Tat Oyi-based candidate therapeutic vaccine: a phase 1 clinical trial in HIV-1 infected patients

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**Tat is a virulence factor of HIV**

Transactivator of transcription (Tat) of HIV-1 is essential for the viral gene expression and productive infection

- Tat protein expression is a first step of the virus life cycle to transactivate its own expression
- It enters the nucleus and then can regulate the expression of host genes which impact the immune system
- Nearly two-thirds of Tat made by infected CD4+ T-cells are secreted into the extra-cellular milieu and the extracellular Tat can be taken up by cells (Rayne EMBO 2010)
- Tat can induce viral expression from latently infected cells

Extracellular Tat acts as toxin by interacting with many cellular pathways in uninfected cells:

- Directly involved in immunosuppression
  - i.e. triggers apoptosis in T-cell, inducing Fas on macrophages
  - Tat Binds to integrins and favors viral entry in T-cell, monocytes and DC
- Directly involved in immunosuppression and pathogenesis
  - i.e. Tat has been shown to play a direct role in inducing neuronal death

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HIV provirus

CD4

LTR
gag pol env
LTR

TAR
RNA

Rev
Nef

Others proteins

Virions assembly

Tat secretion
Why a Tat based vaccine

- Tat is good target for inhibition as the early activator of the retrovirus expression
- Tat Inhibition would block efficiently the viral life cycle and virion release
- It could inhibit the reactivating of latent viral reservoirs
- Few attempts have been made to develop pharmacological inhibitors of Tat

- Neutralization of extra cellular Tat would prevent the binding of Tat to cell membranes and receptors
- To Inhibit the toxic activities of Tat on immune cells and pathways, leading to restoring immune functions or decreasing pathogenicity

- Neutralization of Tat during infection could be an important objective, making Tat a potential therapeutic vaccine candidate.
  - Tat still secreted under ART (S. Mediouni Inf. Dis. 2012)
  - Blocking replication of latent virus in reservoir cells during HAART
Immune response to Tat could protect subjects from disease progression?
Studies suggest that presence of Antibodies to Tat could correlate with slower progression long term survival in human

- Re et al. 1995: Retrospective study in 10 HIV hemophiliac patients followed for 10 years: inverse correlation between Anti Tat Ab and clinical progression
- Re et al. 2001: 44/53 Drug users patients with ART, had Ab to Tat (Tat full length 1-80); inverse correlation between high level of Tat Ab and viral load as measured by HIV RNA.
- Zagury et al 1998: in 182 non progressors NP (26 P-NP) follow up 1-2 years. Inverse correlation (p=0,001) between Tat Ab titers and p24 Ag, CD4 level and clinical signs of progression
- Richardson et al 2003. GRIV cohort: Antibodies to Tat and not to Vpr associated with maintenance of long-term non-progression (epitopes in Tat :N ter, basic domain and C ter Tat). A correlation was observed between anti-Tat IgG titers and cross-reactivity with Tat from diverse viral isolates, including HIV-1 subtype-E (CMU08) and SIVmac251 Tat .
- Rezza et al J. inf Dis 2005. : Ab to native Tat IIIB retrospective study of 252 subjects mean follow up 7,2 years: anti-tat positive subjects (11%) had a 60% lower risk of disease progression.
- Senkaali D,et al. AIDS Res Hum Retroviruses. 2008: No correlation between Tat Ab and disease progression in a cohort in Uganda when using different variants of Tat proteins.
- Bellino et al. Retrovirology 2014. Prospective study in 61 asymptomatic drug-naïve HIV-infected adult volunteers same baseline for CD4, follow up to 42 months, 20/60 with anti Tat Ab, 11 with high persistent levels and 9 with lower level Tat Ab. Inverse correlation was found between high Tat Ab titers and CD4 decline, increase in viral load and start of HAART.

Conflicting results suggest that anti-Tat antibodies when detectable are not necessarily efficient against Tat
CTL to Tat correlate with slower progression?

In 130 children from infected mothers: 9 infected newborns followed for 18 months, 6/9 had CTL of HIV proteins, Tat specific CTL was the earliest immune response and it was associated with better clinical profil (NS).

In 12 seropositives (9 seroconverting) follow up 10 years, frequency of CTL specific for Tat and Rev significantly higher in NP than Rapid Progressors (no correlation with CTL to gag, RT or nef).

In 57 HIV seropositive: CTL to Tat 19% to Rev in 37%. Controllers targeted more CTL epitopes within HIV-1 Tat, compared with the treated individuals (P < 0.03), and responses directed against these epitopes were of significantly higher magnitude in controllers (471 ± 270 SFC/10^6 PBMC vs. 156 ± 71 SFC/10^6 PBMC, P = 0.01).
Tat based vaccine trials in animal models

Earlier Studies of Tat vaccines in animal models did not show protection against high dose or iv challenge:

- Allen et al 2002: Full Tat SIV Mac 239 or tat 28-35 DNA immunization with vaccinia virus induce a CTL specific response but no protection.
- Richardson et al., Silvera et al 2002: Despite specific Ab and cell response in rhesus macaques immunized with unmodified HIV-1 IIIB Tat, SHIV89.6P Tat, and carboxymethylated IIIB and 89.6P Tat toxoids no protection or effect on viral replication.

Several studies reported a reduction in the rate of infection or in the level of viral replication after low dose or intraoral challenges with native Tat vaccination in homologous challenges:

- Pauza et al. 2000. Tat IIIB toxoid or native vaccination attenuated SHIV 86 viral load in 10 Rhesus macaques.
- Cafaro et al. 2010 A retrospective analysis of 112 Mauritian cynomolgus macaques from different preclinical trials, vaccinated (n = 67) or not (n = 45) with Tat (HXB2 tat 86) and challenged with the SHIV-89.6P.

Partial protection could be achieved with multi-component Tat Env vaccines:

- Monini et al 2012. In cynomolgus macaques co-immunized with HIV-1 Tat and Env proteins and challenged intrarectally with a high dose (70 MID$_{50}$) of the R5-tropic SHIV$_{SF162P4cy}$.
- Lakhashe SK et al. 2011. A multi-component vaccine (multimeric HIV-1 gp160, HIV-1 Tat, and SIV Gag-Pol particles) delivered systemically or mucosally and challenged orally or IR with the Heterologous C clade R5-tropic SHIV-1157ip:
  - A sterilizing immunity 2/12 and control of infection (-1 log viral RNA ) was observed in 4/12 immunized rhesus macaques.
  - Protection correlated with strong humoral and cellular immune response to Tat or Gag.
- Bachler BC et al. 2013. Novel biopanning strategy to identify epitopes associated with vaccine protection: only the 6 protected animals had developed antibodies binding a dominant epitopes in Tat.

- Induction of anti-Tat antibodies may be key to achieve protective immunity against HIV.
**Tat vaccine clinical trials**

Immunization with peptides, whole native Tat or Tat toxoid has been reported to induce modest immunogenicity but no evidence of control of HIV replication

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**Tat peptides TUTI-16.**

Tat B cell epitope (Tat 4-12), short peptides with known variant amino acids at variable positions 7, 9 and 12 (plus a promiscuous T helper sequence and a lipopeptide toll-like receptor 2 (TLR2) agonist)

- TUTI-16 in a recent randomized double-blind trial was immunogenic, with high levels of anti-Tat antibodies
- Of 21 immunized subjects, 13 (62%) had HIV rebounds vs. 8 (38%) that remained aviremic after ART cessation, but this distribution was not vaccine-related (p = 0.61)

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**HXB2 Tat ISST-02**

Biologically active HIV-1 Tat protein in 168 HAART patients follow up 3 years

- Vaccination promoted anti-Tat IgM and IgG Abs able to neutralise Tat activity in vitro (Env entry in DC cells)
- Induction T-cell response to Tat epitopes
- Compare to a control cohort: CD4 mean increase was 100 c/µl at year 2 and 3
- A significant reduction of blood proviral DNA was seen after week 72, with Tat (30 μg, 3x), in the group of patients treated with PI inhibitors.

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Goldstein G, Chicca J. Hum Vaccin Immunother. 2012 HIV-1 Tat B-cell epitope vaccination was ineffectual in preventing viral rebound after ART cessation

Enoli F. et al. Retrovirology 2015;12:33. HIV-1 Tat immunization restores immune homeostasis and attacks the HAART-resistant blood HIV DNA: results of a randomized phase II exploratory clinical trial.

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All Tat vaccine were safe, linear epitopes did not induce a protective Ab response, active TatHXB2 immunization could result in some level control of replication
Issues to solve for a Tat vaccine

- Heterogenicity variability of genomes
- Intrinsic moderate immunogenicity of Tat
- Finding the epitopes that could Induce a protective immunity
Huet T, Jazza MC, Brun-Vézinet F, Roelants GE, Wain-Hobson S. AIDS. 1989 A highly defective HIV-1 strain isolated from a healthy Gabonese in a group of pregnant women presenting atypical western blot (no Ab to Env) and stay asymptomatic for at least 2 years.

HIV-1 Oyi variant replicated poorly in PBMC coculture, it was recognised by anti HIV-1Env Ab.

Oyi strains has a mutated Tat protein unable to transactivate HIV transcription (the Oyi strain could be complemented in vitro with a WT Tat gene).

Could it be a defective HIV-1 strain that induces an immune response able to control infection and protect against the disease?
**Tat Oyi has specific mutations**

**Tat variability in the major HIV-1 clades**

<table>
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<th>1</th>
<th>10</th>
<th>20</th>
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<td><strong>TatHXB2</strong></td>
<td>. .................T.............K.............IT.A.................H.N.Q...A...</td>
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<table>
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Up to 28% in sequence variation between the 5 main HIV-1 subtypes
Only C22S: loss of transactivation

**BIOSANTECH**
Tat Oyi has specific immunologic properties (1)

1. **Tat Oyi induces Ab cross-recognizing 5 major Tat variants**
   - Rabbits sera after immunisation with Tat Oyi, Eli and HXB2.
   - anti-Tat Oyi sera had the highest antibody titers
   - anti-Tat Oyi sera were the only one to have a broad antibody response against 5 heterologous Tat variants Subtype A, B, C, D, AE.

2. **Majority of anti-Tat Oyi Abs are directed towards a conserved 3D structure of full length Tat Oyi protein**
   - Western blots showed that non-homologous Tat variants were recognized by antibodies directed against conformational epitopes
   - Tat Oyi if denatured lost its capacity to induce cross reactive Abs

   *(Opi et al: JBC 2002)*

- Full length Tat is neccessary for a vaccine: Rabbit antisera against HXB2 1-100 had a better capacity to neutralize Tat variants (C and D) than HXB2 1-86 antisera *(Opi et al. Vaccine 2004)*

### Titre of Pooled Rabbit Sera Against the Different Variants of Tat (60 Days Post immunis)

<table>
<thead>
<tr>
<th></th>
<th>Anti HXB2</th>
<th>Anti ELI</th>
<th>Anti Oyi</th>
<th>Preimmune</th>
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<td>UG11RP</td>
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<td>16,000</td>
<td>8,000</td>
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<td>8,000</td>
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<td>Eli</td>
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<td>256,000</td>
<td>16,000</td>
<td>500</td>
</tr>
<tr>
<td>CM240</td>
<td>8,000</td>
<td>4,000</td>
<td>128,000</td>
<td>250</td>
</tr>
</tbody>
</table>
3. The tat Oyi induces antibodies against a conserved 3D epitope
A conserved 3D epitope was identified with a mice Monoclonal antibody to Tat Oyi, 7G12
It does recognise the 5 Tat variants : clade A (Ug11RP), clade D (Eli), circulating recombinant form AE(CM240), clade C (96Bw), and clade B (HxB2) (Mediouni et al. JBC 2013)

4. Anti-Tat Oyi Ab against 3D epitope can neutralized Tat activity in vitro (Mediouni et al. JBC 2012 & 2013)
MAb 7G12 was shown to neutralise all Tat variants

➢ Tat Oyi as a novel vaccine candidate would induce an immune response to a 3D epitope not identified previously and able to neutralized the activity of native Tat in vivo.
Antibodies of seropositive patients recognize folded Tat

In sera from 40 HIV-1 infected patients, 19 had anti-Tat Ab:

- 11 recognized Tat peptides and full length
- 8 recognized Tat HXB2 full length exclusively
  These 8 sera recognized Tat Oyi and HXB2

- Dot Blot showed that unfolded Tat was no longer detectable by sera of the second group (n=8) compared to folded Tat. (Mediouni et al. Inf. Dis. 2011)

- Full length Tat is immunogenic and recognised by human sera as a folded protein

- Tat Oyi as a novel immunogen could stimulate an immune response to a unique 3D epitope not previously identified and able to neutralized the activity of native Tat in vivo.
Tat oyi vaccine candidate advantage

The specific feature of Tat Oyi therapeutic vaccine candidate versus other Tat

- It relies on Tat Oyi 3D specific conformation
- The ability to induce an Ab response to conformational epitopes cross reactive to several Tat variants present in the world
- Ability of Ab to 3D epitope to neutralize extracellular Tat
- Immune response against this extracellular HIV target

Did show a protective effect in a Rhesus macaques model of HIV infection
Macaques vaccination with Tat Oyi and SHIV challenge

7 macaques immunized with 100µg full length Tat Oyi (montanide adjuvant). At T0, boost at M1, M2, M3) challenge at M7 with heterologous SHIV BX08 (with 350 AID$_{50}$ IR)

Main Results: 6/7 High titers Ab response to Tat,
1/7 with the highest Tat Ab titer was resistant to infection even after 2$^{nd}$ challenge,
5/6 lower RNA levels than controls (1-2 log) during the chronic phase,
In 7/7 vaccinee Reservoir Cells not detectable at d56 pi.

PBMC co-culture with CD4 macaque cells

- Tat Oyi Vaccinated Macaques
- Control Macaques

Antibodies Titer against GP120

(Watkins et al., Retrovirology 2006)
Preclinical toxicity studies

- No toxicity observed after a 18-months survey in macaques which received four injections of 100μg of Tat Oyi

- No toxicity or loss weight observed in 40 mice which received four intradermal injections of Tat Oyi containing doses of 22μg/50μl and 10μg/50μl solubilized within NaH2PO4 100 mM pH 4.3 and NaCl 9 g/l.
Tat Oyi Clinical trial

A clinical trial phase I/IIa has been started for a therapeutic Tat Oyi vaccine candidate:

- Vaccine candidate: synthetic Tat Oyi full length protein, no adjuvant
- In a double blind study, 48 HIV seropositive under ART
  1: n=12 placebo  
  2: n=12 at 10µg  
  3: n=12 at 33µg  
  4: n=12 at 100µg
- Three intradermal injections of at M0, M1 and M2.
- Follow up 12 months

Criteria:
- Antibodies response to Tat against 5 variants
- HIV RNA level after 1 month ART interruption at M5
In 48 patients, 36 exposed to vaccine candidate

Safety analysis of blinded AEs
• No potential drug interactions identified.
• Most of the reported non-serious AEs and SAEs are common symptoms of HIV infection and are therefore likely attributed to the underlying condition.
  • Non serious AEs: infections, general disorders, gastrointestinal disorders, musculoskeletal disorders, headache
  • The occurrence of injection site pain in four patients which cannot be attributed to the underlying disease or to concomitant medication
  • Only 4 SAEs reported: 2 preimmunisation, 2 that could be attributed to HIV infection

The vaccine candidate with Tat Oyi proved to be safe as no important identified risks during this phase
Thanks

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