PML nuclear body biogenesis and oligomerization-driven leukemogenesis

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Abstract
PML nuclear bodies (NBs), which are increasingly recognized as the central hub of many cellular signaling events, are superassembled spherical complexes with diameters of 0.1–2 μm. Recent studies reveal that RING tetramerization and B1-box polymerization are key factors to the overall PML NBs assembly. The productive RBCC oligomerization allows subsequent PML biogenesis steps, including the PML auto-sumoylation and partners recruitment via SUMO–SIM interactions. In promyelocytic leukemia, the oncoprotein PML/RARα (P/R) inhibits PML NBs assembly and leads to a full-blown leukemogenesis. In this review, we review the recent progress in PML and acute promyelocytic leukemia fields, highlighting the protein oligomerization as an important direction of future targeted therapy.

Keywords: Leukemogenesis, Oligomerization, PML nuclear body, PML/RARα

1. INTRODUCTION
The PML protein (also called TRIM19) is first observed in the 1960s.1,2 PML gene is 53 kb in length and contains nine exons. Through alternatively splicing, at least seven PML isoforms can be observed.3 All of them share a conserved RBCC motif in N-terminus, in which R stands for RING, B for B1- and B2-box, and CC for coiled coil domain. It has been shown that these domains are indispensable for PML nuclear bodies (NBs) biogenesis.4–9 The C-terminus contributes to the specific functions of each PML isoform. For instance, the C-terminal portion of PML-IV (amid 361–633) is critical in p53 regulation.10 In comparison, the C-terminal domains of PML-II and PML-V may contribute to PML-NBs formation.11,12 The NBs-facilitated posttranslational modification is often recognized as the main function of this subcellular complex.13–17 In particular, PML sumoylation is highlighted in acute promyelocytic leukemia (APL) development and targeted therapy. Until today, three sumoylation sites (i.e. K65, K160, K490) are reported in PML. The PML auto-sumoylation and the sumo interaction motif (SIM) in its C-terminus can also contribute to the overall NBs assembly.18,19 In this review we will look into the current understanding of PML NBs biogenesis pathway and their implication in APL pathogenesis and targeted therapy.

1.1. RBCC oligomerization is essential to PML NBs assembly
In the process of PML NBs assembly, the integrity of PML RBCC motif is thought essential.3,7–9 Amongst RBCC domain, the RING tetramerization might precede B1-driven oligomerization.9 Structural-dependent mutations on RING tetramerization and B1-box oligomerization abolish PML NBs formation.8,9 Many TRIMs could facilitate RING-binding E2 enzymes to their nascent substrates, and hence are often recognized as ubiquitin E3 ligases.20 TRIM RING dimerization is critical for ubiquitin conjugating E2 UBC9 (Fig. 1), prompting the hypothesis that high-order assembly of RING domain might be the structural determinants that enable PML/TRIM19 into a sumoylation, but not ubiquitin ligase, E3.3 This is further supported by the recent observation of PML-B1-box polymerization (n > 4).7 Li and coworkers have shown that PML B1-box could form dimer, tetramer, and much higher-order polymer via oligomeric sequence uniquely observed in PML but not other TRIMs (Fig. 1). Indeed, the B1-box oligomerization gives rise to an unexpected K160 lining/concentration that might explain why this lysine position is particularly prone to SUMO long chain. In previous studies,21 it is clear that CC domain can also contribute to RBCC oligomerization. In the recent study,7 via arsenic rescue experiment, a cooperative mechanism has been proposed among RING, B1/2-box, and CC domains. Future study should focus on the mechanism of how RING, B1/2-box, and CC might fold together, leading to the 2D–3D transition and ultimately to PML speckles formation.

1.2. PML NBs PTMs are involved in PML NBs assembly
PML often participates in various cellular signaling via posttranslational modification of itself and partners. Its best-known PTM function is sumoylation. PML could directly
conjugate SUMOs via lysine residues, which endow their ability to interact with partner proteins equipped with a short hydrophobic sequence termed as SIM. Until today, three PML sumoylated sites, that is, K65/K160/K490, are reported. Furthermore, PML and its partners both contain SIM domain. The SUMO–SIM interaction might help the partner proteins sequestration within the NBs. Until today, it has been reported that PML could directly or indirectly interact with at least 120 proteins, including p53, DAXX, SP100, CBP, HDAC, and so on. Of note, the K160 site, which is not critical for NBs formation, is essential for partners recruitment. Furthermore, as demonstrated in P/R transgenic mice, K160 is critical for PML/RARα sumoylation and APL leukemogenesis. In addition to sumoylation, other PTM functions, including phosphorylation and acetylation, are also reported in PML. Like sumoylation, PML phosphorylation is catalyzed by kinases such as ATR, ATM, and CHK2 to assist in DNA damage process. PML acetylation is also reported to happen in the positions of K487 and K505 by the protein acetyltransferase involved in cell apoptosis and cell death regulation. Some anchored PML partners localized in the NBs core could also be modulated by other PTMs (Fig. 1). One of the best-known partners is the tumor suppressor p53. Therefore, PML is often recognized as a critical regulator of p53 activity and p53-mediated cellular processes, such as apoptosis, cell cycle arrest, and DNA repair and senescence, especially through the sequestration of

**Figure 1.** A model for PML nuclear bodies assembly. RBCC-mediated PML oligomerization constitutes the “shell” of PML NBs to recruit UBC9, and hence facilitates PML sumoylation. Sumoylated PML might allow the interaction of partner proteins that contain sumo-interaction-motif (SIM). All these partners’ recruitments ultimately enable the in situ sumoylation and other posttranslational modifications (PTMs).

- RING
- B1
- B2
- CC
- UBC9
- SUMOs
- SUMO sites
- Partner
- Acetyl
- Phosphate

**Figure 2.** The scheme of PML/RARα-driven APL pathogenesis. The oncoprotein PML/RARα drives disruption of PML nuclear bodies while P/R-multimer exerts target genes in deregulation and recruits DBF (RXR, PU.1 and many others) to form hetero-multimers, which in turn might enhance CoR recruitment, leading to abnormal cell arrest.
Mdm2-dependent PTM of p53 and SIRT1-dependent deacetylation of p53.30,31

1.3. Enlightening in oligomerization and tumorigenesis

APL belongs to the M3 subtype of acute myeloid leukemia. The generation of PML/RARα (P/R) oncoprotein and PML NBs abolishment trigger APL pathogenesis.12,33 Primarily, the abnormal P/R expression prompts NBs disruption, leading to nuclear microspeckles formation and losing its pivot properties involved in tumor-suppressive processes, including DNA damage response, cellular apoptosis, senescence, and angiogenesis.25 In recent report, the B1-box-mediated P/R oligomerization is identified as a significant regulator in P/R-driven transactivation and leukemogenesis.9 Supportively, the PML RING–RING and CC–CC interactions are also a significant factor for differentiation arrest and transformation in vivo.8,9,34 All these results highlight the PML oligomerization as an important regulator in leukemogenesis. Furthermore, transcriptional repression due to DAXX recruitment of PML/RARα in a strictly K160-dependent manner and DNA binding factors (DBF)/Cor recruitment by RARα portion are recognized as key contributors in APL development.27,33 PML/RARα could recognize >3000 DNA binding sites via many DBE such as RXR and/or PU.1.13,36 The PML/RARα–DBF complex, which is regulated by multimerization, might cooperate with many corepressor complexes (CoR) such as histone deacetylases (HDACs), polycomb repressive complexes (PRCs), and DNA methyltransferases to exert oligomerization-driven transcriptional repression37–40 (Fig. 2).

Oligomerization-driven tumorigenesis is widely observed in other leukemias such as acute myeloid leukemia (AML), mixed-lineage leukemia (MLL), and acute lymphoblastic leukemia (ALL).41–45 In AML leukemia, the Nervy homology 2 (NHR2) domain in AML1/ETO is shown to form an alpha-helical tetramer. More importantly, NHR2 oligomerization is essential to AML1/ETO’s oncogenic activity, that is, the inhibition of granulocytopoiesis differentiation.42 In ALL leukemia, it has been reported that the diverse oncoproteins, including MLL/GAS7, MLL/AF1P, MLL/GEPRIN, could form oligomerization via coiled coil domain. The disruption of MLL oligomerization precludes leukemogenic transformation.45 In ALL leukemia, the PAX5–PML is shown to repress PAX5 transactivity in heterodimer complex.41 Finally, the tetramerization of stat5, which is frequently associated with various leukemias, is also important for leukemogenesis. The stat5 mutants that target the tetramerization fail to trigger leukemias, reiterating the importance of oligomerization in oncogenic drivers.46

2. CONCLUSION AND PERSPECTIVES

RBCC-mediated oligomerization not only contributes to PML NBs assembly as a premise of the following PML sumoylation and partners recruitment via SUMO–SIM interaction but is also instrumental to APL pathogenesis. PML NBs disruption and the oligomerization of oncoproteins involved in various leukemias suggest that oligomerization could be recognized as an important tumorigenic regulator and valuable direction for future targeted intervention.

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