Review

Human Leukocyte Antigen (HLA) Class I Down-Regulation by Human Immunodeficiency Virus Type 1 Negative Factor (HIV-1 Nef): What Might We Learn From Natural Sequence Variants?

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Abstract: HIV-1 causes a chronic infection in humans that is characterized by high plasma viremia, progressive loss of CD4+ T lymphocytes, and severe immunodeficiency resulting in opportunistic disease and AIDS. Viral persistence is mediated in part by the ability of the Nef protein to down-regulate HLA molecules on the infected cell surface, thereby allowing HIV-1 to evade recognition by antiviral CD8+ T lymphocytes. Extensive research has been conducted on Nef to determine protein domains that are required for its immune evasion activities and to identify critical cellular co-factors, and our mechanistic understanding of this process is becoming more complete. This review highlights our current knowledge of Nef-mediated HLA class I down-regulation and places this work in the context of naturally occurring sequence variation in this protein. We argue that efforts to fully understand the critical role of Nef for HIV-1 pathogenesis will require greater analysis of patient-derived sequences to elucidate subtle differences in immune evasion activity that may alter clinical outcome.
1. Introduction

This review discusses our current knowledge of the Human immunodeficiency virus type 1 (HIV-1) Negative Factor (Nef) protein and its ability to mediate immune evasion through down-regulation of Human Leukocyte Antigen class I (HLA-I) molecules on the surface of virus-infected cells. We highlight recent evidence based on cellular, molecular, and structural biology studies that extends our mechanistic understanding of this important Nef activity. Furthermore, the relevance of naturally occurring Nef sequence variation and its potential impact on protein function and clinical outcome is presented. We emphasize the critical need to examine Nef function using patient-derived viral sequences in order to fully understand the role that Nef plays in HIV-1 pathogenesis.

1.1. HIV-1 Infection and Therapy

HIV-1 causes a life-long infection that is characterized by the rapid destruction of gut lymphoid compartments [1–3] followed by the progressive loss of peripheral blood CD4+ T lymphocytes [4]. The ultimate outcome of infection is a severe decline of the host immune system that is observed clinically as the inability to control opportunistic pathogen infections—a condition that is commonly referred to as acquired immunodeficiency syndrome (AIDS) [4]. Over the past 30 years, a large number of antiviral drugs have been developed that target essential stages in the HIV-1 replication cycle, including enzymatic processes (e.g., reverse transcriptase, protease, and integrase inhibitors) and critical viral:host protein interactions (e.g., CCR5 co-receptor antagonists) [5]. Advances in highly active antiretroviral therapy (HAART) have significantly reduced the burden of HIV/AIDS in regions of the world where these potent drug combinations are available, and HAART is currently our best option to control the spread of HIV [6,7]; however, establishment of latent viral reservoirs and ongoing low-level viremia in treated individuals require that therapy be continued for life. An improved understanding of the mechanisms that allow HIV-1 to persist and cause disease in its human host may uncover new opportunities for clinical or therapeutic intervention [8].

1.2. The Nef Protein

The HIV-1 accessory protein Nef is necessary for viral pathogenesis and progression to AIDS. Its in vivo role was first illustrated in the rhesus macaque model system where a nef-deleted strain of simian immunodeficiency virus (SIV) exhibited reduced viral replication, lower plasma viremia, and attenuated pathogenicity [9]. Nef gene deletion has also been associated with non-progressive HIV-1 infection [10,11]. Several reports have attempted to correlate Nef sequence polymorphisms with clinical outcome [12,13], with mixed results; however, relatively few studies have assessed potential functional impairment of Nef in the context of progressive or non-progressive HIV-1 infection using patient-derived sequences [14–17], and each of these reports examined only a small number of individuals.
HIV-1 Nef is a ~27kd protein that is expressed abundantly during the early stages of viral replication [18]. Nef displays diverse in vitro functions, including the ability to modulate a number of cell surface proteins [19], augment viral infectivity, and enhance viral replication capacity [20,21]. Down-regulation of host cell CD4 [22,23] and HLA-I [24,25] surface molecules are the most extensively studied of Nef’s activities, although some of its functions may share overlapping mechanisms. For example, Nef CD4 down-regulation activity correlates with its ability to enhance viral pathogenesis [26,27]; and lower CD4 expression on virus-infected cells may directly increase viral infectivity [28], virion release [29], viral replication [30], or prevent superinfection [31–33]. Although Nef’s contributions to HIV-1 pathogenesis remain incompletely understood, it has been proposed that progressive disease may require a combination of Nef-mediated functions acting at different times during the infection course [34,35].

1.3. HIV-1 Immune Evasion Strategies

HIV-1 evades host cellular immune responses through Nef-dependent and Nef-independent mechanisms. Nef-mediated down-regulation of HLA-I protects virus-infected cells from recognition by CD8+ T lymphocytes [36], but modulation of other host cell proteins, including CD4, CD88, CD28, CD74 (invariant chain), and HLA class II, may also contribute to Nef-dependent immune evasion [19,37,38]. Nef-independent immune evasion relies on the generation of viral sequence polymorphisms (“escape mutations”) within or near targeted epitopes, resulting in directional evolution of the virus away from immune selection pressure [39,40]. Despite these evasion strategies, CTL may retain antiviral activity, particularly if they recognized viral epitopes that can be presented prior to Nef-induced HLA-I down-regulation [41]. Nef selectively modulates HLA-A and HLA-B alleles through a shared sequence (Y320SQASS326) located in their cytoplasmic tail [42,43], leaving HLA-C allele expression unchanged on the cell surface presumably to counter the innate Natural Killer cell response against HLA-devoid cells [44]. Recent data, however, suggests that Nef-mediated down-regulation of HLA-B is less robust than that of HLA-A [45], which may in part explain the observation that HLA-B alleles tend to be more protective against HIV-1 disease progression in vivo [46].

2. HLA Class I-Mediated Control of HIV-1

2.1. Role of HLA-I in Viral Infection

During the course of viral infection, the cellular proteasome complex degrades viral proteins to produce immunogenic peptide antigens. These cytosolic peptides are transported into the endoplasmic reticulum (ER), captured by HLA-I proteins, and traffic to the cell surface for presentation to circulating antiviral CD8+ cytotoxic T lymphocytes (CTL) (Figure 1). Antigen-specific T cell receptors (TCR) allow a subset of CTL to recognize these “non-self” peptides bound to HLA on the infected cell surface. Following TCR engagement with its HLA/peptide ligand, the CTL forms an “immunological synapse” with the target cell and releases antiviral cytokines and cytotoxic molecules, including perforin and granzymes, to eliminate the infected cell [47].
Figure 1. Presentation of viral peptide antigens by Human Leukocyte Antigen (HLA) class I. Human immunodeficiency virus type 1 (HIV-1) proviral gene expression, including RNA transcription (a) and protein translation (b); generates functional viral proteins (c) as well as truncated or mis-folded proteins that are degraded by the cellular proteasome complex to form short antigenic peptides (d); These peptides are transported from the cytoplasm into the endoplasmic reticulum (ER) (e) where they can be loaded onto HLA-I molecules. Peptide/HLA complexes traffic from the ER through the Golgi and secretory vesicle (SV) network to the plasma cell membrane, where the peptide antigens are presented to circulating cytotoxic T lymphocytes (CTL) (f); The viral Nef protein shuttles HLA molecules located at the cell surface or within the trans-Golgi network into lysosomal compartments (g); where they are degraded. In the absence of Nef-mediated HLA down-regulation, antigen-specific CTL respond to stimulation by releasing cytotoxic molecules, including perforin and granzymes, resulting in elimination of the virus-infected cell (h).

The critical role of HLA-I in control of infection is illustrated by the variety of strategies that viruses have independently developed in order to evade HLA-dependent immune responses [40,48–50]. In the case of HIV-1, Nef down-regulates HLA-A and B alleles on the cell surface and thereby reduces viral peptide presentation to CTL [19,49]. HIV-1 sequences also undergo mutational escape in an HLA-restricted manner, yielding epitope variants that are incompletely or improperly processed, presented and/or recognized by CTL [39,40].

2.2. HLA Class I as a Major Determinant of HIV-1 Pathogenesis

Rates of clinical disease progression during natural HIV-1 infection vary widely, and this has been attributed mainly to differences in HLA-restricted CTL responses [40,51,52]. The association of CTL responses with initial control of plasma viremia during primary HIV-1 infection [53,54], delayed
disease progression [55] and the observation that rhesus macaques depleted of CD8+ cells prior to SIV infection exhibited an inability to control viremia [56] strongly support a critical role for CTL in control of HIV-1 at early times following infection. Over the course of infection, CTL selection pressure results in the generation of viral escape variants [53], and HLA-restricted CTL responses are a major selective force driving viral evolution in an infected host [39]. As a result of immune escape and immune dysfunction, CTL ultimately fail to control infection and the vast majority of individuals’ progress to AIDS in the absence of HAART.

HIV-1 long-term non-progressors (LTNP) and spontaneous controllers have been carefully examined in order to elucidate mechanisms of viral pathogenesis and to identify novel correlates of immune-mediated protection. It has been observed that “protective” HLA-I alleles, most notably B*27 and B*57, are consistently associated with enhanced control of HIV [57]. More recently, genome-wide association studies (GWAS) have identified single-nucleotide polymorphisms (SNPs) in the HLA region on human chromosome 6 as major determinants of lower plasma viral load set point [58,59] as well as the HIV elite controller phenotype [60–62]. Understanding potential differences in the ability of Nef to modulate CTL responses in the context of different HLA alleles or viral peptides is an area of research that merits further attention [45,63].

3. Nef-Dependent Immune Evasion: In Vivo and in Vitro Observations

_in vivo_ observations have clearly suggested that Nef plays a critical role for maintenance of high plasma viremia and for progression to AIDS [9–11]; however, evidence to directly link Nef-mediated HLA-I down-regulation to clinical outcome is more limited. Studies using the SIV-infected macaque model system have observed that Nef mutations that impaired HLA down-regulation activity were restored during the course of infection [64,65], and it was recently shown that high plasma viremia following SIV infection correlated with a high level of _in vivo_ HLA-I down-regulation [66]. Similarly, it has been reported that patient-derived Nef sequences collected during early or chronic HIV-1 infection retained significant ability to down-regulate HLA-I and to evade CTL killing [16,67]; however, other data indicate that this function may be dispensable during very late stage disease [34]. Altogether, these results indicate that HLA down-regulation is a very important _in vivo_ Nef function that is maintained presumably in order to evade ongoing CTL immune pressure on the virus.

_in vitro_ studies from a number of research groups have observed substantial variation in the susceptibility of HIV-infected cells to recognition and killing by CTL using standard co-culture assays [36,68–70]. This result is likely due to the use of different target cells, different CTL clones, and viral strains that may encode Nef variants with differential levels of expression or subtle differences in HLA down-regulation function. In cases where CTL-mediated recognition or killing has been compared directly between HIV strains that encode the _nef_ gene versus variants harboring deletions or non-functional alleles, Nef expression has clearly been shown to confer a protective effect in both cell lines and in primary T cell assays [36,63,70–72]. Notably, even though Nef effectively reduces CTL-mediated killing of virus-infected cells, it may not fully abrogate the ability of responding CTL to produce antiviral chemokines or cytokines [63]. While such incomplete evasion of CTL permits non-cytolytic immune mechanisms to participate in control of chronic infection [73], it
nevertheless allows HIV-1 to establish a persistent infection in the face of robust antiviral host immunity.

4. Nef Structure and HLA Class I Down-Regulation Function

4.1. General Features of Nef and its Domain Structure

Nef serves as an adaptor protein and its various functions appear to utilize non-canonical motifs and interactions to form higher-order complexes with cellular proteins [74]. Despite its small size, Nef engages with multiple and diverse host cell proteins, thereby altering their normal function to favor viral replication. Biochemical characterization of these protein complexes remains difficult, likely due to the small size of Nef, the heterogeneous nature of the complexes that are formed, and the localization of these complexes within lipid membranes.

The protein structure of Nef can be broadly divided into three domains: an N-terminal anchor and flexible loop, a central structured core, and a C-terminal flexible loop [75,76]; and two amino acid positions have been shown to be particularly critical for Nef function. The N-terminal anchor domain of Nef is required for membrane association and localization into detergent-insoluble “lipid rafts” [77], while the central core encodes numerous protein interaction and intracellular trafficking motifs that contribute differentially to diverse Nef functions [78]. The central core domain of Nef adopts a stable tertiary fold, permitting its early characterization using both NMR and X-ray crystallographic methods [79,80]. Our understanding of Nef-mediated HLA down-regulation has been significantly enhanced by the recently reported crystal structure of Nef protein in complex with the MHC-I cytoplasmic domain and the μ1 subunit of the clathrin AP1 complex [76].

4.2. Functional Motifs and Host Proteins Interactions

Myristoylation at glycine residue 2 (G2) allows the Nef protein to localize to lipid membranes, including the inner leaflet of the plasma membranes [81] and aspartic acid residue 123 (D123) is reported to be necessary for protein oligomerization [82]. Recent data also suggest that electrostatic interactions at D123 may be essential for Nef stability [76]. Site-directed mutations at either residue G2 or D123 have been shown to impair nearly all known Nef activities [82,83], however most other mutations indicate that the functional motifs required to down-regulate HLA-I are genetically separable from those required to modulate CD4. Notably, sequences in the central core domain that bind to the clathrin adaptor protein complex AP-2 [84,85], the coatomer protein β-COP [86,87], and the vacuolar membrane ATPase V1H [88,89] are necessary for down-regulation and degradation of CD4 [90], but not HLA-I.

The ability of Nef to down-regulate HLA-I surface expression [24,91,92] has been mapped to three distinct regions of the protein. First, an N-terminal alpha helix (R17ER19M20RRAEPA26) that contains methionine residue 20 (M20) serves an important as a membrane anchor [90,93], and more recently arginine residues 17 and 19 have been shown to form a second β-COP binding motif [87]. This latter observation indicates that the final stages of Nef-mediated CD4 and HLA down-regulation may need to engage the same cellular machinery and lysosomal compartments. Second, an acidic cluster
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(E62EEE65) that binds to the PACS-1 and PACS-2 proteins may enhance Nef localization to the trans-Golgi network (TGN) and/or regulate signaling events that increase the rate of cellular endocytosis [94,95]. Finally, a polyproline (PxxP)3 repeat that includes proline residues 72, 75, and 78 forms an SH3-binding motif that has been shown to interact with Lyn and Hck cellular kinases [90].

4.3. Proposed Mechanisms of HLA Class I Down-Regulation

Two models to explain Nef-mediated HLA-I down-regulation are currently favored, including (1) altered HLA trafficking and (2) enhanced HLA internalization/turnover. These proposed mechanisms are not mutually exclusive, and indeed the two may function collaboratively within the virus-infected cell to ensure robust immune evasion.

In the ‘altered trafficking’ model, interaction between Nef and newly synthesized HLA-I occurs within the secretory pathway, disrupting normal HLA-I transport to the cell surface and redirecting it to endosome/lysosome compartments for degradation. There is strong evidence to support this model, such as the observation that Nef and HLA-I co-localize within the trans-Golgi network rather than at the cell membrane [95]. In addition, direct binding has been demonstrated between Nef and a number of cellular complexes, including AP-1 [96] and β-COP [86]. These complexes are necessary for normal protein transport between the trans-Golgi and endosomes, and mutations in Nef that disrupt these domains dramatically impair HLA-I down-regulation function. Further support comes from biochemical data indicating that the μ1 subunit of AP-1 forms a stable interaction with Nef only in the presence of the HLA-I cytoplasmic tail [97–99]. The interaction between Nef and the AP-1 complex is thought to allow the μ subunit of AP-1 (which typically recognizes YxxØ motifs; where Ø is a bulky hydrophobic residue and x is any amino acid) to associate with a non-conventional sequence (Y320SQA323) in the cytoplasmic tail of HLA-I [42,99,100]. This hypothesis has been validated recently by structural determination of the Nef/μ1/HLA tripartite complex at less than 3-angstrom resolution [76], which captured this complicated interaction ‘in action’. Notably, this new structure demonstrates that a highly cooperative interaction between Nef and μ1 creates a novel binding pocket on μ1 that can accommodate a YxxA motif in HLA-I cytoplasmic tail. More specifically, Nef’s acidic cluster, E62EEE65, provides critical electrostatic interactions with μ1 that stabilize the complex and Nef’s polyproline-rich motif, PxxP, acts as a clamp to secure binding of HLA-I to μ1. These results confirm previous models based largely on biochemical data and provide clear rationale for the role of these Nef sequences in its HLA-I down-regulation activity. Furthermore, the structure by Jia et al. [76] highlights an important role for hydrophobic residues located in Nef’s N-terminal and C-terminal domains. While not participating directly in HLA-I or μ1 binding, Nef residues W13 and M20 anchor the protein core to the plasma membrane and presumably help to position Nef appropriately for optimal interactions with its binding partners. Likewise, residues Y202 and F203 appear to stabilize the HLA-I tail interaction.

In the ‘enhanced turnover’ model, Nef is thought to act through a series of interactions and signaling events to induce clatherin-independent internalization of HLA-I at the plasma cell membrane that requires small GTPases known as ADP-ribosylation factors (ARFs). It is proposed that Nef’s E62EEE65 motif binds to the phosphofurin acidic cluster sorting protein PACS-2 and localizes Nef within the trans-Golgi network (TGN) [94,95]. This allows Nef to bind to Src-family protein kinases.
in the TGN (in particular Hck) [101], triggering a signaling pathway that includes activation of the tyrosine kinase protein ZAP70 and PI-3-kinase, induction of phosphotidylinositol-3-phosphate (PIP₃) on the inner leaflet of the plasma membrane, and PIP₃-mediated recruitment of ARNO [102], that culminates in activation of ARF-6 that results in endocytosis of HLA-I [103]. Several aspects of this model remain controversial, and more recent studies have shown only modest effect of specific ARF-6 inhibitors [104] or a dominant-negative ARF6 mutant [105] on Nef-mediated HLA-I down-regulation.

The unusual functional flexibility of Nef is demonstrated by the fact that both of these HLA-I down-regulation models differ significantly from the mechanism that Nef uses to modulate CD4 expression, reviewed in [74,106]. In that case, direct interactions between Nef, the cytoplasmic tail of CD4, and the AP-2 complex at the plasma membrane [107] results in clathrin-mediated endocytosis of CD4 and its eventual degradation in lysosomes.

5. Natural Variation in Nef Sequence and Implications for Immune Evasion

5.1. Sequence Variability within Described HLA-I Down-Regulation Motifs

Nef is one of the most highly variable HIV-1 proteins; however, the impact of naturally occurring mutations on clinical outcome has rarely been explored in detail [12,108]. This is due, in part, to the fact that many of the motifs associated with its HLA-I down-regulation function are very well conserved in patient-derived sequences [12,109]. Analysis of a panel of 242 HIV-1 subtype B Nef sequence clones obtained from unique individuals by our laboratory (68 from acute infection, 122 in chronic infection, and 52 spontaneous controllers with plasma viral load <50 copies RNA/mL; unpublished data) confirms these previous observations (Table 1). The frequency of the consensus amino acid as well as the Shannon entropy value for each critical residue was determined using web-based tools available at the Los Alamos National Laboratory HIV Sequence Database [110]. Notably, we observed that the Nef residues required for HLA-I down-regulation function—including myristoylation (G2 and S6) and putative stability (W13), proline residues P72, P75, and P78 in the (PxxP)₃ domain, and the aspartic acid residue D123—were essentially unchanged in all patient-derived sequences (all frequencies >99%). While other critical Nef residues displayed less conservation, only glutamic acid residues E63 and E64 were observed to occur in less than 90% of sequences in this cohort of individuals with broad clinical outcome. These changes were mainly conservative substitutions to aspartic acid that are expected to have modest effect on Nef function based on previous studies [111], indicating that population-level variation in the acidic domain may also be limited.

Nef sequences adjacent to known motifs tend to be more variable, but their role in Nef function remains largely unexplored. In addition to P72, P75, and P78, mutations at residues Q73, V74, L76, and R77 in the (PxxP)₃ motif have been shown to alter Nef function [112–114]; however, each of these residues is highly conserved in our patient-derived sequences (Table 1). Greater diversity is observed in the N-terminal motif of Nef, with consensus residues R₈, S₉, V₁₀, and V₁₁ displaying less than 70% identity in our cohort (all entropy scores >1.0; data not shown). Similarly, we observed substantial variability in the C-terminal loop, where Y₂₀₂ and Y₂₀₃ displayed frequencies of ~90%. Finally, a number of Nef codons were shown recently by Lewis et al. [115] to be subject to strong purifying
selection pressure by CTL and to impair Nef’s HLA down-regulation function, including novel polymorphisms N52A, A84D, Y135F, G140R, S169I, and V180E that displayed a range of diversity in our patient sequences. More detailed studies of these regions are warranted to fully explore the possibility that common changes in Nef sequence may affect clinical outcome.

Table 1. Sequence conservation in HIV-1 negative factor (Nef) motifs required for HLA-I down-regulation.

| Nef Domain a | Role | AA | Frequency b | Entropy b | References |
|--------------|------|----|-------------|-----------|------------|
| MG2xxxS6    | Myristoylation | G2 | 100 % | 0 | [116] |
|             |       | S6 | 99.2 % | 0.05 | [116] |
| W13         | Stability (?) | W13 | 100 % | 0 | [76] |
| R17xR19     | β-COP | R17 | 97.5 % | 0.15 | [87] |
|             |       | E18 | 96.7 % | 0.18 | [115] |
|             |       | R19 | 90.5 % | 0.35 | [87] |
| M20         | Stability (?) | M20 | 90.5 % | 0.37 | [93] |
| ?            | Unknown | N32 | 98.8% | 0.08 | [115] |
| E62EEE65    | PACS-1/2 | E62 | 92.1 % | 0.31 | [95,115] |
|             |       | E63 | 75.2 % | 0.72 | [95] |
|             |       | E64 | 88.8 % | 0.48 | [95] |
|             |       | E65 | 91.7 % | 0.35 | [95] |
| P72xxPxR77  | SH3 binding, and (PxxP)3 HLA-I “clamp” | P72 | 100 % | 0 | [76,81,117] |
|             |       | Q73 | 99.6 % | 0.03 | [76,81,117] |
|             |       | V74 | 99.4 % | 0.05 | [76,81,115,117] |
|             |       | P75 | 100 % | 0 | [76,81,117] |
|             |       | L76 | 96.7 % | 0.16 | [76,81,117] |
|             |       | R77 | 100 % | 0 | [76,81,117] |
|             |       | P78 | 99.6 % | 0.03 | [76,81,117] |
|             |       | G83 | 56.2% | 0.73 | [90,115] |
| ?            | Unknown | A84 | 99.2% | 0.05 | [115] |
| D123        | Oligomerization and Stability (?) | D123 | 100 % | 0 | [74,76,82,115] |
| ?            | Unknown | Y155 | 75.2% | 0.64 | [115] |
| ?            | Unknown | G140 | 100% | 0 | [76,115] |
| ?            | Unknown | S169 | 89.7% | 0.47 | [76,115] |
| D175        | Trafficking | D175 | 99.6% | 0.03 | [76,88,115,118] |
| ?            | Unknown | V180 | 99.2% | 0.05 | [76,115] |
| Y202        | Stability (?) | Y202 | 87.6% | 0.39 | [76] |
| F203 c       | Stability (?) | F203 | 9.5% | 0.31 | [76] |
|             |       | Y203 | 90.5% | 0.31 | [76] |
| D123        | Oligomerization and Stability (?) | D123 | 100 % | 0 | [74,76,82,115] |
| ?            | Unknown | Y155 | 75.2% | 0.64 | [115] |
| ?            | Unknown | G140 | 100% | 0 | [76,115] |
| ?            | Unknown | S169 | 89.7% | 0.47 | [76,115] |
| D175        | Trafficking | D175 | 99.6% | 0.03 | [76,88,115,118] |

a: Protein locations based on HXB2 numbering [110]; b: Frequency of consensus residue and Shannon entropy score calculated using 242 clonal Nef sequences collected from unique HIV-1 subtype B-infected individuals from North America (68 acute, 122 chronic, and 52 controllers) and the Entropy-One tool (HIV Sequence Database; [110]); c: Nef used by Jia et al. [76] encoded phenylalanine-203, but tyrosine-203 is prevalent in most sequences.
5.2. Immune-Mediated Attenuation of Nef Function?

HIV-1 Nef is highly targeted by the host immune response during primary infection [119], and a large number of HLA-associated polymorphisms have been identified within or near known CTL epitopes [120]. HLA-mediated immune pressure on Nef drives rapid selection of escape mutations following infection [121,122], and HLA-associated polymorphisms have been identified at approximately half of Nef's 206 residues [120,123]. Indeed, a substantial proportion of natural sequence variation observed in Nef is attributable to immune selection pressure on this protein [120,124]. It has been presumed that the extensive ability of Nef to incorporate sequence changes would allow it to escape from immune pressure with limited consequence for viral fitness. However, while the impact of certain CTL escape mutations on Nef function has been assessed [35,125,126], the broader impact of HLA-restricted pressure on Nef function and viral pathogenesis at the individual or population level has not been clarified, and data to address this important issue are currently lacking.

We have previously reported that HLA-B*35-associated CTL escape mutations R75T and Y85F located in the conserved proline-rich region of Nef can impair HLA-I down-regulation activity [35,126]. Along with other data for A*02 [125], these observations indicate that host immune pressure can alter Nef function in at least some cases and further highlight the need for additional studies to characterize the wide array of Nef sequence variants that are likely to arise within an individual during natural infection. Indeed, population-level analyses have identified a number of HLA-associated polymorphisms in the N-terminal and C-terminal domains of Nef, as well as in sites near other critical Nef residues [120], including several that are selected by the protective allele HLA-B*57. Fully understanding the potential impact of naturally occurring HLA-associated mutations on Nef function and clinical outcome will be an important area for future study.

6. Conclusions

Research advances have significantly improved our understanding of the HIV-1 Nef protein and the mechanisms that it uses to effectively down-regulate HLA class I expression on the surface of infected cells. Molecular and biochemical studies have identified many of Nef's crucial binding partners and have mapped Nef sequence motifs that are required for its function. Structural data have very recently allowed us to visualize the Nef protein in complex with HLA-I and the µ1 subunit of AP-1, validating our current models and identifying potential sites for therapeutic intervention.

Nef-mediated evasion of host immunity is expected to contribute significantly to the establishment and maintenance of persistent HIV-1 infection. However, despite recent progress in the field to understand the cellular mechanisms of Nef-mediated HLA-I down-regulation, our knowledge of Nef's role during HIV-1 disease progression remains poor. To fully elucidate the impact of Nef during natural infection, it will be necessary to extend our current studies of lab-adapted viral isolates to include detailed analyses of patient-derived Nef proteins. Only then will we be able to fully appreciate the consequence of Nef sequence variation on protein function and make important links to clinical outcome.
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Conflict of Interest

The authors declare no conflict of interest.

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