

# A model of N-terminal Cyclin T1 based on FRET experiments

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Human Cyclin T1 is the cyclin partner of kinase CDK9 in the positive transcription elongation factor b (P-TEFb). P-TEFb is recruited by Tat, the transactivator of the human immunodeficiency virus type 1 (HIV-1), to the viral promoter by direct interactions between Tat, Cyclin T1 and the *cis*-acting transactivation-responsive region (TAR) present at the 5'-end of each viral mRNA. At present, no structural data for Cyclin T1 are available. Here, we build a structural model of an N-terminus portion of Cyclin T1 (aa 27–263) based on the X-ray structure of Cyclin H. The model is compared with site directed mutagenesis data from the literature and validated by fluorescence resonance energy transfer (FRET) using Tat as a probe in living cells. This model provides a first step towards the structural characterization of the CDK9–CycT1–Tat–TAR complex, which is crucial for HIV-1 replication and may constitute a promising target for pharmaceutical intervention.

**Keywords:** Cyclin T1; HIV-1; Tat; FRET; Comparative modelling; P-TEFb

## 1. Introduction

The cyclin family of proteins plays a fundamental role in the cell cycle functioning as regulatory subunits of the so-called cyclin dependent kinases (CDK), which act as catalytic subunits [1]. Besides regulation of cell cycle by phosphorylation of a large variety of substrates, Cyclin/CDK complexes are also involved in the control of different cellular mechanisms such as differentiation, apoptosis and, for some members of the family, such as Cyclin H (CycH), Cyclin C (CycC) and Cyclin T1 (CycT1), transcription initiation and/or elongation [2].

The common structural feature of the family is the cyclin box motif. This is a characteristic two-repeats folding motif of ~100 amino acids long [3,4] that constitute a versatile scaffold for mediating protein–protein interactions. Each repeat is composed of five helices, connected by a linker peptide in extended conformation. The helices within each repeat are spatially disposed with the very hydrophobic helix H3 surrounded

by the other four. Remarkably, this motif bears a high degree of structural conservation even at very low primary sequence identity. The relative orientation of the repeats can vary as found for the transcription factor II B (TFIIB) that shows a different arrangement. Cyclin boxes are usually inserted into a protein frame, with additional helical elements found at the N and C termini of the cyclin box. These segments are not conserved among the cyclin family and they likely provide binding specificity for protein–protein interactions.

Cyclin T1 is the cyclin associated to CDK9 in the positive transcription-elongation factor b (P-TEFb), a general elongation co-factor for RNA polymerase II (RNAPII) directed transcription [34]. P-TEFb is able to phosphorylate the carboxy-terminal domain (CTD) of RNAPII, a molecular event associated with increased transcriptional processivity. Initially, CycT1 has been discovered as a cofactor of the human immunodeficiency virus type 1 (HIV-1) Tat transactivator [5]. A tripartite complex is formed between Tat, CycT1 and the *cis*-acting

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transactivation-responsive region (TAR) present at the 5'-end of each viral mRNA. The formation of the P-TEFb-TAR-Tat complex is an essential step towards the assembly of the processive RNAPII machinery at the LTR promoter [6–9]. CycT1 can be grossly sub-divided into two major domains (figure 1(A)). The N terminus of the protein (aa 1–300) shares partial homology with other members of the Cyclin T family (cyclin box) [5]. It contains the Tat recognition motif (TRM) that includes a critical cysteine residue at position 261 present in human CycT1, but absent in its rodent homologue [6,8]. The N terminus domain of CycT1 is also important for the protein–protein associations of the protein with CDK9, NF- $\kappa$ B, CIITA and c-Myc [10–12]. The C terminus of CycT1 is less well characterized and contains a putative coiled-coil region (aa 379–430; coil); a histidine-rich domain (aa 506–530; His) and a carboxy-terminus PEST sequence (aa 700–726). It has been recently shown that

the C terminus of CycT1 is involved in the association with the CTD domain of RNAPII [13,14] and the promyelocytic leukaemia protein, PML [15]. At present, no three-dimensional structure of CycT1 is available. Here, we use bioinformatics tools to model the N terminus (aa 27–263) of CycT1. We build the structural model based on the X-ray structure of CycH, another member of the cyclin family involved in transcription, whose X-ray structure is available. The model fits with all mutagenesis data present in the literature. Furthermore, we take advantage of the complex interactions between the HIV-1 Tat protein and CycT1 *in vivo* to validate the prediction of close proximity between the N and C termini parts of the Cyclin box. Specifically, we measure the fluorescence resonance energy transfer (FRET) between fluorescent proteins genetically fused to the N and C termini of Cyclin T1 and to the C termini of Tat in living cells.

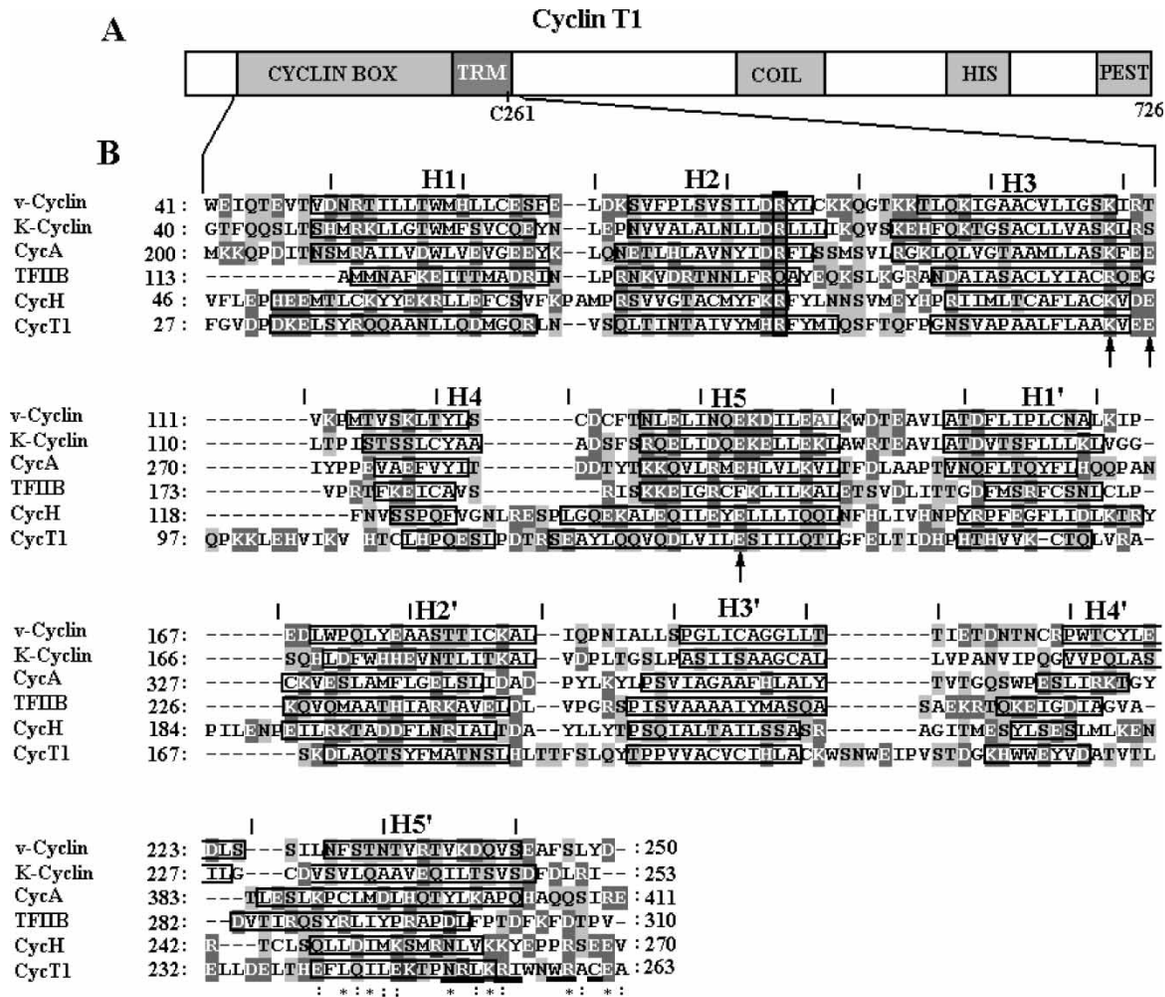


Figure 1. Alignment of Cyclin-boxes. (A) Domains representation of CycT1. (B) Sequence alignment of the cyclin box of CycT1 with the structural templates. The chemical character of the residues is indicated by background colours. Black characters in white background correspond to hydrophobic aminoacids, light grey are polar and dark grey with white characters are charged residues. Black rectangles indicate helical elements as found in crystallographic structures. The conserved arginines that fix the relative orientation of the repeats in all but TFIIB proteins is indicated with a vertical rectangle. Arrows indicate residues whose charge inversion by mutagenesis inhibits CycT1:CDK9 binding. Residues belonging to the Tat recognition motif are underlined. Homology between C-terminal motifs of CycH and CycT1 are indicated with stars for identities and colons for conservative mutations.

## 2. Methods

### 2.1 Bioinformatics

The protocol used for the construction of the structural model of CycT1 (Swiss Prot entry O60563, [5]) was the following:

**2.1.1 Identification of appropriate templates.** We considered a set of five non redundant cyclin proteins whose structure was determined by X-ray crystallography: (i) a viral cyclin (v-cyclin) from herpesvirus saimiri, homolog to type D cyclins, able to specifically activate human CDK6 (PDB code:1BU2) [16]; (ii) a viral D-type cyclin encoded by the Kaposi's sarcoma-associated herpesvirus (K-cyclin, PDB code:1G3N) [17]; (iii) the human Cyclin A (CycA, PDB code:1FIN) [18]; (iv) the general transcription factor II B (TFIIB) [19] (PDB code:1VOL) and (v) the human transcriptional cyclin H (PDB code:1JKW) [20].

**2.1.2 Sequence alignment.** Primary sequences of the above-mentioned cyclins were initially aligned using T-COFFEE (<http://www.ch.embnet.org/software/TCoffee.html>), which was shown to be more efficient in case of low sequence identity [21]. The alignments were then manually refined so as to take into account the secondary structure data.

**2.1.3 Selection of the structural template.** Among the above quoted structural templates, CycH is the only one human protein involved in transcription activation as CycT1. In addition, CycH was also reported to interact with Tat [22]. Indeed, the C-terminal segment of CycH shares high homology with its corresponding part in CycT1 (figure 1) that contains the TRM. For these reasons, CycH was chosen as structural template to coordinate mapping to our target CycT1. The X-ray structure of CycH was determined at a resolution of 2.6 nm with 269 residues traced corresponding to two cyclin box repeats (about 100 amino acids each). Before and after the cyclin box two N and C terminal helical motifs,  $H_N$  and  $H_C$  of 21 and 15 residues long, respectively complete the structure of CycH.

**2.1.4 Construction of the structural models.** Although CycT1 is 726 amino acids long, deletion mutant experiments shown only the N terminus 1–263 [23] amino acids are needed to bind CDK9 and Tat. Our model is based on the structure of CycH from residues 46 to 270. This fragment was mapped to amino acid 27 to 263 of CycT1. Therefore, it covers the necessary protein length relevant for the study of the CycT1 with respect to its functional association with CDK9 and Tat. The construction and refinement of the backbone and side chains of the target protein were performed with the program Modeller V6.2 [24,25].

**2.1.5 Selection of the best model.** The best structural model was based on Modeller's restraints violation. The model selected (figure 2) exhibited a good Ramachandran map, with 93.9% of the residues in most favoured regions, 5.2% in allowed regions and only 0.8% (Ser188 and Thr230) in unfavourable regions.

### 2.2 Experimental methods

**2.2.1 Plasmids and cells.** The expression plasmids pcDNA3-Tat-BFP and pcDNA3-EGFP-CycT1, containing EGFP fused at their 3'- and 5'-ends, respectively, were characterized previously [26]. The plasmid pCycT300-EGFP, carrying EGFP at the 3' end, was obtained by PCR amplification of the N terminus of CycT1 with primers: FW 5'-gc gaa ttc atg gag gga gag agg aag aac-3', RV 5'-gc gga tcc ag gta att tgg acg tta cca-3' and cloning as an EcoRI/BamHI fragment into pEGFP-N1 (Clontech). The plasmid pBFP-CycT300-EGFP, carrying BFP at the 5'-end and GFP at the 3'-end, was obtained by PCR amplification of the previous construct pCycT300-EGFP with primers: FW 5'-gc aag ctt cg gag gga gag agg aag aac-3', RV 3'-gc gtc gac tta ctt gta cag ctc gtc-5' and cloning as a HindIII/Sall fragment into pBFP-C1 (Clontech). To control for correct folding of the fusion proteins we performed transcriptional co-activation assays of the HIV LTR using Tat-EBFP and the various cyclin constructs as described in [27].

The HL3T1 cell line used in this study were cultured in Dulbecco's modified Eagle's medium (DMEM) supplemented with 10% foetal bovine serum.

### 2.2.2 Fluorescence resonance energy transfer (FRET).

The presence of FRET indicates actual protein–protein interaction at distances in the range of the FRET length scale, the Förster radius ( $R_0$ ), defined as the distances at which FRET efficiency is 50%. FRET is exploited *in vivo* by the use of the green fluorescent protein family of fluorophores. For the green fluorescent protein (GFP):blue fluorescent protein (BFP) pair  $R_0 \approx 4$  nm, this implies that simple co-localization of two proteins is not sufficient to yield energy transfer. Epithelial cells (HL3T1) were transiently transfected by the calcium phosphate method with the various expression plasmids fused to the optically-matched fluorescent tags in four-chambers of glass slides. Cells were fixed in 4% paraformaldehyde after 48 h and mounted directly in 70% glycerol for FRET analysis. FRET measurements were carried out as previously described by an epi-fluorescence Axioskop 2 Zeiss microscope mounting a 103 W HBO lamp, a  $100 \times 1.3$  N.A. oil-immersion Plan-Neofluar objective, and Nomarski optics [26,27]. FRET analysis was performed in two steps. Firstly, enhanced green fluorescent protein (EGFP) emission was collected by integrating the fluorescence signal around 520 nm (bandwidth 40 nm) under EGFP excitation at 480 nm (wavelength selection was obtained by 40 nm band-pass filters, excitation power was  $5 \text{ W/cm}^2$ ). Secondly, EGFP emission in the same frequency range was measured

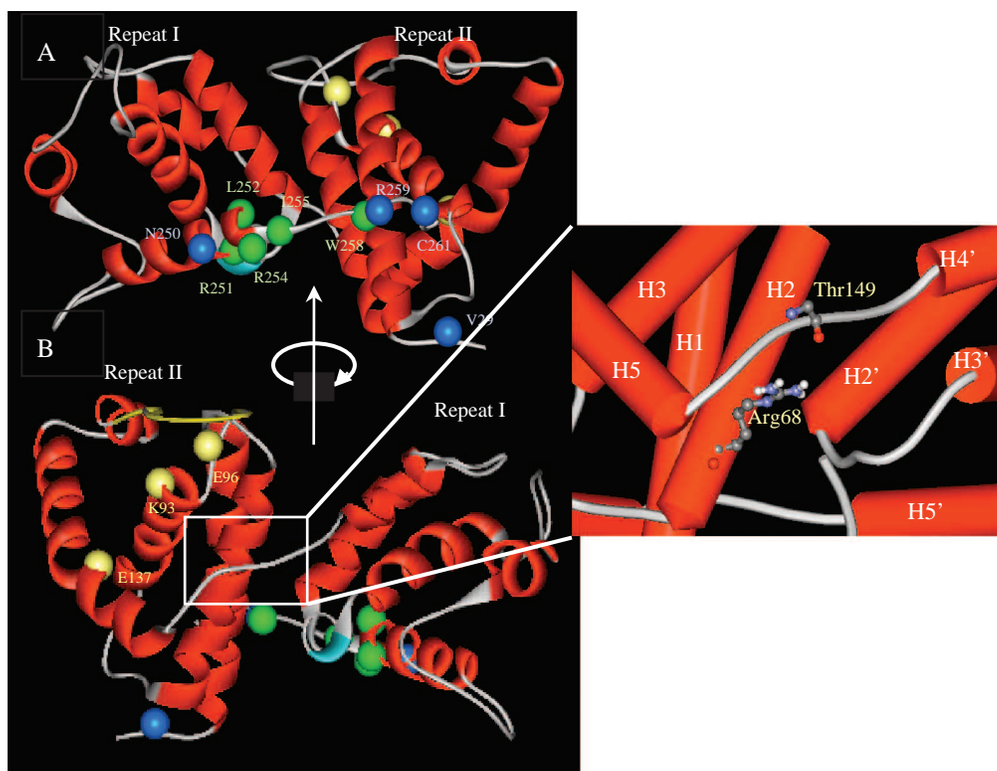


Figure 2. Modelling of CycT. (A) 3D structure of CycT1 as obtained from comparative modelling with CycH. C $\alpha$  of Val29, Asp250, Arg259 and Cys261, which are known to interact directly or indirectly with Tat are shown in light blue with ball representation. C $\alpha$  of Arg251, Leu252, Arg254, Ile255 and Trp258 are shown as green balls. (B) CDK9 binding region (the molecule is rotated 180° with respect to A). C $\alpha$  of Lys93, Glu96 and Glu137, involved in CDK9 recognition are represented in balls and sticks. Glu147 and the loop (marked in yellow) between H3 and H4 that may putatively participate in CDK9 binding are also shown. Inset: a close up showing the conserved interaction between Arg68 and the carbonyl moiety of the Thr149 backbone.

after excitation at 350 nm (power density 2 W/cm<sup>2</sup> and bandwidth 60 nm). Background was detected out of the cell under study for each frame and subtracted from the relevant fluorescent signal. Following this procedure, the ratio between the two measured EGFP emissions (data taken following excitation at 350 nm divided by those at 480 nm) provides the FRET signal. Fluorescence was collected by a PentaMax 512-EFT intensified CCD camera with detection times of the order of 0.1 s (in particular, for data taken under excitation at 350 nm they were five times longer than for those relative to 480 nm excitation). Data acquisition and analysis were performed with the Metamorph software from Universal Imaging Corporation, a subsidiary of Molecular Devices. When evaluating FRET ratios, emission intensities were scaled to take into account the different detection times.

### 3. Results and discussion

#### 3.1 Structural model

**3.1.1 Sequence alignment.** Our alignment with the five templates spanning the respective cyclin box domains is shown in figure 1. We considered a 236 residues long fragment of CycT1 spanning from Phe27 to Ala263 that covers the protein portion involved in HIV-1 Tat and CDK9 binding. As can be observed from figure 1, all the

gaps lie within loopy regions, and better conservation is obtained within the first domain.

As found for other members of the Cyclin family of proteins [4], the percent of identity between CycT1 and the templates is low (table 1). However, the characteristic cyclin box fold is structurally well preserved even at low degrees of sequence conservation. The high hydrophobic character of helix H3 that is surrounded by the other four helices in each repeat is clearly seen from figure 1. The existence of gaps before and after helices H4 and H4', suggest either, the presence in CycT1 of longer loops or helical elements than in all the other cyclins. Site directed mutagenesis experiments on positions Lys93 and Glu96 [28,29], immediately before

Table 1. Percent of sequence identity between CycT1 and the templates obtained from the sequence alignment (figure 1) and root mean square deviation (RMSD) calculated per cyclin repeat and on the whole cyclin boxes. The end to end distance reported correspond to the distance between the first and last residue aligned for each protein.

	CycH	CycA	TFIIB	v-Cyc	K-Cyc
% Identity	14	19	14	9	11
RMSD [Å] (Global)	0.8	1.6	7.7	2	1.5
RMSD [Å] Repeat I	0.5	1.4	1.4	1.3	1.4
RMSD [Å] Repeat II	0.9	1.7	1.5	1.4	1.6
End to end distance [nm]	1.4	2.6	5.6	2	2.3

the first gap, demonstrates that CDK9's binding site is located in this zone, between helix H3 and H4.

The C terminus motifs of CycH and CycT1 align without gaps after H4', at the end of the cyclin box. This segment contains the TRM (residues 250–262) of CycT1. Among all the templates, CycH presents the highest homology with CycT1 (figure 1).

**3.1.2 3D-model.** Based on the alignment of figure 1, a structural model of the protein was built (figure 2) using the structure of CycH as template. The N and C termini of the cyclin box lie on the first repeat, separated 11 Å one from the other. The relative orientation of both repeats is stabilized by the highly conserved residue Arg68 [3], which is in good position to form H-bonds with Tyr49 backbone carbonyl moieties (figure 2). This interaction is conserved in all the templates, with the exception of TFIIB, and was reported to be crucial to fix helix H2 to the linker, and hence, the relative orientation between repeats I and II [20,30]. In fact, the cyclin box of TFIIB presents a glutamine instead of an arginine in the corresponding position. This change provokes the weakness of this strongly stabilizing interaction with the result that the relative orientation of both repeats is different from the rest of the cyclins.

Structural superposition performed on the whole molecules, or considering only one repeat at the time, indicate that the structural dispersion between all the cyclins considered and our model of CycT1 is indeed small (table 1) suggesting that any of the cyclin structures could be used as structural template obtaining practically identical results.

**3.1.3 Consistency with experimental data.** We then investigated the location of the residues of CycT1, which bind to its molecular partners (Tat, TAR and CDK9) and for which mutagenesis data are available in the literature. All the residues reported to participate in CDK9 binding to CycT1 (Lys93, Glu96 and Glu137) [29,30] were solvent-exposed and in close proximity (figure 2). These residues are also highly conserved along the cyclin family [3] (figure 1). Lys93 and Glu96 are located at the end of helix H3, whilst Glu137 is located in the middle of H5. We further notice that other charged residues such as Lys99/100 Glu102 located in the loop connecting H3 with H4 and Glu147 within H5 are also exposed to the solvent and near to the putative CDK9 binding region, so they might also play a role in CDK9 binding.

All the residues known to bind to HIV-1 Tat or TAR [8] (figure 2(A)) resulted exposed to the solvent. Asn250, Arg251 and Leu252 are located in last turn of helix H5', while Arg254, Ile245, Trp258, Arg259 and Cys261 lie in coil conformation.

This conformation is strongly supported by recently reported proteolysis experiments [31] which showed that (i) CycT1 TRM is susceptible to digestion and therefore, is in a flexible conformation. (ii) CycT1 is cleaved after Leu252, indicating that this is the last residue in ordered

(helical) conformation, which is in nice agreement with our theoretical predictions.

Finally, Val29, whose mutation to Leu was reported to render human CycT1 active for equine Tat transactivation in EIAV [32], is also exposed and oriented towards the TRM.

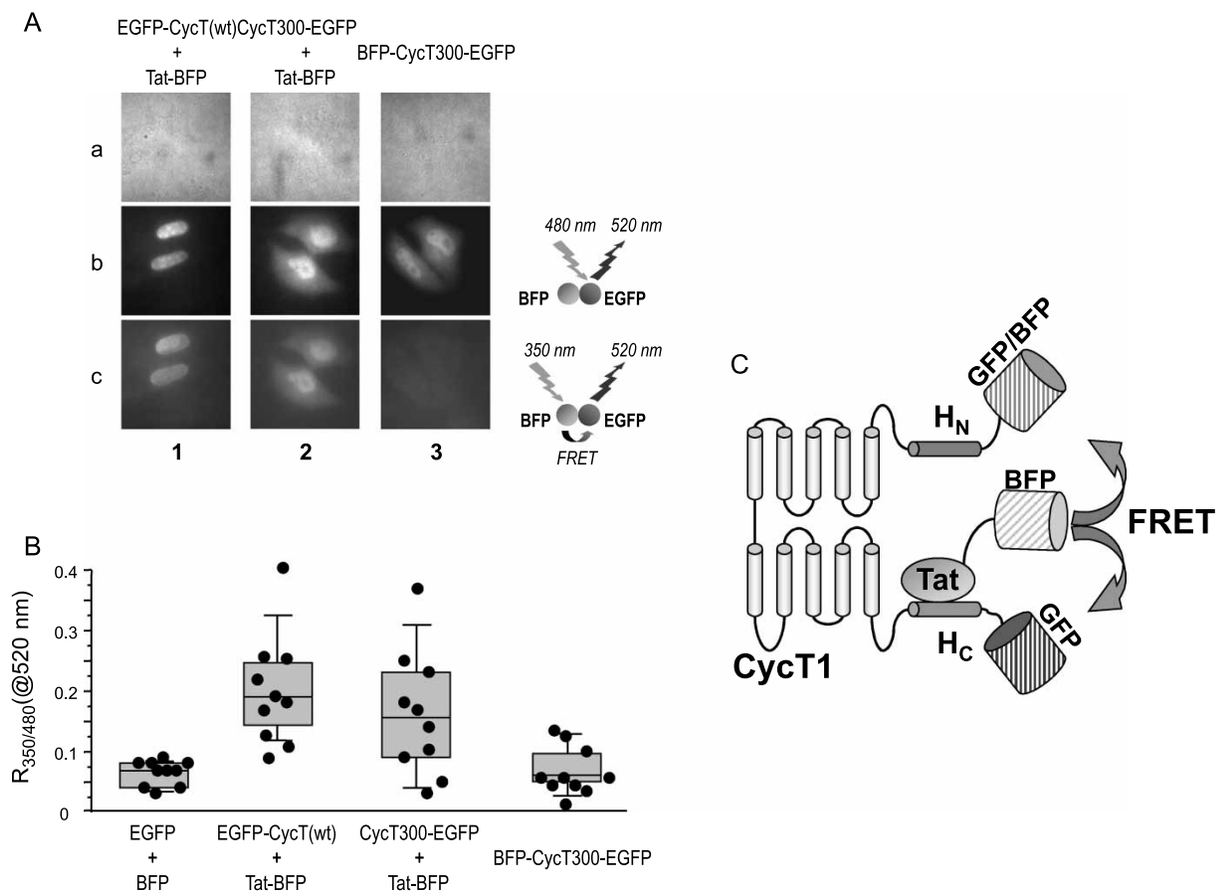
### 3.2 Validation of the model

As stated above, the cyclin box fold is highly conserved even at low identity levels. Therefore, the largest assumption in our model is that in CycT1 the relative orientation of both the cyclin repeats that determine the distance between N and C termini is very similar to that of CycH. To validate this proposal, we performed *in vivo* FRET experiments. FRET has been already successfully used in the last few years to detect CycT1–protein interaction within living cells, using the GFP family of fluorophores [15,26]. Here we used this technique to estimate the proximity between the N and C termini of the protein, which may sensibly vary with the relative orientation of the cyclin repeats.

As a first attempt, we tried to provide an estimation of this proximity by *intra*-molecular FRET in cells co-transfected with plasmids expressing CycT1 tagged at the N-terminus with the BFP and at the C-terminus with EGFP. FRET image analysis of individually transfected cells are shown in figure 3(A), column 3. FRET analysis was performed by comparing EGFP emission at 520 nm (the peak wavelength of EGFP emission), following BFP excitation at 350 nm (figure 3(A), panels in row c), with that following excitation at 480 nm of the same cells (figure 3(A), panels in row b). As it can be seen from figure 3, in this case no FRET signal was obtained.

We then proceed to investigate this proximity by *inter*-molecular FRET, taking advantage of the fact the CycT1's TRM is located around position 261, within the C-terminal helix after the cyclin box. We therefore co-transfected human HL3T1 cells with plasmids expressing Cyclin T1, tagged at the N-termini with EGFP, and Tat tagged at the C-termini with BFP. Most cells transfected with EGFP-Cyclin T1:Tat-BFP (figure 3(A), panel b1) showed the characteristic nuclear punctuated pattern of Cyclin T1 [26,33]. Samples expressing both EGFP-CycT1 and Tat-BFP scored positive for FRET. This suggests that the distance between the N-terminus of CycT1 TRM and the C-terminus of CycT1 is in the range of that of the template (table 1).

Based on these findings, we tested whether fusing GFP at the C terminus of CycT1 provides FRET with Tat-BFP. Cells transfected with CycT300-EGFP:Tat-BFP (figure 3, column 2) showed a nucleo-cytoplasmic distribution of CycT300-EGFP at 520 nm under excitation at 480 nm. The difference of distribution of the protein with respect to the wild-type CycT1 reflected the lack of the carboxy-terminal domain of the protein, which was responsible for the punctuated nuclear localization through binding to the PML protein [15]. EGFP emission at 520 nm, following BFP excitation at 350 nm, scored positive for FRET,



**Figure 3.** FRET analysis *in vivo*. (A) FRET visualization in human cells. The plasmid constructs indicated on top of each column were transfected in HL3T1 cells (a HeLa derivative cell line carrying an integrated LTR CAT cassette). Transfected cells were visualized by transmitted light in Normarski configuration (panels in row a), by excitation at 480 nm and collection at 520 nm, showing direct EGFP excitation (panels in row b), and by excitation at 350 nm and collection at 520 nm, showing EGFP fluorescence after BFP excitation, indicating FRET (panels in row c). (B) FRET quantification. Fluorescent emission at 520 nm from individual cells transfected with the indicated constructs was recorded after excitation at 480 or 350 nm, and integrated intensities over the whole cells were evaluated. Plotted values (indicated by dots) represent the ratio between these two measurements: higher values indicate more efficient FRET between BFP and EGFP. Ten consecutively analysed cells were considered for each transfection; both their individual fluorescence ratios and their percentile box plot distribution are shown. Horizontal lines from top to bottom mark the 10-, 25-, 50-, 75-, 90-th percentiles, respectively. (C) Schematic representation of the putative structure of the N-terminus of Cyclin T1 with respect to the binding of Tat.

substantiating the fact that Tat-BFP was within 4 nm to the N and C termini of CycT1<sub>(1–300)</sub>.

The detailed, quantitative analysis of at least 10 cells expressing for the protein pairs is presented in figure 3B, showing the percentile distribution of FRET values. We consider the signal from cells transfected with EGFP and BFP (background) as baseline.

#### 4. Concluding remarks

The recruitment of P-TEFb by Tat through the specific binding of CycT1 is a crucial step in the replication cycle of HIV-1. However, although a considerable amount of biological data is available, a detailed structural characterization, which may help to rationalize these data and constitute the basis for future experiments, is still lacking.

We present here a theoretical structure of the N terminus of CycT1 based on the structure of its human transcriptional homologue CycH and of other cyclin proteins with known structure. Due to the low level of sequence identity between

target and template (figure 1(B)), the results of the homology modelling are confronted with site directed mutagenesis data present in literature. The residues involved in the CycT1: Tat interaction (from aa 250–261) turned out to be located at the H5'/term-coil-Hc term segment and solvent-exposed (figure 2), in good agreement with proteolysis data [31]. Although we were confident with the gross structural determinants of the cyclin box region of the model, also considering that this fold is highly conserved, we wish to point out that the orientation of the N and C termini remained rather uncertain, depending on the choice of the template (table 1). Thus, FRET experiments were carried out in this work to probe the spatial arrangement of the N and C terminal ends of the protein. By measuring FRET between EGFP-CycT1/Tat-BFP and CycT1-EGFP/Tat-BFP complexes we are able to confirm spatial proximity of these two regions in living cells (figure 3).

In conclusion, the model presented here is consistent with all experimental data up to the best of our knowledge. However, it must be kept in mind that it suffers the obvious limitations and uncertainties of comparative modelling,

particularly at low sequence identity levels. Still, we believe that it may be useful to test working ideas and inspire new biochemical experiments aimed to clarify structural features of protein–protein interactions in which CycT1 is involved.

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