



MEETING ABSTRACT

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Comparison of Tax-1 and Tax-2B post-translational modifications using specific lysine mutants in relation to activation of NF- κ B and intracellular localization

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From 15th International Conference on Human Retroviruses: HTLV and Related Viruses
Leuven and Gembloux, Belgium. 5-8 June 2011

Post-translational modifications of HTLV-1 and HTLV-2 Tax-1 and Tax-2 proteins have been shown to play a critical role in their cellular localization, transactivation and protein interactions. Five of ten lysine residues were found to be major targets for Tax-1 modifications: Lys189(K4); Lys197(K5), Lys263(K6), Lys280(K7) and Lys284(K8), are essential for ubiquitination, while sumoylation takes place on Lys280 (K7) and Lys284(K8). Tax-2 contains four additional lysine residues, namely at position Lys100(K2i), Lys149(K3i), Lys185(K3ii), and Lys356(K10i).

Very few studies have been so far performed on Tax-2 lysine mutants. We have previously demonstrated that Tax-2B is ubiquitinated and sumoylated similarly to Tax-1. To identify the Tax-2 lysine residues which are directly involved in post-translational modifications, we have constructed a series of Tax-2B mutants with substitutions of lysine (K) residues by arginines (R) and analyzed them for NF- κ B and CREB/ATF transactivation, intracellular distribution and extent of ubiquitination and sumoylation. We have found that Tax-2 K7-8R mutant, contrary to its Tax-1 homologue, is only partially affected in its capacity to transactivate NF- κ B pathway, is regularly sumoylated and presents formation of nuclear bodies by confocal analysis. However, Tax-2 mutants with extended (K3ii-8R) and/or total (K1-10iR) mutation rate were severely affected for NF- κ B transactivation and sumoylation. By comparing Tax-2 WT with

mutants K7-8R and K3ii-8R, we observed that the reduction of NF- κ B activity is correlated to a parallel decrease in sumoylation. These results suggest that the target for Tax-2 ubiquitination and sumoylation differs from that described for Tax-1.

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Published: 6 June 2011

doi:10.1186/1742-4690-8-S1-A143

Cite this article as: Turci et al.: Comparison of Tax-1 and Tax-2B post-translational modifications using specific lysine mutants in relation to activation of NF- κ B and intracellular localization. *Retrovirology* 2011 **8** (Suppl 1):A143.

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