Attributes of Host’s Genetic Factors in HIV-1 Pathogenesis

Bechan Sharma*
Professor, Department of Biochemistry, University of Allahabad, Allahabad-211002, UP, India

The human immunodeficiency virus type-1 (HIV-1) is a well-known cause of AIDS leading to the deaths of millions of people all over the world every year. The anti-HIV-1 regimen available as on today have been designed and developed targeting two key viral enzymes such as (1) HIV-1 reverse transcriptase (HIV-1RT) and (2) protease [1,2]. However, increasing incidences of drug failures due to emergence of frequent viral mutations as well as toxicity of these compounds in the patients have led to failures of chemotherapy against AIDS [3,4]. Therefore, research into HIV-1 biology is of paramount importance in order to investigate new targets for design and development of novel chemotherapeutics to inhibit not only the wild type virus but also the drug resistant variants [5]. The ongoing research efforts in this direction suggest that HIV-1 exploits a complex network of many different host cellular factors for the replication of its genome [6,7].

HIV-1 uses host-cellular machinery in order to produce viral genomic material, viral proteins and finally the new mature virions. The hijack and control over host cell processes are mediated by HIV-1 proteins through a complex network of molecular events, including virus-host protein-protein interactions (PPIs) [8]. Therefore, by developing our knowledge of the virus-host interaction network, we can improve our current model of HIV-1 infection and host-cell perturbation and use this information to aid development of new antiviral treatments. One example of a successful antiviral treatment that has come from understanding HIV-host cell interaction is the drug maraviroc [9]. Maraviroc is an entry-inhibitor that binds the CCR5 co-receptor, inhibiting gp120:CD4:CCR5 complex formation and, thus, the entry into the host cell [10,11]. Targeting a host protein in this way demonstrates that the number of possible HIV-1 therapeutic drug targets is not limited to the small viral proteome and that understanding the virus-host interface can lead to the development of novel-acting therapeutic agents [12,13]. The identification of additional cellular cofactors through an independent RNA interference screen has been demonstrated. In fact in last few years, the continued research in this direction has revealed that the cellular factors exhibit dual properties to support as well as to oppose the viral replication; the latter is called as the host restriction factors [14].

After HIV-1 enters the target cell, its cDNA (proviral DNA) integrates into the cellular chromosomes. This process of integration is irreversible in nature and allows HIV-1 to persist in the infected cell in a quiescent or latent stage, which helps the virus to escape from host’s immune surveillance as well as antiviral treatment. It is known that both the virus and the host cellular factors are required for HIV-1 replication and expression. Gradually HIV-1 modifies the cellular environment according to its need of efficient replication and production of viral progeny [14]. The viral regulator protein Tat is required for efficient transcription and elongation of viral transcripts. For this purpose, Tat recruits several cellular proteins to make the chromatin structure accessible for the transcription machinery, to acquire the posttranslational modifications essential for its function, and to produce efficient viral replication. In contrast, the host cell also has several restriction factors to encounter viral replication at different steps. The two accessory proteins encoded by HIV-1 namely Vif and Vpu combat such cellular restrictions and thus significantly contribute in HIV-1 pathogenesis [14].

The different viral proteins which have protein-protein interactions with those of the cellular proteins are as following: The Gag gene of HIV-1 encoded matrix protein (MA) interacts with the cellular Karyopherins, histidyl-tRNA synthetase-like, calmodulin and virion associated nuclear shuttling protein/Nef Associated Factor 1 (VAN/NAF1); the viral capsid (CA) uses host’s Cyclophilin A and retroviral restriction factor Tripartite Motif protein (TRIM5a) for uncoating of the virus and its disassembly. This process of uncoating and disassembly of virions also involves another cellular machinery, Arp2/3 complex. The nucleocapsid (NC) protein of HIV-1 interacts with HP68/RNase L inhibitor and actin; the viral p6 protein interacts with many cellular factors such as Tumor susceptibility gene 101 protein (TSG101), ASK1-interacting protein 1 (AIP1), Neurocural precursor cell-expressed developmentally downregulated 4 (Nedd4) and Ubiquitin. Out of the three pol gene encoded viral enzymes (HIV-1RT; protease and integrase), integrase interacts with integrase interactor 1 (IN11/hsNF5), Lens epithelium-derived growth factor/transcription co-activator p75 (LEDFG/p75), ATR, ATM, Karyopherins, BAF and XRCC5 (Ku autoantigen) [15].

The envelope of HIV-1 is comprised of a lipid bilayer derived from the host’s cell membrane containing certain proteins which mediate receptor binding and membrane fusion. The glycoproteins present at the viral surface recruit cellular CD4 receptor which facilitates docking of HIV-1 with CXCR4 or CCR5 co-receptors allowing internalization of the virus. In addition, HIV-1 also may enter into the CD4+ T-cells via infected dendritic cells (DC) through interaction with DC specific factor, ICAM-3 DC-SIGN. DC-SIGN, a non-integrin, is exclusively expressed by DC, mediates adhesion between DC and resting T-cells via ICAM-3, and is involved in DC-T cell clustering. It is also required for DC-induced proliferation of resting T cells. Upon contact with T-cells, DC helps HIV-1 migrate into T-cells for further infection. Since DC hides HIV-1 as that of macrophages, they evade host’s immune surveillance system and cause failures of vaccines [16].

The accessory proteins encoded by HIV-1 such as Nef recruits PACS-1, ASK-1, PAK, P13-Kinase, lck and VAN/NAF1; Rev interacts with Crm1 and p32; Tat preferably interacts with Cyclin T1; Vpr utilizes karyopherins and Uracil-DNA glycosylase; and the viral Vif interacts
with host’s APOBEC3G (A3G) and helps in its disintegration in order to avoid A3G mediated hypermutation into proviral DNA. The A3G is also involved in uncoating and the disassembly of the virus [17].

The reverse transcription process of HIV-1 genome is essential to synthesize proviral cDNA. The heterodimeric (p66/p51) reverse transcriptase (RT) utilizes cellular tRNA as a primer containing about 18 nucleotides from its 3'-end which are complementary to the viral genomic RNA. It anneals with the primer binding site located at the 5’ terminus of viral LTR region and provides 3'-OH group for synthesis of first (-) single stranded DNA (ssDNA). The other cellular factors involved at different stages of reverse transcription in HIV-1 are actin, tubulin and karyopherin [17,18].

The integrase interactor 1IN1/HSN5, a component of SNF-SWI Complex, facilitates nuclear import of the pre-integration complex (PIC) as well as the integration reaction. IN1 is also needed at the late events of viral life cycle. The binding of human lens epithelium-derived growth factor / transcription co-activator p75 (LEDGE/p75) with integrase is crucial in nuclear import of PIC. HMGal and BAF have also been shown to play key role in this process [17,19,20]. A detailed account of host cell factors that inhibit or assist viral DNA integration processes is reviewed by Sloan and Wainberg [21]. Many of those factors involved in cellular DNA repair processes such as XBP and XPD which are DNA helicases and part of TFII B transcription complex play role in DNA nucleotide excision repair [22]. Ataxia-Telangiectasia Mutated (ATM) is especially critical for repair of DSBs, whereas ATM- and Rad3-related (ATR) kinase appears to be specific for lesions that contain single strands coated with replication protein A (RPA) [19,23].

The P6 domain within HIV-1 gag is involved in budding in association with the host cellular protein, TSG101. HIV-1 release requires TSG101 that sorts proteins into vesicles that bud into multivesicular bodies (MVB) [17,24]. TSG101+PPTAP motif of p6 complex recruits ESCRT III (endosome associated complex required for transport) to the budding site. Similar to TSG101, the multifunctional proteins AIP1/ALIX also bind to the p6 and plays role in pinch-off process. Lysosomal degradation of ubiquitinated receptors is directed by the sequential action of the ubiquitin-binding protein complex composed of Hrs, TSG101 and ALIX with associated proteins. The endocytic sorting machinery is recruited during the budding of RNA viruses, whose Gag proteins are ubiquitinated and have motifs that bind TSG101 and ALIX [25].

Apolipoprotein B m-RNA-editing enzyme-catalytic polypeptide-like 3G (APOBEC3G) can deaminate C to yield U on the minus-strand of viral genome, which causes G-to-A conversion on the complimentary plus strand. The G-to-A hypermutation alters the nucleotide sequences of the viral genome leading to inability in HIV-1 replication. The viral accessory protein, Vif, binds to APOBEC3G to induce proteasome-dependent degradation; thereby it is not incorporated into HIV-1 particles. Thus Vif protects the viral genome by this mechanism. The Vif-mediated ubiquitination / degradation of APOBEC3G involve four critical lysine residues in its C-terminal domain [17,26,27]. However, cellular Cycophilin A (CypA) protects HIV-1 from an unknown antiviral activity in human cells. It regulates the processes of internalization as well as assembly of HIV-1 [28].

The viral accessory protein Nef plays an important role in HIV-1 pathogenesis by interacting with various cellular proteins; one of them being Cyclin K (CycK). CycK acts as a cellular restriction factor and inhibits HIV-1 gene expression and replication in a Nef-dependent manner [29]. Further, the role of cellular protein kinase C (PKC) theta (θ) as another host’s restriction factor opposing the viral replication has been indicated [30]. Unlike the roles of CycK and PKCθ, one of the cellular soluble factor, tumor necrosis factor-alpha (TNF-α) enhances viral replication via activation of nuclear factor-kappa B (NF-κB) [17]. The increasing knowledge of viral protein interactions with host cell factors would be essential for the discovery of new targets that could be used to design new therapeutic strategies [14].

References


