Heat Shock Protein–Based Therapeutic Strategies Against Human Immunodeficiency Virus Type I Infection

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ABSTRACT

Heat shock proteins (hspks) and cyclophilins (CypA) are intracellular chaperone molecules that facilitate protein folding and assembly. These proteins are selectively expressed in cells following exposure to a range of stress stimuli, including viral infection. Hsp species are highly immunogenic, eliciting humoral, cytotoxic T lymphocyte (CTL), and natural killer (NK) cell responses against viruses, tumours, and infectious diseases. This review discusses the roles of stress proteins in immunity and viral life cycles, vis-à-vis the development of Hsp-based therapeutic strategies against human immunodeficiency virus type-1 (HIV-1) infection. Cumulative findings are cited implicating the requirement of CypA in HIV-1 replication and formation of infectious virions. Studies by our group show the upregulated expression of hsp27 and hsp70 during single-cycle HIV infections. These species redistribute to the cell surface following HIV-infection and heat stress, serving as targets for NK and antibody-dependent cellular cytotoxicity. Co-immunoprecipitation and Western blot studies show that hsp27, hsp70, and hsp78 complex with HIV-1 viral proteins intracellularly. Hsp70, hsp56, and CypA are assembled into HIV-1 virions. The ability of hsps to interact with HIV-1 viral proteins, combined with their inherent adjuvant and immunogenic properties, indicates that hsps may serve as vehicles for antigen delivery and the design of vaccines against acquired immunodeficiency syndrome. Infect. Dis. Obstet. Gynecol. 7:80–90, 1999. © 1999 Wiley-Liss, Inc.

KEY WORDS
viral-host interactions; AIDS; cytotoxic T lymphocyte; NK immunity; cyclophilin A

While combination antiviral drug therapies have led to major advances in the management of acquired immunodeficiency syndrome (AIDS), 90% of people infected with human immunodeficiency virus (HIV) worldwide have no access to such treatments. Development of vaccines against HIV type-1 (HIV-1) remains the outstanding challenge in AIDS research.1,2 Generation of AIDS vaccines has been hampered by HIV antigenic diversity, the multiple routes and modes of HIV transmission, and ethical constraints restricting use of whole-killed or live-attenuated viruses. To date, inactivated virus and subunit vaccines have failed to elicit impressive protective cytotoxic T lymphocyte (CTL) or humoral immunity capable of neutralizing a wide spectrum of HIV-1 isolates. This has been attributed to the limited immunogenicity of viral proteins and their failure to encounter major histocompatibility complex (MHC) restriction elements required for T cell recognition.1,2 The immunogenic, carrier, and adjuvant properties of heat shock proteins (hsps) have been exploited as vaccine vectors to elicit protective immunity against cancers and microbial infec-

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tions. This review describes the involvement of hsps in immunity and viral life cycles, vis-à-vis their potential application toward the development of Hsp-based antiviral and vaccine strategies against AIDS.

HEAT SHOCK PROTEINS FUNCTION AS INTRACELLULAR MOLECULAR CHAPERONES

The Hsp family of proteins, representing 2–15 % of total cellular protein, are among the most highly conserved proteins present in procaryotic and eu-caryotic organisms.3–5 Classified by apparent molecular weight (hsp70, hsp60, hsp90, hsp27), hsps were first defined as those proteins selectively synthesized following cellular exposure to temperatures 5 to 10°C above normal.3 Hsps are now referred to as stress proteins since they are induced in response to a vast spectrum of physiological and environmental insults, including viral infection, inflammation, fever, malignant transformation, and cellular exposure to oxidizing agents, cytotoxins and anoxia.5–6

Stress proteins function as chaperone molecules having innate abilities to bind to a broad range of cellular peptides, proteins, and multimeric complexes.6–8 Constitutive Hsp subspecies, referred to as “housekeeping” proteins, direct protein folding, biogenesis, assembly, trafficking, and degradation. Inducible Hsp subspecies rapidly reprogram cellular metabolic, proliferative, and functional events in response to distinct stress stimuli.

Recent advances have revealed the mechanisms of actions of the various Hsp species.7 Hsp70 homologues consist of peptide-binding and ATPase domains that stabilize protein structures in unfolded and assembly-competent states for extended periods of time.4–8 In contrast, mitochondrial hsp60 isoforms form ring-shaped oligomers wherein protein assembly to native states is facilitated.4 Hsp90 species associate with cellular tyrosine kinases, transcription factors, and glucocorticoid receptors, playing suppressor regulatory roles.5–7 Hsp27 proteins suppress protein aggregation, protect against actin polymerization, and represent end components of stress and cytokine kinase (MAPK) cascades.10–13 Binding of the ubiquitin species targets protein complexes for degradation.4–5 Immunophilins, including cyclophilins and FK binding proteins (FKBPs), catalyze cis/trans isomerization of peptidyl-prolyl bonds, key rate-limiting steps in protein folding.3

HSPS PLAY CENTRAL ROLES IN IMMUNE RESPONSE

Besides their biological roles as chaperonins and stress proteins, hsps are extraordinarily immunogenic. Stress proteins have been implicated in host immunity to tumours and viral and microbial infections.14–17 Select Hsp species represent major targets for humoral (B cell) and cell-mediated (αβ and γδ T, natural killer [NK], and lymphokine-activated [LAK] cell) immunity.14–27 The immunogenic properties of hsps have led to considerable interest in their application in vaccine strategies.16–27 Vaccinations with Hsp-peptide complexes have been shown to augment CTL, γδ T, NK, and LAK cell responses against tumour and viral antigens.16–27 Reasons why highly conserved stress proteins play such a prominent role in immunity are still not well understood. Based on in vitro and in vivo studies, several distinct mechanisms wherein hsps induce viral and tumour immunity have been identified.

Stress Proteins Can Act as Classic Antigens

Major portions of the immune repertoire can recognize microbial Hsp and autologous Hsp determinants generated following stress, viral infection, or malignant transformation.14–17 Hsp70 and hsp60 are the predominant antigenic species recognized in many microbial infections, including malaria, leprosy, and tuberculosis. Analysis of immune responses to mycobacterial infections indicates that 20–30 % of the overall CTL and T cell repertoire are directed against Hsp moieties.16–21 Indeed, prophylactic vaccinations with microbial hsps have been shown to generate significant protective immunity against bacterial, fungal, and mycobacterial infections.14–17

Stress proteins are impressively upregulated by both host and pathogen at sites of infection, enhancing Hsp-directed responses. That highly conserved hsps serve as classic foreign antigens is substantiated by correlative links between immunity to foreign microbial hsps and the emergence of Hsp autoimmune reactions in rheumatoid arthritis, systemic lupus, Graves disease, diabetes mellitus, inflammatory diseases, and neurological aging.28 Autoimmune pathology has even been postulated
Fig. 1. Several possible mechanisms wherein hsps may elicit antiviral immunity. 1) Hsps may redistribute to the cell surface in virally-infected cells; 2) Hsp-viral complexes (HIV-VP) may be present on the cell surface; and 3) antigen (Ag)-presenting cells, including macrophages and dendritic cells may incorporate viral peptide-Hsp complexes allowing viral peptides to be expressed in the context of MHC class-I determinants to T-cell receptors (TCR) on CTLs.

to contribute to the ongoing clearance of stressed or infected CD4 cells in AIDS (Fig. 1).23,24

Stress Proteins Can Redistribute to Cell Surface Membranes Following Infection or Transformation

Cytosolic stress proteins, including hsp27 and hsp70, have been shown to relocate to the plasma membrane in transformed, virally-infected or heat-stressed cells.22–26 The de novo distribution of hsps on the plasma membrane specifically targets tumour and HIV-infected cells for elimination by antibody-dependent cellular cytotoxicity (ADCC), NK, LAK and αβ T cell-mediated immunity (Fig. 1).22–26

Hsps Can Serve as Adjuvants and Vaccine Vectors

Hsps can indirectly enhance immune responses to tumours and viral antigens because of their adju-
vant properties. The immunostimulatory properties of the mycobacterial-derived constituents in Freund's adjuvant and Bacillus Calmette-Guérin (BCG) have been attributed to and/or mimicked by adjuvant-free hsp60 and hsp70 carrier proteins.16,17 The strong adjuvant effects of hsps when conjugated to peptides and oligosaccharides have been exploited to stimulate immune responses in a variety of infections.16,17,29–32 Similarly, BCG may be useful as live vaccine vector for HIV-1 constituents.29,30 Mice that have been vaccinated with fused hsp70-HIVp24 complex elicit humoral and CTL immune responses to the HIV-1 capsid proteins for over 1 year, with no corresponding immunity evoked in mice injected with soluble p24 protein.31

**Stress Proteins Can Be Integrally Involved in Antigen Presentation**

Soluble proteins are weakly immunogenic due to their failure to effectively elicit MHC-restricted CTL responses. This has been attributed to the failure of soluble proteins to be incorporated by professional antigen-presenting cells, such as macrophages. In contrast, associations of tumour or viral proteins with hsp70, gp96 or hsp90 moieties stimulate antitumour and antiviral CTL immunity (Fig. 1).19,31–36 It appears that the interactions of hsps with tumour or viral antigens allow for their effective uptake into professional antigen processing cells, such as macrophages, allowing for the processing of antigenic complexes along the MHC class-1 pathway for CTL presentation.

The precise role of hsps is unclear but has been referred to as Hsp-mediated "cross-priming," insofar as the CTL responses are directed against the peptides bound to the hsp and not the hsps themselves.33,34 Hsps appear to serve as antigenic carriers that route exogenous proteins into macrophages, wherein proteins are cleaved and translocated for class 1 MHC presentation. Hsp70, gp96, and hsp90 subspecies, like MHC determinants, can chaperone tumour and viral peptides of 400–2000 Da intracellularly.37 Hsp70 and gp96 moieties are also implicated in the folding of MHC molecules.33–37 Indeed, hsp70 species show considerable homology with MHC class 1 molecules, and hsp70 genes have been localized to the MHC gene complex.34–38 This has led to a postulated role of Hsp moieties in peptide charging of MHC molecules.33–37,39 Immunizations of animals with hsp70 and hsp90 complexes from virally-infected or tumour cells elicit protective antiviral and tumour-specific CTLs and memory cells.35–37,39–46

**STRESS PROTEINS IN THE IMMUNOTHERAPY OF CANCERS, INFECTIOUS DISEASES AND AIDS**

The antigenic, adjuvant, and carrier properties of hsps have heralded their potential application as "magic bullets" in cancer and viral immunotherapy.44–47 Hsp-tumour conjugates and Hsp complexes isolated from cancer cells have been demonstrated to produce tumour-specific CTL responses and protective antitumour immunity in animals.35–37,39–40,42–47 It has been postulated that vaccination with Hsp-peptide complexes purified from tumour cells will 1) maximize the tumour antigenic repertoire in immunized cancer patients; 2) circumvent the requirement to identify and purify unique tumour antigens; 3) eliminate inherent problems of MHC disparity among individuals; and 4) obviate the necessity for adjuvants to stimulate immunity. Although still at the formative stage, Hsp-peptide complexes have been heralded as novel approaches to customize cancer therapeutic vaccines from patients' own cancer cells. Phase 1 clinical trials are currently in progress.44–47

Cell culture and animal model systems have shown that Hsp complexes purified from virally-infected cells can generate viral-specific CTLs and/or protective immunity against simian virus 40, influenza, vesicular stomatitis, and lymphocytic choriomeningitis viral infections.35,41,44 Young et al. are currently exploring the applicability of HIVp24-hsp70 conjugates as vaccines in monkeys.16,30–32,36 Given the vital necessity for an AIDS vaccine, establishing the roles of hsps in HIV replication, assembly, and immunity are warranted.

**THE SPECIFIC ROLE OF STRESS PROTEIN SUBSPECIES IN HIV-1 VIRAL REPLICATION**

Several lines of investigation have implicated stress proteins in viral-host interactions in many DNA and RNA virus infections.5,20,24,48,49 These include demonstrations that:

1) Stress protein expression correlates with levels of viral expression and replication.
2) Stress proteins facilitate rate-limiting steps in viral replication.
3) Hsps affect the folding and assembly of viral intermediates and virions.
4) Incorporation of stress proteins into viruses determines virion infectivity.
5) The presence of stress proteins on the cell surface elicits antiviral immunity.

It is to be expected that the HIV-1 virus may have an obligatory requirement for host stress proteins to facilitate their replication and assembly in CD4 cells. Studies showing involvement of stress proteins in the HIV-1 life cycle will be described, vis-à-vis the potential application of hsps in AIDS antiviral strategies.

MODULATION OF HSP27 AND HSP70 EXPRESSION IN CD4+ CELLS CONSEQUENT TO HIV-1 INFECTION

Viral-induced activations of specific stress genes have been described during latent, productive, and lytic phases of infection with DNA and RNA viruses. It has been shown in many systems that induction of hsp70 and hsp90 genes are among the earliest changes in cellular gene expression following viral infection. Induced expressions of select hsp70 moieties with different viruses suggest viral-dependent hsp70 gene activation.

The HIV-1 virus differentially infects CD4 cells of lymphocytic and monocytic origin through the CD4 and chemokine surface receptors (Fig. 2). Studies in our laboratory have used Northern and Western blot analysis to chart changes in levels of...
Presence of Hsp70 in HIV-1 virions relative to p24 and Pr55gag capsid moieties

A. Blot A probed with αp24 antibodies

B. Blot B probed with αHsp70 antibodies

C. Blot A reprobed with αHsp70 antibodies

Fig. 3. Western blot analysis showing the presence of A) HIV-1 p24 reactive capsid determinants in virion particles; B) hsp70 in purified HIV-1 virions; and C) levels of hsp70 relative to HIV-1 gag proteins.

Hsp mRNA and protein expression in CD4 cells during single-cycle HIV-1 infections. Acute HIV infection of H9, GEM, Jurkat and MT-2 lymphocytic cell lines leads to eight- to 20-fold increases in levels of hsp27 and hsp70 mRNA and protein expression at early stages of infection (1–6 hr) subsequent to viral penetration but prior to viral mRNA synthesis that occurs 36–48 hr postinfection. Initial increases in hsp27 and hsp70 are down-regulated during productive stages of viral synthesis. Elevations in levels of these Hsp species recur at lytic endstages of infection, concomitant to virion release and cytolysis. Induction of hsp27 and hsp70 synthesis are viral dose-related and abrogated by HIV-1 neutralizing antibodies, CD4 antibodies or inactivation of virus. In direct contrast to that observed with hsp27 and hsp70 species, levels of hsp60 and hsp90 mRNA and protein moieties remain unaltered throughout the course of HIV infection. Two-dimensional gel electrophoresis reveals that the hsp70 isoforms induced following HIV infection show distinct isoelectric mobilities from constitutive and heat-inducible hsp70 species.

Studies by a number of investigators have assessed correlative changes in levels of HIV-1 and Hsp gene and protein expression in infected cells. Somewhat contradictory results have been obtained. Stress stimuli, including heat, tumor necrosis factor-α (TNF-α), and interleukin 2 (IL-2), can induce in Hsp and viral gene transcription with similar kinetics. However, heat shock and TNF-α induce states of cellular latency, wherein enhanced Hsp synthesis markedly inhibits overall levels of viral mRNA and protein synthesis. Cyclopentenone prostaglandins (prostaglandins A and J) have also been known to block human T-cell lymphotrophic virus (HTLV-1) and HIV-1 viral replication through induction of an hsp70 stress response. Hyperthermia at 3–4°C above physiological range can block HIV-1 viral replication in vivo and has led to the postulated benefits of localized or systemic hyperthermia in HIV infection.

HSP27 AND HSP70 EXPRESSION ON THE SURFACE OF HIV-1 INFECTED CELLS

Apart from their increased intracellular levels of expression during infection, significant amounts of hsp27 and hsp70 have been shown to redistribute to the cell surface of CD4+ cell lines following heat stress or HIV-1 infection. The enhanced expression of HIV-1 on the cell surface contribute to the enhanced susceptibility of HIV-infected cell lines to NK, LAK, and HIV-specific ADCC cell-mediated cytotoxicity when compared to that observed for their uninfected counterparts (Fig. 1).
This enhanced susceptibility of HIV-infected lymphocytes to natural immunity may contribute to antiviral immunity in vivo. However, it should be noted that basal and IL-2 inducible NK cell-mediated cytotoxicity has been shown to decline progressively with advancing HIV disease.63-68 Indeed, inducible NK cell function is among the most sensitive functional correlates of immunological dysfunction in HIV infection.

**INTERACTIONS OF STRESS PROTEINS WITH HIV VIRAL PROTEINS AND VIRION PARTICLES**

The association of stress proteins with viral intermediates can facilitate their folding, assembly, and morphogenesis with a direct impact on rates of viral production and infectivity. Biosynthetic assembly of the HIV-1 envelope protein glycoprotein 160 (gp160) to its native state has been shown to involve the cellular chaperonins grp78-BiP (the endoplasmic reticulum hsp70 species) and calreticulin.69,70 These stress proteins bind in a specific and transient fashion and are released only after the envelope protein assumes its native conformation.

Our laboratory has used reciprocal co-immunoprecipitation approaches to show that hsp27 and HIV-1 capsid (gag) moieties form intracellular complexes (manuscript in preparation). Similarly, hsp70 can be shown to be complexed with HIV-1 gag precursor moieties that can be precipitated with HIV-1 p24 antibodies.

The strongest evidence that stress proteins are fundamentally required for HIV-1 virion assembly and infectivity come from studies with the chaperonin, cyclophilin (CypA). CypA has been shown to specifically interact with the HIV-1 capsid (gag) protein and is essential for the production of infectious HIV-1 virions.49,71-73 CypA weakens gag-capsid strips and promotes capsid disassembly following HIV-1 penetration into cells.49,74 Studies with cyclosporins show that the failure to incorporate CypA in HIV-1 virions abrogates the infectivity of viruses.49,75 CypA is found to associate with some HIV-1 clades and not others and is generally absent in simian immunodeficiency viruses.76 Levels of HIV-1 virus infectivity and CypA in virions are directly related to cellular levels of CypA expression.77,78 Recent studies indicate that CypA...
and FKBP can also bind to the V3 loop of HIV envelope (env) protein. Indeed, HIV-1 infection of peripheral blood lymphocytes can be blocked by an excess CypA or neutralizing antibodies to CypA. Studies in our laboratory have determined that select Hsp subspecies also associate with HIV-1 virions purified from HIVIIIb-infected (CEM, H9, Jurkat) (manuscript in preparation). HspT0 (inducible) proteins are specifically incorporated into HIV-1 virions (Fig. 3). These findings contradict earlier studies by Bartz et al., who found hsp60 but not hsp70 moieties in polyethylene glycol precipitated viruses isolated from HIV-1 and SIV virions. Using subtilisin extraction procedures, we show that hsp70 is retained in virions while contaminating microvesicular elements are degraded. Thus, hsp60 may be associated with HIV-1 virion assembly intermediates but are not present in purified virions.

Like that reported for CypA, we show hsp70 to be notably absent in SIV virions. We have also isolated capsid virus-like particles from COS-7 cells transfected with vectors producing Pr55 or Pr160 gag capsid precursor proteins. Hsp70, FKBP5 (hsp56), and CypA are found to be present in gag-capsid particles.

**SUMMARY**

Taken together, these results show correlative links in stress protein and HIV replicative pathways in CD4 cells (Fig. 2, Fig. 4). Infection with HIV-1 viruses induces a stress response and stress agonists, such as heat, IL-2 and TNF-α, affect HIV-1 replicative events. Hsp27 and hsp70 stress pathways are induced in CD4 cells concomitant to HIV-1 infection, and these species can distribute on the cell surface of infected cells. Cumulative evidence indicates that hsp70, hsp78, calreticulin, FKBP5, and CypA are among those stress protein subspecies found to be associated with HIV-1 viral proteins or virions.

Having documented existent interactions between stress proteins and HIV-1 viral constituents, it is feasible that Hsp-viral complexes may offer a number of unique vaccine approaches against AIDS. The inherent adjuvant properties of hsp70 may enhance the immunogenicity of HIV-1 proteins and allow exogenously-presented HIV-1 viral proteins to be presented for CTL presentation. Specific HIV subunit vaccines conjugated to hsp70, hsp78, and gp96 proteins may enhance CTL immunity. Immunization with heterogeneous mixtures of Hsp-viral antigenic species may prevent the emergence of CTL escape mutants. Hsp-viral complexes may customize immunogens in distinct HIV-1 infected populations.

Preliminary clinical trials of a number of Hsp-based vaccines against cancer and microbial infections are ongoing. Phase 1 trials for the use of Hsp-24 gag fusion complexes as an AIDS vaccine are in place. Given the difficulty, to date, in generating an AIDS vaccine, investigations into the potential use of Hsp-based AIDS vaccines are warranted.

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