Nanotechnology: A Curative Approach to Combat HIV/AIDS

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ABSTRACT

HIV-AIDS is one of the biggest challenges of the 21st century to cure. However, in the current scenario, various antiretroviral drugs are available which makes the condition chronic rather than worse which manages to increase the survival rate. Antiretroviral drugs are manageable but the bioavailability, lower permeability; poor half-life of the drug limits the potent activity. High dose drug administration leads to higher toxicity which arrays adverse effects and develops resistance against HIV strain. Potent targeting of drugs is lacking due to its instability, chemical degradation, and tissue barrier restriction. The application of nanotechnology to anti-retroviral drug delivery holds the capacity to cure AIDS. The nanotechnology-based efficient delivery system of Nanocarrier (liposomes, dendrimers, nanoparticles, polymeric micelles, nano-emulsion, nanovesicles) plays a major role in drug delivery. Nanocarrier has revolutionized the field of pharmaceutics in the world of drug delivery. This review depicts the nano-based system which is incorporated or encased with ARV drug to increase its efficiency or effectiveness with low adverse effect to abort HIV.

Key Words: Antiretroviral drug, Nanocarrier, HIV, AIDS, Nanotechnology, Drug delivery

INTRODUCTION

Human immunodeficiency virus (HIV), the virus that causes Acquired immunodeficiency syndrome, is one of the world’s most serious public health challenges. According to the statistical report of WHO (Global Health Observatory Data Repository 2018), 37.9 million people are living with HIV infection 1 from which in India, Government of India estimates that about 2.40 million Indians are living with HIV (1.93-3.04 million) with an adult prevalence of 0.25% (2017), while 83% are the in-age group 15-49 years. This virus isolated by Gallo et al. in the USA 2, diversely known as human cell leukaemia virus-III (HCLV-III), human T-lymphotropic virus III (HTLV-III), lymphadenopathy-associated virus (LAV), and ultimately HIV in 1986 by ICTV (International Committee on a taxonomy of virus) 3.

In the late 20th century two types of HIV strain (HIV-1 and HIV-2) categorize, which each evolved from a diverge simian immunodeficiency virus (SIV). HIV-1 is the most common one dispersed across the globe while HIV-2 is more prominent in West Africa. HIV invades from the mucosal layer, damages the immune system, leaving the host body prone to infection with a wide range of bacteria, viruses, fungi, protozoa. HIV infection propagates by the transfer of body fluids due to blood transfusion, organ transplant, sexual contact, from infected mother to offspring. Sexual transmission is one of the main sources to enter into the mucosal surfaces. The female genital tract is the dominant route of heterosexual HIV transmission 4. Sexual transmission through the Rectal route is also a substantial problem that makes it more susceptible to HIV infection due to its physiology 5.

The primary targets of HIV infection are the immune cell i.e. macrophages and dendritic cells which are present in the subepithelial layer of the vagina or cervix mucosa 6. It transmits through the semen or other biological fluids during intercourse which penetrates the stratified squamous epithelium or columnar epithelium of the vagina to infect the target cell. HIV possesses the glycoprotein called gp120 as mention in Figure 1., on the surface of the viral coat which binds to the transmembrane protein receptor CD4 or chemokine receptor CCR5, CXCR4 of the T-helper lymphocytes and infects the cell 7.
The HIV infects the host cell via endocytosis it fuses with the cell membrane and releases its viral RNA into the host cytoplasm it undergoes reverse transcription succeeded by the integration of proviral DNA into the host chromosomes. Subsequently, it creates new viral particles budding out from it to infect the other cells. It infects macrophages and depletes the amount of CD4+ cells which is the distinctive feature of the HIV infection. HIV Viruses utilizes different genes(a) structural genes (three) — gag, pol, and env (b) six regulatory genes — tat, rev, nef, vif, vpr, and vpu, to enhance its productivity and hijack the cellular mechanism of the host cell to generate its progeny to infect the other cells. After successful infection viral particles reside into the main anatomical site such as macrophages, bone marrow, lymph node, spleen, lung, and central nervous system (CNS). When it resides in the CNS it causes significant loss of neural networks and ultimately leads to serious complications such as HIV Associated Dementia (HAD). When the patient is not treated well it will die within 5-10 years.

In the current scenario, various Antiretroviral drugs are available in the market with a different combination which depends upon the stage of infection. High Activity Antiretroviral Therapy (HAART) is employed to treat AIDS/HIV. It had been launched in 1996 and involves a combination of at least three antiretroviral (ARV) drugs. This therapy has been used to extend the life span of HIV-infected patients. However this therapy is used to manage the infection with some extend, but total recovery is not yet been achieved because these ARV drugs have some limitation such as moderate water solubility, low target specificity, low half-life time reactivity, poor blood-brain barrier permeability, poor bioavailability is one of the major problems. The ARV drug works on the principle of blocking and inhibiting mechanisms, based on the stage of the HIV cycle. Reverse transcriptase inhibitor blocks the activity of reverse transcriptase enzyme which prevents the conversion of viral RNA to DNA. Nucleoside analogue reverse transcriptase inhibitors, including zidovudine, lamivudine, stavudine, abacavir, emtricitabine, zalcitabine, dideoxycytidine, didanosine, tenofovir disoproxil fumarate, and didanosine get to incorporate in between the nucleotide chain at the time of reverse transcription which leads to chain termination, whereas non-nucleoside analogue reverse transcriptase inhibitors, including etravirine, delavirdine, efavirenz, and nevirapine block the binding efficiency of the reverse transcriptase enzyme. In the market, various kinds of drug regimens to treat AIDS are available or have been prescribed. But to deal with the selection of a correct combination of ARV drugs is a difficult task because due to various factors like drug properties, drug resistance status, patients’ acceptance reactivity, drug costing, Drug toxicity, or any other drug adverse effect. HAART is prescribed for a lifelong duration and any disobedience leads to a drastic increase in the viral load in the infected patient. The major drawback of ARV drug is the shorter duration availability in the bloodstream of the body so that the viral particle in the site of the reservoirs like CNS, Lymph nodes, and lungs are less exposed to the drug, so to eradicate the viral particle higher doses of the drug are required for a prolonged period, which develops resistance in HIV strain. The reservoir also contains latently infected cells which include CD4+ T-cell, monocytes/ macrophages lineage which contains integrated transcription silencing provirus within their genome which can reinfect the patient due to activation of the proviral genome. To overcome such kind of problems and limitations nano-based drug delivery system, nanomedicines and various nano approaches play a major role in the drug efficiency, drug reactivity, drug target specificity, lowering the toxicity and adverse effect of the drug and various major challenges which current ARV drug face in the current scenario.

The nanobiotechnological world which emerges out with the great advance technologies with greater efficiency due to its nanoscale mechanism ranges (1-100nm). Nanotechnology has revolutionized the field of pharmaceutics in the world of drug delivery. The basic principle is to modulate pharmacokinetics of the chemical molecule which becomes worthy to eradicate the HIV from the body without harming from its consequences. It also enhances the biodistribution and bioavailability of the drug to expose the virus particle for a prolonged period with its greater target specificity. In an enclosed nano-system that carries anti-HIV drug its absorption, excretion, distribution, and metabolism are not governed by the drug but by the physical, chemical properties of the nano-system which is based on its size and electric charge. Application of nanotechnology to ARV drug delivery holds the capacity to cure AIDS because it possibly avails the drug at the reservoir site and also increases the half-lives of the drug. This review depicts about the nano-based system such as liposomes, dendrimers, nanoparticles, polymeric micelles, nano-emulsion, nanovesicles, and peptide self-assemble nano-drug which are incorporated or encased with ARV drug to increase its efficiency or effectiveness with low

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**Figure 1:** The structure of HIV.
adverse effect to abort HIV from an infected person and also reactivate the latent HIV particle from its reservoir site to prevent further future infection.

**Current available strategies for the treatment of HIV- AIDS based on its life cycle**

Currently, available ARV drugs work on the principle of inhibition of the life cycle of HIV which blocks or inhibits the activity or virulence factor of the viruses. HIV viruses require the receptor molecules to enter the Host cell based on viral antigen-receptor interaction. Host cells contain a CD4 receptor molecule and one of the major co-receptor molecules (CCR5 or CXCR4) on surface 24. Viral antigen called spike protein (gp120) recognizes this receptor and mediates the viral infection with the cell membrane of the host cell with the involvement of the viral fusion protein (gp41) 25. The ARV class of drug target this process to prevent the entry to the host cell. HIV entry involves three complicated processes: (1) An attachment step that requires CD4 receptor binding, (2) Co-receptor binding, (3) fusion process. Chemical or drug which acts against this process is represented in Table 1.

**Table 1:** Represents chemical components or drug and its description of attachment inhibitor, co-receptor binding and fusion inhibitor

<table>
<thead>
<tr>
<th>Chemical component or drug</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cyclodextrinsulfate, dextran sulfate.</td>
<td>Non-specific blocking of HIV virion [26].</td>
</tr>
<tr>
<td>Cyanovirin-N (Cellegy Pharmaceuticals)</td>
<td>It binds to a conserve high manose carbohydrate region of gp120 which prevent attachment 27.</td>
</tr>
<tr>
<td>PRO-2000 (Indevus Pharmaceuticals), Naphthalenesulfonate polymer.</td>
<td>It binds non-specifically to the CD4 receptor to prevent interaction of antigen and receptor. (Microbicides) 46.</td>
</tr>
<tr>
<td>Glycyrrhizin.</td>
<td>A triterpenoid saponin isolated from liquorice root inhibits HIV-1 replication by partially inhibiting viral adsorption to CD4+ cells 36.</td>
</tr>
<tr>
<td>Amphotericin, lentinan, suramin, aurantricarboxylic acid, glycyrrhizin sulfate.</td>
<td>It interferes with the nonspecific viral attachment process 39.</td>
</tr>
<tr>
<td>PRO 542 (Progenics Pharmaceuticals)</td>
<td>Recombinant tetrameric antibody CD4-TgG2 which targets the CD4 binding site on gp120 by mimicking the CD4 receptor 40.</td>
</tr>
<tr>
<td>TNX355</td>
<td>A humanized IgG4 monoclonal antibody against CD4, which binds to the D2 domain of CD4 and arrest CD4-induced post-binding conformational changes 30.</td>
</tr>
<tr>
<td>CADA (cyclotriadisulfonamide)</td>
<td>CADA function by a specific CD4 down-modulating potency 31. As a primary mode of action, it inhibits the interaction of CD4-gp120 32.</td>
</tr>
<tr>
<td>BMS-488043</td>
<td>Co-receptor binding inhibitor.</td>
</tr>
<tr>
<td>AOP-RANTES and NNY-RANTES</td>
<td>It competes with gp120 to bind the CCR5 co-receptor and inhibit HIV infection 33.</td>
</tr>
<tr>
<td>PRO-140 (Progenics Pharmaceuticals)</td>
<td>It is a murine anti-CCR5 monoclonal antibody that binds a complex epitope spanning multiple extracellular domains on CCR5 34.</td>
</tr>
<tr>
<td>TAK-799, TAK-652 (Take-da Chemical Industries)</td>
<td>It inhibits the binding to CCR5 co-receptor 35.</td>
</tr>
<tr>
<td>Maraviroc (MVC, UK-427,857, Pfizer, Inc.), Vicriviroc (SCH-D, SCH-417,690, Schering-Plough), Ancriviroc (SCH-C, SCH-351,125, Schering-Plough)</td>
<td>It is selective CCR5 antagonist with potent antiviral activity against all CCR5-tropic HIV-1 viruses 36.</td>
</tr>
<tr>
<td>KRH-1636 (Kureha Chemical Industries)</td>
<td>It is an arginine-based CXCR4 antagonist with efficient anti-HIV-1 activity 37.</td>
</tr>
<tr>
<td>AMD3100 (AnorMED, Inc., now Genzyme Corporation)</td>
<td>It is a low molecular weight bicyclic analogue with potent anti-X-4 HIV variant activity 38.</td>
</tr>
<tr>
<td>Fusion inhibitors</td>
<td>Enfuvirtide (T20, Fuzeon, Trimeris/Roche)</td>
</tr>
<tr>
<td>Enfuvirtide</td>
<td>It is a 36-amino acid synthetic peptide that imitates the HR2 region (residues 127–162 in C-terminal) of gp41. It prevents the formation of the six-helix bundle structure that is crucial as the energy source for the fusion process 39.</td>
</tr>
<tr>
<td>TRI-999 and TRI-1144 (Trimeris)</td>
<td>TRI-999 and TRI-1144 (Trimeris) are oligopeptide fusion inhibitors 40.</td>
</tr>
<tr>
<td>(Betulinalamino)alkanoic acid, C-28 aminoalkanoic acid &amp; its derivatives.</td>
<td>It modulates antiviral fusion potency 41.</td>
</tr>
</tbody>
</table>

After fusion to the plasma membrane, viral protein p24 gets introduced into the host cell cytoplasm 42. While migrating towards the nuclear pore reverse transcription of the viral RNA takes place inside the core. Inside the viral core, RNA transformed into DNA with the help of reverse transcriptase by incorporating nucleotide triphosphates 43 from the cytoplasm and ARV drug target this mechanism to combat viral DNA synthesis by blocking or terminating the process of reverse transcription. Based on this inhibition, it divided
into two classes non-nucleoside reverse transcriptase inhibitors (NNRTI) and nucleoside reverse transcriptase inhibitors (NRTI) as shown in Table 2.

### Table 2: Represents Market available Drugs and its brand name of NNRTI and NRTI.

<table>
<thead>
<tr>
<th>Non-nucleoside reverse transcriptase inhibitors (NNRTI)</th>
<th>Generic names</th>
<th>Abbreviations</th>
<th>Brand name</th>
<th>Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Efavirenz</td>
<td>EFV</td>
<td>Sustiva®</td>
<td>Bristol-Myers Squibb</td>
<td></td>
</tr>
<tr>
<td>Etravirine</td>
<td>TMC125</td>
<td>Intelen®</td>
<td>Tibotec</td>
<td></td>
</tr>
<tr>
<td>Nevirapine</td>
<td>NVP</td>
<td>Vire-mune®</td>
<td>Boehringer-Ingelheim</td>
<td></td>
</tr>
<tr>
<td>Delavirdine</td>
<td>DLV</td>
<td>Rescriptor®</td>
<td>ViiV Health care.</td>
<td></td>
</tr>
<tr>
<td>Rilpivirine</td>
<td>TMC278</td>
<td>Edurant®</td>
<td>Tibotec</td>
<td></td>
</tr>
<tr>
<td>Doravirine</td>
<td>MK-1439</td>
<td>Pifeltro®</td>
<td>Merck &amp; Co.</td>
<td></td>
</tr>
</tbody>
</table>

Nucleoside Reverse transcriptase inhibitors (NRTI)

<table>
<thead>
<tr>
<th>Didanosine</th>
<th>ddI</th>
<th>Videx®</th>
<th>Bristol-Myers Squibb</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zalcitabine</td>
<td>ddC</td>
<td>HIVID®</td>
<td>Roche</td>
</tr>
<tr>
<td>Lamivudine</td>
<td>3TC</td>
<td>Epivir®</td>
<td>GlaxoSmith-Kline</td>
</tr>
<tr>
<td>Abacavir</td>
<td>ABC</td>
<td>Ziagen®</td>
<td>GlaxoSmith-Kline</td>
</tr>
<tr>
<td>Zidovudine</td>
<td>AZT</td>
<td>Retrovir®</td>
<td>GlaxoSmith-Kline</td>
</tr>
<tr>
<td>Tenofovir</td>
<td>TDF</td>
<td>Vemlidy®</td>
<td>Gilead Sciences</td>
</tr>
<tr>
<td>Stavudine</td>
<td>d4T</td>
<td>Zerit®</td>
<td>Bristol-Myers Squibb</td>
</tr>
<tr>
<td>Emtricitabine</td>
<td>FTC</td>
<td>Emtriva®</td>
<td>Bristol-Myers Squibb</td>
</tr>
<tr>
<td>Entecavir</td>
<td>ETV</td>
<td>Baraclude®</td>
<td>Bristol-Myers Squibb</td>
</tr>
</tbody>
</table>

After completion of reverse transcription viral coat deconstruct and import pro-viral DNA to the nucleus. Integrase mediates the insertion of viral DNA into the host chromosome and hijacks the cellular machinery to produce viral mRNA \(^{45}\). Integrase inhibitor target this mechanism to prevent viral DNA insertion into the host. Integrase inhibitor drug represented in Table 3.

### Table 3: Represents integrase inhibitors which are available in the market.

<table>
<thead>
<tr>
<th>Integrase inhibitors</th>
<th>Generic names</th>
<th>Abbreviations</th>
<th>Brand name</th>
<th>Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Raltegravir</td>
<td>RAL</td>
<td>Isentress®</td>
<td>Merck</td>
<td></td>
</tr>
<tr>
<td>Dolutegravir</td>
<td>DTG</td>
<td>Tivicay®</td>
<td>ViiV Health care.</td>
<td></td>
</tr>
<tr>
<td>Elvitegravir</td>
<td>EVG</td>
<td>Vitekta®</td>
<td>Gilead Sciences</td>
<td></td>
</tr>
</tbody>
</table>

After insertion of the viral DNA, it starts towards the processes of virion generation. Viral mRNA transcript translates protein and splice into mature protein by the protease \(^{46}\). Some classes of ARV drugs as follows in Table 4. which interfere’s with the protease enzyme which prevents the maturation of the protein processing. Virus particles are reconstructed at the cell membrane by the recruitment of the freshly synthesized viral proteins. Two viral genomic-RNA with its enzyme and regulatory protein is packed into the viral capsid. Assembled virus particles are liberated from the cell surface by a process of budding.

### Table 4: Represent protease inhibitors with its Brand name and Manufacturer.

<table>
<thead>
<tr>
<th>Protease inhibitors</th>
<th>Generic names</th>
<th>Abbreviations</th>
<th>Brand name</th>
<th>Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Saquinavir</td>
<td>SQV</td>
<td>Invirase®</td>
<td>Roche</td>
<td></td>
</tr>
<tr>
<td>Atazanavir</td>
<td>ATV</td>
<td>Reyataz®</td>
<td>Bristol-Myers Squibb</td>
<td></td>
</tr>
<tr>
<td>Indinavir</td>
<td>IDV</td>
<td>Crixivan®</td>
<td>Merck</td>
<td></td>
</tr>
<tr>
<td>Nelfinavir</td>
<td>NFV</td>
<td>Viracept®</td>
<td>Agouron Pharmaceuticals</td>
<td></td>
</tr>
<tr>
<td>Fosamprenavir</td>
<td>FPV</td>
<td>Lexiva®</td>
<td>ViiV Health care.</td>
<td></td>
</tr>
<tr>
<td>Ritonavir</td>
<td>RTV</td>
<td>Norvir®</td>
<td>Abbvie</td>
<td></td>
</tr>
<tr>
<td>Darunavir</td>
<td>DRV</td>
<td>Prezista®</td>
<td>Abbvie</td>
<td></td>
</tr>
<tr>
<td>Tipranavir</td>
<td>TPV</td>
<td>Aptivus®</td>
<td>Boehringer Ingelheim</td>
<td></td>
</tr>
<tr>
<td>Amprenavir</td>
<td>APV</td>
<td>Agenerase®</td>
<td>GlaxoSmith-Kline</td>
<td></td>
</tr>
</tbody>
</table>

Based on different stages of HIV infection, the patient receives a combinational drug treatment as shown in Table 5. , which increase the inhibitory or viricidal effect on the HIV life cycle.
Table 5: Represents different ARV drug combination with its brand name.

<table>
<thead>
<tr>
<th>Drug combination</th>
<th>Brand name</th>
<th>Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lopinavir + Ritonavir</td>
<td>Kaletra™</td>
<td>Abbott Labs</td>
</tr>
<tr>
<td>Lamivudine + Zidovudine</td>
<td>Combivir™</td>
<td>GlaxoSmithKline</td>
</tr>
<tr>
<td>Abacavir + Lamivudine</td>
<td>Trizivir™</td>
<td>GlaxoSmithKline</td>
</tr>
<tr>
<td>Tenofovir + Emtricitabine</td>
<td>Truvada™</td>
<td>Gilead Sciences</td>
</tr>
<tr>
<td>lamivudine + tenofovir-id- isoproxifumarate</td>
<td>Temixys™</td>
<td>Celltrion</td>
</tr>
</tbody>
</table>

**Nanotechnology-based drug delivery system to treat HIV-AIDS infection**

Being of nano-size the nanomaterial behaves differently than the conventional drug because due to its size reduction and have their therapeutic effect within the living system. The employment of nanotechnology platforms for the delivery of drugs is the trailblazing of medication in various areas of disease treatment. Using nanotechnology, it’s become possible to appreciate enhanced delivery of poorly water-soluble drugs, focused delivery of medicines to specific cells or tissues, and intracellular delivery of macromolecules. Nanocarriers offer several benefits, such as the protection of drugs against degradation, drug specificity, and delivery of biochemical molecules, such as proteins, peptides, oligopeptides, and oligonucleotides. Nanocarriers also utilize to overcome the limitation of therapeutic applications such as drug delivery challenges, drug bioavailability, drug conformation stability, physicochemical stability, increased permeability, tissue clearance of the drug, cellular uptake, reduction of immunogenic response. A brief overview of the nano-system is represented in Figure 2.

**Liposomes**

A liposome is a small microscopic vesicle that is made up of phospholipid bilayers which are usually encircling by the aqueous core as mention in Figure 3. This is advantageous to carry hydrophilic drugs by entrapping inside the core and while the hydrophobic drug is incorporated in the lipid bilayer.

![Figure 3: This picture represents the structural arrangement of liposomes.](image-url)

The size of the liposomes can be 25nm to several microns which gives the advantage of permeability. In their preparation, natural or synthetic phospholipids along with the cholesterol and additionally lipids, protein, or peptide fragments are used. Liposomes when entering into the living system it recognizes as the foreign particle which is engulfed by the mononuclear phagocytic cells such as macrophages, so liposomes represent favourable carrier for the anti-HIV drug to the infected cell. Thus, liposomes can enhance the efficiency of the anti-HIV drug by lowering its side effects. Three types of liposomes are there which include, small uni-lamellar vesicles, large uni-lamellar vesicles, and multi-lamellar vesicles. In their native form liposomes are grabbed by the reticuloendothelial system and quickly clear from the circulation. This property was studied in vivo mice modal by using drug Zidovudine. The Zidovudine drug is an amphiphilic compound, with the use of a liposomal drug delivery system, it results in greater efficiency and a longer half-life in the model. The liposomal loaded system of zalcitabine (2’,3’-dideoxycytidine, ddc) had been studied on a mouse macrophage cell line by Makabi-Panzuet al. which shows the high intracellular uptake due to anionic charge on the liposomes. Oussoren et al. synthesized the liposome of the phosphorylated form of zalcitabine and evaluated them in the murine acquired immunodeficiency syndrome model which arrays chemical stability, better retention entrapment, and reduced viral load in mononuclear phagocyte system in both spleen and bone marrow. Various in vitro and in vivo studies were performed by Jain et. al. by entrapping ARV drugs such as acyclovir, indinavir, zidovudine, and lamivudine into the permuted liposomal system which shows high efficacy of transdermal flux compare to free drug. It indicates better permeability of liposomal formulation. The liposomal system of zidovudine shows 12 folds higher concentration as compared to control in 24 hours at the targeted site. There was
higher accumulation at the reticuloendothelial organ system after the introduction of zidovudine loaded elastic liposomes trans-dermally. Further, zidovudine loaded surface engineered liposomes were formulated by Kauret. al for lymphatic targeting. The surface of liposomes was orchestrated by the charges and site-specific ligand to intensify localization to lymphatics, prominently to lymph nodes and spleen. The engineered liposomes charged particle was created using Stearylamine, Dicetyphosphate, and mannose conjugate. Evaluating these three substances, fluorescent microscopy reveals better localization of mannose conjugate than the negative or positive charged liposomes. Liposomes are speedily phagocytosed by the macrophages so to increase prolonged circulation time and bioavailability of the drug its surface is modified with the hydrophilic molecule such polyethene glycol. Clayton et al. created pegylated liposomes with targeting ligand derived from HIV gp120 directed monoclonal antibody F10 and demonstrated as the novel approaches to combat HIV-1. This nano-immuno-liposomes shows greater and longer antiviral activity than the free drug or drug encapsulating nontargeted liposomes. Saiyedet al. prepared magnetic liposomes which contain azidothymidine 5'- triphosphate, the average size of these magnetic liposomes 150nm it is prepared using phosphatidylycholine and cholesterol with a magnetite loading efficiency of 54% and 45.3%. It is studied to check the transmigration across an in vitro blood-brain barrier model and monocyte mediated transport by applying an external magnetic field. The outcome results in apparent permeability of magnetic azidothymidine liposomes were the 3 folds better than the free azidothymidine.[58] Dubey et al. experimented with liposomal Indinavir in the J774.A1 macrophage cell line and modified plain liposomes with covalently coupled bd-1-than-nopyranoside with 1,2-Bis(dimethylphosphino)ethane (DMPE) to generate mannosylated-DMPE (Man- DMPE) and further conjugated with choline and cholesterol to make upgraded mannosylated liposomes. Flow cytometric analysis reveals a better result than the plain liposomes. Cell-derived liposomes show greater and efficient targeting. It is made from the cytoplasmic membranes of the cell expressing CCR5, the human receptor for gp120 that is mainly found on the surface of HIV infected cells and HIV-virion this exhibit significant 60% reduction in the viability in the HIV infected model cell due to binding and nullifying the infectivity. The liposomes are also incorporated with the soluble-CD4, a soluble form of peptide ligand of gp120 shows efficient targeting of infected cells. Pegylatedsaquinavir (SVQ) liposomes were less cytotoxic and sustained drug release efficiency in cell viability assay of Jurkat T- cell. The immune-liposomes loaded with the Heparin-activated serine protease inhibitor antithrombin III (hep-AIII), injected into the nonhuman primate system model. The Result shows the greater than 1(10) log gradual decrease in the plasma viral load which concludes hep-AIII as salvage or alternative agent for HIV strain resistant to standard ARV drug. Hence this critical literature review depicts how liposomes play a significant role in drug encapsulation and work as an effective nanocarrier for drug delivery for the elimination of HIV drug.

**Dendrimers**

A dendrimer is made up of dendrons, a small branching unit that contains interior and periphery end group, it is a polymeric nanostructure composed of several branching units in a layer by layer pattern which characterizes size, growth and the microenvironment within it, as represented in Figure 4.

![Dendrimer](Image)

**Figure 4:** This picture represents the dendrimer structure with its core-shell for ARV drug entrapment.

Dendrimer contains space lying inside the dendron which can be used for the drug entrapment, targeted drug-releasing, protection against degradation from the surrounding environment, specific targeting. The size of the dendrimer is less than 100nm with less polydispersity and higher functionality it is like a conventional polymer with the branched unit in 3-Dimensional architecture. The core can be synthesized by Ammonia and ethylene diamine surrounding by the highly branched repeating units such as polyether, porphyrins, poly-amido-amines, polyphenyl, and polyamine acids. The properties of the core-shell are chiefly based on the multivalent surfaces which contain targeting moieties or functional group so this allows tagging ARV drugs or dendrimer itself can act as the drug after attaching synthetic functional group along with the synthetic peptide polymer which hinders with HIV viruses. A diverse array of dendrimers can be synthesized according to the biological entity such as the HIV antigen for its interaction by doing subtle modification of branching unit type, linker, dendrimer generation, moieties, and surface. The Anti-HIV drug Efavirenz loaded with tuftsin-conjugated fifth-generation poly (propylene imine) (T5PP) dendrimer which shows the prolonged effect of a drug in 24h, negligible cytotoxicity, and cellular uptake 34.5 times higher than the free drug in vitro to the infected macrophages. The 2G-NN16 amino-terminated carbosilane dendrimer use to deliver siRNA to HIV infected astrocytes in vitro which shows less cytotoxicity with controlled drug release. The 2G-NN16/siRNA-dendripplexes successfully cross the blood-brain barrier model revealed by in vitro transcy-
Nanoparticles have increasingly experimented with targeted ARV drug delivery to achieve modulated pharmacokinetics, improved efficacy, decrease in systemic toxicity, and adverse effect. Mainly three types of nanoparticles are employed for anti-HIV curative.

**Polymeric nanoparticles**

A polymeric nanoparticle can be generated as per a desirable approach for targeted delivery of ARV drugs. Various polymers are used for the construction of anti-HIV polymeric nanoparticles such as poly (lactic acid) (PLA), poly (lactic-co-glycolic acid) (PLGA), poly(alkyl) cyanoacrylates,poly (ethylene glycol-co-(lactic-glycolic acid)), poly(caprolactone), and poly(methyl) methacrylate. PLA and PLGA have been approved by the FDA for human use. A diverse drug can be incorporated in these polymers based on their hydrophilicity or hydrophobicity and release characteristics can easily be modified based on the requirements. The zidovudine loaded polyvinylpyrrolidone (PVP)/stearic acid (SA)-polyethylene glycol (PEG) nanoparticles (PSNPs) were developed by emulsification–solvent evaporation method. And studied in vitro murine neuro-2a and HeLa cells which demonstrate significantly, improvement in cellular internalization, stable colloidial suspension, better cellular uptake, increased half-life, no cytotoxicity.

Shah and Amiji developed saquinavir-loaded poly (ethylene oxide)-modified poly(epsilon-caprolactone) (PEO-PCL) nanoparticulate system using the solvent displacement process. The cellular uptake and biodistribution of PEO-PCL were analyzed in vitro into THP-1 human monocyte/macrophage (Mo/Mac) cell line which ends in higher accumulation of drugs than the aqueous form. The electromagnetic interference within the permeability of nanoparticles loaded SQV studied on the human brain microvascular endothelial cells. Here nanoparticles are used as poly butyl cyanoacrylate (PBCA), methyl methacrylate-sulfo-propymethacrylate (MMA-SPM) to study the blood-brain barrier human model which arrays higher permeability across the blood-brain barrier in PBCA nanoparticle than MMA-SPM nanoparticle. Destache et al. studied ARV drug releases from poly (DL-lactide-co-glycolide) NPs in BALB/c mice and compared with free intraperitoneal injected ARV drug such as ritonavir, lopinavir, and efavirenz on mice, which result in higher availability of the drug up to 28 days as compared to conventional one. Furthermore, more tenofovir- loaded chitosan-based nanoparticles fashioned by the Meng et al. to maximize its mucoadhesion. By decreasing the size from 900 nm to 188nm of nanoparticle it shows non-cytotoxicity to vaginal epithelial cell line with increased 6% to 12% mucoadhesion. Hence polymeric nanoparticle shows efficient drug delivery to combat HIV.
Solid lipid nanoparticles (SLN) and Nanostructured lipid carriers (NCL)

SLN are small microscopic system composed of physiological lipid which emerges to form solid nanoparticles on aqueous surfactant solution (size 50 to 1000nm). SLN offers a great opportunity to attach ARV drugs because due to its miniature size, high drug loading capacity, slow degradation of lipid matrices, large surface area reactivity. SLN also provides sustained release minimizing drug toxicity, dosing frequency, and fluctuation in plasma drug levels. SLN shows biphasic drug release due to its structure, initial burst due to its surface adsorption, and slow-release from its lipid core due to gradual degradation.

SLN of protease inhibitors name atazanavir were produced by Chattopadhyay et al. to check permeability and studied on the human brain microvascular cell line (hCMEC/D3) a blood-brain barrier model which successfully results from a higher accumulation of drug by endothelial cell monolayer than the aqueous drug solution with apparent permeability across the barrier membrane. Zidovudine palmitate-loaded SLN develops by Heiat et al. which accommodate trilaurin as the lipid core with a combination of dimyristoyl phosphatidylglycerol which results in neutral charge.

Later it is modified with polyethylene glycol moieties and higher phospholipid content to the surface which results in improved plasma circulation with increases half-life of the drug. The various modification was done to change the surface moieties of the SLN with different methods to achieve higher drug accumulation and significant permeability across the blood-brain barrier. Lopinavir SLN was modified using a hot self-nano-emulsion technique which includes a hot isotropic mixture of stearic acid, poloxamer, and polyethylene glycol in water with rapid cooling which results optimize bioavailability compare to bulk lopinavir. In perfusion experiment, a high amount of positively charged or negatively charges SLN will result into a high cortical cerebrospinal volume which loses the integrity of brain barrier in rats. This tends to make attention that high amount of surface charges modification in SLN will increase the adverse effect on health.

NCL is the fashioned or tailored SLN with solid lipid matrix incorporated with liquid lipids with various fatty acid chain in compromised organized crystalline structure which offer higher drug accommodation capacity. NCL is made up of low toxic lipid molecules that provide hydrolytic and oxidative stability. It also shows the biphasic drug release potential with a liquid lipophilic surface which contains drug and solid core with a higher melting point which allows drug release by diffusion and matrix erosion. Lots of NCL have been manufactured in the current scenario to treat HIV. Advance formulated NCL, prepared by Kasongoet al., is the mixture of Precirol ATO 5 and trans cutol loaded with didanosine nanostructured lipid carrier manufactured using hot high-pressure homogenization method dried using liquid nitrogen and then dried material were pass through 200µm membrane which results into solid lipid particles in surfactant solution which results in high relative stability for 2 months at 25°C and increased encapsulation efficiency.

Inorganic nanoparticles

This class of nanoparticles includes metal elements like iron, gold, silver, titanium, and silica which nowadays employed in anti-cancer therapeutics, molecular labeling of biomarkers, theranostics approaches, bioimaging, biosensor, Nanoparticles of a noble metal such as gold, silver, and platinum have been formulated using diverse methods such as chemical bioreduction, hard template, solution-phase synthesis, Gas-phase deposition, and sol-gel. The silver nanoparticle is getting more prominent attention due to its anti-microbial and anti-viral effects against hepatitis B, herpes simplex virus, respiratory syncytial virus, monkeypox virus, and HIV-1 in vitro including clinical isolates and resistant strains. Silver nanoparticles can bind to the gp120 protein and prevent viral entry, it also inhibits the CD4 mediated viral fusion and interfere in post-entry stages of the invasion of the HIV life cycle. The Silver nanoparticles provoke higher antiviral efficiency and therapeutic index as compared to silver ion salts of sulfadiazine.

Another potent gold nanoparticle arrays anti-HIV activity, when conjugated with the HIV inhibitors such as TAK-779 and SDC-1721 which results in better anti-HIV activity than its free form. Inorganic nanoparticles possess limitations such as cytotoxicity, DNA damage, causes cellular apoptosis as evaluated by membrane leakage assay and LDH assay. Furthermore, gene chip (microarray) analysis stipulated an induction of a large number of genes, particularly, stress associated genes coding metallothionein and heat shock protein. This review suggests formulating such inorganic nanoparticles which is less toxic to the mammalian cell and must be enhanced with modified advanced hybrid technology to achieve greater efficiency.

Polymeric micelles

Polymeric micelles are the nano-engineered product of block polymer and having core-shell just like surfactant-based micelles and have been utilized for improving permeability, aqueous solubility, protection towards chemical degradation, controlled drug release, offer surface modification due to hydrophobic moieties as mention in Figure 6.

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Polymeric micelles are architected as hydrophobic core and hydrophilic shell which allows entrapping anti-HIV drugs based on their polarity. Additionally, surface properties of polymeric micelles such as hydrophilic block can be modified by docking antibodies or another chemical ligand specific for receptors present on the diseases like HIV-AIDS. A lot of pharmaceutical scientists have formulated polymeric micelles loaded ARV drug such as lamivudine conjugated with stearic acid-g-chitosan oligosaccharide micelles by esterification method which results in pH-dependent drug release, low cytotoxic effect, higher cellular uptake against hepatitis B virus-infected HepG2.2.15 tumour cells. Polymeric micelles of Efavirenz evinces significant absorption rate and 3 folds increase in the pharmacokinetic parameters of C-max values from 1789, and 2657ng/ml to 2856, and 7056ng/ml in single-dose between 20 and 80mg/kg given to healthy adult volunteers. Hence polymeric micelles show the greater ability for Anti-HIV drug formulation.

Figure 6: This pictorial representation shows polymeric nanoparticle for drug entrapment and shows targeting moieties for focused ARV drug delivery.

Nanocrystals

Nanocrystal drug itself could be a nano-size drug particle that may be disseminated in aqueous or non-aqueous media. Drug nanocrystals are often formulated using methods that supported a top-down approach or a bottom-up approach. Top-down approaches like media milling and high-pressure homogenization are the foremost preferred methods for the generation of nanocrystals because of their amenability for large-scale production. Nanocrystal drug provides longer colloidal stability, prolong, and persistent targeting due to increased surface area. Different platform technologies such as Nano-Crystal®, Nanopure®, and NANOEDGER® have been flourished for the formulation of drug nanocrystals. Transformation of the pure drug to nanoscale particle leads to increased parameters of pharmacokinetics, bioavailability, surface area reactivity. Nanoscale engineering of pure drugs is developed because of the intensely hydrophobic drug that is strenuous to deliver as a solution intravenously or drugs exhibiting dissolution rate-limited oral bioavailability. With the help of the media milling technique, Baret et al. formulated Rilpivirinenanocrystals of 200, 400, and 800 nm were injected in mice and dogs by intramuscular and intra-subcutaneous routes and their pharmacokinetic behaviour was monitored. Experimentally, all treatment results in significant detectable rilpivirine levels up to 90 days in dogs and 3 weeks in mice indicating their efficacy in long-term prophylaxis of HIV. The author’s study also indicates that the intra-subcutaneous route of administration shows stable plasma concentration while intramuscular shows peak level concentration as well as higher clearance. Rilpivirine concentrations were also detected in lymphoid tissues throughout the treatment, stimulating the uptake of nanocrystals by macrophages.

Hence this review suggests that nanocrystal drugs show better potentiality to treat HIV. However, the route of administration matters in nanocrystal drug delivery.

Miscellaneous- Nanoparticle utilize for (LRA) latency-reversing agents

ARV drugs play a major role in managing the HIV infectious condition; However, it is not possible to cure HIV-AIDS because due to its latency. HIV infected cells such as macrophages or monocytes which reside in the bone marrow, cerebrospinal fluids, lymph nodes which are considered as the reservoir of the human body, LRA are not able to available at this site due to its low bioavailability. Nanotechnology plays a better role in transporting LRA to the reservoir site by peptide self-assemble nanoparticle loaded with Panobinostat. Panobinostat is a highly potent hydroxamic acid Histone deacetylase inhibitor that binds to the specific target of the DNA to reactivate the silenced proviral genome; it is used in the clinical development of multiple myeloma. Experimentally it shows better blood-brain barrier permeability, effective reactivation of silenced proviral genome in vitro, increased activation of peripheral blood mononuclear cells from latently infected patients ex vivo, increased cellular drug uptake more accurate than the free Panobinostat alone. Therefore, this strategy exhibits that nanotechnology can aid to improve the activation of latent HIV, and this study lays a foundation for further enhancement of LRA delivery systems to combat silenced HIV proviral genome.

CONCLUSION

In this literature review, we have discussed the potentiality of a nanotechnology-based drug delivery system to provide an empirical application for anti-HIV therapy. Nanotechnology-based system development of ARV drugs offers wide and efficient targeted drug delivery with modulated pharmacokinetics, a higher therapeutic index as demonstrated by in vitro and animals in vivo studies. These nano-systems provide prolong drug circulation, high bioavailability, drug stability, better permeability, bioaccumulation in known reservoir sites for HIV. It also demonstrated the application of ARV nanocarriers to deliver drugs across the blood-brain barrier and other tissue to kill HIV. Based on the HIV lifecycle, di-
verse nanocarriers are surface modified with different moiety to prevent viral fusion with intended ARV drug delivery. The majority of works done in the field of nanocarrier ARV drug delivery system incorporate a single ARV agent. So, this review tends to notice about the multidrug delivery system which involves a combination of drugs can lead to tremendous efficacious treatment and downgrading of resistance profiles. Hence, nanotechnology provides a multifunctional system for scaling up therapeutic approaches with innovative formulation to fulfill diverse biological requirements.

Acknowledgement: The authors wish to express their acknowledgement to the I/C Principal TrilokAkhani PIAS, Parul University & Dr Iivala Anand Shaker, Professor & HOD Dept. of Biochemistry, PIMSR, Parul University for their constant help and support throughout in preparing this manuscript. The authors are also grateful to publishers of all those articles, journals and books from where the literature for this article has been reviewed and discussed.

Conflict of Interest: Nil

Source of Funding: Nil

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Baghel et al.: Nanotechnology: A curative approach to Combat HIV-Aids


