Target-Directed Azide-Alkyne Cycloaddition for Assembling HIV-1 TAR RNA Binding Ligands

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In memory of Professor Rolf Huisgen

Abstract: The highly conserved HIV-1 transactivation response element (TAR) binds to the trans-activator protein Tat and facilitates viral replication in its latent state. The inhibition of Tat–TAR interactions by selectively targeting TAR RNA has been used as a strategy to develop potent antiviral agents. Therefore, HIV-1 TAR RNA represents a paradigmatic system for therapeutic intervention. Herein, we have employed biotin-tagged TAR RNA to assemble its own ligands from a pool of reactive azide and alkyne building blocks. To identify the binding sites and selectivity of the ligands, the in situ cycloaddition has been further performed using control nucleotide (TAR DNA and TAR RNA without bulge) templates. The hit triazole-linked thiazole peptidomimetic products have been isolated from the biotin-tagged target templates using streptavidin beads. The major triazole lead generated by the TAR RNA presumably binds in the bulge region, shows specificity for TAR RNA over TAR DNA, and inhibits Tat–TAR interactions.

Human immunodeficiency virus type-1 (HIV-1) contains a cis-acting regulatory element called TAR RNA located upstream of the transcriptional start site (+1 to +59). TAR RNA forms a stable hairpin structure with a hexanucleotide loop and a three base bulge (UCU). A trans-activator protein Tat binds to the bulge region of the TAR RNA and activates HIV-1 replication by stimulating the elongation efficiency of RNA pol II.[3] The interaction between TAR and Tat is essential for viral replication and growth; however, mutated TAR shows a reduced affinity for Tat and is unable to replicate efficiently.[4] Various strategies have been developed to inhibit viral replication, such as targeting the HIV-1 protease and the long terminal repeat (LTR) region. Although numerous classes of small molecules have been reported,[5,6] mutated virus makes them ineffective over time.[7] However, the TAR sequence and its heterogeneity are highly conserved throughout the evolution of the virus (Figure 1a).[8] Therefore, the TAR RNA can be targeted for an anti-HIV therapeutic strategy because of its structural integrity and central role in HIV-1 replication.

Herein we demonstrate the rapid discovery of HIV-1 TAR-Tat inhibitors by TAR RNA guided in situ cycloaddition of azide and alkyne fragments. The in situ cycloaddition was also performed with a TAR RNA without the UCU bulge (TAR RNA w/b; w/b = without bulge) and a TAR DNA (Figure 1). The target-guided synthetic (TGS) approach, commonly known as an in situ click reaction, accelerates the discovery of novel pharmaceutical molecules through the involvement of the biological target itself in the selection process.[10] This bio-orthogonal approach has been demonstrated previously using enzymes and a few nucleic acid targets,[11] however, TAR RNA has not been employed as a target to date. The in situ click approach is a unique process which relies on target-guided Huisgen 1,3-dipolar cycloaddition of alkyne and azide residues to produce five-membered nitrogen-containing triazole heterocycles.[12] Triazoles are stable under oxidative or reductive conditions as well as to basic or acidic hydrolysis. Furthermore, they actively participate in hydrogen bonding and π–π stacking interactions with biological targets.[13]

Both alkynes and azides are energetic species and there is a high kinetic barrier for cycloaddition. The kinetic barrier can be overcome by bringing the reactants into proximity with a biological template or with a catalyst.[14] The biological target samples various combinations of the alkynes and azides and synthesizes the best binders with high efficacy. Given the dynamic folding topology of RNA structures, the in situ...
cycloaddition approach could be a potential strategy for generating selective RNA-binding ligands. Various antibiotics and different classes of ligands have been reported previously for targeting TAR RNA. Herein, TAR RNA has been employed for the first time as a template to generate its specific binder that could efficiently inhibit Tat–TAR interactions. For the in situ cycloaddition, we designed and synthesized a library of alkyn (1a-d) and azide (2a-k) building blocks (Figure 2a, see also Schemes S1–S3 in the Supporting Information). Alkynes 1a–c consist of a different number of five-membered thiazole heterocycles, prepared by iterative amide coupling. The thiazole ring system and peptides are privileged scaffolds in medicinal chemistry. Alkyne 1d is considered to have pharmacological activities because of its carbazole ring system. Our azide library 2a–k consists of aliphatic and aromatic functional groups such as amines (2a, 2e, 2f, and 2h), a carboxylic acid (2b), a nitro (2d), an ester (2g), an amino acid (2h), a carboxamide (2j), and a guanosine azide (2k).

The combinations of these building blocks could lead to 88 possible regioisomers (anti-(1,4)- and syn-(1,5)-triazoles).

The multicomponent reaction consisting of 4 alkynes and 11 azides was stirred in the presence of TAR RNA to generate potential hit ligands. As a model system for our study, we used a 5′-biotin-tagged 29 nt TAR RNA (Figure 1a). The biotin-tagged RNA can be easily separated from the reaction mixture along with the hit triazole products by magnetic separation using streptavidin magnetic beads (Figure 3). For the in situ RNA-templated reaction, the alkynes and azides were incubated in a 1:4 ratio together with the target TAR RNA (10 µM) in 20 mM sodium cacodylate, 180 mM NaCl, and 10 mM MgCl₂ buffer at pH 7.4 and 25 °C for 72 h. To identify the selective leads for the TAR RNA, control experiments were performed with biotinylated TAR RNA w/b and biotinylated TAR DNA (Figure 1b,c).

The hit products were characterized by HPLC-MS analysis, which showed the formation of two triazole products 3ca (cycloadduct of alkyne 1c and azide 2a) and 3ba (cycloadduct of alkyne 1b and azide 2a) in a ratio of 41:59 by the TAR RNA (Figure 4a(ii)). Since the triazole product 3ca was also formed in the presence of the control TAR DNA (Figure 4a(iii)), the triazole lead 3ba was identified as the selective lead for TAR RNA. In addition, triazole product 3aa (cycloadduct of alkyne 1a and azide 2a) was selectively formed using the control TAR RNA w/b (Figure 4a(iii)). The TAR RNA leads 3ca and 3ba were not formed by the TAR RNA w/b, thus suggesting an interaction between these ligands and the UCU bulge region of the TAR RNA. We further performed in situ cycloaddition of individual alkynes with the azide library in the presence of the TAR RNA and controls (Figure S2b). HPLC and ESI-MS analysis showed the reactions yielded similar lead compounds (Figure 4b, see also Figure S2a).

Hit triazole products 3aa, 3ba, and 3ca were prepared from their respective alkyne and azide fragments under Cu-catalyzed conditions (Figure S2b). The products were obtained in high yields as anti-(1,4-triazole) regioisomers. Comparison of the HPLC traces of the RNA-templated products with the authentic 1,4-triazole samples (Cu-catalyzed products) revealed that the RNA targeting reactions generated predominantly the anti-(1,4-triazole) isomers.

The yields of the lead anti-triazoles were further determined by time-dependent cycloaddition of the corresponding alkyne (1a, 1b, and 1c) and azide (2a) fragments in the presence of the biotin-tagged target and control DNA and RNA targets (Figure S4). Alkyne 1b reacted with azide 2a in the presence of TAR RNA to form anti-3ba in 62% yield (Figure S3a). Similarly, lead anti-3ca (from 1c and 2a) was obtained in 54% yield with TAR RNA (Figure S3b). However, in the presence of TAR DNA, anti-3ca was formed in 42% yield. Lead anti-3aa (from 1a and 2a) was formed in 51% yield in the presence of the control TAR RNA w/b.

Analysis of the binding properties of the hit triazole products against the target TAR RNA and controls were performed using isothermal titration calorimetry (ITC) studies (Figure 5a,b, see also Figure S5). Ligand 3ba showed an exothermic binding interaction with TAR RNA with a dissociation constant (K₅) of 0.49 µM. In comparison, ligand 3ca showed an approximately 15-fold lower binding affinity, with K₅ = 7.2 µM for the TAR RNA (Table 1). Both ligands were found to bind to TAR RNA with a 1:1 stoichiometry (N) and the ΔG values were found to be −9.54 kcal mol⁻¹ for 3ba and −7.01 kcal mol⁻¹ for 3ca. Moreover, 3ba interacted neither with the TAR RNA w/b nor with the TAR DNA. However, 3ca exhibited exothermic binding interactions with TAR DNA, with K₅ = 11.1 µM and ΔG = −6.76 kcal mol⁻¹. Ligand 3aa showed exothermic binding interactions with TAR RNA w/b, with K₅ = 9.4 µM and ΔG =
Table 1: Binding interactions of triazole leads (3aa, 3ba, and 3ca) with TAR RNA, TAR DNA, and TAR RNA w/b.

| RNA and DNA | Hit ligands | $K_d$ [μM] | $ΔG$ [kcal mol$^{-1}$] | $F/F_0$ | $K_d$ [μM] | DC50 [μM] |
|-------------|-------------|-----------|-----------------------|--------|-----------|-----------|
| TAR RNA     | 3aa         | –         | –                     | 1.6    | –         | –         |
|             | 3ba         | –         | –                     | 5.0    | 0.54      | 1.5       |
|             | 3ca         | –         | –                     | 5.2    | 6.7       | 10.6      |
| TAR DNA     | 3aa         | –         | –                     | 1.7    | –         | –         |
|             | 3ba         | –         | –                     | 1.5    | –         | –         |
|             | 3ca         | –         | –                     | 4.3    | 10.7      | –         |
| TAR RNA w/b | 3aa         | 9.4       | –6.23                 | 3.8    | 11.3      | –         |
|             | 3ba         | –         | –                     | 1.8    | –         | –         |
|             | 3ca         | –         | –                     | 1.7    | –         | –         |

[a,b] ITC and fluorescence experiments were conducted in 20 mM sodium cacodylate, 180 mM NaCl, and 10 mM MgCl$_2$ at pH 7.4. 3aa: $λ_{em}$ = 280 nm, $λ_{ex}$ = 435 nm; 3ba: $λ_{em}$ = 280 nm, $λ_{ex}$ = 393 nm, and 410 nm; 3ca: $λ_{em}$ = 280 nm, $λ_{ex}$ = 435 nm. $K_d$ values are determined from [a] ITC and [b] fluorescence titrations ($K_d$ = ±5%).

The modeling study revealed that the lead compound 3ba showed an improved binding with $K_d = 0.54$ μM compared to 3ca with $K_d = 6.7$ μM (Table 1). Moreover, ligand 3ca also displayed considerable binding interactions with the control TAR DNA and showed a 4.3-fold increase in fluorescence intensity and $K_d = 10.74$ μM (Figure S6a). In comparison, 3ba didn’t result in a significant increase in the fluorescence intensity in the presence of TAR DNA and TAR RNA w/b (Table 1), which is in agreement with the ITC results. In addition, 3aa exhibited a 3.8-fold increase in fluorescence intensity with TAR RNA w/b with $K_d = 11.3$ μm, and didn’t interact with the TAR RNA or the control TAR DNA (Figure S6b).

We further assessed the ability of the lead anti-triazole ligands to displace the Tat peptide from a preformed complex between fluoresceinated carboxytetramethylrhododendrin (TAMRA) labeled Tat peptide and fluorescein amidite (FAM) labeled TAR RNA by a Förster resonance energy transfer (FRET) based assay (Figure 6c). In the absence of ligands, the association of Tat with TAR RNA results in an efficient quenching of the FAM dye. The addition of the ligand may displace the Tat peptide, thereby resulting in an increase in the fluorescence emission of the FAM dye (Figure 6c, see also Figure S7). The displacement of Tat from TAR is expressed as DC50, which reflects a ligand’s ability to displace 50% of the bound Tat peptide from the TAR RNA. Ligand 3aa showed a lower DC50 value (1.5 μM; Figure 6b) than ligand 3ca (DC50 = 10.6 μM), which indicates 3ba has a higher affinity for TAR RNA. However, ligand 3aa wasn’t able to displace 50% of the bound Tat peptide from the TAR RNA (Table 1), which suggests that the ligand 3ba may displace the peptide by binding to the UCU bulge of TAR RNA. The biophysical experiments—ITC, fluorescence, and displacement studies—are thus in agreement with the in situ cycloaddition results.

Molecular docking of 3ba and 3ca with the target RNA (PDB: 1ANR) was performed with the AutoDock program v4.1 to elucidate the structure–activity relationship of the ligands with the RNA structure (Figure 7, see also Figure S8). The modeling study revealed that anti-3ba could bind to the bulge region of the target while anti-3ca interacted with the major groove region of the target RNA. The presence of a -CH$_2$ linker provided added flexibility to the first thiazole

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ring of anti-3ba, thereby making it favorable for interacting with the bulge region of the TAR RNA. The water-soluble chain of anti-3ba interacted with the phosphate backbone of the other strand, thus making the interaction even more stable ($\Delta G = -9.12 \text{ kcal mol}^{-1}$). Anti-3ca, on the other hand, exhibited a planar crescent-shaped structure which stacks between the major groove of the TAR RNA ($\Delta G = -6.85 \text{ kcal mol}^{-1}$) and TAR DNA. The bulge region further
widens the groove (in TAR RNA and DNA) and makes the stacking interactions more favorable but nonspecific for anti-

3ca.

In summary, in situ click chemistry has been successfully applied to discover novel thiazole-based peptidomimetics as HIV-1 TAR-Tat inhibitors. The TAR RNA itself acts as a template for the cycloaddition and promotes the formation of a thiazole peptidomimetic 1,4-triazole lead that shows high selectivity for TAR RNA. Isothermal calorimetric and fluorescence studies reveal that a thiazole peptidomimetic with two thiazole rings and flexible methylene units strongly interacts with the bulge region of TAR RNA and effectively inhibits the Tat-TAR RNA interactions. Further studies are currently underway to apply this method for the discovery of novel TAR RNA inhibitors and to investigate their antiviral activities.

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Conflict of interest

The authors declare no conflict of interest.

Keywords: click chemistry · cycloaddition · HIV-1 TAR RNA · peptidomimetics · Tat peptide

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