De novo assembly of a PML nuclear subcompartment occurs through multiple pathways and induces telomere elongation

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Summary
Telomerase-negative tumor cells use an alternative lengthening of telomeres (ALT) pathway that involves DNA recombination and repair to maintain their proliferative potential. The cytological hallmark of this process is the accumulation of promyelocytic leukemia (PML) nuclear protein at telomeric DNA to form ALT-associated PML bodies (APBs). Here, the de novo formation of a telomeric PML nuclear subcompartment was investigated by recruiting APB protein components. We show that functionally distinct proteins were able to initiate the formation of bona fide APBs with high efficiency in a self-organizing and self-propagating manner. These included: (1) PML and Sp100 as the constituting components of PML nuclear bodies, (2) telomere repeat binding factors 1 and 2 (TRF1 and TRF2, respectively), (3) the DNA repair protein NBS1 and (4) the SUMO E3 ligase MMS21, as well as the isolated SUMO1 domain, through an interacting domain of another protein factor. By contrast, the repair factors Rad9, Rad17 and Rad51 were less efficient in APB nucleation but were recruited to preassembled APBs. The artificially created APBs induced telomeric extension through a DNA repair mechanism, as inferred from their colocalization with sites of non-replicative DNA synthesis and histone H2A.X phosphorylation, and an increase of the telomere repeat length. These activities were absent after recruitment of the APB factors to a pericentric locus and establish APBs as functional intermediates of the ALT pathway.

Key words: Alternative lengthening of telomeres, DNA repair, DNA recombination, Promyelocytic leukemia nuclear bodies

Introduction
Telomeres, the ends of the linear chromosomes, contain repetitive DNA sequences [in humans (TTAGGG)ₙ, typically 3–20 kb in length] that are organized into a specialized nucleoprotein complex. This structure protects the telomeres from being processed as a DNA double-strand break (DSB) by the DNA repair and recombination machinery of the cell (de Lange et al., 2006; Verdu and Karlseder, 2007). Telomeres shorten with every cell division owing to the incomplete replication of the lagging strand and additional exonucleolytic activities. Upon reaching a critical length cellular senescence is induced (Collado et al., 2007). In most tumor cells the reverse transcriptase telomerase is reactivated and the telomere repeat sequences can be extended, thereby allowing unlimited proliferation. However, some immortalized cell lines and 10–15% of cancer cells use an alternative lengthening of telomeres (ALT) mechanism for the maintenance of their telomere repeats (Henson et al., 2002). This pathway involves DNA repair and recombination processes (Dunham et al., 2000). ALT-positive cells are characterized by the association of telomeric DNA with promyelocytic leukemia nuclear bodies (PML-NBs), forming ALT-associated PML-NBs (APBs) (Henson et al., 2002). PML-NBs are mobile nuclear subcompartments present in most mammalian cells and have been implicated in a variety of cellular functions including apoptosis, senescence, tumor suppression, transcription, antiviral response and DNA replication and repair (Bernardi and Pandolfi, 2007; Dellaire and Bazett-Jones, 2004; Lallemand-Breitenbach and de The, 2010; Takahashi et al., 2004). APBs colocalize with DNA repair and recombination proteins, and a number of models for the molecular mechanisms have been proposed to explain the role of APBs in the ALT pathway (Cesare and Reddel, 2010; Draskovic et al., 2009; Henson et al., 2002; Jiang et al., 2007). However, it is not clear whether a functional link between APB formation and telomere lengthening exists. To address this issue, we investigated the de novo formation of APBs. Protein components of APBs were recruited to telomeres tagged with stable integrations of bacterial lac operator DNA sequence (lacO) repeats in the ALT-positive human osteosarcoma U2OS cell line (Jegou et al., 2009). This system allowed elucidating the APB assembly process after enriching one factor and the dissection of the interaction network that leads to APB formation and the recruitment of DNA repair and recombination factors. Furthermore, we show that the de novo formation of APBs induced the elongation of telomeric repeats in a DNA repair based synthesis process. These results demonstrate that APBs are indeed functional intermediates in the ALT pathway and identify them as potential targets for the treatment of ALT-positive tumors.

Results
Recruitment of PML and Sp100 to lacO-labeled telomeres leads to the de novo assembly of bona fide APBs
The ALT-positive U2OS cell line F6B2, which has three stable integrations of bacterial lac operator (lacO) repeats (referred to as
telomeric lacO arrays) directly adjacent to the telomeres of chromosomes 6q, 11p and 12q (Jegou et al., 2009), was transfected with a bacterial LacI repressor fused to a GFP-binding protein (GBP) (Rothbauer et al., 2006; Zolghadr et al., 2008). The LacI construct with (GBP–LacI–RFP) or without (GBP–LacI) an additional red fluorescent mRFP1 marker was used to recruit GFP- or YFP-tagged proteins and interacting factors to the telomere-associated lacO arrays (Fig. 1A). As described in previous studies, the GBP domain binds with high affinity to GFP with an equilibrium dissociation constant of 0.23 nM (Rothbauer et al., 2006). Thus, this system is equivalent to the use of direct fusion constructs of LacI with the protein of interest (Kaiser et al., 2008; Soutoglou and Misteli, 2008; Tumbar et al., 1999). Accordingly, the recruitment of GFP–PML through GBP–LacI–RFP in the F6B2 cell line resulted in colocalization of GFP–PML with three telomeres (Fig. 1B). In order to address whether tethering PML to the lacO-labeled telomeres leads to the assembly of APB-like structures at these sites, the presence of the main structural components of PML-NBs was analyzed. For this, we used the PML III splicing variant that itself appears to have no specific interactions with shelterin proteins (e.g. TRF1 and TRF2), as opposed to PML IV (corresponding to PML 3 in the arabic numbering scheme) for which binding to TRF1 has been reported (Yu et al., 2009). Whereas PML IV showed a behavior similar to that of PML III in control experiments (data not shown), the use of PML III allowed us to separate the initial telomeric recruitment event (provided in our system by lacO and GBP–LacI) more clearly from the other protein–protein interactions of PML.

PML-NBs are composed of PML and Sp100 proteins that carry post-translational small ubiquitin-like modifier (SUMO) modifications and organize in a spherical shell (Bernardi and Pandolfi, 2007; Lang et al., 2010; Shen et al., 2006). Accordingly, we investigated whether PML or Sp100 recruitment would result in the accumulation of other components (Fig. 2). GFP-tagged PML protein was efficiently bound to the lacO arrays through GBP–LacI–RFP. This triggered the subsequent recruitment of endogenous Sp100 to these sites with an efficiency of 100% (Fig. 2A). The reverse experiment, tethering GFP–Sp100 to the lacO arrays, also induced the formation of PML-NBs because endogenous PML was detected at all GFP–Sp100-positive telomeric lacO arrays (Fig. 2B). Furthermore, the recruitment of GFP–PML increased the presence of endogenous SUMO isoforms to more than 90% (93 ± 9% for SUMO1, 98 ± 11% for SUMO2 and SUMO3, Fig. 2C,D). By contrast, transfection of only GBP–LacI–RFP or GBP–LacI–RFP together with the isolated GFP domain did not lead to a significant enrichment of endogenous PML or Sp100 at these sites (Fig. 2E; supplementary material Fig. S1A). Likewise, co-transfecting GFP-LacI without the GBP domain together with GFP–PML did not target PML to the lacO arrays (data not shown). The residual degree of colocalization in the control cells is likely to reflect the presence of endogenously formed APBs at the three tagged telomeres, as well as random superposition of the two signals in the same optical section of the confocal images.

The analysis of the de-novo-assembled APB-like structures by confocal laser scanning microscopy (CLSM) revealed a cap-like organization of PML protein around telomeric repeats (Fig. 1B; supplementary material Fig. S1) that was indistinguishable from that of the de-novo formed complexes of PML nuclear bodies (APBs) at the telomeres. (A) Schematic representation of the experimental approach. The U2OS cell clone F6B2 employed in this study has three integration sites of the lacO arrays, adjacent to the telomeres of chromosomes 6q, 11p and 12q (Jegou et al., 2009). Different GFP-tagged proteins were recruited to the lacO arrays through a fusion of LacI repressor to a high-affinity GFP-binding domain (GBP) and a red fluorescent protein domain (GBP–LacI–RFP). Endogenous interaction partners of the GFP-labeled protein were then identified by subsequent immunostaining and evaluation of the colocalization of the fluorescence signals by confocal laser scanning microscopy (Rothbauer et al., 2006; Zolghadr et al., 2008). (B) The F6B2 U2OS cell line was co-transfected with GFP–PML and GBP–LacI expression vectors. Through binding of GBP–LacI to the lacO repeat sequences, GFP–PML was recruited to these sites. Staining of the telomeric repeats TTA(G)3 with a Cy3-labeled PNA probe revealed the colocalization of telomeres with the GFP–PML signal in the confocal images (see arrows). This indicates the formation of bona fide APBs at the three telomere sites of chromosomes 6q, 11p and 12q. Scale bar: 10 μm.
of endogenous APBs (Jegou et al., 2009; Lang et al., 2010). Two telomere signals within one APB could only be distinguished in a very small fraction (0.3 ± 0.1%) of the de-novo-formed telomeric PML subcompartments evaluated here. For endogenous APBs in U2OS cells this number was even lower at ∼0.10 ± 0.04% (six out of 5803). These results were obtained with an advanced automated three-dimensional image analysis of confocal three-dimensional

stacks of endogenous APBs in U2OS cells visualized by immunostaining against PML and TRF2 using the method described previously (Osterwald et al., 2011; Wörz et al., 2010).

Together, our results indicate that the artificial enrichment of GFP–PML at lacO-tagged telomeres led to the assembly of bona fide APBs (defined as PML-NBs at telomeres) with respect to their structural composition.

Fig. 2. Formation of a de novo APB by recruitment of PML and Sp100 protein. Cells were co-transfected with GBP–LacI–RFP and the indicated GFP constructs. This resulted in the tethering of the GFP-tagged protein to the three lacO-labeled telomeres. Association of the main APB components, PML, Sp100 and SUMO, was detected by immunostaining the endogenous proteins and evaluating the colocalization of the two fluorescence signals in optical sections obtained by confocal laser scanning microscopy imaging. (A) Recruitment of GFP–PML yielded 100% colocalization with endogenous Sp100 compared with 24 ± 5% in the absence of GFP–PML (P < 0.0001). (B) GFP–Sp100 displayed 100% colocalization with endogenous PML compared with the control value of 32 ± 5% in the absence of GFP–Sp100 (P < 0.0001). (C, D) GFP–PML induced 93 ± 9% (endogenous SUMO1) and 98 ± 11% (endogenous SUMO2/3) colocalization compared with 32 ± 5% and 24 ± 4% in the control experiments, in which GFP–PML was absent (P < 0.0001 for both analyses). (E) Recruitment of the isolated GFP domain. This led to a colocalization of only 19 ± 5% with endogenous PML, which did not significantly differ from the 20–30% background observed in the control transfection with only GBP–LacI–RFP (P = 0.48). Scale bars: 10 μm. All error bars show the s.d.
Fig. 3. See next page for legend.
SUMO1 interactions are essential for APB assembly

Because impairment of sumoylation disrupts PML-NB formation, and sumoylated telomeric proteins are crucial for the formation of APBs in ALT-positive cells (Potts and Yu, 2007; Shen et al., 2006), we investigated the effect of tethering the SUMO domain to the lacO-tagged telomeres. Recruitment of GFP-tagged SUMO1, SUMO2 or SUMO3 constructs was clearly sufficient to initiate the formation of APBs, as judged by the degree of colocalization of the three GFP-SUMO variants with endogenous PML, endogenous Sp100 and endogenous Rad51, i.e. 80–85% (Fig. 3A–C), 60–80% (supplementary material Fig. S2A–C) and 40–50% (supplementary material Fig. S2D–F), respectively. Structural APB components, such as PML and Sp100, are subject to sumoylation and, at the same time, contain SUMO-interacting motifs (SIMs) (Hecker et al., 2006; Knipscheer et al., 2008; Shen et al., 2006). Thus, in these experiments the effect of GFP-SUMO1 that was covalently conjugated to its target proteins and non-covalent interactions through the SIMs could not be distinguished (supplementary material Fig. S3). Accordingly, we investigated SUMO constructs that could not be conjugated to other proteins. The covalent attachment of SUMO occurs after cleavage of its C-terminus, exposing a Gly-Gly motif that becomes bound to a lysine residue of the target protein (Geiss-Friedlander and Melchior, 2007; Muller et al., 2001). The YFP–SUMO1ΔC7, GFP–SUMO2ΔC4 and the GFP–SUMO3ΔC13 mutants lack the C-terminal double-glycine motif and can no longer be attached to target proteins (Ayaydin and Dasso, 2004; Lin et al., 2006; Mukhopadhyay et al., 2006). Tethering the SUMO1ΔC7 mutant to the lacO-labeled telomeres resulted in APB formation with an efficiency that was similar to that of the conjugable wild-type SUMO1 construct (77±6% compared with 82±10% colocalization with endogenous PML, Fig. 3A,D). Thus, the interaction of an isolated SUMO1 domain with the SIMs of other proteins was sufficient for the APB nucleation event. By contrast, the non-conjugable mutants of SUMO2 and SUMO3 were significantly less efficient in this respect, yielding colocalization with endogenous PML of 46±5% (GFP–SUMO2ΔC4) and 45±7% (GFP–SUMO3ΔC13) (Fig. 3E,F).

To test whether SIM–SUMO interactions are indeed essential for the de novo APB assembly, the YFP–SUMO1ΔC7(-) variant was evaluated. This variant was constructed by replacing the residues Val38 and Lys39 with alanine residues. These residues are part of the D-strand of SUMO1, which is crucial for SIM binding, as shown in several studies (e.g. Perry et al., 2008; Song et al., 2005).

Accordingly, YFP–SUMO1ΔC7(-) can neither be conjugated to another protein nor bind to a SIM. As shown in Fig. 3G, tethering this construct to the telomeres did not increase colocalization with endogenous PML over background levels. Thus, a SIM interaction with SUMO1 is a central component of APB nucleation. This conclusion is in line with the behavior of yet another type of SUMO construct, namely a C-terminally tagged GFP fusion of SUMO3 (Fig. 3H). This fusion protein appeared to be mostly resistant to cleavage of the C-terminus during the maturation process because it was not conjugated (supplementary material Fig. S3). Interestingly, this variant was also unable to induce de novo APB assembly upon telomere recruitment, which might be due to interference of the C-terminal GFP tag with SIM binding.

De novo APB formation can be induced with high efficiency by recruiting the shelterin components TRF1 and TRF2, and recombination factor NBS1, but not Rad9, Rad51 or Rad17

Other known APB components were tested for their capability to induce the assembly of PML-NBs when recruited to telomeres. First, the telomere repeat binding factors 1 and 2 (TRF1 and TRF2, respectively), were investigated (Fig. 4A,B). Both TRF1 and TRF2 bind to telomeric repeats and are therefore present in endogenous APBs. Tethering these factors resulted in a strong increase of colocalization with endogenous PML, with TRF2 being somewhat more efficient than TRF1 (85±7% colocalization with GFP–TRF2 and 70±8% with GFP–TRF1, Fig. 4A,B).

Because APBs are characterized by the presence of several DNA repair and recombination proteins, the propensity of such proteins to drive PML-NB assembly at lacO-tagged telomeres was examined. The recombination factors NBS1 and Rad51, as well as the DNA repair factors Rad9 and Rad17, were tethered to the lacO arrays as GFP fusions (Fig. 4C–F; Table 1). Recruitment of NBS1, which is a central component of the Mre11–Rad50–Nbs1 (MRN) repair and recombination complex, increased endogenous PML levels at the lacO telomeres with a high efficiency to 83±9% (Fig. 4C) (Jiang et al., 2005; Jiang et al., 2007; Wu et al., 2003). Rad51 is a central player in DSB repair through homologous recombination and is also involved in normal telomere function, presumably by promoting t-loop formation (Verdun and Karlseder, 2007; West, 2003). Furthermore, Rad51 is present in APBs (Yeager et al., 1999). The recruitment of GFP-tagged Rad51 led only to a small increase of endogenous PML at these telomeres to 40±4% (Fig. 4D). The Rad9 and Rad17 proteins are part of the RFC–Rad17–9–1-1 complex that participates in the DNA damage response, plays a role in telomere stability and is a component of APBs (Nabetani et al., 2004; Pandita et al., 2006; Parrilla-Castellar et al., 2004). Enriching GFP–Rad9 at the lacO-labeled telomeres resulted in subsequent recruitment of endogenous PML of 59±6% (Fig. 4E). In contrast to the other investigated proteins, recruitment of GFP–Rad17 did not initiate the assembly of PML-NBs (Fig. 4F).

The composition of de novo APBs is indistinguishable from endogenous APBs

To assess whether the de novo-assembled APBs also contain endogenous proteins involved in DNA repair and recombination, we investigated their composition by immunostaining (Fig. 5; Table 1). NBS1, Rad17 and Rad9, all bona fide components of functional APBs (Jiang et al., 2007; Jiang et al., 2009; Nabetani et al., 2004; Wu et al., 2003), were enriched between twofold
(NBS1, Fig. 5A) and approximately fourfold (Rad9, Fig. 5B) after GFP–PML recruitment. Thus, our de novo assembly approach resulted in PML-NBs associated at telomeres that were equivalent to APBs in terms of their protein composition by all criteria reported for endogenous APBs in the literature.

**APB components can be assembled efficiently at a pericentric lacO integration site by targeting PML, TRF1, TRF2 or NBS1 to this locus**

In order to examine whether the assembly of APB proteins requires the telomeric location of the lacO array, the U2OS cell clone F42B8 was investigated; this clone has one lacO array insertion at the pericentric region of chromosomes 2p (supplementary material Fig. S4) (Jegou et al., 2009). These cells showed a higher level of colocalization of endogenous PML with the lacO arrays (44 ± 6%) compared to the telomeric lacO sequences (supplementary material Fig. S5A). This is in line with findings that described the colocalization of PML-NBs with pericentric heterochromatin (Everett et al., 1999; Luciani et al., 2006) and the previously reported interaction of heterochromatin protein 1 (HP1) with PML-NBs, which is also enriched at these sites (Hayakawa et al., 2003). Whether LacI-bound lacO loci per se have an increased propensity for association with PML-NBs is not clear. According to a previous study, LacI–GFP-tagged lacO arrays display no preferred colocalization with PML-NBs in clones of the human HT-1080 sarcoma cell line for integration sites at 13q22, 5p14, 3q26.2, 13p and 1q11 (Chubb et al., 2002). By contrast, investigations of a baby hamster kidney cell line with a lacO array at unknown genomic localization showed an association of PML-NBs with this region (Tsukamoto et al., 2000). Recruiting GFP–PML to the pericentric lacO array enriched endogenous SUMO isoforms to an extent that was similar to that measured for the telomeric lacO loci. (supplementary material Fig. S5B,C). Furthermore, TRF1 and TRF2 were similarly efficient in the subsequent recruitment of endogenous PML to the pericentric locus as they were at telomeric sites (Fig. 4A,B; supplementary material Fig. S6A,B). A comparable result was obtained when tethering NBS1 to the pericentric lacO array (supplementary material Fig. S6C), whereas GFP-tagged Rad51 could not initiate PML-NB formation at this locus (supplementary material Fig. S6D). We then tested whether the accumulation of endogenous APB maker proteins at pericentric regions upon GFP–PML recruitment was different. Remarkably, the protein composition of the nuclear bodies induced by recruitment of GFP–PML to the pericentric lacO arrays revealed that all factors were enriched under these conditions to a similar or even higher degree than at the telomeric sites (Fig. 5, right-hand panels). Thus, the de-novo-assembled nuclear bodies at the pericentric chromatin locus had an APB-like protein composition.

**APB assembly can be induced by the MMS21 SUMO E3 ligase and occurs in two steps**

The SUMO E3 ligase MMS21 induces the sumoylation of several telomere-repeat-associated proteins, such as TRF1, TRF2 and Rap1, in ALT-positive cells and thereby supports APB formation (Potts and Yu, 2007). In order to investigate the role of MMS21, we first tested for the presence of endogenous MMS21 at the lacO-labeled telomeres after GFP–PML recruitment (Fig. 6A). MMS21 was highly enriched upon tethering PML at these sites, resulting in an increased level of colocalization from 19 ± 3% to 68 ± 7%. Interestingly, the nuclear bodies formed de novo at the pericentric lacO array contained endogenous MMS21 at similar levels to those detected at the telomeric loci (79 ± 9% as opposed to 28 ± 5% in the GFP control, Fig. 6B).

Next, we sought to test whether the presence of MMS21 at telomeres is sufficient to initiate APB formation in terms of recruiting endogenous PML. To this end, GFP–MMS21 was bound to the telomeric lacO sequences. We observed that GFP–MMS21 was highly efficient in promoting APB assembly as it increased the colocalization between telomeric lacO sites and endogenous PML from 19 ± 5% to 86 ± 9% (Fig. 6C). Notably, tethering the GFP–MMS21 to the pericentric lacO sites also increased the percentage of colocalizing endogenous PML from 44 ± 6% to 95 ± 10%, which suggests that other sumoylation targets or interaction partners might exist in addition to telomere-associated proteins (supplementary material Fig. S6E). Next, we addressed the question of whether the GFP–MMS21-induced targeting of endogenous PML protein to the telomeric lacO sites was accompanied by the enrichment of the DNA repair factor Rad9. The enrichment of endogenous PML and Rad9 at these sites was evaluated by immunofluorescence (Fig. 6D). Remarkably, 35 ± 4% of the GFP–MMS21-bound telomeres colocalizing with PML did not contain Rad9 (Fig. 6D1,D3,E). By contrast, no colocalization of endogenous Rad9 with GFP–MMS21 was detected without the simultaneous presence of PML. To compare this result with the situation for native APBs, we investigated the PML:Rad9 ratio at telomere repeats identified with GFP–TRF2. The vast majority (98%) of endogenous APBs (defined as colocalization of PML and TRF2) contained Rad9, and only 0.9% of the telomeres marked by TRF2 had Rad9 but no PML (Fig. 6E). On average, we detected 54 ± 11 telomeres per cell, of which 8 ± 3 were associated with APBs. Endogenous APBs were found in almost every cell of the asynchronous cell population, in contrast to previous reports (Yeager et al., 1999). We note that our CLSM-based detection also included relatively small colocalization spots, as discussed in further detail elsewhere (Osterwald et al., 2011). In summary, the fully assembled functional endogenous APBs contained both PML and Rad9, which is in line with previous work showing PML colocalizing with almost all Rad9 foci in U2OS cells (Nabetani et al., 2004). In the de novo assembly process initiated by recruitment of GFP–MMS21, however, a two-step process was revealed: tethering MMS21 to the telomere led to the concomitant assembly of the PML, Sp100 and SUMO network, presumably through sumoylation of target proteins. Subsequently the DNA recombination and repair factor Rad9 was recruited, as apparent from the 35 ± 4% of telomeres with colocalization of GFP–MMS21 and endogenous PML that did not contain endogenous Rad9 (Fig. 6D,E).

**Recruitment of PML induces DNA repair synthesis at telomeric, but not at pericentric, sites**

Because the ALT mechanism involves DSB repair and recombination processes, we investigated whether APB assembly induced these activities. First, we probed de-novo-formed APBs for the presence of the phosphorylated form of the histone variant H2A.X (γH2A.X), a molecular marker for DSB repair and a component of APBs (Cserei et al., 2009; Ismail and Hendzel, 2008; Nabetani et al., 2004). Indeed, an 11 ± 7% higher γH2A.X colocalization was found, which is indicative of an increased activity in DSB repair processes (Fig. 7A). By contrast,
Fig. 4. Initiation of APB formation by shelterin and DNA repair and recombination proteins. Confocal images of cells that were co-transfected with GBP–LacI–RFP (column 1), the indicated GFP fusion protein (column 2; the merge of the RFP and GFP signal is shown in column 3) and immunostained for endogenous PML protein to determine APB formation (column 4). The colocalization of the GFP signal at telomeric lacO arrays with the immunofluorescence of endogenous PML at these sites (column 5) yielded 19 ± 5% in the control, in which an isolated GFP domain was recruited. The propensity of proteins to induce APB formation when recruited to the telomeres as GFP fusions was evaluated in terms of colocalization with endogenous PML. This yielded values of (A) GFP–TRF1, 70 ± 8% ($P<0.0001$); (B) GFP–TRF2, 85 ± 7% ($P<0.0001$); (C) NBS1–GFP, 83 ± 9% ($P<0.0001$); (D) Rad51–GFP, 40 ± 4% ($P<0.0005$); (E) GFP–Rad9, 59 ± 6% ($P<0.0001$); and (F) GFP–Rad17, 31 ± 6% ($P=0.10$). Scale bars: 10 μm. All error bars show the s.d.
no significant enrichment of the γH2A.X signal was detected when GFP–PML was tethered to the pericentric lacO arrays (Fig. 7A, right-hand panel).

Second, we tested for non-replicative DNA synthesis with a 5-bromo-2'-deoxyuridine (BrdU) pulse labeling after transfection of the cells with GBP–LacI–RFP and GFP–PML, and subsequent staining with an anti-BrdU antibody. To differentiate the BrdU signal did not significantly differ from the background level (Fig. 7A, right-hand panel). Taken together, these results indicate that the de-novo-assembled nuclear bodies consist of an APB-like protein composition independent of the chromosomal site of assembly. However, these nuclear bodies have to be assembled at telomeres to induce DNA repair processes, as detected here by H2A.X phosphorylation and BrdU incorporation.

### De novo APBs induce telomere repeat extension

In order to directly evaluate changes in telomere repeat length associated with de novo APB formation, fluorescence in situ hybridization (FISH) with a Cy3 fluorescently tagged peptide nucleic acid (PNA) probe against the telomere repeat sequence was conducted (Fig. 8) (Jegou et al., 2009). Owing to the heterogeneity of telomere repeat length in ALT-positive cells, a substantial number of the chromosomal ends were too short to display a detectable telomere repeat PNA-Cy3 signal. The fraction of these telomeres was determined at several time points after transfecting F6B2 cells with GBP–LacI, and either GFP–PML or a GFP-only control. The recruitment of GFP–PML to the telomeric lacO arrays led to an increase of the detectable TTA(G)3 signal at these sites that increased over time from 57±7% (12 hours) to 81±9% (96 hours after transfection). These experiments, a telomere signal was counted if it comprised >0.025% of the total PNA-Cy3 intensity in a given nucleus (Fig. 8B, left-hand panel). The telomere repeats were also examined at the pericentric lacO arrays. Notably, there was no significant change of the telomeric repeat signal when GFP–PML was targeted to the telomeres (Fig. 8B, right-hand panel). This suggests that the observed increase of the TTA(G)3 signal at the lacO-labeled telomeres can be, indeed, attributed to an extension of the telomere repeats as opposed to a PML-mediated association with another telomere repeat sequence. Next, a quantitative analysis of the TTA(G)3 signal intensity distribution was conducted. This revealed the appearance of a fraction of telomeres (~20%) with an increased normalized repeat length of 3.4±0.8% when GFP–PML was targeted to the telomeres (Fig. 8C). Finally, the images were inspected to determine whether the increase of telomere repeat signal at the de-novo-formed APBs was due to an induction of clustering of two or more telomeres. Within the resolution of the CLSM images, only two out of 604 (i.e. 0.3%) of the complexes showed a telomere intensity signal distribution indicative of the association of two telomeres.  

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### Table 1. APB de novo assembly and recruitment of endogenous proteins

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<th>Protein</th>
<th>Initiation of APB formation by GFP fusion protein*</th>
<th>Recruitment of endogenous protein to de novo APBs†</th>
<th>Initiation of APB-like structure assembly by GFP fusion protein*</th>
<th>Recruitment of endogenous protein to APB-like compartment†</th>
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<td>Rad51</td>
<td>+</td>
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<td>Rad17</td>
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<td>γH2A.X</td>
<td>n.d.</td>
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<td>GFP</td>
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The measured degree of colocalization at the lacO loci was: +++ (>80%; +++, >50%; +, >20%; –, no significant enrichment over background (P>0.05); n.d., not determined; wt, wild type.

*The indicated GFP fusion proteins were bound to the lacO arrays through GBP–LacI–RFP. APB formation was evaluated by immunostaining for endogenous PML at these sites, except for PML where endogenous Sp100 was measured.

†The de novo APB formation was induced by tethering GFP–PML (except for PML when GFP–Sp100 was used). Recruitment of endogenous proteins colocalizing with the GBP–LacI–RFP signal was detected by immunofluorescence.

‡The same antibody was used for detecting both the endogenous isoforms SUMO2 and SUMO3 simultaneously.
Discussion

Here, we have investigated the assembly mechanism of APBs and their function in the ALT pathway by recruiting protein components of APBs to lacO-tagged telomeres in U2OS cells (Fig. 1). As described previously, the structure of the PML-NB component of APBs is determined by PML and Sp100 proteins, in conjunction with their sumoylation, to mediate the non-covalent binding of the two proteins through their SIMs (Bernardi and Pandolfi, 2007; Lang et al., 2010; Shen et al., 2006).

The shelterin components TRF1 and TRF2 were highly capable of inducing the formation of de novo APBs after enrichment at the telomeric lacO arrays (Fig. 4A,B). This is consistent with their requirement for APB formation in previous reports (Jiang et al., 2007). The results obtained here suggest that the amount of TRF1 and/or TRF2 accessible for protein–protein interactions or post-translational modifications, particularly sumoylation, can be a limiting factor for APB assembly at endogenous telomeres. We note that the recruitment of TRF1 and TRF2 to the lacO arrays also allowed us to target very short telomeres, which presumably lack parts of the shelterin complex. Enrichment of TRF1 and/or TRF2 at these telomeres could provide the required additional interaction surface for APB formation. Surprisingly, TRF2 was somewhat more efficient than TRF1 in recruiting endogenous PML, although a direct interaction between TRF1 and PML IV in the context of APB formation has been reported recently (Yu et al., 2009). However, the antibody used in the present study recognizes all PML isoforms so that a specific recruitment of PML IV might not be detected in our assay.

With our experimental system we were able to dissect the role of the three different paralogues, SUMO1, SUMO2 and SUMO3 (Fig. 3). Intriguingly, a non-conjugable SUMO1 mutant was found to be highly efficient in triggering the assembly of APB proteins, whereas mutated SUMO2 and SUMO3, which could not be conjugated to a target protein, showed only a moderate propensity to initiate this process. We speculate that the modification of telomeric proteins with SUMO1 (as mimicked in our experiments by the tethering of a non-conjugable SUMO1 mutant or the MMS21 SUMO E3 ligase) would be sufficient to initiate the formation of an APB through recruitment of SIM-containing APB components. This conclusion is further corroborated by our findings that SIM interactions of non-conjugable SUMO1 are crucial for efficient APB nucleation, in line with previous reports (Bernardi and Pandolfi, 2007; Shen et al., 2006). Moreover, our recent high-resolution three-dimensional analysis of PML-NBs revealed that the SUMO1 modification is localized preferentially in the spherical shell of PML and Sp100 protein, whereas the SUMO2 and/or SUMO3 modification was found also in the interior of PML-NBs (Lang et al., 2010). This finding points to
functional differences between the SUMO isoforms as observed here, too. Thus, we propose that PML binds SUMO1 directly through its SIM, whereas SUMO2 and SUMO3 are more weakly or indirectly bound by the main PML-NB components.

The SUMO1 modification of target proteins, as an initiating factor for APB formation, could be mediated by the MMS21 SUMO E3 ligase. This enzyme is responsible for sumoylation of the shelterin components TRF1, TRF2 and Rap1 in ALT cells.

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**Fig. 6.** See next page for legend.
and can auto-sumoylate itself (Andrews et al., 2005; Potts and Yu, 2005; Zhao and Blobel, 2005). In support of this model, recruitment of MMS21 was found to initiate APB formation as efficiently as the SUMO1 domain (Fig. 3A-D; Fig. 6C), indicating that MMS21 promotes APB assembly through a recruitment (indicated by arrows in D1, magnifications are shown in D2 and D3), whereas endogenous Rad9 was found only at a fraction of these sites (filled arrows in D1, a magnification is shown in D2). (E) Upper panel: Quantification of PML and Rad9 colocalization after GFP–MMS21 tethering (n=182 lacO-tagged telomeres). Lower panel: Analysis of endogenous APBs, identified by transfection of GFP–TRF2, in U2OS cells revealed that 16 ± 1% of telomeres colocalized with PML and 15 ± 1% with Rad9. A total of 2.0 ± 0.3% telomeres were associated with only PML, and 0.9 ± 0.2% with only Rad9 (n=1722 telomeres). Note the different scale of the y-axis. Scale bars: 10 µm (A–C,D1); 0.5 µm (D2,D3). All error bars show the s.d.

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have been found to be required for recruiting the DNA repair factors Mre11, Rad50 and Brc1 to APBs (Naka et al., 2002; Wu et al., 2003; Zhu et al., 2000). This process seems to be tightly regulated in the endogenous environment as we observed a higher enrichment of endogenous NBS1 after recruiting GFP–PML to a pericentric site, which might lack specific inhibitory mechanisms. In addition to its DNA repair and recombination activity, NBS1 could target PML-NB assembly to certain telomeres at which it is enriched. This might lead to a DSB-repair-mediated elongation at these sites. Furthermore, the strong accumulation of PML protein after tethering of NBS1 to the pericentric lacO array supports an ALT independent relationship between DSB signaling and PML-NB formation, as suggested previously (Dellaire and Bazett-Jones, 2004; Dellaire and Bazett-Jones, 2007). Thus, the persistent deposition of NBS1 at a chromatin locus might mimic a DNA DSB situation (Soutoglou and Misteli, 2008).

The homologous recombination (HR) factor Rad51 plays an important role in HR-mediated DSB repair as it forms nucleoprotein filaments on single-stranded DNA to promote the pairing of homologous strands and strand exchange. It was one of the first recombination factors to be described as an APB component in ALT cells (Yeager et al., 1999). Interestingly, recruitment of this factor to the telomeric lacO arrays promoted APB formation only weakly (Fig. 4D). This is consistent with a previous report showing that siRNA-mediated knockdown of Rad51 in U2OS cells does not lead to a disruption of APBs (Potts and Yu, 2007). With respect to the assembly mechanism, this suggests a classification of APB proteins into those that are capable of initiating the assembly and others that are only recruited subsequently. This model also accounts for the results obtained upon testing the repair factors Rad9 and Rad17. For these proteins, the accumulation of endogenous proteins in the de-novo-formed APBs was higher (Fig. 5B,C) than the increase in the level of endogenous PML proteins when GFP–Rad9 and GFP–Rad17 were enriched at the telomeres (Fig. 4E,F). Thus, both Rad9 and Rad17 are less efficient as initiation factors for APBs but readily assemble at these sites once these complexes are formed. This view is supported by our finding that Rad9 binding to telomeres correlated with the presence of PML but not vice versa (Fig. 6). We conclude that during APB formation, the assembly of structural nuclear body core components PML and Sp100 precedes the subsequent binding of DNA repair and recombination factors according to the mechanism depicted in Fig. 9. The comparison of de-novo-assembled nuclear bodies after recruitment of GFP–PML to a pericentric lacO array revealed that the essential features of the assembly process were independent of the chromosomal site. SUMO1, SUMO2 and SUMO3, NBS1, Rad9, Rad17 and MMS21 were enriched at both the telomeric and the pericentric sites to similar levels. This points to a self-organization process for the genome-associated structural PML-NB subcompartment through stochastic interactions of the constituting components, as opposed to a...
defined sequential order of binding events (Fig. 9) (Dinant et al., 2009; Hancock, 2004; Hebert and Matera, 2000; Matera et al., 2009; Misteli, 2007; Wachsmuth et al., 2008). This aspect of APB formation is very similar to that reported for Cajal nuclear bodies (Kaiser et al., 2008). It is supported by experiments on the dissociation and reassembly of PML-NBs through varying the degree of molecular crowding in their environment (Hancock, 2004). In addition to previous findings on the self-organizing properties of nuclear bodies, we propose that the APB protein interaction network is stabilized by a feedback and propagation mechanism that comprises: (1) the MMS21 SUMO E3 ligase and possibly other E3 ligases; (2) the post-translational sumoylation of PML, Sp100, telomeric proteins TRF1, TRF2 and Rap1, and MMS21 itself; and (3) the SUMO-interacting domains of PML and Sp100 (Fig. 9). In contrast to the initiating proteins, other APB proteins, such as Rad9, Rad17 and Rad51, are incorporated later, in conjunction with the phosphorylation of the H2A.X histone variant. Thus, a preassembled subset of APB components is required for the subsequent binding of other factors. This feature of the sequential assembly mechanism has been reported previously for the recruitment of Sp100 and Daxx to early G1 PML-NBs and the prior binding of the MRN complex to telomeres before APB assembly (Chen et al., 2008; Jiang et al., 2007).

The APBs assembled here by recruiting essential structural components of APBs, such as PML, Sp100 and SUMO1, to the telomere-associated lacO arrays were indistinguishable from their endogenous counterparts with respect to protein composition and structural organization according to all criteria reported previously. This allowed us to address the question of whether this nuclear subcompartment has an essential function within the ALT pathway. To this end, we evaluated the presence of the phosphorylated H2A.X histone variant, as a molecular marker for DSB repair, as well as non-replicative DNA synthesis through the incorporation of BrdU into the DNA (Fig. 7). We found that the de-novo-assembled APBs were positive for these two hallmarks of DNA repair. Presumably these activities are coupled to the DNA damage response pathway, as previous work has shown that BrdU incorporation in APBs is dependent on the phosphoinositide 3-kinase (PI3K)-like kinases ATR and ATM (Nabetani et al., 2004). Finally, we showed by quantitative FISH that the recruitment of GFP–PML to the telomeric lacO arrays led to an increase in the telomere repeat length at these sites (Fig. 8). A fraction of the de-novo-formed APBs (10–15%) was competent in inducing telomere extension during an ~24 hour time period. This number is consistent with the result that not all of the de novo APBs contained the complete set of DNA repair and recombination factors investigated here (Fig. 5; Table 1). A longer incubation of the cells further increased the percentage of functional APBs so that ~30% of the APBs had telomere extension activity after 96 hours (Fig. 8B, left panel). As discussed above, recruitment of PML to a non-telomeric site led to the formation of a nuclear body that contained all of the tested APB proteins. However, this nuclear subcompartment was non-functional with respect to H2A.X phosphorylation and the non-replicative synthesis of telomeric DNA as there was no significant difference in BrdU incorporation and the detected telomere repeat signal (Fig. 7; Fig. 8B right panel).

Previously, a study of artificially enlarged telomere–PML-NB complexes, which form upon transfection of ALT cells with a mutated form of the herpesvirus ICP0 protein, showed that these APB structures promote the association of multiple telomeres (Draskovic et al., 2009). Our structural analysis of de-novo-assembled and endogenous APBs in U2OS cells by
conventional CLSM and high-resolution 4Pi microscopy, here and elsewhere, revealed a cap-like structure of PML protein formed around a single telomere signal in U2OS cells (Fig. 1B; supplementary material Fig. S1) (Jegou et al., 2009; Lang et al., 2010). Two separate telomere signals were only distinguishable for a small fraction of APBs (0.1-0.3%). Thus, under our experimental conditions, we did not find evidence that telomere clustering could explain the observed increase of the telomere repeat signal at the de novo-formed APBs. Furthermore, although all APB marker proteins were present at the pericentric lacO arrays after PML recruitment, no additional binding of telomeres or extrachromosomal telomeric repeat (ECTR) DNA was detected. Thus, an association of ECTRs with PML and TRF2 protein in APBs, proposed previously on the basis of sucrose gradient fractionation of nuclear components after DNA damage induction and sonication, was not apparent in our system (Fasching et al., 2007). The de novo assembly of PML protein at the lacO-tagged telomeres induced the binding of other APB components and led to H2AX phosphorylation, BrdU incorporation and an increase in the telomere repeat DNA signal. Accordingly, we conclude that the formation of bona fide APBs promotes the extension of the telomere repeat sequence by a DNA-repair-coupled synthesis process. Our study does not provide information on the nature of the telomere repeat template used for synthesis, i.e. whether it is intra- or inter-chromosomal or an APB-associated ECTR. Furthermore, given the large number of partially contradicting results in the literature, it is well conceivable that different telomerase-independent mechanisms for telomere repeat extension exist. In addition, certain cellular conditions could lead to the formation of telomeric PML-NBs that are incapable of promoting telomere extension. For example, APB-like colocalizations of PML-NB and telomeres have been detected in telomerase-negative human cancer cell lines upon exogenous expression of the telomerase RNA component, but other characteristics of the ALT pathway were missing (Pickett et al., 2009). Likewise, a human cell line that maintains telomeres in the absence of telomerase but without the formation of APBs has been described (Cerone et al., 2005). Thus, it will be important to further dissect the exact combination of protein factors that are sufficient to trigger telomere extension in APBs and to investigate the effects of their presence or absence. We anticipate that the experimental approach introduced here, in conjunction with RNA interference (RNAi)-based three-dimensional confocal microscopy high-content screening for genes involved in APB formation (Osterwald et al., 2011), will provide a valuable approach for subsequent studies. It will allow us to precisely identify all protein components that are sufficient to form a telomeric PML-NB subcompartment structure, as well as the additional factors needed to induce telomere extension at these sites. This will serve to select protein targets for inhibiting telomere extension, and thus cell proliferation, in tumors that make use of the ALT pathway.

Materials and Methods

Protein constructs

The cDNAs encoding TRF1, SUMO1, SUMO2, SUMO3, MMS21, Rad51, Rad9 and Rad17 were obtained from the DKFZ Genomics and Proteomics Core Facility and cloned into pcDNA-DEST53 (N-terminal GFP-tag) and pcDNA-DEST47 (C-terminal GFP tag) expression vectors (Invitrogen). Constructs for GFP-PML III and GFP–TRF2 were as described previously (Jegou et al., 2009). The other constructs were kindly provided as indicated: GFP–PML IV [Peter Hemminger, FLI Jena, Germany (Weidtkamp-Peters et al., 2008)], GFP–CenpA [Stephan Diekmann, FLI Jena, Germany (Hemmerich et al., 2008)], GFP–Sp100 and NBS1–2GFP (Thomas Hofmann, DKFZ, Heidelberg, Germany), pEYFP-SUMO1AC (Frauke Melchior, ZMBH Heidelberg, Germany). The non-SIM-interacting mutant pEYFP-SUMO1AC7(-) was created by site-directed mutagenesis to convert Val38 and Lys39 into alanine residues. Non-conjugable GFP–SUMO2AC4 and GFP–SUMO3AC13 constructs were created from the corresponding pcDNA-DEST53-SUMO2/3 vectors by site-directed mutagenesis. The first glycine codon of the C-terminal Gly-Gly motif with a stop codon. The fluorescence three-hybrid system for recruiting GFP-tagged proteins to lacO arrays through GBP–LacI and GBP–LacI–RFP was provided by Chromotek (Munich, Germany).

Cell culture work, immunostaining and PNA FISH

The U2OS cell clones F6B2 and F42B8 were cultured and transfected as described previously (Jegou et al., 2009). Cells were typically fixed 24 hours after transfection with 4% paraformaldehyde in PBS buffer. For the analysis by immunostaining, cells were washed and permeabilized for 5 minutes with ice-cold 0.1% (v/v) Triton X-100 solution in PBS. After three PBS washes, cells were blocked for at least 15 minutes with 10% goat serum in PBS, the solution was removed, and the cells were incubated with appropriate dilutions of specific antibodies against γH2A.X (1:100, rabbit, Millipore), NBS1 (1:200, NB100-143, Novus Biologicals), PML (1:150, PG-M3, Santa Cruz Biotechnology), Rad9 (1:100, M-389, Santa Cruz Biotechnology), Rad17 (1:200, H-300, Santa Cruz Biotechnology), Sp100 (1:200, AB1138, Chemicon), SUMO1 (1:100, FL-101, Santa Cruz Biotechnology) or SUMO2/3 (1:200, rabbit, Abcam). For immunofluorescence of MMS21, cells were fixed with 1% paraformaldehyde, and permeabilization and blocking was conducted in 0.2% (v/v) Triton X-100 and 3% BSA in PBS for 20 minutes, and the antibody was incubated in the same buffer (1:500, Abnova, NSMCE2 MaxPab, B01). For 5-bromo-2’-deoxyuridine (BrdU) staining, cells were seeded, transfected and incubated for 1 or 2 days. Then 100 μM BrdU (Sigma-Aldrich) was added to the medium for 2 to 4 hours, the cells were fixed, permeabilized with 0.2% (v/v) Triton X-100 in PBS, denatured with 1.5 M HCl for 30 minutes and then stained with an antibody against BrdU (1:50, B44, BD Biosciences). After incubation with primary antibodies the coverslips were washed with PBS containing 0.002% (v/v) NP40. The appropriate secondary antibodies conjugated to Alexa Fluor 488 or Alexa Fluor 633 (Molecular Probes) were diluted according to the manufacturer’s instructions in PBS, applied to the cells and incubated for 30-60 minutes. After another PBS wash, the coverslips were mounted with Vectashield (Vector Laboratories) or Prolong Gold antifade reagent (Molecular Probes) both containing DAPI. For telomere PNA FISH, cells were grown on a slide or coverslip, transfected, incubated for the indicated time, washed with PBS and fixed with 4% paraformaldehyde. After permeabilization with 0.2% (v/v) Triton X-100 in PBS, cells were dehydrated by a series of ethanol washes (70%, 85% and 100% ethanol), air-dried and a Cy3-labeled (CCCTAA)3 PNA probe (Dako, Glostrup, Denmark) was added. Then, samples were denatured at 80°C for 3 minutes and hybridization was conducted for at least 3 hours at 30°C. Slides were then washed consecutively with 70% formamide in 10 mM Tris-HCl pH 7.4, 2 × SSC, 0.1 × SSC at 55°C and 0.25% (v/v) Tween-20 in 2 × SSC. In order to enhance the GFP signal, immunofluorescence was conducted as described above using an antibody against GFP (1:500, ab290, Abcam). FISH experiments on metaphase chromosomes were conducted as described before using 200 ng of a Cy3-labeled oligonucleotide probe hybridizing to the lacO sequence (Jegou et al., 2009).

Confocal fluorescence microscopy, image analysis and statistical evaluation

Fluorescence images were acquired with different Leica TCS SP5 confocal laser scanning microscopes. Optical sections with spacing of 0.3 μm along the z-axis were recorded. Fluorescence intensities in the different color channels were analyzed on the individual z-slices. Cells with appropriate expression levels of the fluorescent cells were chosen. Spots were counted as colocalizing if the signal at the telomere was at least twofold above the background and comprised at least two pixels with a size of 200 nm. The percentage of lacO arrays with colocalization was determined with the indicated value giving the number of lacO arrays evaluated. Error bars were calculated as ±1σ, which yields the standard deviation for a Poisson distribution. Data obtained from the image analysis represent averages from at least three independent experiments. In the figures, maximum intensity projections of the image stacks are shown. In order to determine whether the percentages of colocalization after recruiting the proteins of interest were significantly different from the ones obtained in the controls, the Fisher’s exact test was used to calculate P-values.

Western blotting

A total of 5 × 10^6 F6B2 cells were seeded and transfected with GFP–SUMO3–GFP, incubated for 24 hours, washed with PBS, incubated with ice-cold RIPA buffer for 30 minutes at 4°C and centrifuged at 4°C. The supernatant was subjected to SDS-PAGE (12% gels) and, after blocking with 3% BSA in PBS, subjected to western blot analysis with an antibody against GFP (ab290, Abcam) according to the manufacturer’s protocol.
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