Research

Optimization of the doxycycline-dependent simian immunodeficiency virus through in vitro evolution

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Abstract

Background: Vaccination of macaques with live attenuated simian immunodeficiency virus (SIV) provides significant protection against the wild-type virus. The use of a live attenuated human immunodeficiency virus (HIV) as AIDS vaccine in humans is however considered unsafe because of the risk that the attenuated virus may accumulate genetic changes during persistence and evolve to a pathogenic variant. We earlier presented a conditionally live HIV-1 variant that replicates exclusively in the presence of doxycycline (dox). Replication of this vaccine strain can be limited to the time that is needed to provide full protection through transient dox administration. Since the effectiveness and safety of such a conditionally live virus vaccine should be tested in macaques, we constructed a similar dox-dependent SIV variant. The Tat-TAR transcription control mechanism in this virus was inactivated through mutation and functionally replaced by the dox-inducible Tet-On regulatory system. This SIV-rtTA variant replicated in a dox-dependent manner in T-cell lines, but not as efficiently as the parental SIVmac239 strain. Since macaque studies will likely require an efficiently replicating variant, we set out to optimize SIV-rtTA through in vitro viral evolution.

Results: Upon long-term culturing of SIV-rtTA, additional nucleotide substitutions were observed in TAR that affect the structure of this RNA element but that do not restore Tat binding. We demonstrate that the bulge and loop mutations that we had introduced in the TAR element of SIV-rtTA to inactivate the Tat-TAR mechanism, shifted the equilibrium between two alternative conformations of TAR. The additional TAR mutations observed in the evolved variants partially or completely restored this equilibrium, which suggests that the balance between the two TAR conformations is important for efficient viral replication. Moreover, SIV-rtTA acquired mutations in the U3 promoter region. We demonstrate that these TAR and U3 changes improve viral replication in T-cell lines and macaque peripheral blood mononuclear cells (PBMC) but do not affect dox-control.

Conclusion: The dox-dependent SIV-rtTA variant was optimized by viral evolution, yielding variants that can be used to test the conditionally live virus vaccine approach and as a tool in SIV biology studies and vaccine research.
Background

More than 20 years after the identification of human immunodeficiency virus (HIV) as the causative agent of AIDS, an effective HIV/AIDS vaccine remains elusive. All vaccine candidates thus far tested in human efficacy trials have failed to prevent HIV infection or suppress the viral load. In the experimental model system of pathogenic simian immunodeficiency virus (SIV) in macaques, live attenuated virus vaccines have proven to be much more effective than other AIDS vaccine approaches. For example, 95% of the Indian rhesus macaques immunized with a live attenuated SIV demonstrated a viral load suppression of more than 3 logs (compared to unvaccinated animals) upon challenge with a wild-type SIV, whereas such protection was observed in only 7% of macaques immunized with other vaccine strategies [1]. In most of the studies, SIV was attenuated through deletion of one or several accessory functions from the viral genome (reviewed in [1-4]). Although the majority of macaques vaccinated with such deletion variants of SIV can efficiently control replication of pathogenic challenge virus strains, the attenuated virus could revert to virulence and cause disease over time in some vaccinated animals [5-8]. Similarly, some of the long-term survivors of the Sydney Blood Bank Cohort infected with an attenuated HIV-1 variant in which nef and long terminal repeat (LTR) sequences were deleted, eventually progressed to AIDS [9]. An HIV-1 Δ3 variant with deletions in the vpr, nef and LTR sequences regained substantial replication capacity in long-term cell culture infections by acquiring compensatory changes in the viral genome [10]. These results underline the evolutionary capacity of attenuated SIV/HIV strains, which poses a serious safety risk for any future experimentation with live attenuated HIV vaccines in humans.

Evolution of the attenuated vaccine virus upon vaccination is due to the persistence of the virus and ongoing low-level replication. The error-prone viral replication machinery can facilitate the generation and accumulation of mutations in the viral genome that improve replication and pathogenicity. To minimize the prospect of such undesired evolution of the vaccine strain, we and others previously presented a unique genetic approach that exploits a conditionally live HIV-1 variant [11-15]. In our HIV-rtTA variant, the Tat-TAR regulatory mechanism that controls viral transcription was inactivated by mutation of both the Tat protein and the TAR RNA element, and functionally replaced by the components of the Tet-On system for inducible gene expression [16]. The rtTA gene encoding a synthetic transcriptional activator was inserted in place of the nef gene, and the corresponding tet-operator (tetO) DNA binding sites were inserted into the LTR promoter. Since the rtTA protein can only bind tetO and activate transcription in the presence of doxycycline (dox), HIV-rtTA replicates exclusively when dox is administered.

Upon vaccination with this virus, replication can be switched on temporarily and controlled to the extent needed for induction of the immune system by transient dox administration. Upon long-term in vitro passage of the initial HIV-rtTA variant on T cells, the virus acquired additional modifications in both the rtTA and tetO components that significantly improved replication [17-22]. This designer HIV-rtTA was thus optimized through in vitro virus evolution, resulting in a dox-dependent variant that replicates in vitro in T cell lines and ex vivo in human lymphoid tissue [23]. In addition, we constructed an HIV-1 variant that depends not only on dox for gene expression, but also on the T20 peptide for cell entry [24].

To evaluate the safety and effectiveness of such a conditionally replicating virus as a candidate AIDS vaccine, a dox-dependent SIV variant is needed that can be tested in macaques. Moreover, such an SIV variant may be an ideal tool to study the immune correlates of vaccine protection, since both the level and duration of virus replication can in principle be controlled by dox administration. Such studies may reveal the critical information needed for the design of an HIV vaccine that is safe and equally effective as a live attenuated virus. Based on our experience in developing HIV-rtTA, we recently constructed a similar dox-dependent SIVmac239 variant [25]. Surprisingly, inactivation of the Tat protein was not allowed in the SIV-rtTA context, even though gene expression was transcriptionally controlled by the incorporated Tet-On system. This result suggests that Tat has additional essential functions in SIV replication in addition to its role in the activation of transcription. The Tat-positive SIV-rtTA variant replicated in a dox-dependent manner in T cell lines, but not as efficiently as the parental SIVmac239 strain. We anticipated that SIV-rtTA could evolve to a better replicating variant and therefore initiated multiple cultures. We did indeed identify modifications in the U3 and TAR regions that significantly enhance SIV-rtTA replication in T cell lines and macaque peripheral blood mononuclear cells (PBMC). Importantly, these modifications do not affect dox-control. These evolved SIV-rtTA variants should allow future in vivo studies in macaques.

Results

In vitro evolution of the dox-inducible SIV-rtTA variant

We recently described the construction of a dox-dependent SIVmac239 variant in which the natural Tat-TAR mechanism of transcription control was replaced by the dox-inducible Tet-On gene expression system (Fig. 1A). In this variant, the bulge and loop sequences in stem-loop 1 (SL1) and stem-loop 2 (SL2) of TAR are mutated (TARm; substituted nucleotides marked in a gray circle in Fig. 1B), which prevents the binding of Tat and precludes Tat-responsiveness of the LTR promoter. Furthermore, this virus carries the gene encoding the rtTA transcriptional
Figure 1
Evolution of the dox-inducible SIV-rtTA variant. (A) In the SIVmac239-based SIV-rtTA variant, the Tat-TAR regulatory mechanism was inactivated through mutation of TAR (TARM), and functionally replaced by the dox-inducible Tet-On regulatory system through the introduction of the gene encoding the rtTA transcriptional activator protein at the site of the nef gene and two dox-responsive tet operator (tetO) elements between the NFKB and Sp1 sites in the U3 promoter region [25]. The TAR mutations and tetO elements were introduced in both the 5’ and 3’ LTR. (B) The TAR RNA element of SIV-rtTA can fold a branched hairpin structure with three stem-loop domains (SL1-3). The mutations that had been introduced in SL1 and SL2 to inactivate TAR, are encircled in gray (SL1: +27U-A, +28U-A, +34C-A, +36G-U; SL2: +62U-A, +68C-A, +70G-U). Upon long-term culturing of SIV-rtTA in PM1 cells, additional nucleotide substitutions are observed in TAR. The number of the culture in which the mutation is observed is shown (#), with the asterisk (*) indicating the transient presence of the mutation. (C) Alternative folding of the SL1 domain can result in a 6-bp spacer between the bulge and loop sequences. However, this spacer extension slightly reduces TAR stability (ΔG_{6bp} = -67.5 kcal/mole; ΔG_{5bp} = -65.2 kcal/mole). Alternative folding of the +63A-G mutated TAR RNA results in a 6-bp bulge-loop spacer in SL2 but does not affect TAR stability (ΔG_{6bp} = ΔG_{5bp} = -67.5 kcal/mole). Formation of an A+63-U+78 base pair in the +78C-U mutant results in a similar 6-bp bulge-loop spacer in SL2 and increases the stability of this TAR variant (ΔG_{6bp} = -65.2 kcal/mole; ΔG_{5bp} = -65.8 kcal/mole). (D) TAR can fold an alternative extended hairpin structure in which the SL1 and SL2 sequences fold a large stem-loop structure. The introduced and acquired mutations are shown as in B.
activator protein at the position of the nef gene and two
dox-responsive tet operator (tetO) elements between the
Nfkb8 and Sp1 binding sites in the U3 promoter region
(Fig. 1A). Dox induces a conformational change in the
rtTA protein that triggers binding to the tetO sites and acti-
vation of transcription from the downstream start site. In
the absence of dox, rtTA cannot bind to the tetO sites and
viral gene expression is not activated. Since transcription
is critically dependent on dox, this SIV-rtTA variant replic-
cates exclusively in the presence of dox. As the TAR muta-
tions and tetO elements were introduced in both the 5'
and 3' LTR, they are stably maintained in the viral prog-
cy.

We demonstrated that SIV-rtTA replicates in a dox-
dependent manner in PM1 T-cells, but not as efficiently as
the wild-type SIVmac239 variant [25]. Since macaque
studies will likely require an efficiently replicating variant,
we set out to optimize SIV-rtTA through in vitro viral evolu-
tion. We therefore started 13 cultures of the Tat-positive
SIV-rtTA variant in PM1 cells and passaged the virus onto
fresh cells at the peak of infection when massive syncytia
were observed. The cultures were maintained for up to
234 days. The period between infection and the appear-
ance of syncytia decreased over time and we could reduce
the volume of the virus inoculum that is needed to start a
new infection. These observations indicate that the repli-
cation capacity of the virus had improved and we ana-
alyzed the proviral genome present in these long-term
cultures. This analysis revealed that the virus steadily main-
tained the introduced TAR mutations, rtTA gene and tetO
elements, but acquired additional mutations in the LTR
region (Fig. 2). We observed one or several nucleotide
substitutions in the TAR sequence in all 13 cultures. In
eight of these cultures, additional nucleotide substitutions
or deletions were present in the Sp1 sites, which are
located between the tetO sites and the TATA box.

**Mutations in TAR affect RNA structure**

We observed an A-to-G substitution at TAR position +63
in seven independent cultures (Fig. 1B). The high fre-
quency may indicate that this change is an important evolu-
tionary route toward improved replication. This
substitution may induce a base pairing rearrangement in
SL2 by formation of a G+63-C+78 base pair, resulting in a 6-
bp spacer between the bulge and loop domains (Fig. 1C).
Remarkably, we observed a C-to-U substitution at position
+78 in two other cultures that has the same impact on the
TAR structure, as it also allows the formation of a 6-bp
bulge-loop spacer through A+63-U+78 base pairing in SL2
(Fig. 1C). In fact, the mutated SL1 can also form a 6-bp
spacer between the bulge and loop domains (Fig. 1C),
although analysis of the thermodynamic stability with the
Mfold RNA folding software [26,27] revealed that this
spacer extension slightly reduces TAR stability ($\Delta G_{5\text{ bp}} = -$
67.5 kcal/mole; $\Delta G_{6\text{ bp}} = -67.2$ kcal/mole). Another
remarkable mutation is seen at position +21 in three cul-
tures. This G-to-A mutation destabilizes the lower stem of
SL1 by generating an A+21-C+49 mismatch but it creates a 7-
nt sequence (C/1A/GC) at the start of the SL1 sequence
that is repeated at the start of SL2. Nearly all other nucle-
otide substitutions were observed in individual cultures.
These mutations seem to destabilize the TAR structure by
either replacing a G-C base pair by a less stable G-U base
pair, or by causing a base pair mismatch (Fig. 1B).

Recently, Pachulska-Wieczorek et al. showed that HIV-2
TAR can fold an alternative secondary structure in addi-
tion to the classical branched hairpin (BH) structure with
SL1, SL2 and SL3 [28]. In this extended hairpin (EH)
structure, the SL1 and SL2 sequences fold a single,
extended stem-loop structure. SIVmac239 TAR, which is
very similar to HIV-2 TAR, and the mutated SIV-rtTA TAR
may also co-exist in comparable BH and EH forms (Fig.
1B and 1D, respectively). At first glance, the individual
TAR mutations observed in SIV-rtTA upon prolonged cul-
turing seem to either stabilize the EH structure by creating
more stable base pairs (e.g. replacement of a G-U base pair
by a more stable A-U base pair) or destabilize this
structure by creating mismatches or less stable base pairs (e.g.
replacement of a G-C base pair by a G-U base pair). Since
the equilibrium between the BH and EH conformers may be
essential in viral replication, we used Mfold RNA anal-
ysis to estimate the thermodynamic stability of the BH
and EH structures for the wild-type (TAR\textsuperscript{wt} in SIVmac239),
mutated (TAR\textsuperscript{m} in SIV-rtTA) and evolved TAR sequences
(Table 1). The difference between these AG values
($\Delta G_{\text{BH\text{-}EH}}$) reflects whether the BH form is more stable
and favored ($\Delta G_{\text{BH\text{-}EH}} < 0$) or the EH form ($\Delta G_{\text{BH\text{-}EH}} > 0$).
This analysis revealed that TAR\textsuperscript{wt} is more stable in the
EH form ($\Delta G = -68.2$ kcal/mole) than in the BH form ($\Delta G
= -65.3$), yielding a $\Delta G_{\text{BH\text{-}EH}}$ of 2.9 kcal/mole. The bulge
and loop mutations that we introduced in TAR\textsuperscript{m} to pre-
vent Tat trans-activation stabilize the BH form and desta-
bilize the EH structure. As a result the $\Delta G_{\text{BH\text{-}EH}}$ is reduced to
-3.6 kcal/mole. The most frequent +63A-G substitution
does not affect the stability of the BH structure but par-
tially restores the stability of the EH form, resulting in a
$\Delta G_{\text{BH\text{-}EH}}$ of -1.2 kcal/mole. Most of the other nucleotide
substitutions reduce the stability of the BH structure and
at the same time stabilize the EH structure. As a result, the
$\Delta G_{\text{BH\text{-}EH}}$ of these TAR elements is increased to values
between -2.0 to 6.3 kcal/mole. In cultures 4, 9 and 10, the
virus accumulated multiple TAR mutations that resulted
in a gradual increase in the $\Delta G_{\text{BH\text{-}EH}}$. In cultures 1 and 5,
such a gradual increase through the accumulating muta-
tions is not observed, but the virus acquired additional
mutations in the Sp1 region. These results suggest that the
bulge and loop mutations that we introduced in SIV-rtTA
shifted the BH-EH equilibrium into the direction of the
**Figure 2**

**SIV-rtTA acquires additional mutations in the U3 and TAR region upon long-term culturing.** SIV-rtTA was cultured with dox in PM1 cells for up to 234 days. Cellular proviral DNA was isolated from 13 independent cultures at different times and the LTR region was subsequently PCR amplified and sequenced. The number of the culture (#) and the day of sampling are indicated on the left. The -90 to +130 U3/R region is shown with +1 indicating the transcription initiation site. The Sp1 and TATA box are shown in grey. The mutations that were introduced in TAR to abolish Tat responsiveness are underlined. The additional nucleotide substitutions and deletions (Δ) observed in the SIV-rtTA cultures are indicated.
BH form, and that nucleotide substitutions selected during virus evolution reduce this preference for the BH form or even restore the preference for the EH structure. The only exceptions are the +46C-T and +72G-A mutations observed in culture 12, which only marginally affect the BH and EH stability. The virus in this culture did however acquire an additional nucleotide substitution in the Sp1 sites, which may have improved replication.

To demonstrate that the introduced and acquired mutations do indeed affect TAR folding, we analyzed the electrophoretic mobility of in vitro transcribed RNAs corresponding to TARwt, TARm and the evolved +21G-A, +63A-G and +78C-U variants. The RNAs were denatured by heat, renatured in the presence of MgCl2 and subsequently analyzed by denaturing and non-denaturing polyacrylamide gel electrophoresis. All RNAs migrate similarly on a denaturing polyacrylamide gel, as expected based on their identical size (Fig. 3A). In contrast, TARm migrates slower than TAR wt on the non-denaturing gel (Fig. 3B). Since branched RNA conformers migrate slower than extended molecules, the observed migration pattern is in agreement with a predominant EH structure of TARwt under these conditions, as previously shown by Pachulska-Wieczorek et al. [28], and a BH structure of TARm. The +21G-A, +63A-G and +78C-U TAR RNAs show the fast wild-type migration capacity, which demonstrates that these mutations restore EH folding of TAR in this in vitro assay.

SIV-rtTA expresses the wild-type Tat protein but the mutations introduced in TAR prevent binding of Tat and activation of transcription [25]. One possibility is that the acquired TAR mutations restore Tat binding. We therefore performed an Electrophoretic Mobility Shift Assay (EMSA) to analyze the effect of the +21G-A, +63A-G and +78C-U changes on Tat binding. In the absence of Tat, all in vitro transcribed TAR RNAs migrate similarly on the EMSA gel (Fig. 3C). Upon incubation with Tat, TARwt efficiently shifts into a slower migrating Tat-TAR complex. This Tat-TAR complex is not observed for TARm, demonstrating that the introduced TAR mutations do effectively prevent Tat binding. The +21G-A, +63A-G and +78C-U substitutions do not restore Tat binding.

Mutations in U3 and TAR do not affect promoter activity
In addition to the mutations in TAR, SIV-rtTA acquired mutations in the U3 region upon long-term culturing (Fig. 2). We observed a G-to-A substitution in one of the four G-rich Sp1 sites in six cultures. Furthermore, a 1-nt deletion in one of the Sp1 sites and a 14-nt deletion that affects two Sp1 sites were observed once. Since the U3 and TAR mutations may affect SIV-rtTA promoter activity, we re-cloned the evolved LTR sequences into an LTR promoter-luciferase reporter construct. We made constructs with the +21G-A, +63A-G or +78C-U TAR mutation. The +63A-G mutation was also combined with the G-to-A substitution (mSp1) or 14-nt deletion in the Sp1 sites (ΔSp1), exactly as it appeared at day 115 in culture 3 and at day 104 in culture 1, respectively.

To test the dox responsiveness of these SIV-rtTA promoters, these plasmids were co-transfected with an rtTA-expressing plasmid into C33A cervix carcinoma cells. After two days of culturing with 0 to 1000 ng/ml dox, we measured the intracellular luciferase level, which reflects gene expression (Fig. 4A). The original SIV-rtTA promoter was inactive in the absence of dox and its activity gradually increased with an increasing dox level. All evolved promoter variants showed a similar low activity without dox and a similarly high activity with dox, which demonstrates that the acquired U3 and TAR mutations do not sig-

### Table 1: Nucleotide substitutions affect the stability of the branched hairpin (BH) and extended hairpin (EH) conformation of TAR.

|          | ΔGBH | ΔGEH | ΔΔGBH-EH | culture |
|----------|------|------|----------|---------|
| TARwt (SIVmac239) | -65.3 | -68.2 | 2.9     |         |
| TARm (SIV-rtTA) | -67.5 | -63.9 | -3.6    | 312, 251, 3115, 481, 6124, 7136, 1328 |
| +63A-G       | -67.5 | -63.9 | -2.0    | 1133    |
| +21G-A       | -62.5 | -66.6 | -4.1    | 4324    |
| +78C-T       | -65.8 | -65.7 | -0.1    | 512, 1017 |
| +78C-T +44T-C| -63.7 | -64.1 | 0.4     | 5198    |
| +72G-A       | -62.5 | -64.2 | 1.7     | 948     |
| +21G-A +78C-T+99C-T | -58.4 | -63.6 | 5.2     | 5124    |
| +21G-A +74G-A | -58.8 | -63.4 | 4.6     | 9130    |
| +5G-A +59A-T | -63.3 | -69.6 | 6.3     | 10136   |
| +73T-C       | -66.5 | -65.1 | -1.4    | 11147   |
| +73T-C +49C-T| -64.2 | -64.9 | 0.7     | 8222    |
| +46C-T +72G-A | -67.6 | -63.6 | -4.0    | 12120   |

a ΔG values (kcal/mole) as determined with the Mfold RNA analysis software. b ΔΔGBH-EH = ΔGBH-ΔGEH. c Culture in which the mutation is observed (see Figure 2), with the day of earliest detection in superscript.
Acquired mutations in TAR restore secondary structure but not Tat binding. In vitro transcribed TAR RNA corresponding to the wild-type SIVmac239 (TARwt), SIV-rtTA (TARm) and the evolved +21G-A, +63A-G and +78C-U variants was denatured by heat, renatured in the presence of MgCl₂ and analyzed on a denaturing gel (A) and on a non-denaturing gel (B). Under these non-denaturing conditions, branched hairpin (BH) RNA conformers migrate slower than extended hairpin (EH) molecules [28]. (C) Binding of SIV Tat to TAR was analyzed in an Electrophoretic Mobility Shift Assay (EMSA). TAR RNA was incubated with 0 or 100 ng Tat protein (indicated with - and +, respectively) and analyzed on a non-denaturing gel. The position of unbound TAR RNA and TAR-Tat complex is indicated.

Figure 4
U3 and TAR mutations do not affect dox and Tat responsiveness of the SIV-rtTA promoter. (A) To assay dox responsiveness, C33A cells were transfected with LTR-promoter/luciferase reporter constructs corresponding to the original and evolved SIV-rtTA variants and an rtTA-expressing plasmid. After two days of culturing with 0 to 1000 ng/ml dox, the intracellular luciferase level, which reflects promoter activity, was measured. The error bar represents the standard deviation (SD) for 3 to 8 experiments. (B) To assay Tat responsiveness, C33A cells were transfected with the promoter/luciferase plasmids and 0 to 50 ng SIV Tat-expressing plasmid. Two days after transfection, the promoter activity was analyzed by measuring the intracellular luciferase activity. The error bar represents the SD for 2 to 4 experiments. (C) 293T cells were transfected with the SIV-rtTA proviral constructs and cultured for two days with or without dox. Virus production was quantified by measuring the CA-p27 level in the culture supernatant. The error bar represents the standard deviation for 2 experiments.
nificantly affect the basal and dox-induced promoter activity.

To test the Tat responsiveness of the new SIV-rtTA promoters, we transfected C33A cells with the promoter/luciferase plasmids plus 0 to 50 ng SIV Tat-expressing plasmid [25] and measured the luciferase level after two days (Fig. 4B). Neither the original SIV-rtTA construct nor the evolved variants responded to Tat. Only the control construct with a wild-type SIVmac239 TAR sequence showed increased activity with an increasing amount of Tat. Thus, the acquired U3 and TAR mutations do also not restore Tat responsiveness, which is in agreement with the inability of the evolved TAR RNAs to bind Tat (Fig. 3C).

**Evolved U3 and TAR sequences improve SIV-rtTA replication**

To determine the effect of the acquired U3 and TAR mutations on virus production and replication, we introduced the evolved LTR sequences into the SIV-rtTA genome. The mutations were introduced in both the 5' and 3' LTR of the SIV-rtTA plasmid, such that they are stably inherited in the viral progeny. The SIV-rtTA constructs were transfected into 293T cells and after two days of culturing with or without dox, virus production was quantified by measuring the CA-p27 level in the culture supernatant (Fig. 4C). The original and new SIV-rtTA variants showed a similarly high level of virus production with dox and a similarly low level without dox. These results demonstrate that the acquired U3 and TAR mutations do not significantly affect dox-dependent viral gene expression and virus production, which is in agreement with the results of the promoter activity assays (Fig. 4A).

To evaluate the replication capacity of the SIV-rtTA variants, PM1 T-cells were transfected with 5 μg of the proviral plasmids and cultured in the presence and absence of dox (Fig. 5A). None of the SIV-rtTA variants replicate in the absence of dox, which is in agreement with their dox-dependent promoter activity. In the presence of dox, the new variants with either the +21G-A, +63A-G or +78C-U TAR mutation replicate more efficiently than the original SIV-rtTA, which demonstrates that these TAR mutations significantly improve viral replication. The +63A-G mSp1 and +63A-G ΔSp1 variants seem to replicate with a similar efficiency as the +63A-G variant. However, comparison of the replication capacity of these variants upon transfection of 1 μg of the proviral plasmids revealed that the Sp1-mutated variants replicate more efficiently (Fig. 5B). This result demonstrates that the acquired Sp1 mutations further improve SIV-rtTA replication. The original SIV-rtTA did not show any replication within the time frame of this experiment, which illustrates that the replication capacity of the new variants has increased significantly. Despite this large improvement, the new SIV-rtTA variants did not

**Figure 5**

**U3 and TAR mutations improve SIV-rtTA replication.** (A) PM1 T-cells were transfected with 5 μg of the original (grey symbols) or LTR-mutated SIV-rtTA proviral plasmid (black symbols) and cultured with or without dox (closed and open symbols, respectively). Virus replication was monitored by measuring the reverse transcriptase level in the culture supernatant. (B) Cells were transfected with 1 μg SIV-rtTA or SIVmac239 proviral plasmid and cultured with dox (SIV-rtTA variants) or without dox (SIVmac239).
replicate as efficiently as wild-type SIVmac239, which was included in this experiment for comparison.

To demonstrate that the acquired mutations do not selectively improve viral replication in the human PM1 T cells that were used in the evolution study, we next assessed the replication capacity of the SIV-rtTA variants in primary PBMC isolated from cynomolgus macaques (Fig. 6A). For comparison, we included the wild-type SIVmac239 and the SIV-rtTA-mTat variant in which Tat is inactivated by a Tyr-55-Ala mutation [25]. Upon infection, cells were cultured with or without dox. In the absence of dox, none of the SIV-rtTA variants showed any replication, while SIVmac239 replicates efficiently (not shown). SIV-rtTA-mTat does also not show any replication in the presence of dox, which is in agreement with previous observations in T cell lines and indicates that SIV-rtTA requires Tat for a non-transcriptional function in the viral life cycle. The original Tat-positive SIV-rtTA replicates poorly in the PBMC upon dox administration, whereas the new variants in which we introduced the U3 and TAR changes replicate much more efficiently. However, these viruses do not replicate as efficiently as wild-type SIVmac239. Similar results were obtained when replication of the +63A-G, +63A-G mSp1 and +63 A-G ΔSp1 variants was tested in PBMC isolated from rhesus macaques (Fig. 6B). Also in these cells, the new SIV-rtTA variants replicated to much higher levels in the presence of dox than in its absence, although with somewhat delayed replication kinetics when compared to SIVmac239. These studies suggest that the evolved LTR sequences significantly improve SIV-rtTA replication in macaque PBMC. Importantly, the Sp1 and TAR mutations do not affect dox-control in these primary cells.

**Discussion**

In this paper, the optimization of the conditionally live SIV-rtTA variant through viral evolution is described. We recently constructed this dox-dependent SIVmac239 variant by replacing the natural Tat-TAR mechanism of transcription control by the dox-inducible Tet-On regulatory mechanism. Although the original SIV-rtTA variant replicates in T cell lines and in primary macaque PBMC, it replicates poorly when compared with the parental SIVmac239 [25][Figs. 5 and 6]. Upon long-term culturing, the virus acquired several mutations in the TAR and U3 region. These mutations significantly improve viral replication, but do not affect dox control. We thus generated novel SIV-rtTA variants that replicate efficiently and in a dox-dependent manner in both T-cell lines and primary macaque PBMC.

We previously used virus evolution to optimize a similarly constructed dox-dependent HIV-1 variant. Upon long-term culturing, this HIV-rtTA variant acquired several mutations in the rtTA and tetO components of the introduced Tet-On system, which considerably improved viral replication [17-19,21]. These optimized rtTA and tetO components were used for the construction of SIV-rtTA and these elements were stably maintained upon evolution of this virus. Unlike HIV-rtTA, SIV-rtTA further
improved its replication capacity through additional mutations in the TAR and Sp1 region.

For the construction of SIV-rtTA, both the bulge and loop domains in TAR were mutated to prevent binding of Tat and trans-activation of transcription. Interestingly, the acquired nucleotide substitutions in TAR upon SIV-rtTA evolution do not restore the wild-type bulge and loop sequences. The frequently observed changes at positions +63 and +78 do however allow the formation of a 6-bp spacer between the bulge and loop domains in SL2 (Fig. 1C). This is remarkable since trans-activation by HIV-2 Tat, which is very similar to SIV Tat, is optimal with a bulge-loop spacing of 6 bp [29]. However, we demonstrate that the evolved TAR elements do not bind Tat and that transcription from the modified SIV-rtTA promoter is not activated by Tat. We also frequently observed a G-to-A nucleotide substitution at position +21, which creates a 7-nt repeat at the start of SL1 and SL2. If this sequence would bind a transcription factor, either as LTR DNA or TAR RNA, duplication of the motif could increase promoter activity. However, the +21 substitution did not affect the low basal promoter activity in the absence of dox or the high induced activity in the presence of dox. Similarly, the other TAR and U3 mutations do not affect the transcription process.

In silico RNA folding analysis and in vitro RNA mobility assays revealed that the acquired TAR mutations do affect the structure of this RNA element. As previously proposed for HIV-2 TAR [28], the TAR motif of SIVmac239 and SIV-rtTA may fold alternative structures: the classical branched hairpin (BH) structure with SL1, SL2 and SL3 (Fig. 1B) and an extended hairpin (EH) structure in which the SL1 and SL2 sequences form a single, extended stem-loop structure (Fig. 1D). We demonstrate that the wild-type TAR in SIVmac239 favors the EH form. The bulge and loop mutations that we had introduced in SIV-rtTA shift the equilibrium toward the BH form. Interestingly, nearly all mutations observed in the evolved variants partially or completely restored the wild-type situation in which the EH form is favored. Although the role of the EH TAR conformation and the possible EH-BH riboswitch in the SIV life cycle has yet to be resolved, these results suggest that a proper EH-BH equilibrium is important for efficient viral replication.

Interestingly, alternative folding of the leader RNA has also been proposed for HIV-1. In this case however, the TAR structure is identical in the alternative conformations. The energetically favored structure of the HIV-1 leader is formed by a long-distance interaction (LDI) between the sequences around the polyadenylation site and the dimerization initiation signal (DIS) [30]. In the alternative structure, termed the branched multiple hairpin (BMH) conformation, both the polyadenylation and DIS motifs fold a stem-loop element. Mutations that affect the equilibrium between the dimerization-incompetent LDI structure and the dimerization-prone BMH structure significantly affect HIV-1 replication [30-33]. Our recent studies with HIV-rtTA showed that HIV-1 TAR can be truncated, deleted or replaced by a non-related stem-loop element when not required for the activation of transcription, which demonstrates that TAR has no additional essential role in HIV-1 replication [34]. However, destabilization of TAR blocked replication, which can possibly be explained by unwanted pairing of free nucleotides in the destabilized TAR structure with downstream leader sequences, thereby affecting the LDI-BMH equilibrium [35]. Thus, although TAR is not a functional domain of the LDI-BMH conformational switch in HIV-1, it can indirectly affect this function. In analogy with these HIV-1 studies, it cannot be excluded that the bulge and loop mutations introduced in SIV-rtTA caused misfolding of the leader RNA. These mutations may change the local TAR folding or generate a new sequence with complementarity to downstream sequences, which could result in an interaction between TAR and other leader domains. The additional TAR mutations in the evolved variants may prevent this interaction and thus restore viral replication. Although further analyses will be needed to understand this misfolding scenario, it is supported by our recent observation that precise truncation of structural TAR domains is compatible with SIV-rtTA replication (manuscript in preparation).

We demonstrated that SIV-rtTA requires wild-type Tat protein for replication in T-cell lines [25] and primary macaque PBMC (this study), although gene expression is controlled by the incorporated Tet-On system. These results suggest that Tat has additional functions in the SIV replication cycle in addition to its role in the activation of transcription. For this reason, the SIV-rtTA variant used in this study encodes the wild-type Tat protein. Reversion of the bulge and loop mutations in TAR, which had been introduced to prevent Tat binding and trans-activation of transcription, would restore the Tat-TAR mechanism of transcription control. However, this evolution route would require multiple nucleotide substitutions, which is not likely to occur. Indeed, we never observed restoration of the Tat-TAR axis in numerous long-term cultures of SIV-rtTA. Nevertheless, the likelihood of this unwanted evolution route can be further reduced by introducing novel mutations in Tat that would inactivate the first function (activation of transcription) but not the second function (currently unknown). However, such Tat mutations remain to be identified. Alternatively, this evolution route can be blocked by the complete or partial deletion of TAR (e.g. only SL1 and SL2), as we recently showed that the
complete removal of TAR in HIV-rtTA does not significantly affect replication [34].

The optimization of SIV-rtTA through viral evolution resulted in new dox-controlled variants that replicate efficiently in the PM1 T cell line and in primary PBMC from cynomolgus and rhesus macaques. These novel SIV-rtTA variants may be good candidates to study the efficacy and safety of a conditionally live virus as AIDS vaccine in macaques. Furthermore, this virus may be an ideal tool to study the immune correlates of protection if the level and duration of replication in vivo can be stringently controlled by dox administration. Such studies may reveal crucial information needed for the design of a safe and effective HIV vaccine.

Methods
Viruses and cells
We previously described the construction of the SIV-rtTA plasmid encoding the dox-dependent virus that is based on the SIVmac239 isolate [36][GenBank accession M33262] and contains the wild-type Tat gene (pSIV-rtTA-Tatwt in [25]). The plasmid pSIVmac239 encodes the full-length SIVmac239 isolate [37].

Human embryonal kidney (HEK) 293T and cervical carcinoma C33A cells [38] were cultured in 2-cm² wells and transfected with 1 µg SIV-rtTA or SIVmac239 DNA by calcium phosphate precipitation, as previously described [19]. PM1 T cells [39] were cultured and transfected by electroporation [25]. PBMC were isolated from cynomolgus and rhesus macaques [41] and cultured as described for the PM1 cells [25]. Cells were activated with 2 µg/ml PHA (cynomolgus macaques PBMC) or anti-CD3 monoclonal antibody (rhesus macaques PBMC) for two days prior to infection with C33A or 293T produced virus, and maintained with 100 units/ml recombinant IL-2 following infection. Cells were cultured with 1000 ng/ml dox (Sigma D-9891) when indicated. The virus level in the culture medium was determined by CA-p27 ELISA (SIV-Env3, Invitrogen). The NotI-NarI fragment of the TA-clone was digested with EcoRI (position 10584) and XhoI (position 10981), and used to replace the corresponding 3'-LTR fragment in pBS-3'SIV-rtTA [25]. The NheI-XhoI fragment (GTTAGGGCTTTAATCGTAGGGATTTTCCTGCTTCG, nt 10991–10959) for amplification, and SIV-Env5 (GTTAGGGCTTTAATCGTAGGGATTTTCCTGCT, nt 8723–8777) and tTA-Rev4 (GAAGAGGGCTTTAATCGTAGGGATTTTCCTGCT, nt 9882–9865) for sequencing.

For the cloning of the evolved U3 and TAR sequences into the 5’ LTR of SIV-rtTA, the proviral DNA was PCR amplified with primers SIV-LTR4 (AGGTACCGGCCGCCGCGCTCTGGAAAGGATTTTAATCGTAGGGATTTTCCTGCT, nt 1–36) and SIV-LTR5 (ATGGACGTCTCGAGTCGCATGCTAGGCGCCAATCTGCTAGGCGCCAATCTGCT, nt 9681–9700) for sequencing. For the analysis of Tat-coding exon 2 and the 5’-half of rtTA, we used primers tTA3 (GTGTGCGGCCGCCGCGCTCTGGAAAGGATTTTAATCGTAGGGATTTTCCTGCT, nt 8691–8712) and tTA-Rev3 (TGAAATCGAGTTTCTCCAGGCCCATATGA, nt 9291–9901) for amplification, and SIV-Env5 (GTTAGGGCTTTAATCGTAGGGATTTTCCTGCT, nt 8723–8777) and tTA-Rev4 (GAAGAGGGCTTTAATCGTAGGGATTTTCCTGCT, nt 9882–9865) for sequencing.

For the introduction of the evolved U3 and TAR mutations into the 3’ LTR of SIV-rtTA, the proviral DNA was PCR amplified with primers SIV-LTR4 (AGGTACCGGCCGCCGCGCTCTGGAAAGGATTTTAATCGTAGGGATTTTCCTGCT, nt 1–36) and SIV-LTR5 (ATGGACGTCTCGAGTCGCATGCTAGGCGCCAATCTGCTAGGCGCCAATCTGCT, nt 9681–9700) for sequencing. For the analysis of Tat-coding exon 2 and the 5’-half of rtTA, we used primers tTA3 (CTGTGCGGCCGCCGCGCTCTGGAAAGGATTTTAATCGTAGGGATTTTCCTGCT, nucleotides 9681–9700 in pSIV-rtTA-Tatwt) and SIV-LTR3 (ATGGACGTCTCGAGTCGCATGCTAGGCGCCAATCTGCTAGGCGCCAATCTGCT, nt 9709–9727), SIV-LTR8 (AAAGGCTCTAAACGCAAGCCCA, nt 10944–10926) and SIV-LTR10 (GAAAGGCTTTAATCGTAGGGATTTTCCTGCT, nt 10832–10812) for sequencing. For the analysis of Tat-coding exon 2 and the 5’-half of rtTA, we used primers SIV-LTR1 (AGGTACCGGCCGCCGCGCTCTGGAAAGGATTTTAATCGTAGGGATTTTCCTGCT, nt 8691–8712) and tTA-Rev3 (TGAAATCGAGTTTCTCCAGGCCCATATGA, nt 9291–9901) for amplification, and SIV-Env5 (GTTAGGGCTTTAATCGTAGGGATTTTCCTGCT, nt 8723–8777) and tTA-Rev4 (GAAGAGGGCTTTAATCGTAGGGATTTTCCTGCT, nt 9882–9865) for sequencing.

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We used standard molecular biology procedures for all manipulations and plasmids were propagated in either E. coli TOP10 (for TA-cloning; Invitrogen), DH5α (pBS-3'SIV-rtTA and LTR-luc plasmids) or Stbl4 (pSIV-rtTA-
Tat<sup>wt</sup> constructs; Invitrogen). All constructs were verified by sequence analysis.

**Promoter activity assay**

To determine dox-responsiveness of the promoter-luciferase constructs, C33A cells were transfected with 20 ng LTR-luciferase<sup>firefly</sup> plasmid, 0.4 ng rtTA-expressing plasmid pCMV-rtTA<sub>F86Y A209T</sub> [19], 0.5 ng pRL-CMV (Promega) and 980 ng pBluescript. This pBluescript was added as carrier DNA, and pRL-CMV, in which the expression of renilla luciferase is controlled by the CMV immediate early enhancer/promoter, was co-transfected to allow correction for differences in transfection efficiency. The cells were cultured after transfection for 48 hours with 0–1000 ng/ml dox. Cells were lysed in Passive Lysis Buffer and firefly and renilla luciferase activities were determined with the Dual-Luciferase assay (Promega). The expression of firefly and renilla luciferase was within the linear range and no squelching effects were observed. The promoter activity was calculated as the ratio of the firefly and renilla luciferase activities, and corrected for between session variation [44]. To determine Tat-responsiveness of the promoter-luciferase constructs, C33A cells were transfected with 20 ng LTR-luciferase<sup>firefly</sup> plasmid, plus 0–50 ng SIVmac239 Tat-expressing plasmid pcDNA3-SIV-Tat<sup>wt</sup> [25]. 0–50 ng pcDNA3 (empty expression vector, total amount of Tat-expressing plasmid and pcDNA3 was kept at 50 ng), 0.5 ng pRL-CMV and 950 ng pBluescript. Cells were cultured for 48 hours and luciferase activities were subsequently measured.

**Tat binding and TAR conformer assay**

<sup>32</sup>P-labeled TAR transcripts were produced as described previously [25]. In brief, the TAR region in SIV-rtTA LTR-luciferase plasmids was amplified by PCR with a 5' primer encoding the T7 promoter sequence directly upstream of the +1 position. The DNA products were in vitro transcribed with the MEGashortscript T7 transcription kit (Ambion). The TAR RNA transcripts were dephosphorylated with calf intestine alkaline phosphatase and 5'-end labeled with the KinaseMax kit (Ambion) in the presence of 1 μl [γ-<sup>32</sup>P]-ATP. The labeled transcripts were purified on a denaturing 8% acrylamide gel.

For the Tat binding assay, <sup>32</sup>P-labeled TAR RNA (200 counts/s) was denatured in 10 μl water for 1 min at 85°C followed by snap cooling on ice. After addition of 10 μl 200 mM KCl, 100 mM Tris-HCl (pH 8.0), 5 mM MgCl<sub>2</sub>, TAR RNA was renaturated at room temperature for 15 min. After adding 4 μl of non-denaturing loading buffer (30% glycerol, bromophenol blue), 10 μl of the sample was analyzed on a non-denaturing 8% acrylamide gel (in 45 mM Tris, 45 mM Borate and 0.1% Triton X-100) at 450 V at 4°C, or on a denaturing 4% acrylamide gel (in 45 mM Tris, 45 mM Borate and 0.5 mM EDTA) at 450 V at room temperature. The gels were dried and analyzed with a PhosphorImager. The thermodynamic stability of TAR RNA (nt +1 to +124) was determined with the MFold RNA folding program (version 3.2) at the Rensselaer Polytechnic Institute bioinformatics web server [26,27].

**Competing interests**

The authors declare that they have no competing interests.

**Authors’ contributions**

ATD and BB designed the viral replication and evolution studies and drafted the manuscript; BK and AH performed the replication, evolution and promoter analysis experiments; MC and MO carried out the Tat binding and TAR conformer assays; BK, MPa, NA, FY, MPi and JL performed the PBMC experiments. All authors have read, revised and approved the manuscript.

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References

1. Koff WC, Johnson PR, Watkins DI, Burton DR, Lifson JD, Hasenkrag KJ, McDermott AB, Schultz A, Zamb TJ, Boyle R, Desrosiers RC: HIV vaccine design: insights from live attenuated SIV vaccines. Nat Immunol 2006, 7:19-23.

2. Johnson RP, Desrosiers RC: Protective immunity induced by live attenuated simian immunodeficiency virus. Curr Opin Immunol 1998, 10:436-443.

3. Mills J, Desrosiers R, Rud E, Almond N: Live attenuated HIV vaccines: proposal for further research and development. AIDS Res Hum Retroviruses 2000, 16:453-461.

4. Whitney JB, Ruprecht RM: Live attenuated HIV vaccines: pitfalls and prospects. Curr Opin Infect Dis 2004, 17:17-26.

5. Baba TW, Jeong YS, Penninck D, Bronson R, Greene MF, Ruprecht RM: Pathogenicity of live, attenuated SIV after mucosal infection of neonatal macaques. Science 1995, 267:1820-1825.

6. Baba TW, Liska V, Khimani AH, Ray NB, Dailey PJ, Penninck D, Bronson R, Greene MF, McClure HM, Martin LN, Ruprecht RM: Live attenuated, multiply deleted simian immunodeficiency virus type 1 vaccine in infant and adult macaques. Nat Med 1999, 5:194-203.

7. Chakrabarti LA, Metzner KJ, Ivanovic T, Cheng H, Louis-Virelizier J, Connor RL, Cheng-Mayer C: An attenuated simian immunodeficiency virus strain, SIVmac239, protects macaques against a lethal HIV-1 challenge. Proc Natl Acad Sci USA 2001, 98:2363-2368.

8. Whatmore AM, Cook N, Hall GA, Sharpe S, Rud EW, Crane MP: Repair and evolution of nef in vivo modulates simian immunodeficiency virus virulence. J Virol 1995, 69:1751-1753.

9. Churchill MJ, Rhodes DI, Learmont JC, Sullivan JS, Wesselingh SL, Cooke IR, Deacon NJ, Gorry PR: Longitudinal analysis of human immunodeficiency virus type 1 nef/long terminal repeat sequences in a cohort of long-term survivors infected from a single source. J Virol 2000, 74:1047-1052.

10. Berkhout B, Verhoef K, van Wamel JLB, Back B: Genetic instability of live-attenuated HIV-1 vaccine strains. J Virol 1999, 73:1138-1145.

11. Verhoef K, Marzio G, Hillen W, Bujard H, Berkhout B: Strict control of human immunodeficiency virus type 1 replication by a genetic switch: Tet for Tat. J Virol 2001, 75:979-987.

12. Smith SM, Khoroshev M, Marx PA, Orenstein J, Centlivre M, Berkhout B: Construction of a doxycycline-dependent simian immunodeficiency virus type 1 vaccine by controlling both gene expression and cell entry. J Virol 2005, 79:3855-3858.

13. Zhou X, Vink M, Berkhout B, Das AT: Modification of the Tet-On regulatory system prevents the conditional-live HIV-1 variant from losing doxycycline control. Retrovirology 2006, 3:87.

14. Koseleva Y, Ito Y, Lima RG, Grivel JC, Das AT, Berkhout B, Margolis LB: Depletion of CD4 T lymphocytes in human lymphoid tissue infected ex vivo with doxycycline-dependent HIV-1. Virol 2004, 328:i-6.

15. Das AT, Baldwin CE, Vink M, Berkhout B: Improving the safety of a conditional-live human immunodeficiency virus type 1 vaccine by controlling both gene expression and cell entry. J Virol 2005, 79:3855-3858.

16. Mathews DH, Sabina J, Zuker M, Turner DH: Expanded sequence dependence of thermodynamic parameters improves prediction of RNA secondary structure. J Mol Biol 1999, 288:911-940.

17. Sali A: Mfold web server for nucleic acid folding and hybridization prediction. Nucleic Acids Res 2003, 31:3406-3415.

18. Pauchańska-Włochorek K, Purzycka KJ, Adamiaik RW: New, extended hairpin forms of cDNA TAR RNA point to the structural polymorphism at the 5' end of the HIV-2 leader RNA. Nucleic Acids Res 2006, 34:2984-2997.

19. Berkhout B, Jeang KT: Detailed mutational analysis of TAR RNA: critical spacing between the bulge and loop recognition domains. Nucleic Acids Res 1991, 19:6169-6176.

20. Berkhout B, Back B: Two alternating structures for the HIV-1 leader RNA. RNA 2001, 7:143-157.

21. Berkhout B, Ooms M, Beerens N, Huthoff H, Southern E, Verhoef K: In vitro evidence that the untranslated leader of the HIV-1 genome is an RNA checkpoint that regulates multiple functions through conformational changes. J Biol Chem 2002, 277:19967-19975.

22. Ooms M, Huthoff H, Russell R, Liang C, Berkhout B: A riboswitch regulates RNA dimerization and packaging in human immunodeficiency virus type 1 virions. J Virol 2002, 76:10814-10819.

23. Abbaskin M, Ooms M, Haasnoot PGC, Berkhout B: The HIV-1 leader RNA conformational switch regulates RNA dimerization but does not regulate mRNA translation. Biochem 2005, 44:9058-9066.

24. Das AT, Harwig A, Vrolijk MM, Berkhout B: The TAR hairpin of human immunodeficiency virus type-1 can be deleted when not required for Tat-mediated activation of transcription. J Virol 2007, 81:7742-7748.

25. Vrolijk MM, Ooms M, Harwig A, Das AT, Berkhout B: Destabilization of the TAR hairpin affects the structure and function of the HIV-1 leader RNA. Nucleic Acids Res 2008. Advance Access published June 27.

26. Regier DA, Desrosiers RC: The complete nucleotide sequence of a pathogenic molecular clone of simian immunodeficiency virus. AIDS Res Hum Retroviruses 1990, 6:1221-1231.

27. Vink M, Berkhout B: Role of CD8(+) lymphocytes in control of simian immunodeficiency virus infection and resistance to rechallenge after
transient early antiretroviral treatment. J Virol 2001, 75:10187-10199.

42. Cline AN, Bess JW, Piatak M Jr., Lifson JD: Highly sensitive SIV plasma viral load assay: practical considerations, realistic performance expectations, and application to reverse engineering of vaccines for AIDS. J Med Primatol 2005, 34:303-312.

43. Maudru T, Peden KW: Adaptation of the fluorogenic 5′-nuclease chemistry to a PCR-based reverse transcriptase assay. Biotechn 1998, 25:972-975.

44. Ruijter JM, Thygesen HH, Schoneveld OJ, Das AT, Berkhout B, Lamers WH: Factor correction as a tool to eliminate between-session variation in replicate experiments: application to molecular biology and retrovirology. Retrovirology 2006, 3:1-8.