

Article

# AML-related NPM mutations drive p53 delocalization into the cytoplasm with possible impact on p53-dependent stress response

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**Simple Summary:** Nucleophosmin (NPM) is one of the most abundant nucleolar proteins and its mutations frequently occur in acute myeloid leukemia (AML). The mutations cause aberrant cytoplasmic localization of mutated protein (NPMmut) and frequently mediate dislocation of NPM interaction partners. Tumor suppressor p53 is known to interact with NPM in response to genotoxic stress and its cytoplasmic localization is an unfavorable prognostic factor in cancers. This study aims to characterize the NPM-p53 interaction and to elucidate effect of the NPM mutations on p53 localization and expression in live cells. In addition, cellular dynamics of NPMmut and p53 after treatment with nuclear export inhibitor Selinexor is described and the mechanism of the Selinexor action proposed. Our results contribute to better knowledge of leukemogenic potential of NPM mutations and assist in the search for targeted AML therapy.

**Abstract:** NPM interaction with tumor suppressor p53 is a part of a complex interaction network and considerably affects cellular stress response. An impact of characteristic AML-associated NPM mutations on interaction with p53 has not been investigated yet, although consequences of NPMmut-induced p53 export to the cytoplasm are important for understanding of leukemogenic potential of these mutations. We investigated p53-NPM interaction in live HEK-293T cells by FLIM-FRET and in cell lysates by immunoprecipitation. Results were confirmed in leukemia cell lines. eGFP lifetime-photoconversion was used to follow redistribution dynamics of NPMmut and p53 in Selinexor-treated cells. We confirmed the p53-NPMwt interaction in intact cells and newly documented that this interaction is not compromised by the NPM mutation causing displacement of p53 to the cytoplasm. Importantly, the interaction was not abolished for non-oligomerizing NPM variants with truncated oligomerization domain, suggesting that oligomerization is not essential for interaction of NPM forms with p53. Inhibition of the nuclear exporter XPO1 by Selinexor caused expected nuclear relocalization of both NPMmut and p53. However, significantly different return rates of these proteins indicate nontrivial mechanism of p53 and NPMmut cellular trafficking. We suggest that the altered p53 regulation in cells expressing NPMmut offers a new target for AML therapy.

**Keywords:** p53; nucleophosmin; mutation; acute myeloid leukemia; FLIM-FRET; Selinexor; photoconversion

## 1. Introduction

Nucleolus is a subnuclear compartment with multiple cellular roles. Sensing cellular stress is one of them [1]. During the stress response nucleoli become more condensed and nucleolar proteins partially translocate to the nucleoplasm. Nucleolar phosphoproteins, such as nucleophosmin (NPM) or nucleolin (NCL), participate in the stress response via complex interaction network [2]. The translocated proteins can stabilize tumor suppressor

p53 either by direct interaction [3-6] or by interaction with proteins regulating intracellular p53 levels [7]. NPM is an abundant nucleolar phosphoprotein with numerous functions in many important cellular processes, including the p53-dependent response to genotoxic stress [1, 8]. Thanks to its chaperone function NPM interacts with p53 as well as with other tumor suppressors, such as pRb or p14Arf. While the interaction with p53 occurs via region close to the C-terminus of NPM [3], interactions with pRb and p14ARF take place via the N-terminal NPM region that is responsible also for NPM-oligomerization [9, 10].

Ubiquitin ligase HDM2 targets the p53 for degradation by proteasome [11]. In response to stress, NPM competes with the HDM2 for binding to p53, which protects p53 from ubiquitination. Alternatively, NPM binds directly to HDM2 causing release of p53 from the ligase [12, 13]. The regulatory interaction network including HDM2, p53, p14Arf and nucleolar phosphoproteins allows for rapid stabilization and activation of p53 in response to genotoxic and pathogenic stimuli affecting the DNA stability [14, 15]. The localization of the proteins engaged in the p53 degradation/stabilization balance is affected by their mutual interaction. The p14Arf is sequestered to nucleolus by binding to NPM, where it is kept away from p53 [16, 17]. Protein p53 was also observed to be pulled to the nucleolus by overexpressed NPM [3]. Another nucleolar phosphoprotein NCL does not possess appropriate nucleolar localization sequence (NoLS) [18, 19] and its nucleolar localization is supposed to be mediated by its association to NPM [19]. Independently of its NoLS, attachment of NPM to the nucleolus has been recently ascribed to the affinity of its C-terminal region to G-rich nucleic acids (Mitrea et al., 2016), which applies also for NCL.

To our knowledge, any systematic investigation of the p53-NPM interaction in cells under conditions other than the genotoxic stress has not been performed yet. In this work we therefore focus on the p53-NPM interaction in intact cells under standard physiological conditions. Protein tagging by fluorescent proteins makes possible monitoring of the cellular dynamics and protein proximity directly in live cells [20-22]. FLIM utilizing resonance energy transfer (FRET) between the tagged proteins is a highly sensitive tool for this purpose [23, 24]. Here we characterized p53-NPM interaction both in cell lysates and live cells using methods previously established for the detection of NPM oligomers, i.e. fluorescence confocal microscopy [25], fluorescence lifetime imaging (FLIM) [23] and immunochemical methods [24].

A mutation of the *NPM1* gene resulting in the altered C-terminus of NPM and aberrant localization of mutated NPM (NPMmut) to the cytoplasm appears in approximately 50% AML with normal karyotype [26-28]. The original NoLS of wild type NPM (NPMwt) is highly compromised in NPMmut and strong nuclear export signal (NES) for the XPO1 exporter appears at the altered C-terminus [29, 30] in addition to the two NESes already present in its N-terminal domain [10]. The most frequent AML-related mutation type A gives rise to mutated protein (NPMmutA, NPMmut) lacking tryptophans W288 and W290 [28]. For comparison, the mutation of type E retains NPM's W288, which causes that nucleolar localization of the mutated protein (NPMmutE) is partially preserved [31]. The interaction of NPM monomers within the oligomer and its interaction with p14Arf were shown to persist when NPM is mutated in its C-terminus [29]. In cells expressing NPMmut, interacting proteins NPMwt and p14Arf become partially dislocated to the cytoplasm due to their binding to NPMmut [32]. In analogy, other NPM-interacting proteins, such as p53, are also candidates for dislocation via the interaction with NPMmut. The dislocation should interfere with their proapoptotic activity, which could lead to uncontrolled cell division [9]. On the other hand, the interaction of NPM with NCL, taking place through AA187-241 region of the NPM molecule [19], is inhibited by the NPM mutation and NCL is therefore not translocated to the cytoplasm together with NPMmut [25]. Since p53 was found to interact with a domain near the C-terminus of NPM (AA186-259 or AA242-269, respectively [3, 4]), one could expect that the p53-NPM interaction was affected by this mutation as well. The detailed mechanism and role of the p53-NPMmut interaction in the leukemogenesis is unknown so far. The main part of this article therefore

investigates impact of the NPM mutation on the p53-NPM interaction and presence of this interaction is documented both in live cells and cell lysates.

Cytoplasmic localization of p53 plays an adverse role in the cell cycle regulation [33] and it was reported to launch apoptosis via interaction with mitochondrial proapoptotic factors [34]. On the other hand, displacement of the p53 from its mainly nuclear localization could lead to aberrant growth signals resulting in cancer development [35]. The proteasome-mediated p53 degradation can also be affected by the p53 translocation [36]. Therapeutic restoration of the p53 functionality therefore seems to be a promising anti-cancer approach [37]. As NPM mutations are mutually exclusive with p53 mutations in AML, the return of wild-type p53 to the nucleus could re-establish its function. The blockade of exportin 1 (XPO1) could be a strategy, since gradual p53 relocation to the nucleoplasm owing to the XPO1 blockade should help to stabilize the p53 population in uncontrollably proliferating cells and lead to re-activation of p53-dependent apoptosis pathway.

The XPO1 mediates outward protein transport from the nucleus [38]. The transport depends on recognition of specific leucine-rich NESes [39], two of which were identified also in p53 [40, 41]. Leptomycin B is a typical drug blocking XPO1. As this drug is too toxic for the clinical use [42], other drugs belonging to the group of selected inhibitors of the nuclear export (SINE) attract attention nowadays [43]. These drugs, such as Selinexor, were reported to reactivate p53 by restoring its nuclear accumulation via the XPO1 blockade and, consequently, its proper regulatory function [35]. Since NPMmut is also an XPO1-client, it is not surprising that Leptomycin B or Selinexor have been reported to induce relocation of NPMmut to the nucleus [44, 45]. Selinexor has been already tested both for treatment of cancers associated with impaired p53 and for the AML with mutated NPM [46]. In the second part of this work, we address p53 interactions and trafficking in HEK-293T cells and leukemia cell lines treated with Selinexor. Our results elucidate mechanism of p53 and NPMmut co-translocation in live cells.

## 2. Materials and Methods

### *Cell cultivation*

Adherent cell line HEK-293T was kindly provided by dr. Š. Němečková (Department of Immunology, Institute of Hematology and Blood Transfusion). Leukemia cell lines MV4-11, OCI-AML2, OCI-AML3, and KASUMI-1 were purchased from DSMZ (Germany). The cells were cultivated in growth media with fetal bovine serum (FBS), glutamine and antibiotics (all from Sigma-Aldrich) according to manufacturers' recommendation: MV4-11 in RPMI-1640/10% FBS, OCI-AML2 and OCI-AML3 in alpha-MEM/20% FBS, KASUMI-1 in RPMI-1640/20% FBS and HEK-293T in DMEM/10% FBS. Selinexor (Selleckchem) was added into the cell culture from 10mM stock solution to final concentrations and for times specified in the text.

### *Plasmid construction and cell transfection*

Plasmids for expression of fluorescently tagged proteins were constructed by standard molecular cloning techniques. Preparation of plasmids for NPMwt and NPMmut expression was described in [47], preparation of plasmids for  $\Delta$ 117wt in [24]. For cloning, DNA fragments corresponding to specific *NPM1* and *TP53* transcript variants (RefSeq. NM 002520.7 and NM 000546.6, respectively, from NCBI database) were PCR-amplified from cDNA library (Jurkat cells, Origene) or from plasmid pBI-p53 wt/EGFP (Plasmid #16543, Addgene), respectively, using extended primers and subsequently they were subcloned to vectors peGFP-C2 or pmRFP1-C2 (originally Clontech) using XhoI and BamHI unique restriction sites (Thermo Scientific) and T4 DNA ligase (NEB). Plasmids for  $\Delta$ 117mut were prepared according to [24, 47] with primers used for construction of the single mutants to obtain the desired double mutant.

The constructed plasmids were amplified in *E. coli* competent and purified with the PureYield Plasmid Miniprep System (Promega). HEK-293T cells were seeded to the cell density of  $1 \times 10^5$ /ml 24h prior transfection and then transfected with jetPrime transfection reagent (Polyplus transfection) according to the manufacturer's protocol. Growth

medium was replaced 4h after the transfection and cells were further grown for 20-40h prior analysis. NPM and p53 constructs appearing in this study are listed in Table 1.

**Table 1.** List of constructs. All protein constructs are tagged on their N-terminus, symbol "/" denotes mixture of constructs in text of the article.

Construct	Abbreviation	Construct	Abbreviation	Construct	Abbreviation
eGFP_NPMwt	G_NPMwt	eGFP_Δ117NPMwt	G_Δ117wt	eGFP_p53wt	G_p53
mRFP1_NPMwt	R_NPMwt	mRFP1_Δ117NPMwt	R_Δ117wt	mRFP1_p53wt	R_p53
eGFP_NPMmut	G_NPMmut	eGFP_Δ117NPMmut	G_Δ117mut		
mRFP1_NPMmut	R_NPMmut	mRFP1_Δ117NPMmut	R_Δ117mut		

#### *Cell lysis*

As described previously [25], cells were washed with PBS and lysed in Laemmli sample buffer (SB, 50mM Tris pH 6.8, 2% SDS, 100mM DTT, 10% glycerol), boiled at 95°C for 5 min, centrifuged at 200.000g/4°C for 4h and the supernatant was stored at -20°C.

#### *Western blotting*

Five to ten microliters of each sample were subjected to SDS-PAGE and transferred into PVDF membrane (BioRad). Mouse monoclonal antibodies against β-actin, GFP, dsRed, NCL, FBL and NPM (clone 3F291, for endogenous NPM), were from Santa Cruz Biotechnology. All mouse primary antibodies were used at a dilution 1:100-1:500. Rabbit monoclonal antibody against p53 (Abcam) and rabbit polyclonal antibodies against NPM (Abcam) and NPMmut (pab50321, Covalab) were used at 1:1.000-1:2.000 dilution. Anti-mouse and anti-rabbit HRP-conjugated secondary antibodies were purchased from Thermo Scientific and used at concentrations 1:10.000-1:50.000. ECL Plus Western Blotting Detection System (GE Healthcare) was used for chemiluminescence visualization and evaluation by G-box iChemi XT4 digital imaging device (Syngene Europe). Alternatively, SuperSignal West Atto Ultimate Sensitivity Substrate (ThermoFisher Sci) was used for highly sensitive detection of co-immunoprecipitated proteins. Images included in Figures are always representative of at least three independent experiments.

#### *Co-immunoprecipitation Trap assay*

Transfected cells expressing fluorescent proteins were processed after 40h-incubation. GFP-, RFP- and p53-Trap\_A system (Chromotek) were used following the manufacturer's instructions as described in [30]. Briefly, eGFP/mRFP1-expressing adherent cells were washed with in ice-cold PBS and scrapped from dish. Alternatively, suspension cells of leukemia cell lines were collected by centrifugation and washed with PBS. The cell pellet was lysed in the lysis buffer (10mM Tris/Cl pH7.5, 150mM NaCl, 0.5 mM EDTA, 0.5% NP-40, protease and phosphatase inhibitors) on ice for 30 min and centrifuged at 20.000g/10min/4°C. The lysate was applied on the Trap\_A beads and rotated for 1h at 4°C. Then the beads were pelleted and extensively washed in the diluting buffer (10mM Tris/Cl pH7.5, 150mM NaCl, 0.5mM EDTA), resuspended in 2xSDS-sample buffer (100 mM Tris pH 6.8, 4% SDS, 200 mM DTT, 20% glycerol), boiled for 10 min and centrifuged at 2.500g/2min/4°C. Supernatant was stored at -20°C until used for SDS-PAGE.

#### *Live cell imaging*

Cells were grown on glass bottom Petri dish (Cellvis). Fluorescence experiments were carried out at 37°C after sealing the Petri dish with parafilm to prevent CO<sub>2</sub> leakage. The subcellular distribution and co-localization of eGFP- and mRFP1-fused NPM variants were observed under the confocal laser scanning microscope FV1000 (Olympus Corporation). One Petri dish was typically imaged for 1 hour and data were processed by FluoView FV10-ASW 3.1 and ImageJ-Fiji software.

*FLIM - data acquisition and analysis*

FLIM experiments were carried out on the inverted IX83 microscope with FV1200 confocal scanner (Olympus). The microscope was equipped with cell-cultivation chamber (Okolab) and FLIM add-on comprising picosecond semiconductor lasers, fiber-coupled GaAsP hybrid detectors, and TimeHarp 260PICO TCSPC detection electronics (all PicoQuant, Berlin, Germany). Specifically, cellular FLIM experiments were performed with a UPLSAPO 60x NA 1.2 water immersion objective (Olympus), eGFP fluorescence was excited at 485 nm by the LDH-DC-485 laser head (PicoQuant) and emission decays were collected on the pixel-by-pixel basis using combination of 560 nm short-pass dichroic and Semrock 520/35 bandpass filter in the detection path. To avoid pile-up, the data collection rate at brightest pixels was kept below 5% of the laser repetition rate. Acceptor photobleaching was done by the 562 nm cw laser. eGFP lifetime-photoconversion (PC) was performed as described previously [48]. Selected ROI was scanned by intense 488 nm light till eGFP emission decreased to about 20-30% of its initial value. Depending on the sample, the power of the argon 488 nm cw-laser line was set between 0.5 and 1.0 mW at the back-aperture of the objective in order to complete the lifetime-conversion within 1–2 min. All experiments were performed at 37°C.

FLIM data were analyzed using the SymPhoTime64 software (PicoQuant). The lifetime images were generated in the SymPhoTime64 by the “fast-FLIM” approach when mean pixel lifetimes were calculated by a method of moments [49]. Specifically, the lifetime  $\tau_{fast}$  was determined as the difference between the barycentre of the fluorescence decay and the time-offset  $t_{offset}$  at the steepest growth of the decay curve at each pixel:

$$\tau_{fast} = \frac{\sum I_i t_i}{\sum I_i} - t_{offset} \quad (1)$$

where  $I_i$  stands for the emission intensity at time  $t_i$ . Least-squares reconvolution was applied for accurate analysis of cumulative decays from larger ROIs (cell, nucleus, nucleolus). Fluorescence of eGFP was typically assumed to decay bi-exponentially [50, 51] according to the formula:

$$I(t) = a_1 \cdot e^{-t/\tau_1} + a_2 \cdot e^{-t/\tau_2} \quad (2),$$

where  $\tau_i$  and  $a_i$  are lifetime components and corresponding amplitudes, respectively. The intensity-weighted mean fluorescence lifetime was calculated as:

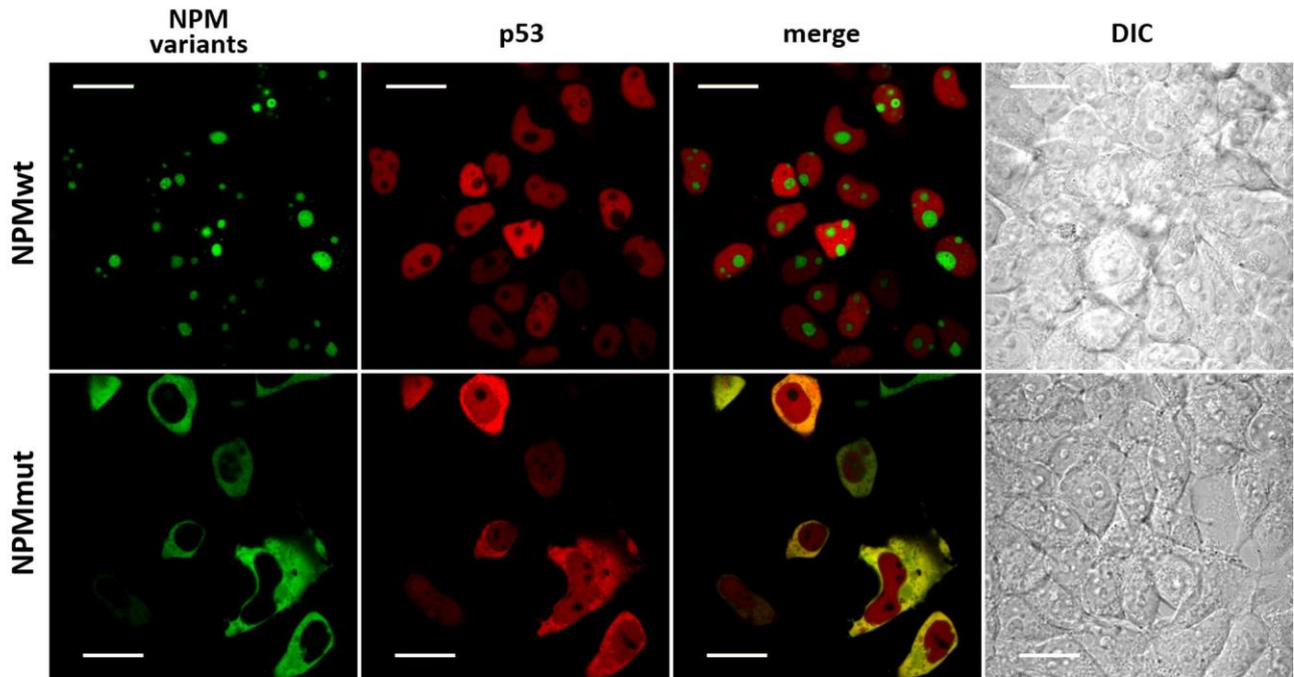
$$\tau_{mean} = \sum f_i \cdot \tau_i, \quad f_i = a_i \tau_i / \sum a_i \tau_i \quad (3),$$

where  $f_i$  are intensity fractions of the  $i$ -th lifetime component.

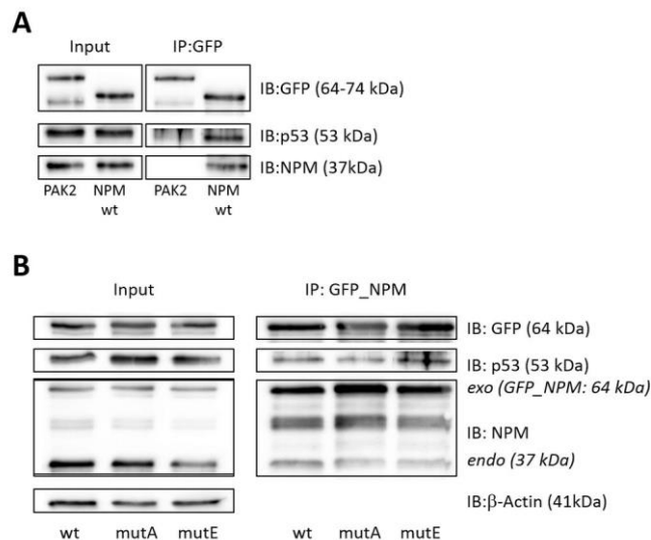
**3. Results**

The C-terminal mutation drives NPM translocation to the cytoplasm. We investigated interaction of fluorescently tagged NPMwt/mut and p53 in transfected HEK-293T cells, similarly as previously done for interaction of NPM variants in mixed oligomers [30]. As seen in Figure 1, exogenous p53 is markedly delocalized to the cytoplasm of HEK-293T cells co-expressing NPMmut. We observed extensive translocation of mRFP1\_p53 (R\_p53) to the cytoplasm when co-expressed with eGFP-tagged NPMmut (G\_NPMmut). This is in the clear contrast with p53 localized mainly in the nuclei in cells co-expressing eGFP-tagged NPMwt (G\_NPMwt). Although the effect of co-expression on the p53 translocation is clearly observable in both tagging combinations, Figure S1, it is more obvious for combination G\_NPMmut/R\_p53. The observed translocation suggests that p53 interacts with NPM regardless the C-terminus mutation. We confirmed this finding by co-immunoprecipitation as done for endogenous NPM in mixed NPM oligomers [30], i.e. NPMwt and NPMmut were labeled with eGFP, transfected into HEK-293T cells and immunoprecipitated with the GFP-Trap. Then co-immunoprecipitated endogenous p53 was detected by immunoblotting. Indeed, the interaction between p53 and NPMwt, as well as between p53 and NPMmut was proven, Figure 2. The endogenous p53 was clearly

detected in all precipitates, regardless NPM mutation type. Alternative experiments documenting the interactions used p53-Trap for endogenous p53 immunoprecipitation and combination of GFP/RFP Traps for immunoprecipitation from cells co-expressing R\_NPMwt and G\_p53. Results presented in supplemental Figure S2 bring evidence that the p53-NPM interaction is not inhibited by the mutation of the NPM C-terminus.



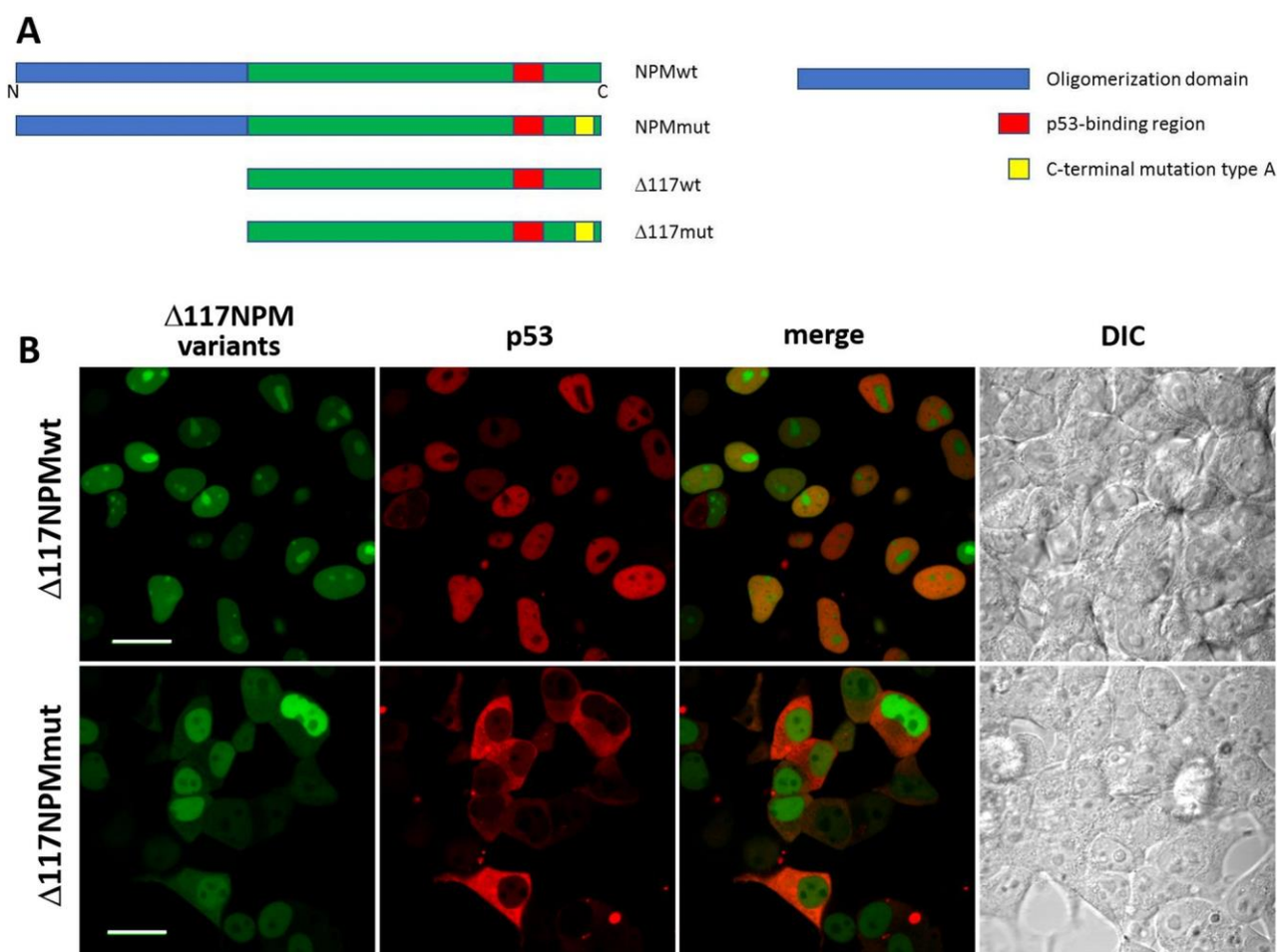
**Figure 1.** Localization of R\_p53 co-expressed with eGFP-labeled NPM variants in HEK-293T cells. Upper row - R\_p53/G\_NPMwt co-expression, lower row - R\_p53/G\_NPMmut co-expression, first column - eGFP fluorescence, second column - mRFP1 fluorescence. Bar indicates 20  $\mu$ m.



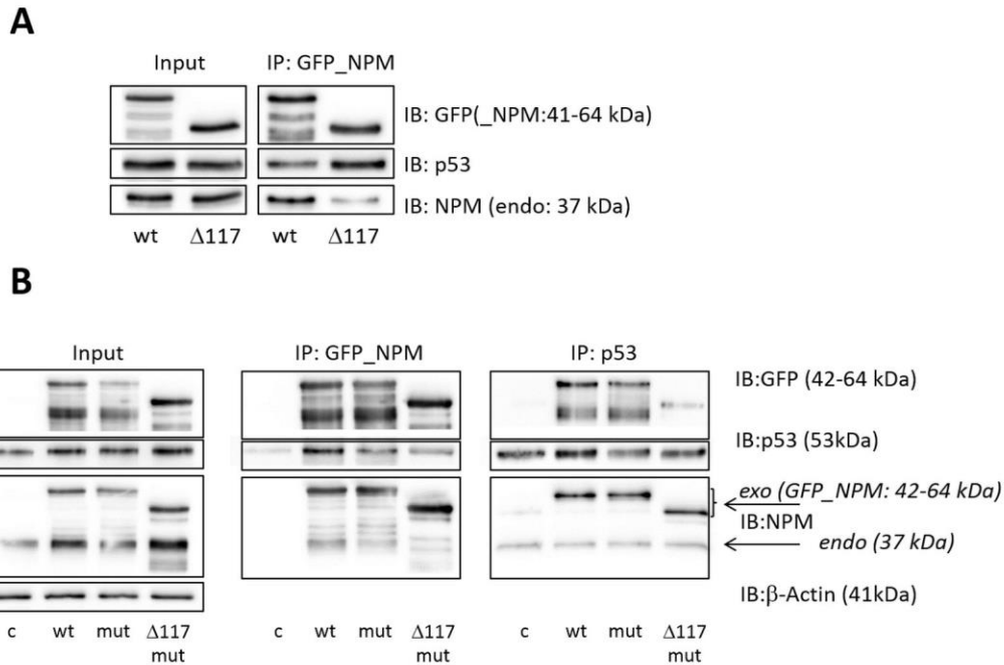
**Figure 2.** Interaction of p53 with NPM variants in lysates of HEK-293T cells. **(A)** Interaction of p53 with NPMwt. Cells were transfected with G\_NPMwt or G\_PAK2 (non-interacting control of a comparable size). Endogenous NPM as well as p53 co-precipitated only with NPMwt; **(B)** Interaction of p53 with NPMmut. Cells were transfected with eGFP-labeled NPMmutA, NPMmutE or NPMwt for comparison. Endogenous NPM as well as p53 co-precipitated in all samples. The samples are identical with those used in [25] for assessment of the interaction between NPM and nucleolin.

Further we investigated the role of NPM oligomerization in the p53 delocalization and NPM-p53 interaction. As the oligomerization of NPM is mediated by its N-terminal domain (AA1-117) [52, 53], we truncated the N-terminus to obtain NPMwt and NPMmut

variants with compromised oligomerization,  $\Delta 117$ wt [24] and  $\Delta 117$ mut, respectively, Figure 3. Similarly to  $\Delta 117$ wt, we found  $\Delta 117$ mut being localized mainly in the nucleus, nevertheless, with substantial cytoplasmic fraction caused by the C-terminal mutation. At the same time, extensive translocation of R\_p53 to the cytoplasm was observed when co-expressed with eGFP-tagged  $\Delta 117$ mut (G\_ $\Delta 117$ mut). The observation strongly suggests an existence of interaction between p53 and  $\Delta 117$ mut, similar as for NPMmut. The effect of p53 co-expression with  $\Delta 117$ mut on the p53 translocation to the cytoplasm is comparable in both tagging combinations, Figure S3. To independently confirm the p53- $\Delta 117$ mut interaction, we performed immunoprecipitation experiments presented in Figure 4. Consistently with our previous results [24], we detected slightly weaker interaction between NPMwt and  $\Delta 117$ wt than between NPMwt monomers. Concurrently, endogenous p53 was pulled down with  $\Delta 117$ wt more efficiently than with NPMwt, suggesting that the interaction between p53 and  $\Delta 117$ wt might be even stronger than between p53 and NPMwt. In  $\Delta 117$ mut precipitates the endogenous p53 was detected as well, but in a lower extent, Figure 4B. Endogenous NPMwt did not precipitate with  $\Delta 117$ mut at all, which indicates inhibited oligomerization of this construct. Presented results document that the interaction capability of NPMwt with p53 is independent of the NPM oligomerization status, which implicates that monomeric NPM is able to interact with p53. However, the lower extent of exogenous p53 detected in  $\Delta 117$ mut precipitates indicates lower binding affinity for the double mutant.

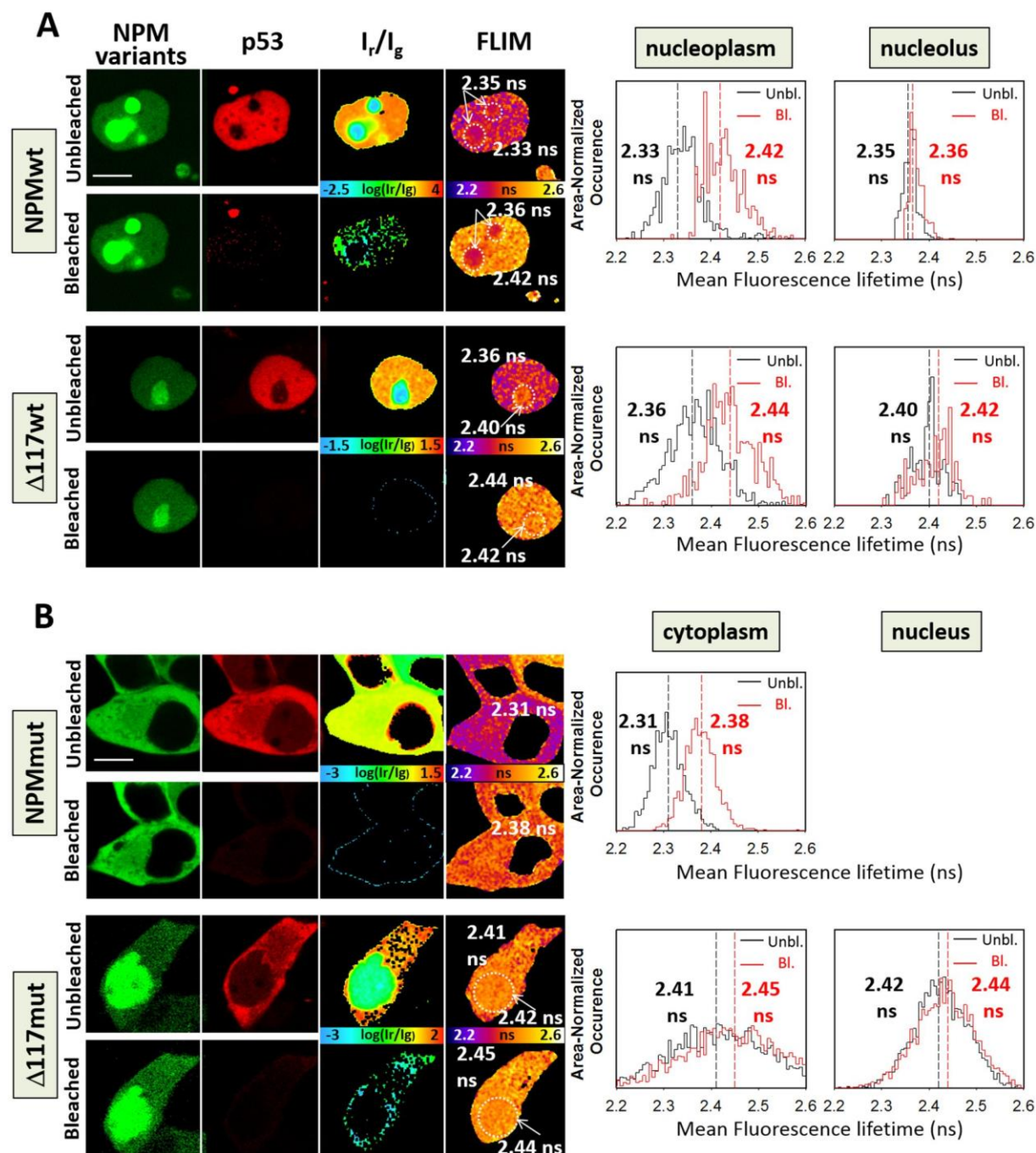


**Figure 3.** Localization of R\_p53 co-expressed with eGFP-labeled NPM variants with truncated oligomerization N-terminus (AA1-117) in HEK-293T cells. (A) Comparison of used NPM variants; (B) Upper row - R\_p53/G\_ $\Delta 117$ wt co-expression, lower row - R\_p53/G\_ $\Delta 117$ mut co-expression, first column - eGFP fluorescence, second column - mRFP1 fluorescence. Bar indicates 20  $\mu$ m.



**Figure 4.** Interaction of p53 with  $\Delta 117$  variants of NPM in lysates of HEK-293T cells. **(A)** Interaction of p53 with  $\Delta 117$ wt. Cells were transfected with G\_ $\Delta 117$ wt or G\_NPMwt for control. Levels of co-precipitated endogenous NPM and p53 differ between the samples; **(B)** Interaction of p53 with  $\Delta 117$ mut. Cells were transfected with G\_NPMmut, G\_ $\Delta 117$ mut or G\_NPMwt for comparison. Untransfected cells (c) served as a precipitation control. Levels of co-precipitated proteins were investigated by immunoblotting.  $\beta$ -Actin expression served as a loading control.

For investigation of the p53-NPMmut interaction in live cells we utilized FLIM-FRET, which visualizes fluorescence lifetime across the cellular structures. FRET reliably detects proximity of the donor and the acceptor on the molecular scale and presents itself by shortening of the donor fluorescence lifetime [54, 55]. eGFP and mRFP1 fused to p53 and NPM were used as the donor and the acceptor, respectively. The shortening of the eGFP lifetime ( $\tau_{GFP}$ ) in the presence of the acceptor-labeled protein then indicates molecular complex formation. To unambiguously prove the presence of the energy transfer, the acceptor was photobleached, which cancels FRET. Consequently,  $\tau_{GFP}$  should increase [56]. Interaction of R\_p53 with G\_NPMwt and G\_NPMmut is presented in Figure 5. Images from experiment with the G\_NPMwt/R\_p53 combination are presented in the first two rows of Figure 5A. Presence of FRET between G\_NPMwt and R\_p53 is clearly documented in the nucleoplasm by the increase of  $\tau_{GFP}$  from 2.33 to 2.42 ns upon the acceptor photodestruction. Nucleoli without sufficient concentration of the acceptor serve as an alternative internal control and FRET does not occur there, as expected. R\_p53 co-expressed with G\_ $\Delta 117$ wt displays very similar behavior (Figure 5A, lower two rows). Compared to the mainly nucleolar G\_NPMwt [24], the amount of G\_ $\Delta 117$ wt in the nucleoplasm is enhanced and FRET between G\_ $\Delta 117$ wt and R\_p53 is clearly present there, as  $\tau_{GFP}$  increases from 2.36 to 2.42 ns upon the acceptor photobleaching. Nucleolus with a low acceptor concentration serves again as the internal control and FRET is not detected there. Lifetime histograms of the G\_NPMwt-R\_p53 and G\_ $\Delta 117$ wt-R\_p53 pairs highly resemble each other indicating similar interaction. In agreement with the immunostaining, the lack of the NPM oligomerization domain does not evidently prevent the interaction between p53 and NPM monomers.

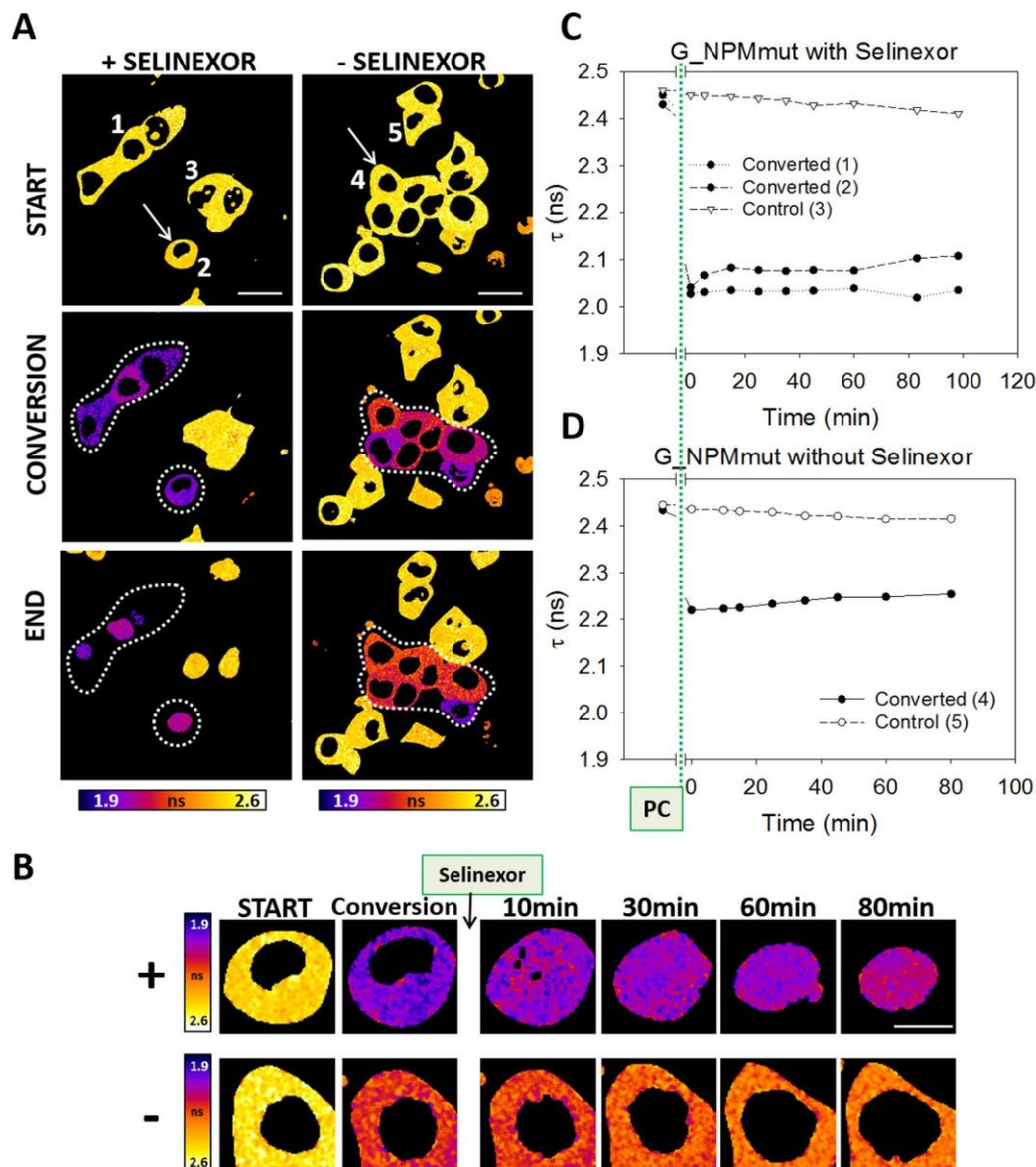


**Figure 5.** Interaction of R\_p53 with eGFP-labeled variants of NPM in live HEK-293T cells examined by FLIM-FRET. (A) Interaction of p53 with NPM variants with unmodified C-terminus. Upper two rows - co-expression of R\_p53 with G\_NPMwt, lower two rows - co-expression of R\_p53 with G\_Δ117wt; (B) Interaction of p53 with NPM variants bearing mutation at the C-terminus. Upper two rows - co-expression of R\_p53 with NPMmut, lower two rows - co-expression of R\_p53 with G\_Δ117mut. Columns from left to right represent localization of the green fluorescence  $I_{green}$  (NPM variants), red fluorescence  $I_{red}$  (R\_p53), ratio  $I_{red}/I_{green}$ , and fluorescence lifetime maps of eGFP before (row Unbleached) and after (row Bleached) mRFP1 photobleaching. Histograms contain pixel-based lifetime distributions from the nucleolus and nucleoplasm before (black line) and after (red line) the mRFP1 photobleaching. Bar represents 10  $\mu$ m.

Interaction between G\_NPMmut and R\_p53 is assessed in the first two rows of Figure 5B. FRET was detected in the cytoplasm where the donor lifetime increases from 2.31ns to 2.38 ns after the acceptor destruction. In agreement with Figure 1, G\_NPMmut is almost absent in the nuclei, which prevents FRET measurements there. The last two rows in Figure 5B document the cellular interaction between R\_p53 and G\_Δ117mut. We can see that

R\_p53 is strongly delocalized to the cytoplasm while the nucleus is depleted. The opposite is true for G\_Δ117mut, since the lack of NESes from oligomerization domain suppresses cytoplasmic transport of Δ117mut relatively to NPMmut and the protein stays mainly in the nucleus. In contrast to the immunostaining, we were not able to prove FRET between G\_Δ117mut and R\_p53 neither in the cytoplasm nor in the nucleus of live cells. As R\_p53 is translocated to the cytoplasm, there is not enough acceptor in the nucleus to observe FRET, which is clearly visible in a lifetime histogram in the last column of Figure 5B. Although the acceptor is in a significant excess in the cytoplasm and FRET should be detected, lifetime histograms show only minor change there before and after the photo-bleaching. The observed discrepancy might be explained by rather low binding affinity of p53 to G\_Δ117mut, as suggested by the low amount of endogenous p53 co-immunoprecipitating with G\_Δ117mut, compared to G\_NPMmut and G\_Δ117wt. Binding conditions and concentrations in live cells might be therefore unfavorable for the detection of trace amounts of the complex by FRET. Alternatively, the critical Forster distance could be somehow larger for this particular protein combination, e.g. due to an unfavorable orientation factor of the labels [57]. The weak interaction then might be missed by FRET. The intriguing observation shown in the last two columns of Figure 5 indicates a nontrivial mechanism of NPM/p53 cellular co-trafficking when both NPM termini are altered.

Further we investigated effect of Selinexor on the p53 and NPMmut relocalization back to the nucleus. Since both NPM and p53 are clients of XPO1, we tried to verify whether their Selinexor-induced nuclear import is synchronous in a complex, as could be assumed from the p53-NPMmut interaction. To monitor the relocalization, we performed time-lapse FLIM measurement utilizing the lifetime-based photoconversion (PC) of eGFP [48]. The method allows marking a selected population of eGFP-labeled protein by a brief scan by intense 488nm light, which irreversibly converts eGFP to a form with a shorter emission lifetime. Since the photoconverted and unconverted tags differ in the lifetimes, diffusion and cellular redistribution of the highlighted protein can be easily followed by FLIM. This allows quantification of the converted and unconverted eGFP fractions across the cell by the localized lifetime analysis. First we monitored the effect of Selinexor on the relocalization of G\_NPMmut (Figure 6). After the PC, the FLIM image was measured and then 4 μM Selinexor was added. Rows of Figure 6A from the top to bottom show FLIM images before the PC, just after the PC and 80 min later. The photoconverted areas are marked by the dashed line. We can notice significant difference in the G\_NPMmut localization after the Selinexor treatment (first column). While G\_NPMmut stays in the cytoplasm in the control sample (second column), the addition of Selinexor induced marked relocalization of G\_NPMmut to the nucleus that was finished in about 60 min. Figure 6B demonstrates this process in more detail for selected cells in the presence and the absence of Selinexor, cell no. 2 and 4, respectively (panel A). Quantitative evolution of mean eGFP lifetimes ( $\tau_{\text{GFP}}$ ) in individual cells is depicted in Figures 6C and 6D. It can be immediately seen that the addition of Selinexor has negligible effect on the time evolution of  $\tau_{\text{GFP}}$  of eGFP tagging NPMmut either in the control or in the photoconverted cells (compare differences between panels C and D). The small rise in  $\tau_{\text{GFP}}$  in the photoconverted cells (closed symbols) is a consequence of synthesis and maturation of fresh eGFP with longer lifetime which gradually elevates the measured lifetime. As apparent from Figure 6C, D, this rise of about 50ps in 80 min is almost insignificant. All G\_NPMmut is relocalized to the nucleus at that time.



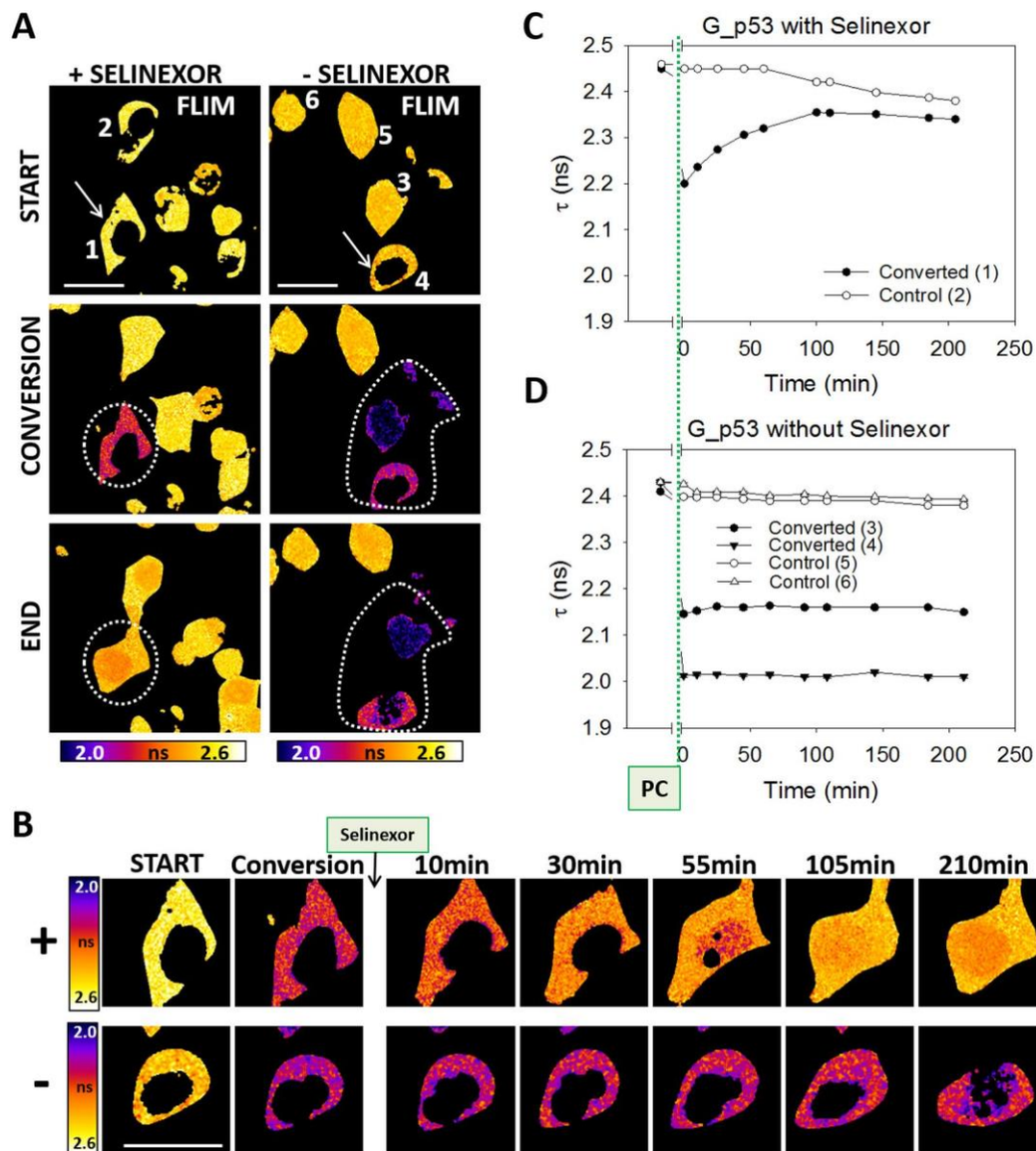
**Figure 6.** Effect of Selinexor on localization and lifetime of G\_NPMmut in live HEK-293T cells. (A) Lifetime images of G\_NPMmut in the presence and absence of 4  $\mu$ M Selinexor before (the 1<sup>st</sup> row), immediately after (the 2<sup>nd</sup> row), and 80 min after the photoconversion (PC) (the 3<sup>rd</sup> row). The photoconverted area is bordered by the dashed line. Bar represents 20  $\mu$ m; (B) Detailed time evolution of G\_NPMmut lifetime maps in single cell with photoconverted eGFP in the presence (upper row, cell no. 2) and in the absence (lower row, cell no. 4) of Selinexor. The visualized cells are marked by arrows in panel A. The nuclear signal was negligible in the absence of Selinexor. A minute increase in  $\tau_{\text{GFP}}$  is noticeable both in the presence and absence of Selinexor. Bar represents 10  $\mu$ m; (C, D) Time evolution of  $\tau_{\text{GFP}}$  in the photoconverted cells (1, 2, 4) and in control cells (3, 5) in the presence (C) and absence (D) of Selinexor. Cells used for integration of the plotted signal are marked in the panel A.

Then we performed similar relocalization experiments with cells expressing G\_p53 accompanied by R\_NPMmut to ensure cytoplasmic localization of G\_p53, Figure 7. We selected cells with marked amount of G\_p53 in the cytoplasm, photoconverted those cells and mapped evolution of  $\tau_{\text{GFP}}$ . Figure 7A depicts FLIM images of the photoconverted and control cells in the presence and absence of Selinexor. The process is shown again in more detail in Figure 7B (cells no. 1 and 4 from panel A, marked by arrows). Compared to G\_NPMmut examined in Figure 6, the G\_p53 nuclear relocation is slower and incomplete within 3 hours. The relatively slow return of G\_p53 to the nucleus was accompanied by a faster, significant increase of  $\tau_{\text{GFP}}$  upon the application of Selinexor. This is best seen in panels C and D where evolution of  $\tau_{\text{GFP}}$  averaged over individual cells is shown. We can see that while control cells without Selinexor exhibit almost no lifetime change, Figure 7D,

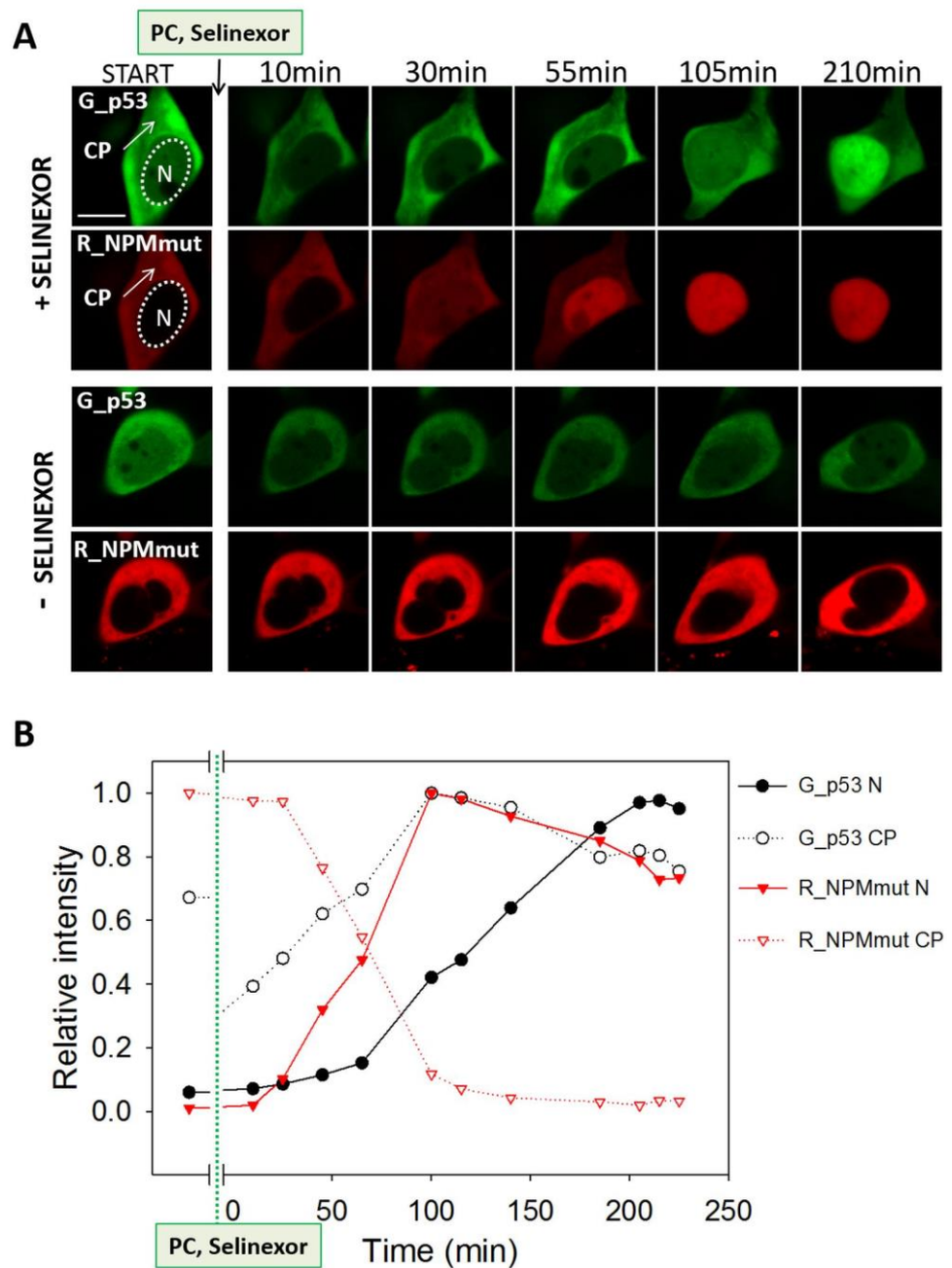
the addition of the drug induces significant increase of  $\tau_{\text{GFP}}$  in the photoconverted cell (cell no. 1, panel A), Figure 7C. Since the eGFP photoconversion was shown to be irreversible and fluorescence lifetimes stable in time [48], the observed increase in  $\tau_{\text{GFP}}$  should be caused either by *de novo* synthesis of fresh long-lifetime G\_p53 or by degradation of the short-lifetime photoconverted protein. The process reaches its plateau at about 2h with the characteristic rise half-time ( $T_{1/2}$ ) of about 60 min. Some decrease in  $\tau_{\text{GFP}}$  of the control unconverted cell in the presence of Selinexor (cell no. 2, panel A) is likely caused by a gradual reimport of eGFP-tagged p53 to a dense nucleus, where eGFP exhibits slightly lower lifetime clearly visible at the end of the experiment, Figure 7A. Such eGFP lifetime shortening in a dense environment modulating the refraction index has been commonly observed [48, 58].

Selinexor-induced synthesis/maturation of fresh G\_p53 supposedly causing the observed increase in  $\tau_{\text{GFP}}$  should be accompanied by an increase in the emission intensity. Indeed, increasing fluorescence of exogenous G\_p53 was observed for both converted and unconverted cells upon the Selinexor addition, as documented in Figures 8 and S4, respectively. In Figure 8, relocalization dynamics of G\_p53 and R\_NPMmut in photoconverted cells from Figure 7 (cells no. 1 and 4) is shown in detail. Upper rows of Figure 8A display a gradual relocalization of G\_p53 and R\_NPMmut after the addition of Selinexor. It can be seen that while both proteins gradually relocalize to the nucleus, relocalization of R\_NPMmut appears faster. Lower two rows containing control data without Selinexor do not indicate any evolution of the protein amount and distribution. Time evolution of fluorescence signal in the nucleus and cytoplasm of the Selinexor-treated cell from Figure 8A is shown in Figure 8B. From the intensity increase in the nucleus we can conclude that NPMmut re-localizes to the nucleus about twice faster than p53, half-rise times  $T_{1/2}$  being of about 60 and 120 min for NPMmut and p53, respectively. Such asynchronous behavior indicates, that NPMmut and p53 may respond independently to the nuclear export inhibition. Interestingly, despite outflux of p53 from the cytoplasm, photoconversion-reduced cytoplasmic intensity of G\_p53 rises in a response to Selinexor. The observation is fully consistent with the increased  $\tau_{\text{GFP}}$  documented in Figure 7C. Kinetic behavior is also rather similar being completed in about 2h. Similar rise in the fluorescence intensity was observed also for unconverted cells. Figure S4 demonstrates that the cytoplasmic emission quickly rises from the non-reduced pre-treatment level and later is followed by a decrease caused by p53 outflux to the nucleus, Figure S4, B (open circles). Presented results strongly support conclusion that Selinexor upregulates *de novo* synthesis of G\_p53 in live cells. In agreement with another drug from the SINE family, KPT-185 [59], we can attribute the observed p53 up-regulation to the Selinexor-induced stabilization and activation of p53 [60].

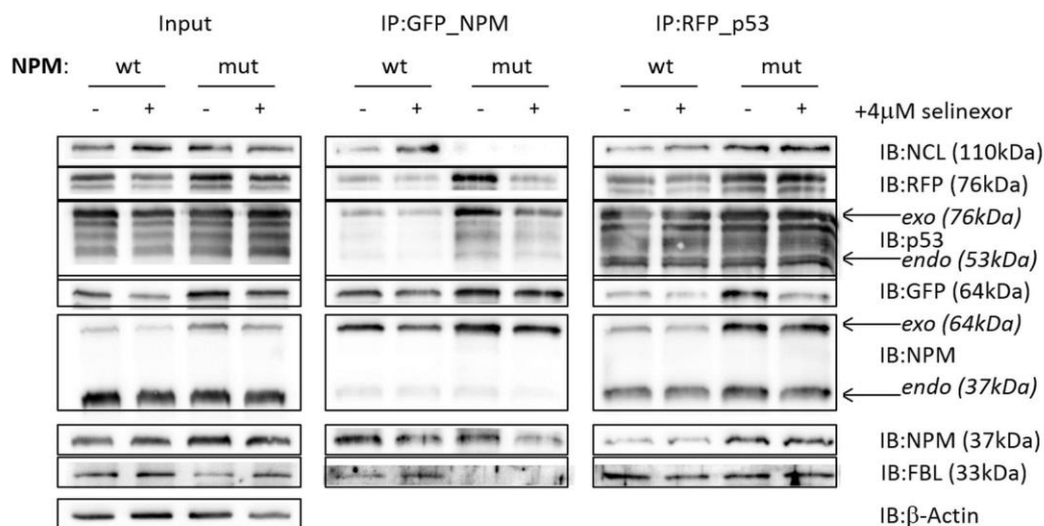
Effect of Selinexor on the p53-NPMwt/mut interaction was further tested by immunoprecipitation. We performed GFP/RFP immunoprecipitation from Selinexor-treated cells co-transfected with G\_NPMwt or G\_NPMmut, and with R\_p53, Figure 9. We found comparable amounts of the p53/NPM complexes in all samples irrespectively of the presence of Selinexor. The result suggests that the complex forms independently of the Selinexor treatment. Selinexor had no effect on NPMwt interaction with NCL as well, which was used as a control for immunoprecipitation. In accordance with our previous findings [25], NCL was not detected in NPMmut precipitates. Moreover, NCL was found in all p53 precipitates confirming interaction of these two proteins [5]. Interestingly, identical results as for NCL were obtained also for another nucleolar protein, fibrillarin (FBL). Specifically, FBL interaction with NPMwt was inhibited by the NPM mutation and FBL interacted with p53, Figure 9. These findings are to the best of our knowledge novel.



**Figure 7.** Effect of Selinexor on localization and lifetime of G\_p53 co-expressed with R\_NPMmut in HEK-293T cells. **(A)** Lifetime images of G\_p53 in the presence and absence of 4  $\mu$ M Selinexor before (the 1st row), immediately after (the 2nd row), and 210 min after the PC (the 3rd row). The photoconverted area is bordered by the dashed line. Bar represents 20  $\mu$ m; **(B)** Detailed time evolution of G\_p53 localization and  $\tau_{GFP}$  maps in single cell with photoconverted eGFP in the presence (upper row, cell no. 1) and absence of Selinexor (lower row, cell no. 4). The cells are marked by arrows in panel A. The nuclear signal was negligible in the absence of Selinexor. Bar is 20  $\mu$ m; **(C, D)** Time evolution of  $\tau_{GFP}$  in the photoconverted (1, 3, 4) and control (2, 5, 6) cells in the presence **(C)** and absence **(D)** of Selinexor. Cells used for integration of the plotted signal are marked in the panel A.



**Figure 8.** Effect of Selinexor on the fluorescence intensity and localization of G\_p53 and R\_NPMmut co-transfected in HEK-293T cells. (A) Intensity images of G\_p53 co-expressed with R\_NPMmut at different times upon the addition of 4  $\mu$ M Selinexor (upper two rows, cell no. 1). The control experiment without Selinexor is presented in the lower two rows (cell no. 4). Depicted are cells from Figure 7B. Bar represents 10  $\mu$ m; (B) Fluorescence intensity changes of G\_p53 (circles) and R\_NPMmut (triangles) in the nucleus (close symbols, full lines, N) and the cytoplasm (open symbols, dash lines, CP) of the cell from the upper two rows of the panel A. Areas used for integration of the fluorescence signal are separated by the dotted line in panel A. The initial irreversible photoconversion of G\_p53 reduced eGFP intensity to about 45% of its initial value before the addition of Selinexor (open circles).

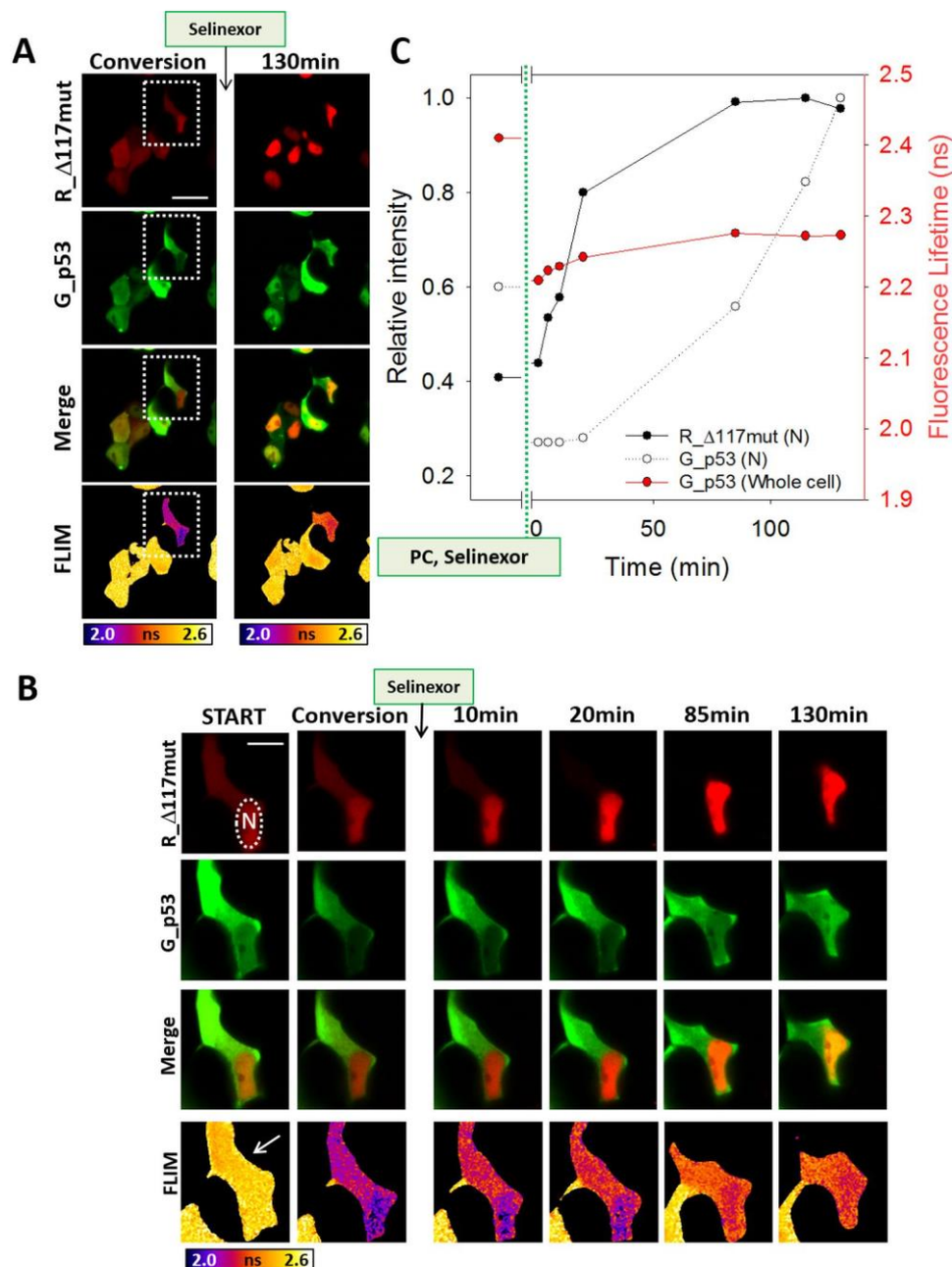


**Figure 9.** Effect of Selinexor on interaction of p53 with NPM variants in lysates of HEK-293T cells. The cells were co-transfected with G\_NPM variants (wt, mut) and R\_p53 in duplicates. Selinexor (4 μM) was added to one sample of each variant 2h before the GFP/RFP-precipitation. Levels of co-precipitated proteins were investigated by immunoblotting. β-Actin expression served as a loading control.

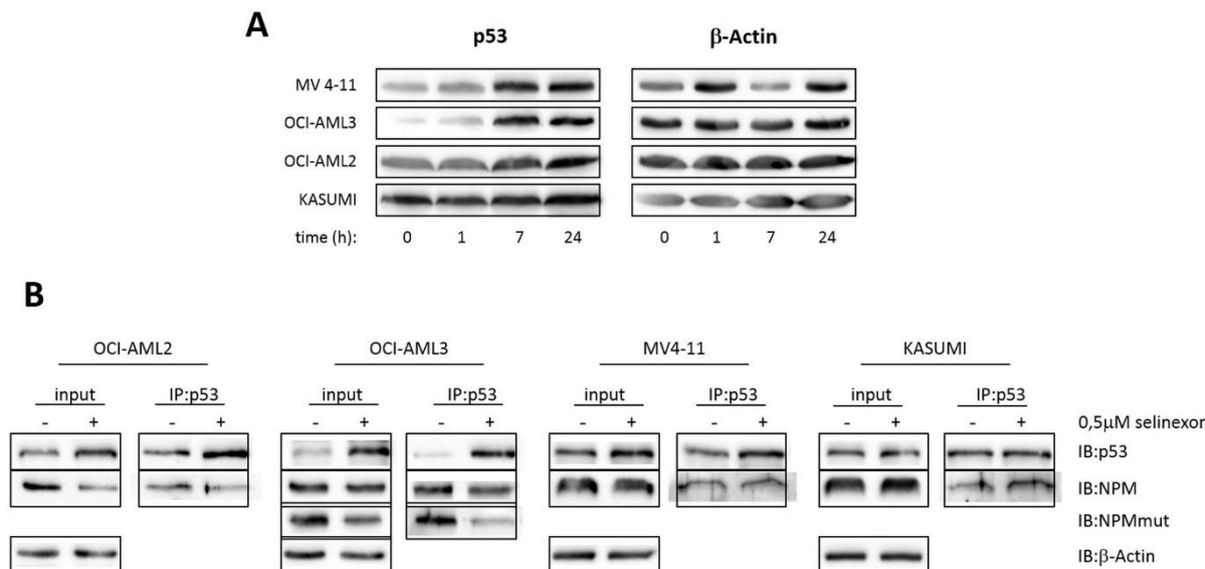
The asynchronous reimport of the complex components back to the nucleus was observed in transfected HEK-293T cells also when p53 was translocated to the cytoplasm by  $\Delta 117$ mut, see Figure 10. The panel A shows, from the top to bottom, intensity images of G\_p53, R\_ $\Delta 117$ mut, their merge, and FLIM images just after the PC and 130 min later. Relocalization dynamics in the photoconverted region is shown in detail in Figure 10B. Rapid relocation of R\_ $\Delta 117$ mut in the nucleus, gradual increase of G\_p53 nuclear intensity as well as slight increase of  $\tau_{\text{GFP}}$  in the photoconverted area can be seen upon the addition of Selinexor. Based on the nuclear rise in the R\_ $\Delta 117$ mut emission intensity depicted in Fig. 10C, it seems that  $\Delta 117$ mut relocalizes back to the nucleus even faster than NPMmut shown in Figure 7, with  $T_{1/2}$  of about 20-30 min. Nevertheless, it should be noted that the relative amount of  $\Delta 117$ mut in the cytoplasm was lower than for NPMmut prior the Selinexor addition, compare Figure 8A and Figure 10B. It is also evident that similarly to NPMmut, the double mutant  $\Delta 117$ mut re-localizes to the nucleus much faster than p53. Increase in  $\tau_{\text{GFP}}$  of G\_p53 integrated across the whole cell does not exhibit so marked elevation in presence of  $\Delta 117$ mut, suggesting lower Selinexor-induced upregulation compared to cells with NPMmut. Mechanistic reasons for this difference are unknown at the moment.

The HEK-293T cell line is suitable for simulations with fluorescently tagged proteins thanks to its highly effective transfectability. Nevertheless, its p53 signaling may be different from that in leukemia cells. We therefore performed immunoprecipitation experiments on a set of leukemia cell lines, Figure 11. Our choice of the cell lines was representative of AML-characteristic mutations. OCI-AML2 cells carry DNMT3A (R635W) mutation, OCI-AML3 line contains NPM1 and DNMT3A (R882C) mutations, and MV4-11 cells have FLT3-ITD mutation. All these lines express wild type p53. The fourth tested line, KASUMI-1, carries the p53 mutation (R248Q) causing high p53 expression in intact cells. We detected Selinexor-induced increase of p53 levels in cell lysates of all the cell lines expressing p53wt, Figure 11A. Despite the rise in p53 even in cell lines lacking NPMmut, the p53 expression increases faster in OCI-AML3 containing NPM with the mutated C-terminus. To confirm persisting NPM-p53 interaction observed in Selinexor treated cells transfected with the exogenous FP-labeled proteins, we performed similar experiments with our panel of AML-related cell lines, Figure 11B. Due to the higher sensitivity of AML cells to the treatment, the concentration of Selinexor was lowered 8-fold to 0.5 μM and the exposure of cells to the drug was prolonged from 2h to 24h. In agreement with the results on HEK-293T cells, NPM co-precipitated with p53 in all examined cell lines irrespectively

of the Selinexor presence. In the OCI-AML3 cell line, NPMmut was also detected with a mutant-specific antibody in the p53 precipitates.



**Figure 10.** Cellular dynamics of G\_p53 and R\_Δ117mut upon Selinexor treatment. (A) Overview of localization, intensity and lifetime changes of co-expressed G\_p53 and R\_Δ117mut in HEK-293T cells after addition of 4 μM Selinexor. eGFP within the region bordered by the dashed line was irreversibly photoconverted to the low-lifetime form before the Selinexor addition. Bar represents 30 μm; (B) Detailed time evolution of the photoconverted region from panel A. Nucleus (N) is marked by the dashed line. Bar represents 10 μm; (C) Intensity change of G\_p53 (open circles) and R\_Δ117mut (closed circles) at the nucleus of the cell from Panel B. Signal for the lifetime change was integrated from the whole cellular area (red circles).



**Figure 11.** Effect of 0.5  $\mu$ M Selinexor on interaction between p53 and NPM in AML-derived cell lines. (A) p53 expression at indicated times after the Selinexor addition; (B) p53-precipitation 24h after the Selinexor addition. Control samples without Selinexor were collected at the same time. Levels of co-precipitated NPM were investigated by immunoblotting.  $\beta$ -Actin expression served as a loading control.

#### 4. Discussion

A presence of interaction between NPM and p53 was reported in stressed cells almost twenty years ago [3]. Since then, the interaction between purified recombinant proteins was confirmed by surface plasmon resonance and regions responsible for the interaction were specified in GST-pulldown assays [4]. As the C-terminus of NPM (AA242-269) mediates the interaction with p53 [4], it would not be surprising, if the interaction was disrupted by the AML-associated NPM mutation located at the same terminus. However, no data confirming this hypothesis have been published yet. In this work we bring several lines of evidence that both p53-NPMwt and p53-NPMmut interactions occur under physiological conditions in live cells.

We have reported consistently with others that due to the heterooligomer formation with NPMmut, a fraction of NPMwt is delocalized into the cytoplasm [29, 30]. Similarly, confocal microscopy revealed extensive p53 delocalization into the cytoplasm of cells co-expressing NPMmut, Figure 1, which suggests interaction between p53 and NPMmut. The complex formation was therefore investigated by co-immunoprecipitation using GFP Trap beads. The experiments have confirmed already published p53-NPMwt interaction [3, 4]. Moreover, they newly proved interaction between p53 and C-terminally mutated variants of NPM that are characteristic for AML, Figure 2. The discovery of the p53-NPMmut interaction appears rather surprising in the light that NCL, which also interacts with C-terminal NPMwt region (AA187-241) [19], was not detected in the complex with NPMmut [25]. Interaction of p53 with both NPMwt and NPMmut was further unambiguously confirmed in live cells, using FLIM-FRET, Figure 5. Subsequently, p53 was found also in the G $\Delta$ 117wt-precipitates, which suggests that the NPM oligomerization domain is unessential for its binding to p53, see Figure 4A. In fact, even higher amount of p53 co-precipitated with  $\Delta$ 117wt than with NPMwt. FRET between G $\Delta$ 117wt and R $\Delta$ p53 confirmed this finding in live HEK-293T cells, Figure 5A. We conclude that neither the wild-type C-terminus nor the oligomeric NPM form is necessary for p53 binding.

We therefore tested how the double mutation affects formation of p53/NPM complex. Massive cytoplasmic localization of fluorescently labeled p53 was observed when co-expressed with  $\Delta$ 117mut in live HEK-293T cells, as shown in Figure 3. Surprisingly, the cells abundantly exhibited p53 in the cytoplasm, despite the prevalingly nuclear localization of  $\Delta$ 117mut. The result was the same for both eGFP and mRFP1 labeling combination, see Figure S3. This observation seems not to be completely in line with the finding that lower

p53- $\Delta$ 117mut binding was detected by co-immunoprecipitation, Figure 4. This suggests that the p53/ $\Delta$ 117mut complex is formed, nevertheless its  $K_D$  is likely higher and lower amount of the complex is therefore detected by immunoprecipitation. This conclusion is supported also by FLIM results in Figure 5B, where only insignificant FRET stemming from a small fraction of the R\_p53/G\_ $\Delta$ 117mut complex in the cytoplasm was detected on a high background of free uncomplexed G\_ $\Delta$ 117mut donor. Due to the very low amount of R\_p53 in the nucleus, the complex cannot be detected there by FLIM-FRET, either.

Both p53 and NPM are client proteins of nuclear exporter XPO1. However, only in cells expressing NPMmut, p53 is ubiquitously detected in the cytoplasm. As NPM is supposed to form complex with p53 [3, 4] and we found this complex also in intact cells, we thus expect that p53 and NPMmut are simultaneously co-exported out of the nucleus in this form of p53/NPMmut complex. Our data suggest that even weaker binding of p53 to  $\Delta$ 117mut is sufficient to facilitate translocation of p53 to the cytoplasm. However, an alternative transport mechanism involving ternary complex with some other NPM- and p53-interacting partner, cannot be ruled out at this moment.

The translocation of p53 to the cytoplasm implicates important consequences for its turnover and function. In intact cells, p53 protein level is tightly controlled by numerous posttranslational modifications and it is kept at low level by proteasome. However, stable p53 production from *TP53* transcript allows for rapid reaction on stress stimuli by stopping the protein degradation and triggering its transcription activity. A regulation of the p53 expression, stability and activity takes place mainly in the nucleus and the p53 delocalization likely causes a loss of its function [33, 61]. Targeting the cytoplasmic p53 and restoration of its nuclear localization could be therefore a promising alternative for the therapy of the AML with NPMmut. An inhibition of XPO1 by Selinexor [37, 62] seems to be an auspicious choice [35].

When the XPO1 exporter is blocked by Selinexor, NPMmut is no longer exported to the cytoplasm and nuclear export/import equilibrium is changed favoring relocalization of NPMmut into the nucleus [45]. In co-transfected cells treated with Selinexor, NPMmut thus cannot drive p53 translocation to the cytoplasm in the p53/NPMmut complex. At the same time, p53, as another XPO1 client, cannot be exported to the cytoplasm and returns to the nucleus via importins [63]. Indeed, we observed relocalization of both NPMmut and p53 to the nucleus upon addition of 4 $\mu$ M Selinexor to transfected HEK-293T cells, Figures 6 and 7. The observed Selinexor-induced increase of p53 expression is in accordance with the activation of p53-dependent apoptosis after Selinexor treatment [60, 62]. Importantly, p53 and NPMmut responded independently and exhibited different dynamics upon Selinexor addition. Figures 8 and S4 demonstrate that the redistribution of NPMmut after application of Selinexor is considerably faster than that of p53. Hence, it seems evident that p53 and NPMmut do not have to be synchronously co-transported to the nucleus. Our results indicate that cytoplasmic complexes of p53 with C-terminally-mutated NPM variants are weaker and rather dynamic. Exchanging complex constituents and rapid equilibration could allow for independent nuclear import of p53 and NPMmut variants. Abolishment of the p53-NPM interaction by Selinexor is therefore not required. The detection of the p53/NPMmut complex in Figure 9 is fully consistent with the outlined mechanism.

Cytoplasmic mislocalization of p53 requires presence of NPMmut. Targeting of the NPMmut-p53 interaction might therefore facilitate restoration of the p53 proapoptotic and regulative function in the nucleus. Nevertheless, strong translocation of p53 to the cytoplasm was observed in the presence of  $\Delta$ 117mut when lower amount of the p53-NPM complex is expected to be formed in the nucleus independently of the Selinexor, owing to the documented weaker p53- $\Delta$ 117mut interaction. The faster reimport of  $\Delta$ 117mut to the nucleus, compared to that of NPMmut where the interaction is stronger, is therefore consistent with higher cytoplasmic pool of free  $\Delta$ 117mut ready for import. The observed asynchronous reimport of p53 and NPMmut suggests that an induction of the p53 nuclear translocation rather than a disruption of the p53-NPMmut interaction could stabilize p53 and restore its cellular function. Therapeutic boosting of the p53 reimport could be the

right target, since p53 is also a subject of ubiquitinylation and fast degradation in the cytoplasm [36].

The therapeutic effect of Selinexor is supposed to be mediated by restoration of the nuclear localization of p53. This could happen to some extent in all cells, however, the effect should be augmented in cells with mislocalized p53. Significant induction of p53 is visible in Figure 11A for the OCI-AML3 cell line expressing NPMmut. The cells display very low basal levels of p53, likely caused by its high degradation when mislocalized in the cytoplasm. Fast p53 induction in the presence of Selinexor then could be partially explained by its restored nuclear localization and lower degradation rate there. However, due to the very low p53 concentration in intact cells, the supposed NPMmut-dependent translocation of p53 to cytoplasm of OCI-AML3 cells cannot be checked by immunostaining. Extensive induction of p53 shown in Figure 11B accompanied by slightly lowered NPMmut level in Selinexor-treated OCI-AML3 suggests that this exclusive response could be related to the presence of NPMmut. We conclude that the p53 induction might explain positive therapeutic effect of Selinexor in treating AML with NPMmut, possibly in combination with other p53-inducing agents, such as mdm2/p53 inhibitors, e.g. nutlin-3a [37].

## 5. Conclusions

We examined importance of p53-NPM interaction for accumulation of p53 in the cytoplasm. The interaction was found to be unaffected by the AML-related mutation of the NPM C-terminus. We proved that the p53-NPMmut interaction persists also when the oligomerization of NPM is compromised by the deletion of its N-terminal domain (AA1-117). Finally, the cytoplasmic localization of exogenous p53 was found also in the presence of double mutant  $\Delta$ 117mut co-expressed in live HEK-293T cells. To the best of our knowledge, the NPMmut-mediated cytoplasmic translocation of p53 has not been reported yet. We documented that in response to the Selinexor treatment, both p53 and NPMmut returned to the nucleus, however, with significantly different kinetics. We concluded that p53 and NPMmut do not have to be co-transported to the nucleus as stable complexes. They rather form weaker short-lived complexes allowing rapid exchange of its constituents and independent nuclear import.

**Supplementary Materials:** The following are available online at [www.mdpi.com/xxx/s1](http://www.mdpi.com/xxx/s1), Figure S1: Localization of p53 co-expressed with NPMmut in HEK-293T cells shown for two tagging combinations; Figure S2: Interaction between NPMwt and p53 in cell lysates of transfected HEK-293T cells; Figure S3: Localization of p53 co-expressed with  $\Delta$ 117mut in HEK-293T cells shown for two tagging combinations; Figure S4: Effect of Selinexor on the fluorescence intensity and localization of G\_p53 and R\_NPMmut co-transfected in HEK-293T cells. Supplementary Material 2 contains original Western Blot images.

**Author Contributions:** Conceptualization, A.H., D.S., P.H. and B.B.; methodology, A.H., D.S., P.H. and B.B.; formal analysis, A.H., D.S., P.H. and B.B.; investigation, A.H., D.S., P.O., P.R., P.H. and B.B.; resources, A.H., D.S., P.H. and B.B.; data curation, A.H., D.S., P.H. and B.B.; writing—original draft preparation, A.H., D.S., P.H. and B.B.; writing—review and editing, A.H., D.S., P.H. and B.B.; funding acquisition, P.H. and B.B. All authors have read and agreed to the published version of the manuscript.

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**Data Availability Statement:** The data presented in this study are available in Supplementary Materials, see above.

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