.

Many inhibitory receptors, including PD-1, can negatively regulate T cell receptor (TCR) signaling via immunoreceptor tyrosine-based inhibitory motifs (ITIM) or immunoreceptor tyrosine-based switch motifs (ITSM) found in their cytoplasmic tails (17). Upon binding ligands, ITIM/ITSM domains within the cytoplasmic tails of inhibitory receptors are phosphorylated and recruit Src homology region 2 domains within the cytoplasmic tails of inhibitory receptors are phosphorylated and recruit Src homology region 2 containing phosphatases (SHP-1 and SHP-2). Overall, PD-1/PD-L1 signaling pathway can regulate exhaustion phenotype by suppressing TCR signaling (139), inducing T cell suppressor genes (140) and by reducing T cell motility (141). TIGIT uses a similar strategy to negatively regulate T cell function (142). However, inhibitory receptors can engage more than one suppressive mechanism to attenuate T cell functions. In contrast to ITIM signaling, LAG-3 is known to function through KIEELE motifs located at its relatively short intracellular tail to negatively regulate cell cycle progression (143). Tim-3 also utilizes non-canonical inhibitory mechanisms that are distinct from, and complementary to, PD-1 (144). High surface expression of Tim-3 often correlates with severely exhausted T cell subsets during chronic infections (145, 146). The inhibitory receptor CTLA-4 functions by outcompeting CD28 stimulatory receptor by binding to their common ligands CD80 or CD86 to suppress T cell functions (147). Uniquely, CTLA-4 can utilize trans-endocytosis; a mechanism of capturing and removing common ligands from the surface of an antigen presenting cell (APC), thus making them unavailable for stimulatory receptor binding (148). All these inhibitory receptors can employ non-overlapping mechanisms of T cell suppression, making their functional role in promoting T cell exhaustion rather diverse and complex.

### T Cell Exhaustion in Cancer

Immunosuppressive factors found in the tumor microenvironment and the tumor-antigen load greatly influence the degree of cancer-mediated T cell exhaustion. Similar to T cell exhaustion in chronic infections, tumor-infiltrating CD8+ T cells display attenuated effector functions including impaired cytokine secretion and sustained high surface expression of inhibitory receptors (PD-1, CTLA-4, Tim-3, LAG-3, and others) (76, 149–151). However, exhausted T cells in cancer show subtle differences in their gene expression profiles from infection mediated T cell exhaustion. For example, tumor-specific CD8+ T cells derived from a late stage melanoma cancer model showed overexpression of several genes involved in cell cycle regulation, DNA repair and immune responses which was comparatively different from gene expression profiles derived from EBV-specific and CMV-specific exhausted CD8+ T cells (152). These differentially expressed genes were related to inhibitory receptors. Accordingly, CD160 and several other inhibitory receptors were not co-expressed in tumor-specific exhausted CD8+ T cells compared to virus-specific exhausted CD8+ T cells. Some inhibitory receptors like BTLA are upregulated in exhausted tumor-specific CD8+ T cells and not in exhausted virus-specific CD8+ T cells (76). These distinct gene expression profiles of multiple inhibitory receptors suggest different underlying mechanisms governing receptor upregulation in chronic viral infections and cancer mediated exhaustion. As such, the differential expression of inhibitory receptors may shape the extent of T cell exhaustion in each scenario and provide a molecular signature that will help to diagnose diseases.

### Tumor Microenvironment

Despite several overlapping functional and phenotypic features found in exhausted T cells induced by chronic viral infections or cancer, the progression of cancer mediated T cell exhaustion is not fully understood. This is partly because of the complexity presented by the tumor microenvironment. The surrounding environment of a developing tumor is comprised of stroma (containing fibroblasts, immune cells, and extracellular matrix) (77), blood vessels, infiltrating inflammatory cells and a number of cells associated with host tissues (Figure 1). The cellular environment inside the tumor is not homogenous throughout the cancer (153–155). Hence, the tumor microenvironment (TME) is continuously evolving with tumor progression. Tumor-infiltrating lymphocytes, such as cytotoxic and regulatory T, B and natural killer (NK) cells, associated M2 macrophages (TAM) (156, 157), infiltrating dendritic cells...
changes on cancer cells and abrogated IFN presentation or downregulation of PD-L1 driven by epigenetic immunoeediting are the loss of tumor-associated antigen (TAA) immunotherapy treatments in some clinical settings (162). Thus, decision-making, which has been poorly discussed in the context their nature and can trigger unique signaling pathways for cellular molecular forces at play during these encounters differ greatly in

The biophysics of T cell-APC encounter is highly dynamic. The receptor landscapes on both sides of the immunological synapse.

impaired T cell responses would follow changes in the immune landscape could therefore abort T cell responses and disrupt host immune regulatory mechanisms. In this section we highlight the studies that attempt to unravel the link between changes in biophysical properties of the membrane and T cell dysfunction.

Immunoeediting comprises complex adaptive mechanisms where cancer reduce its immunogenicity to evade recognition and destruction of selected clones (162, 165). Examples of immunoeediting are the loss of tumor-associated antigen (TAA) presentation or downregulation of PD-L1 driven by epigenetic changes on cancer cells and abrogated IFNγ—a key regulator of antigen process and presentation—delivered by tumor-infiltrating lymphocytes (TIL) in the TME (166–169), which can lead to incomplete elimination and persistence of adapted tumors becoming clinically evident (169). Insufficient TAA presentation poses a challenge for adaptive cytotoxicity by driving immunological ignorance (170, 171). The poor immunogenicity of transformed cells that escaped recognition can further promote insufficient activation of T cells, evident by unsuccessful immunotherapy treatments in some clinical settings (162). Thus, the selective advantage acquired by evasive cancer cells with impaired T cell responses would follow changes in the immune receptor landscapes on both sides of the immunological synapse. The biophysics of T cell-APC encounter is highly dynamic. The molecular forces at play during these encounters differ greatly in their nature and can trigger unique signaling pathways for cellular decision-making, which has been poorly discussed in the context of T cell dysfunctions.

**BIOPHYSICAL LANDSCAPE OF DYSFUNCTIONAL T CELLS**

T cell signaling and the cascade of events that follow T cell activation require close encounter of two cell membranes: the plasma membrane of the T cell and the APC. The dynamic and heterogeneous nature of the membrane environment, composed of different types of lipids, receptors and ligands makes the process of T cell-APC conjugation complex to understand. The close contact between the two opposing membranes is accompanied by the formation of unique structural features that promote information transfer through receptor–ligand interactions. The interface between a T cell and an APC is known as the immune synapse and its formation involves spatial redistribution of surface receptors and ligands to facilitate the initiation of immune responses (172). T cells continuously form membrane protrusions known as filopodia or microvilli which help them probe the surface of APCs and to sense biophysical properties in the surrounding environment (Figure 2A). Formation of these structural features require extensive membrane remodelling assisted by cytoskeleton rearrangements (178). Moreover, due to relatively high cell motility and relatively slow diffusion rates of engaged receptors and ligands, both cells experience pulling-pushing and shear forces. Recent studies have attempted to quantify these mechanical forces during immune synapse formation (179, 180). It is generally thought that these forces enable T cells to probe the surrounding environment and execute effector functions at optimal levels. Changes to this biophysical landscape could therefore abort T cell responses and disrupt host immune regulatory mechanisms. In this section we highlight the studies that attempt to unravel the link between changes in biophysical properties of the membrane and T cell dysfunction.

**Structure and Functions of Immune Synapses**

The immunological synapse is crucial for T cell activation and is sometimes referred to as an activation synapse. Immune receptors, signaling molecules, cytoskeletal components and cell organelles all participate in the formation of the immune synapse (181). In cytotoxic T cells (CTLs), the immune synapse is also the interface in which cytolytic granules are delivered to the target cells and in these cases they have also been referred to as cytotoxic synapses or lytic synapses (182).

**Immunological Synapses for Activation**

When forming activation synapses, key signaling molecules congregate to form a distinct sub-synaptic domain known as the supramolecular activation cluster (SMAC) (Figure 2B). The central region of SMAC (cSMAC) primarily contains TCRs and tyrosine kinases which are crucial for the initiation of TCR signaling. The peripheral SMAC (pSMAC) surrounds the cSMAC and contains integrins like LFA-1 which binds to ICAM-1 expressed on APCs, facilitating the adhesion of T cell to APC. Under some circumstances, T cell activation is achieved via TCR microclusters and without the need of classic immune synapse formation (183, 184). It has also been proposed that the cSMAC serves as the site for signal termination and receptor recycling (185). One of the main functions of activation synapse is the initiation and amplification of TCR signaling. Upon TCR
binding to its cognate pMHC, a cascade of signaling events take place leading to T cell activation, proliferation and execution of effector functions. TCR-pMHC ligation also triggers substantial structural alterations in the membrane (186). The function of filopodia is broad including crucial roles in cell-cell adhesion and cell migration (177). Immune synapses. Formation of immune synapses are important steps in T cell activation and executing T cell effector functions through cytotoxicity. When forming activation synapses, signaling receptors (in T cells) and ligands (in APCs) spatially segregate into a bull’s eye-like structure forming the supra-molecular activation cluster that is separated into central (cSMAC—red), peripheral (pSMAC—yellow) and distal (dSMAC—green) regions, where each zone preferentially recruit different signaling receptors to initiate T cell signaling. Cytotoxic synapses are formed when a CTL encounters a target cell. Cytotoxic synapses differ from activation synapses in the recruitment of lytic granules to the synaptic cleft with the help of Golgi apparatus and the microtubule organising centre (MTOC). Delivery of lytic granules and exocytosis of granule contents are highly depended on calcium influx. Mechanosensing. T cells are constantly being subjected to mechanical stresses when undergoing kinapse formation with APCs. TCRs display mechanosensing properties by exerting pulling-pushing and shear forces on pMHC molecules on APCs. When a TCR engage with cognate-antigen peptide, a catch bond is formed, which triggers conformational changes in the TCR-CD3 complex and initiate T cell signaling. Conversely, TCR encounter with a non-cognate-antigen peptide results in a slip bond, where an exponential decay of bond lifetime is observed with increasing force. Slip bonds fail to trigger TCR signaling.

The Vav family of proteins are involved in modulating these cytoskeletal changes at the immune synapse. As shown by previous studies, Vav1 mediates downstream signaling in T cells via PLCγ1 and TCR-induced calcium flux (189–191). The absence of Vav1 affects the stability of the TCR signaling clusters and impair both calcium flux and MAP kinase phosphorylation (192). By activating RHO GTPases such as RAC1 and CDC42 (193), Vav1 is implicated in series of events facilitating Wiskott-Aldrich syndrome protein (WASP) and WASP-family verprolin-homologous protein-2 (WAVE2) to activate actin-related protein 2 and 3 (ARP2/3), which leads to polymerisation and

**FIGURE 2** T cell biophysical landscape. (A) Membrane protrusions. T cells continuously form actin-rich membrane protrusions known as filopodia or microvilli which help them to sense biophysical properties in the surrounding environment. Microvilli are involved in early T cell activation (173) and where signaling components including TCR and TCR-associated signaling molecules get accumulated (174, 175). The tip of a microvilli is zoomed in to illustrate the accumulation of T cell signaling molecules. Compared to microvilli, filopodium membrane projections are larger in size ranging from 10-40 µm in length in different cells (176). The function of filopodia is broad including crucial roles in cell-cell adhesion and cell migration (177). (B) Immune synapses. Formation of immune synapses are important steps in T cell activation and executing T cell effector functions through cytotoxicity. When forming activation synapses, signaling receptors (in T cells) and ligands (in APCs) spatially segregate into a bull’s eye-like structure forming the supra-molecular activation cluster that is separated into central (cSMAC—red), peripheral (pSMAC—yellow) and distal (dSMAC—green) regions, where each zone preferentially recruit different signaling receptors to initiate T cell signaling. Cytotoxic synapses are formed when a CTL encounters a target cell. Cytotoxic synapses differ from activation synapses in the recruitment of lytic granules to the synaptic cleft with the help of Golgi apparatus and the microtubule organising centre (MTOC). Delivery of lytic granules and exocytosis of granule contents are highly depended on calcium influx. (C) Mechanosensing. T cells are constantly being subjected to mechanical stresses when undergoing kinapse formation with APCs. TCRs display mechanosensing properties by exerting pulling-pushing and shear forces on pMHC molecules on APCs. When a TCR engage with cognate-antigen peptide, a catch bond is formed, which triggers conformational changes in the TCR-CD3 complex and initiate T cell signaling. Conversely, TCR encounter with a non-cognate-antigen peptide results in a slip bond, where an exponential decay of bond lifetime is observed with increasing force. Slip bonds fail to trigger TCR signaling.
accumulation of actin filaments at the immune synapses (194, 195).

Cytotoxic Immune Synapses
Cytotoxic synapses are crucial in executing T cell effector functions. One important difference between a cytotoxic synapse and an activation synapse is the recruitment of lytic granules to the synaptic cleft (Figure 2B). CTLs exert their cytotoxicity by first binding to the target cell and then releasing lytic granules containing perforin and granzymes via exocytosis, and finally detaching from the target cell (196). Similar to activation synapses, attachment to the target cell is primarily mediated by LFA-1 which also aids the formation of SMAC (197). Importantly, the pSMAC has been implicated in stabilising the cytotoxic synapses in CTLs, as the disruption of pSMAC formation results in impaired target cell lysis (198). During CTL mediated target cell lysis, granules containing cytotoxic enzymes (lytic granules) are recruited to the immune synapse with the help of the Golgi apparatus and the microtubule organising centre (MTOC). In NK cells, it was shown that dynein, a cytoskeletal motor protein is responsible for the transport of lytic granules to the MTOC and then MTOC polarises to deliver lytic granules to the synaptic cleft (199). Targeted delivery of lytic granules and exocytosis of granule contents are highly depended on calcium influx (200). Following the detachment from the target cell, CTLs are capable of effectively killing multiple targets sequentially (201).

Immune Synapse Dysfunctions in Chronic Diseases
Chronic diseases are often associated with the phenotype of T cell exhaustion. The root cause for a number of chronic diseases stems from the inability of T cells to form functional immune synapses with the target cells, leading to impaired T cell activation resulting suboptimal immune responses (202, 203). Understanding the mechanisms that induce impairments in immune synapse formation is an important step in developing effective therapeutics that can reverse T cell exhaustion by restoring effector functions.

Disruption of T cell and target cell contact during T cell activation or T cell-mediated cytotoxicity can impair the formation of functional immune synapses. This can be detrimental in terms of maintaining immunity against pathogens and cellular malignancies. Leukocyte adhesion disorder (LAD) is a classic example of a chronic disease caused by defective expression (LAD type-I) or activation (LAD type-III) of cell adhesion molecules, primarily β-2 integrins like LFA-1 (204). Since LFA-1 plays a crucial role in the assembly of immune synapses, LAD patients often have recurrent bacterial infections due to their compromised immune system (205). Similarly, defects in WASP family of proteins directly affect actin mediated cytoskeletal rearrangement during immune synapse formation. This was shown in Wiskott-Aldrich Syndrome where CTLs lose their cytotoxicity (206). In a rare type of non-Hodgkin lymphoma known as anaplastic large cell lymphoma (ALCL), it has been shown that WASP and WASP-interacting-protein (WIP) are expressed in low amounts (207). In WASP knockout (KO) mice, fast onset of tumor growth has been observed (208). In the same study, the metastatic rate of B16 melanoma was shown to be higher, indicating an overall loss of T cell tolerance towards the cancer. However, somewhat contradictory observations were made in mouse breast carcinoma, where the metastatic spread was decreased in the absence of WASP (209), suggesting differing roles for WASP in cancer progression depending on the cancer model. Interestingly, addition of exogenous IL-2 was able to rescue the cytotoxicity of WASP KO NK cells by restoring their ability to form immune synapses (210, 211). In fact, IL-2 treatment is commonly used as an immunotherapy treatment in the attempt to promote T cell proliferation and restore or enhance T cell effector functions (212). WASP is one of many proteins in which irregular protein expression can lead to immune dysfunctions because of cytoskeletal organisation defects during immune synapse formation. A growing number of putative proteins including Dock8, RAC2, RHOH, CORO1A, ACTB and many others have been implicated in modulating actin-dependent cytoskeleton organisation to promote efficient T cell activation. Their individual functions have been reviewed elsewhere (213).

Failure to deliver lytic granules to the synaptic cleft leave CTLs with impaired cytotoxicity and reduced pathogen clearance. The continual stimulation from the innate immune system together with dysfunctional adaptive immune responses can result in systemic inflammation which is detrimental to the host homeostasis. This is readily observed in herpes viral infections, particularly Epstein–Barr virus (EBV) and cytomegalovirus (CMV). These viruses with their lifelong latency in the host may result in persistent antigenic-stimulation mediated by the innate immune system (214). This has been linked to hemophagocytic lymphohistiocytosis (HLH), a life-threatening syndrome presented with attenuated killing capacity of T cells and NK cells (215).

In cancer models, much of the evidence for defects in the formation of immune synapses comes from haematological malignancies (15, 216, 217). For instance, one report demonstrated that CD8+ and CD4+ T cells are unable to form proper immune synapses with chronic lymphocytic leukemia (CLL) cells, which hindered anti-tumor activity (15). The authors show that when a healthy T cell encounters CLL-B cells, F-actin polymerisation was suppressed and there was impaired recruitment of key adhesion and signaling molecules to the immune synapse of the T cells (15). These observations agree with other cancer models where tumor-infiltrating lymphocytes (TIL) showed similar defects in actin polymerisation (218, 219). The exact mechanism behind tumor-induced immune synapse defects in T cells is not yet clear. However, immuno-modulatory drugs such as lenalidomide are shown to be effective in reversing actin polymerisation defects in patients with follicular lymphoma (216).

Human immunodeficiency virus type-1 (HIV-1) also induces signaling dysfunctions in CD4+ T cells as a part of its viral pathogenesis. The abundantly expressed viral protein Nef plays a central role in impairing the immune synapse formation in HIV-
1 infected T cells (220). Nef achieves this by hijacking the host membrane protein trafficking machinery to promote the spread of infection (221). HIV-1 infected T cells showed poor cell spreading, suggesting a Nef-dependent inhibition of actin polymerisation. In parallel, a reduced recruitment of TCR-CD3 complex, Lck and other actin polymerisation-related proteins to the immune synapses was observed (222). Interestingly, Nef sequesters Lck away from TCR-CD3 complex, in both the presence or absence of CD4, and slows down TCR internalisation (220), thereby arresting TCR recycling and downregulation following TCR-pMHC ligation. This leads to accumulation of TCRs on the cell surface, resulting in T cell hyperactivation which is readily observed in untreated HIV-infection (223, 224). Previous reports also show that Nef has downstream effects on transcription factors like NFAT and NF-kB which are important to execute T cell mediated immune responses (225, 226).

From T cell-APC conjugation to T cell activation and cytolytic granule trafficking to targeted cytotoxicity, these chronic diseases highlight the importance of each stage in immune synapse formation for the execution of optimal T cell immune responses.

**T Cell Mechanosensing**

The initial contact between T cell and APC demonstrated by TCR binding to its cognate antigen-peptide triggers downstream signaling events that would activate T cells to efficiently execute their effector functions. However, the outcome of TCR signaling is largely impacted by mechanical forces applied to the TCR-pMHC complex (227, 228). Essentially, exogenous forces applied to TCR-pMHC interactions are transmitted into the cell as biochemical signals through mechanotransduction, the process which describes how physical perturbation experienced by receptors are translated into chemical signals (229). Conversely, biochemical signals generated by the cell is being translated into mechanical forces that are exerted on the surrounding environment.

Attempts to *ex vivo* activate and expand T cells utilising soluble anti-CD3 antibodies have largely failed (228, 230–232). TCR triggering *ex vivo* can be achieved by immobilising CD3 complex activating antibodies on rigid surfaces such as beads or tissue culture plates, as evidenced by the increased Ca^{2+} influx and phosphorylation of ZAP70 in T cells (233, 234). One explanation for why surface attached antibodies induce activation but those in solution do not is that surface association mimics mechanical forces created by the movements of synaptic membranes on TCR-pMHC complexes and suggests that TCR signaling cannot be initiated unless pulling-pushing stresses or shear forces are applied to the complex (Figure 2C) (234, 235).

Early studies showed that mechanical forces at the piconewton (pN) range applied through pMHC-coated beads were enough to induce Ca^{2+} influx and ERK phosphorylation in T cells (236). However, when similar forces were applied to CD28, CD62L, or ICAM-2 no significant increase in Ca^{2+} influx was observed (234). Since these mechanical forces involving antigen recognition by T cells operate at pN range, it poses technical challenges when elucidating them in biological systems. Recently, using biomembrane force probe (BFP) a number of studies have shown the threshold for cognate TCR-pMHC interaction to be at the scale of ~10 pN (237, 238). In addition to TCRs, other mechanosensors such as Piezo1 contributes to optimal T cell signaling (232). However, the extent of Piezo1 involvement in TCR signaling and T cell responses is yet to be elucidated. As the mechanical forces applied on TCR-pMHC require some level of rigidity from both biological supports (i.e. membranes), the stiffness of APC is expected to influence T cell responses. Stimulating substrates with anti-CD3 and anti-CD28 antibodies on supports with relatively high rigidity drive greater productions of IFN-γ, TNF-α and IL-2, up-regulation of activation markers and proliferative capacity compared with softer substrates (180, 228, 239–241). Furthermore, CTLs increase the stiffness of APC by stretching the synaptic region to modulate the speed of the perforin pore formation and consequently promote faster target cell lysis (242). These studies highlight the importance of fine tuning the rigidity of stimulating cultures for optimal T cell response which has implications in adoptive T cell immunotherapies (239, 240, 243, 244). One may also question whether subtle mechanical changes in TCR-pMHC affinity can trigger specific signal transduction pathways and affect downstream T cell responses.

Forces applied on TCR-pMHC complex can affect their bond lifetime in unexpected ways. For instance, catch bonds, where pulling forces applied to the bond, increases its bond lifetime (245), have been described for several receptor–ligand interactions (Figure 2C) (246, 247) and recently also for TCR-agonistic peptide MHC interactions (237, 238, 248). In contrast, antagonistic peptides form slip bonds characterized by short lifetimes (Figure 2C) (237). Independent reports using BFP (237, 238, 249, 250) and optical tweezers (248) on cell systems that express transgenic TCRs and cell-free experiments also using optical tweezers (248, 251) have demonstrated that while agonistic-peptides can increase TCR-pMHC binding lifetime, antagonistic-peptides tends to reduce it. These studies further demonstrated that catch bonds reach their maximum lifetime under a mechanical force in the range of ~10 pN and the lifetime of slip bonds decreases exponentially with increased mechanical force.

Another crucial aspect of T cell mechanosensing is to understand how TCR and cognate-pMHC binding events get translated into biochemical signals in T cells that are specific to the antigenic peptide. One theory suggests a conformational change of the TCR-CD3 complex during pMHC binding that would dislodge and release the cytoplasmic tails of CD3 from the inner leafllet of the plasma membrane. This would expose immunoreceptor tyrosine-based activation motifs (ITAMs) to get phosphorylated by tyrosine kinases Lck and Fyn. Accordingly, a TCR specific antigenic peptide would expose the cytoplasmic tails of CD3 for a longer period and permit more efficient phosphorylation by tyrosine kinases to generate a stronger signal to transduce (252, 253). A theory based on kinetic segregation model explains a local disruption of the kinase-phosphatase balance during TCR-pMHC binding is sufficient
to generate a productive signal as TCRs get phosphorylated by the segregation of phosphatase like CD45 (254, 255). Here, catch bonds (formed with cognate ligands) or slip bonds (formed with non-cognate ligands) formed during TCR-pMHC interactions (Figure 2C) may generate differential segregation patterns that would then be translated to a strong or weak signal, respectively (250).

In summary, regulating mechanosensing capacity of TCR is crucial to recognise and translate mechanical cues into cell signals during T cell activation and in execution of immune responses.

**T Cell Mechanosensing in Chronic Diseases**

T cells constantly patrol and migrate to different tissue compartments in search of cognate-antigens. This tissue migration involves continuous changes in T cell morphology driven by actin polymerisation which impose considerable mechanical force at the cellular level. During the contact between T cell and APC, the role played by mechanical forces in mediating immune responses are now becoming clear. The highly dynamic interactions between the extracellular matrix (ECM) and actin cytoskeleton is directly linked with translating mechanical cues from the environment into cell signals. Substrate stiffness is a mechanical cue that is implied to regulate number of cellular functions including proliferation, migration and differentiation (256–258). T cells are exposed to a range of substrate stiffnesses during their lifespan as stiffness values change substantially in different cells that T cells encounter. For example, while skeletal muscles have a stiffness in the range of ~10 kPa (259), elastic modulus of human bones may vary from 7–25 GPa (260).

A number of studies reported that reduced stiffness in cancer cells as a mechanism of promoting their growth independent of ECM stiffness (261, 262). Since optimal T cell responses require surfaces or biological membranes with relatively high rigidity, by reducing surface stiffness, cancer cells can effectively evade immune detection and subsequent cytotoxicity. Concurrently, a local increase in ECM stiffness is associated with disease progression (263). It has been reported that cancer cells are able to modify their surrounding ECM stiffness in order to promote metastasis (264). In fact, ECM associated adhesion proteins are known to play a vital role in different stages of cancer metastasis which overall influence the invasiveness of a cancer (265). In some cases, cancer cells are shown to synthesise their own ECM proteins to promote metastasis (266). Additionally, ECM stiffness influence the outcome of desmoplastic response (i.e. pervasive growth of dense collagen stroma around a tumor) associated with tumors (267). For instance, desmoplasia in pancreatic and breast cancer promote tumor progression and results in poor prognosis (268, 269). In mammary tumors, lysyl oxidase enzyme is linked to remodelling and increasing ECM stiffness as the inhibition of this enzyme reduced tissue stiffening and delayed tumor progression (270). Several studies have demonstrated a correlation between collagen density, a primary component of the ECM, and the infiltrative capacity of T cells into tumor islets (271–273). Densely packed collagen fibres are suggested to obstruct T cell entry into the tumor microenvironment, and overall reduce their proliferative and cytotoxic capacity (273). Recently, a study using confocal microscopy coupled with optical tweezers was able to track changes in biophysical properties of cancer cells in a multicellular 3D breast cancer model (274). The study was able to identify a stiffness gradient decreasing outward from the core of the growing tumor, suggesting cancer cells with softer biophysical characteristics are likely to be located at the periphery of the tumor. Moreover, this also implies that T cell mediated cytotoxicity become less efficient at the edge of the tumor, thereby increasing the invasiveness of the tumor at the periphery. Whether modulation of cell stiffness is a reliant mechanism for immune evasion during chronic infections is yet to be determined. Overall, understanding T cell responses to mechanical cues such as substrate stiffness may become crucial to understand the biophysical landscape of exhausted T cells.

**TCR Diversity in Immune Responses**

In adaptive immunity, the engagement of TCR with pMHC molecules plays a pivotal role in shaping the overall immune responses against foreign pathogens, malignancies and allergens. When TCRs recognise and bind to their cognate antigen, it triggers intracellular signaling pathways that activate the expression of multiple genes linked to several effector functions in T cells. Hence, the quality and magnitude of T cell effector functions are linked to the strength and quality of TCR-pMHC interactions. Primarily, the strength of these interactions are measured by TCR affinity to its antigen (275). T cell signaling is a complex function that involves the affinity of TCR-pMHC interactions, coreceptor binding and co-stimulatory and co-inhibitory signal integration (276), but in general, affinity of TCR-pMHC can predict the sensitivity of a T cell to a specific antigen (277). TCR affinity also dictates selective polyclonal expansion of antigen-specific T cells during immune responses.

Rearrangement within the variable regions of the TCR and thymic selection generates an immune repertoire of T cells with differing antigen specificities. During an infection, the expansion of T cell clones specific to a small number of immunodominant antigens can skew the immune repertoire (278). This form of TCR bias can be influenced by multiple factors ranging from thymic selection to initial immune response to an antigen (278). Overall, the adaptive immune system is shown to maintain a diversified population of antigen-specific T cell clones with varying affinities which possess the capacity to clonally expand and form memory T cells. In some cases a bias T cell clonal expansion, either towards high or low affinity has been observed (279, 280). During the acute phase of an infection, early models of clonal selection have shown that antigen-specific T cells with higher affinity are selectively expanded from the polyclonal T cell population to mediate immune responses and proceed to become memory T cells to retain acquired immunity (281–283). This selective enrichment of antigen-specific high-affinity clones has been described as a form of affinity maturation of T cells (284). During persistent antigen exposure, however, the profile of memory T cell clones shifts towards a low-affinity repertoire (16, 279, 285). Hence, distinct affinity profiles for antigen-specific T cells are generated and maintained during
acute and chronic phase of an infection. Elucidating the underline mechanisms behind this differential clonal expansion under different phases of antigen exposure proven to be beneficial in developing therapeutic interventions aimed at restoring T cell immune responses in chronic diseases.

Measuring TCR Affinity

TCR affinity to its cognate antigen-MHC complex is generally reported as the ratio of $k_{on}$ and $k_{off}$ rates, $K_D$: the equilibrium dissociation constant. In simple terms TCR-pMHC interactions with low $K_D$ values (i.e. high affinity interactions) are typically associated with longer binding dwell times. Due to rapid dissociation (high $k_{off}$) of TCR-pMHC complex, it is often difficult to determine the affinity of TCR-pMHC interactions using conventional kinetic measurements. Generally, TCR-pMHC affinity is in the scale of micromolar range (1 to 300 µM), albeit with considerable variability (275, 286). Among several approaches which have been useful in characterising TCR-pMHC binding affinity, surface plasmon resonance (SPR), two-dimensional micropipette adhesion frequency assay (2D-MP) (287, 288) and pMHC multimer-binding have demonstrated their wide applicability in physiological settings (Figure 3).

The usage of SPR to determine TCR-pMHC affinity dates back three decades (290, 291). SPR measures a signal that correlates to a change in mass on a sensor-surface (i.e. sensor chip) where the binding partner, in this case pMHC molecules are immobilized on the sensor-surface and TCR molecules flown over to bind (Figure 3A). SPR offers much lower sample requirement and versatility over earlier techniques like isothermal titration calorimetry (ITC), but with some drawbacks which have been reviewed elsewhere (292, 293). While SPR provides the means of directly measuring TCR-pMHC binding affinity, in physiological settings, the TCR is also attached to a surface where several other receptor-ligand interactions would take place simultaneously. Moreover, SPR would not account for the forces introduced with bystander TCRs, auxiliary receptors, cell adhesion molecules and membrane fluctuations which overall can modify TCR-pMHC interactions.

**FIGURE 3** | Measuring TCR affinity. (A) Surface plasmon resonance (SPR). SPR measures the equilibrium dissociation constant ($K_D$) of TCR-pMHC interactions in which pMHC is immobilized on a sensor surface and TCR molecules are injected in a continuous flow. Binding of TCR to pMHC results in a change of mass on the sensor surface and is recorded in a sensogram which is then used to calculate $K_D$. (B) Micropipette adhesion assay. This technique uses two probes, one that is stationary which contains a red blood cell (RBC) attached to a functionalized glass bead to act as the adhesion force transducer and a mobile force probe bearing a T cell coupled to a piezotranslator. During adhesion-retraction cycles carried out by the mobile probe, the deformation of the RBC, displacement of the glass bead and the force generated in each cycle is recorded. (C) Multimer staining. This technique enhances the binding avidity of TCR-pMHC by increasing the valency of the interaction, results in more stable multimeric TCR-pMHC complexes for efficient labelling and detection. To date numerous forms of pMHC multimers have been reported which includes tetramers, pentamers, octamers, and dextramers (289).
binding affinity. Hence, measuring functional TCR-pMHC binding affinity in native cell membrane environment would be more physiologically relevant (288).

When predicting TCR-pMHC affinity in the context of T cell-APC interactions, the 2D micropipette adhesion assay (2D-MP) technique has proven to be useful. In 2D-MP, a human red blood cell (RBC) decorated with the ligand of interest acts as a sensor for measuring adhesion kinetics with the cell of interest expressing the cognate receptor (Figure 3B) (294). A micromanipulation device is used to bring these two cells into close proximity, in a tightly controlled environment enabling receptor–ligand binding. These binding events are captured as the degree of deformation of RBC membrane when the T cell is pulled away. These adhesion cycles were repeated and then translated into a binding curve which allows the calculation of binding kinetics for a given receptor–ligand interaction in two-dimensional space. By measuring affinity in the native membrane environment of a receptor, this method provides more physiologically relevant binding kinetics that have more applicability in cell biology. A modified version of 2D-MP known as the fluorescence biomembrane force probe (fBFP) uses osmolarity adjusted human red blood cell attached to a functionalized glass bead to act as the adhesion force transducer (Figure 3B) (237). This technique combined with single-molecule force spectroscopy and fluorescence microscopy enables the measurement of singular receptor–ligand binding event kinetics. Both 2D-MP and fBFP have measured much faster off-rates for TCR-pMHC interactions [30–8,300 fold faster (287)] than reported by 3D kinetic measurements derived from SPR. It should be noted that rapid 2D off-rates obtained for TCR-pMHC do not necessarily correlate with the rapid off-rates in 3D, indicating kinetics of TCR-pMHC interactions including off-rates and antigen-peptide affinity differ substantially from 2D to 3D space. A major drawback of these techniques is their requirement for highly specialized equipment to measure cellular level kinetics. This limits their usage in predicting population level kinetics during T cell-APC interactions.

Multimers of pMHC are the most commonly used method in identifying antigen-specific T cells from a polyclonal population. Due to their low affinity, monomeric pMHC are ineffective as a labelling probe in detecting antigen-specific T cell clones from a pool of other T cells. Multimer technology overcomes this by increasing the valency of TCR-pMHC interaction by multimerizing pMHC complexes to increasing avidity, which results in more stable multimeric TCR-pMHC complexes for efficient labelling and detection (295). The pMHC multimers can be in the form of tetramers, pentamers or octamers (Figure 3C) (296, 297). Recently, multimer labelling has been shown to introduce biases towards detection of high-affinity TCR-pMHC interactions and underestimation of interactions with low-affinity. This may distort the overall view of antigen-specific T cell diversity in a polyclonal population (298). Therefore, multimer binding intensity does not necessarily correlate with the functional responses produced by antigen-specific T cell population (299, 300).

It is evident that each affinity measurement technique in isolation overlooks the clonal diversity of T cell immune repertoire which is crucial to understand the full extent of immune responses during a disease. When combined, these techniques would resolve the shortcomings of affinity biases in predicting antigen-specific polyclonal T cell diversity.

T Cell Affinity Repertoire in Chronic Diseases

TCR affinity and signaling strength in response to a specific antigen sets the threshold for clonal selection to execute immune responses. Based on the premise that high-affinity TCR-pMHC interactions leads to efficient T cell activation, high-affinity T cell clones have a selective advantage over other clonotypes in mediating primary and secondary immune responses during the acute phase of a disease (281, 283, 301). This observed lack of affinity diversity becomes reduced when the disease progresses into a chronic phase. Mounting evidence suggest that this enhanced diversity in T cell clonal affinity is due to the recruitment of low-affinity TCR expressing T cells in the immune repertoire (280, 302–304). This distribution pattern of affinity clones of antigen-specific T cells can differ between disease models, and how this diversity is maintained to produce life-long immune responses is still under investigation. Moreover, screening affinity diversity in the T cell immune repertoire is challenging as current detection methods are suboptimal in identifying the full breadth of clonal diversity. Excluding low-affinity clones from the measured T cell repertoire will underestimate the full capacity of functional responses exerted by the immune system during chronic diseases.

A number of studies have demonstrated the effectiveness of low-affinity antigen-specific T cell clones in mediating immune responses, combating infections and preventing tumor progression similar to high-affinity counterparts (279, 280, 300, 305). Based on CD4+ T cell responses to six different LCMV-antigens, Martinez et al. reported limited correlation between TCR affinity and dominance in clonal expansion (280). Moreover, both high and low-affinity T cell clones possess similar proliferative capacity (306, 307) and phenotypic characteristics (302) which all together challenge the prerequisite of high-affinity TCR-pMHC interaction dominance in driving clonal expansion and mediating immune responses. However, the strength of TCR-pMHC ligation may determine the magnitude of clonal expansion and the onset of contraction phase. This was demonstrated in Listeria monocytogenes infection model using altered peptide ligands with varying affinities (302). Despite the observed similarities in rapid proliferation rates in low and high-affinity antigen-specific T cell clones, Zehn et al. showed that weaker ligand interactions lead to early onset of contraction of T cell proliferation (302). In the same study, early appearance of low-affinity T cell clones in the blood stream after antigen-stimulation suggest a role for TCR-pMHC affinity in modulating the kinetics of T cell migration. Moreover, the reduced memory T cell expansion during successive challenge of a weak ligand/antigen indicates a correlation with the strength of recall stimulus and memory T cell responses.

Despite numerous attempts undertaken to predict the affinity diversity of T cells in different disease models, most clonal diversity observations of immune responsive T cells come from unrelated models of acute and chronic diseases. It would be highly
relevant to demonstrate the evolutionary trajectory of antigen-specific T cell affinity using longitudinal observations during acute and chronic phase of the same disease model. Using two LCMV infection models, Andargachew et al. showed that the overall affinity diversity of CD4+ T cell clones were similarly maintained throughout acute and chronic antigen exposure along with effector and memory T cells showing similar affinity distribution patterns in both phases (308). Their study accompanied 2D-affinity kinetic measurements derived from 2D-MP assay. During the transition of effector to early memory CD4+ T cells, both acute and chronic LCMV infection showed an increased functional avidity. However, the half-maximal effective concentration (EC_{50}) for IFN-γ and IL-2 production was much lower for acute-LCMV CD4+ T cells compared to chronic exhausted CD4+ T cells, which indicates a higher antigen sensitivity and functional avidity for CD4+ T cells in the acute phase of LCMV infection. This study also drew parallels between chronic and acute-LCMV with their selective recruitment of low-affinity T cell clones into the immune repertoire.

More recently, a longitudinal study used memory T cell inflation to illustrate the evolutionary trajectory of cytomegalovirus (CMV)-specific CD8+ effector memory T cell affinity during acute and chronic phase antigen exposure (16). T cell inflation is described as the atypical accumulation of memory T cells in blood and peripheral tissues in response to persistent low-level antigen exposure (309). Using both human and mouse CMV models, Schober et al. analyzed TCR affinity distribution among the CD8+ T cell population. The TCR-pMHC dissociation rate (k_{off}) measurements obtained from real-time fluorescence microscopy (310) conclusively demonstrated that T cells with lower TCR affinity were enriched in the inflationary CD8+ T cell pool compared to the acute phase. Moreover, in-depth analysis of CMV-specific TCR repertoire obtained from the mouse model showed clones underrepresented in the acute phase of the infection (medium to low-affinity clones) were recruited at higher proportions to the immune repertoire at later stages via clonal succession (311). The authors also suggest that recruitment of low-affinity T cell clones compensated the loss of functional avidity provided by high-affinity clones which become senescent at late stages of the infection. This form of “reversed-affinity maturation” could be an important adaptation of the immune system to maintain life-long effective T cell responses against persistent viral infections which particularly exhibit low antigen expression levels. In the long-run, selective expansion of low-affinity T cell clones may provide an effective strategy in generating lasting pathogen control along with reduced immunopathology.

Whether the above mechanisms prove to be effective in regulating clonal diversity in T cell exhaustion related chronic diseases remains to be explored. T cells with high-functional avidity often exhibit exhaustion phenotypes under high levels of antigen exposure (312, 313). This can increase the probability of pathogen escape from the immune system, leaving low-affinity clones of antigen-specific T cells to take up the task of pathogen clearance or maintain a life-long host-pathogen equilibrium with reduced immunopathology. It should be noted that low-affinity T cell clones can also become functionally exhausted as reported by several tumor models (314, 315).

The affinity repertoire of effector T cells during cancer is less well understood than in chronic infections. Previous studies suggest that TILs with high-affinity are more likely show exhaustion markers albeit having superior control in eliminating tumor cells compared to their low-avidity counterparts (312, 316, 317). A growing number of studies have recognized a distinct role for low-avidity TILs in tumor clearance. Studies have shown low-avidity TCR interactions with tumor antigens activate tumor-specific T cells in a similar manner to high-avidity TCR interactions with the tumor antigen (317, 318). Moreover, with prolong exposure to the tumor, both clonotypes showed exhaustion markers including sustained upregulation of inhibitory receptors, with higher degree of exhaustion observed for high-avidity tumor-specific T cell clones (317). Another study using adoptive transfer of OT-I (high-avidity) and OT-3 (low-avidity) transgenic tumor-specific CD8+ T cells was able to demonstrate that OT-3 T cells were able to mediate tumor regression in pancreas with minimum autoimmunity, contrast to OT-1 T cells which in addition to the rapid eradication of the tumor, caused autoimmune diabetes in the mouse model (314). These studies suggest a necessary role for low-avidity tumor-specific T cells in anti-cancer immune responses. Thus, elucidating various mechanism underlying the expansion of T cell affinity repertoire during cancer progression is important to understand cancer immune surveillance and more complex immune-oncology concepts like immunoediting. Immunoediting of malignant cells can generate slightly variable neoantigen presented to the existing TCR repertoire in the TME causing decreases in the overall TCR affinities and T cell-cytolytic responses, and consequently tumor evasion (319–321).

**Other Factors Influencing T Cell Affinity Diversity**

Numerous other internal and external factors including the expression level of co-stimulatory/inhibitory receptors in T cells, co-stimulatory/inhibitory ligands on APC, and the dose and density of antigen presentation by APCs play a key role in shaping the functional avidity of antigen-specific T cell immune repertoire. For example, the CD27/CD70 mediated co-stimulation in T cells has been shown to lower the threshold of TCR activation to respond to low-affinity antigens, which promotes to generate a higher degree of memory T cell clonal diversity (322). Conversely, higher expression of B7-1 along with ICAM-1 and LFA-3 are linked to selectively enriching T cell clones with high functional avidity (323, 324). De novo expression of B7-1 by anti-myeloma cellular vaccines improves cytotoxicity and helper-dependent memory formation of subdominant CD8+ T cell clones by avoiding tolerogenic effects (320). The antigen density presented by APCs during T cell priming and during infections has been shown to influence the functional avidity of immune responsive T cell populations (325). For instance, higher antigen density on APCs can compensate for low-affinity TCR-pMHC interactions. In human melanoma model it was demonstrated that low antigen doses presented by dendritic cells (DCs) produce melan-A-specific CD8+ T cells with high functional avidity which had lower dependence on CD8 coreceptors (326). B cells in infection models like Friend virus (FV) are linked to efficient priming and subsequent expansion of T cells with low functional avidity,
overall diversifying CD4+ T cell immune repertoire (327). Moreover, the degree of B cell activation correlates with B cell mediated clonal expansion of low-avidity CD4+ T cells. Indeed, antigen specific B cells have long been speculated to drive clonal diversity in CD4+ T cells (328).

CONCLUSION

Dysfunctional T cells are distinct from effector and memory T cells based on their functionality, metabolic activity, and epigenetic makeup. Recent findings strengthen the link between dysfunctional T cells and the progression of chronic diseases, thus, unravelling potential mechanisms behind the functional impairment of T cells with changes to its immune receptor and biophysical landscape during disease progression. In T cell exhaustion, the sustained upregulation of inhibitory receptors becomes a key feature that modifies the immune receptor landscape of T cells. Hence, these receptors have become primary targets in developing checkpoint blockade therapies aimed at restoring effector functions in non-responsive T cells. Although this has shown much clinical success in managing the progression of chronic diseases, there remains to be several limitations which hinder its wide applicability. Acquired resistance is one of the emerging challenges faced by checkpoint blockade therapy and may be overcome by combinatorial therapeutic strategies. The effectiveness of these therapeutic approaches in rescuing terminally exhausted T cells remains to be explored.

Apart from alterations in immune receptors expression profile, understanding changes to the cellular physiology of T cells during disease progression has become increasingly relevant to elucidate factors that promote T cell dysfunctions. Throughout their lifetime T cells are subjected to a myriad of mechanical forces experienced during cell migration, cell–cell interactions or exerted by the surrounding ECM. It is now becoming clear that these forces have important roles in T cell activation and the resultant effector functions, and may also play a central role in T cell dysfunctions. The concept of mechanotransduction as a mechanism of regulating cell behavior and function is not new (329), however, to understand this process at a subcellular level by means of measuring these infinitesimal forces require hypersensitive tools. As highlighted in this review, we have discussed the application of biophysical tools that can measure the strength of TCR-pMHC interactions as a proxy to predict the quality and magnitude of T cell mediated immune responses. Other techniques, including traction force microscopy (TFM) (180, 330), micro-pillar array detectors (mPADS) (331) and DNA-based molecular tension sensors (332, 333) have demonstrated to be useful in measuring mechanical forces experienced by TCRs in vivo. Importantly, DNA-based tension gauge tether was able to map mechanical forces during T cell activation modulated on a nanoparticle surface (332). This technique has been useful in determining T cell force threshold to distinguish functionally relevant mechanical forces from those do not trigger T cell activation, thus, providing a “fidelity checkpoint” for antigen discrimination (332). Utilization of these techniques to exploit single-molecule biomechanics of immune receptors may become useful in elucidating more complex cellular interactions faced by T cells such as found in the TME. Further, with the aid of these advanced biophysical tools, it is possible to develop new class of immunotherapies that aims to revamp T cell effector functions by recalibrating the mechanical force threshold of T cells to trigger more effective anti-tumor immune responses against the stiffness gradient of a growing tumor.

Lately, the usage of engineered chimeric antigen receptor (CAR) T cells as an effective immunotherapy has been useful in treating several cancer models (334–336). Importantly, previously observed non-classical immune synapses formation in CAR T cells correlates with the rapid recruitment of lytic granules to the synaptic cleft and killing the target cells much faster than classic CTLs (184). These unique functional features can be utilized to improve immunotherapy treatments against solid tumors. However, the potency of CAR T cells has been limited by several factors including T cell exhaustion (337, 338). The early constructs of CARs possessed affinities in the range of nano molar scale, rendering high-affinity interactions between CAR and antigens (339). These interactions are much stronger than physiologically relevant affinities displayed by TCR and pMHC, leading to off-target toxicities. This emphasises the role of affinity modulation in immunotherapy, which also becomes useful in designing prophylactic vaccine strategies to develop lasting immunity against pathogens. Immunotherapies aimed at treating patients in the acute phase of an infection should utilize the proliferative capacity of high-avidity T cell clones to achieve pathogen clearance. Accordingly, when an acute infection exacerbates into the chronic phase where pathogen clearance becomes inefficient, immunotherapies should make use of low-avidity clones to promote lasting host–pathogen equilibrium which delivers minimal immunopathology.

The importance of T cell biomechanics and how they differ between T cell subtypes needs further investigation. So far, much of the evidence of biomechanical influence in mediating T cell immune responses comes from expansion of clonal avidity of antigen-specific T cells during infections or cancer. Overall, in depth understanding of the biophysical properties behind mediating optimal immune repose may help to identify broader principles governing T cell dysfunctions in chronic diseases and present new and improved avenues to develop clinical interventions in the future.

AUTHOR CONTRIBUTIONS

SG and NP reviewed the relevant literature and wrote the manuscript. JG and KG provided critical feedback and helped shape the manuscript. All authors contributed to the article and approved the submitted version.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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