Promyelocytic Leukemia Protein (PML) and Daxx Participate in a Novel Nuclear Pathway for Apoptosis

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Abstract

The promyelocytic leukemia protein (PML) gene of acute promyelocytic leukemia (APL) encodes a cell growth and tumor suppressor essential for multiple apoptotic signals. Daxx was identified as a molecule important for the cytoplasmic transduction of the Fas proapoptotic stimulus. Here, we show that upon mitogenic activation of mature splenic lymphocytes, Daxx is dramatically upregulated and accumulates in the PML nuclear body (NB) where PML and Daxx physically interact. In the absence of PML, Daxx acquires a dispersed nuclear pattern, and activation-induced cell death of splenocytes is profoundly impaired. PML inactivation results in the complete abrogation of the Daxx proapoptotic ability. In APL cells, Daxx is delocalized from the NB. Upon retinoic acid treatment, which induces disease remission in APL, Daxx relocalizes to the PML N Bs. These results indicate that PML and Daxx cooperate in a novel NB-dependent pathway for apoptosis and shed new light in the role of PML in tumor suppression.

Key words: apoptosis • PML • Daxx • nuclear body • acute promyelocytic leukemia

Introduction

The promyelocytic leukemia protein (PML) gene is involved in reciprocal chromosomal translocations with the retinoic acid receptor α (RAR α) locus that are specifically found in almost 100% of cases of acute promyelocytic leukemia (APL), a distinct subtype of myeloid leukemia (1–6). This translocation leads to the production of a PML-RAR α chimeric oncoprotein, which is thought to interfere with both PML and RAR/RX receptor X (RXR) pathways (5, 6).

PML is a RING finger IFN-inducible protein typically concentrated within discrete speckled nuclear structures called PML nuclear bodies (N Bs) (5–9). PML acts as a cell growth and tumor suppressor, antagonizing initiation, promotion, and progression of tumors of various histological origins and, particularly, of the lympho-hemopoietic compartment (10). PML is essential for the ability of Fas, TNF-α, ceramide, and IFN to induce programmed cell death and is required for DNA damage-induced apoptosis in mitogen-activated splenic lymphocytes (11).

In APL cells, the presence of PML-RAR α leads to the delocalization of PML from the N Bs and its accumulation into aberrant nuclear microspeckled structures where PML, RXR, and the PML-RAR α oncoprotein colocalize. Upon RA treatment, which induces the degradation of the oncoprotein, the terminal differentiation of the APL blasts and the complete albeit transient remission of the disease, PML and the other NB proteins reacquire their natural nuclear localization (5, 9, 12–14). Thus, the deregulation of the PML NB pathway by PML-RAR α may play a crucial role in tumorigenesis.

In agreement with the critical role of PML in apoptosis, PML-RAR α has a marked antiapoptotic function, and this activity depends, at least in part, on its ability to deregulate PML function (11, 15).

Daxx was originally identified as an adaptor molecule that binds to the death domain of the Fas receptor (16). Daxx has also been found to localize in the nucleus (17, 18).
However, its nuclear function is unknown. Daxx acts as a proapoptotic molecule that can enhance Fas-mediated apoptosis, as well as the Fas-mediated activation of the c-Jun N H\textsubscript{2}-terminal kinase (JNK) pathway through its interaction with the apoptosis signal-regulating kinase 1 (ASK1) (19, 20). Targeted disruption of the Daxx gene in the mouse results in early embryonic lethality, defining an important developmental role for this protein (18).

Here, we show that PML and Daxx participate in a novel nuclear pathway for lymphocyte apoptosis. PML is essential for both the proper localization of Daxx into the NBs and its proapoptotic ability. The deregulation of this pathway results in impaired lymphocyte activation-induced cell death providing a further basis for the increased susceptibility to lymphoma observed in PML-/- mice. We also find that in APL cells, Daxx is delocalized from the NB or 100-mm plates 24 h before transfection. Transfections with reporter under the control of a Gal4-responsive upstream activating sequence (UAS), and colonies were grown on synthetic dropout (SD)-Trp, SD-Leu, or SD-Trp/-Leu/-His plates. After 3 d, the primary keratinocytes and splenocytes were prepared as described previously (11, 21). N B4 cells (American Type Culture Collection) were grown in RPMI 1640 medium with 10% FBS. Passage 1 keratinocytes and Cos-1 cells were transferred into 6-well dishes or 100-mm plates 24 h before transfection. Transfections with plasmid DNA were carried out using the SuperFect™ reagent (Qiagen) following the manufacturer’s instructions.

Expression Vectors. Full-length Daxx cDNA was obtained by reverse transcription PCR from mouse testis total RNA, and subcloned into pcDNA3.1 vector (Invitrogen). Mouse Daxx and human PM L cDNAs were also subcloned in pEGFP (green fluorescent protein [GFP]) and pEBFP (blue fluorescent protein [BFP]) vectors (Clontech), respectively. Mouse Daxx was also subcloned in pcDNA3.1 together with a 6×His tag, His(6×)-LacZ, pcDNA3.1 was purchased from Invitrogen.

Immunoprecipitation. After transfection with plasmid DNA, Cos-1 cells were washed in PBS, and cell lysates were prepared by adding 1 ml of ice-cold E1A buffer (250 mM NaCl, 50 mM Hepes pH 7.0, 0.1% NP-40, 5 mM EDTA) supplemented with a complete protease inhibitor cocktail (Boehringer Mannheim). Either a mouse anti-PML mAb (1:50-1:100 dilution, sc-966; Santa Cruz Biotechnology), a mouse anti-Daxx mAb (1:400 dilution, sc-8043; Santa Cruz Biotechnology), or a rabbit polyclonal anti-Daxx antibody (1:400 dilution, sc-7152; Santa Cruz Biotechnology) was added to 50 μl of lysate, and the solution was incubated at 4°C overnight. For the endogenous immunoprecipitation, mouse primary splenocytes were Con A activated as above for 3 d and washed in PBS. Cell lysates were prepared by adding 1 ml of ice-cold E1A buffer. Either the rabbit polyclonal anti-PML (1:500) or the mouse anti-Daxx mAb (1:400) was added to the lysate, and the solution was incubated at 4°C overnight. Protein A-agarose beads (Sigma Chemical Co.) were added and incubated for 1 h with gentle rocking before pelleting by centrifugation. The beads were washed three times in E1A buffer and twice with PBS, and the immune complexes were released from the protein A-agarose beads by boiling in sample buffer for 5 min. Cell lysates and immunoprecipitation products were analyzed by Western blot.

Preparation of Subcellular Fractions. Subcellular fractionation was performed as described (22). In brief, cells were lysed in hypotonic buffer (5 mM Tris pH 7.4, 5 mM KCl, 1.5 mM MgCl\textsubscript{2}, 0.1 mM EGTA [pH 8.0], 1 mM dithiothreitol, 10 μg/ml aprotinin, 10 μg/ml leupeptin, 10 mM benzamidine, 1 mM PM SF, 0.2 mM Na\textsubscript{3}VO\textsubscript{4}) at 4°C for 30 min. After homogenization (20 strokes with a Dounce homogenizer B pestle), samples were centrifuged (2,000 g for 5 min) to isolate the nuclei, which were lysed in E1A buffer. Nuclease N P-40-insoluble fractions were lysed directly in SDS-sample buffer. The resulting supernatant was considered as cytosolic fraction.

Yeast Two-Hybrid Assays. Yeast two-hybrid assays were performed using MATCHMAKER Two-Hybrid System 2 (Clontech). In brief, full-length Daxx and PML were subcloned into pAS2-1 or pACT-2 vector, pAS2-1-Daxx and/or pACT-2-PML were transformed into yeast strain Y187 containing a LacZ reporter under the control of a Gal4-responsive upstream activating sequence (UA5), and colonies were grown on synthetic dropout (SD)-Trp, SD-Leu, or SD-Trp/-Leu/-His plates. After 3 d, the colonies were inspected for blue color development following the manufacturer’s protocol. To quantitate the protein–protein interactions, β-galactosidase activity of yeast transformants (three to four clones per transformation) grown in liquid culture was determined using o-nitrophenyl β-d-galactopyranoside (ONPG) as substrate, as per the manufacturer’s instructions (Hoffmann-La Roche).

Purification of B Cells. B and T splenic subpopulations were stained with FITC-conjugated anti-B220 and anti-CD90 antibodies (PharMingen) for 15 min and then washed and incubated with magnetic beads conjugated with an anti-FITC antibody (Miltenyi Biotec) for 5–10 min. After two washes, cells were separated on LS+ MACS separation columns (Miltenyi Biotec). The purity of the various subpopulations was ~95% as assessed by flow cytometric analysis of cells stained with an FITC-conjugated anti-B220 and anti-CD90 antibodies (PharMingen).

Apoptosis Analysis. Primary keratinocytes and splenocytes were harvested and cytospun onto glass slides. Cells were fixed with 4% paraformaldehyde in PBS for 30 min at room temperature and permeabilized in 0.1% Triton X-100, 0.1% sodium citrate for 1 min at 4°C. After 1 h in blocking solution (PBS with
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10% goat serum) at room temperature, cells were incubated with an anti-His mAb (Penta, 1:200; Qiagen). Cells were then stained with a Texas red-conjugated anti–mouse antibody (1:200) diluted in the terminal deoxynucleotidyl transferase–mediated dUTP nick end labeling (TUNEL) reaction mixture (Roche) for 1 h at room temperature. Cells recognized by the anti-His antibody were scored for TUNEL positivity and nucleus condensation as apoptotic cells. In addition, apoptosis was scored by staining with propidium iodide and by evaluation of subdiploid DNA content by FACS® analysis.

Results

Daxx Localizes to Discrete Nuclear Regions upon Splenocyte Activation. We attempted to identify molecules that could interact with PML in transducing the apoptotic stimulus in mitogen-activated splenocytes. To this end, we searched for proteins that colocalize with PML in the NBs of splenocytes before and/or after Con A activation. Cytospins of splenocytes obtained from PML1/1 mice were prepared either immediately or after 3 d of culture in the presence of Con A. Subsequently, the cells were subjected to confocal microscopy, we observed that while at the steady-state Daxx was localized in the cytosol, where it showed a diffuse localization pattern, in Con A–activated splenocytes Daxx was mainly localized in discrete speckled nuclear regions (Fig. 1 A). The vast majority of the cells displayed a speckled nuclear pattern with ~5.5 ± 3.2 dots per cell (Fig. 1 A). Similar results were obtained using a polyclonal Daxx antibody (not shown).

We also observed that the immunofluorescence staining of Daxx in splenocytes became much stronger after Con A activation. To investigate whether this could be the result of upregulated Daxx expression after Con A treatment, we studied levels of Daxx expression in splenocytes before or after Con A activation by Northern and Western blot analyses (Fig. 1, B and C). After 72 h of Con A treatment, the mRNA level of Daxx was markedly upregulated, suggesting that Con A can induce Daxx expression at the transcription level (Fig. 1, B and C). In addition, we analyzed the protein levels of Daxx in the cytoplasmic and nuclear soluble and insoluble fractions. We observed that before Con A activation, Daxx was detected at very low levels and exclusively in the cytoplasm (Fig. 1 C). Localization of Daxx in the cytosol was also observed in Jurkat T lymphocytes and murine hepatocytes (our unpublished observations; and Weinberg, R.A., personal communication). By contrast, upon Con A activation Daxx was strongly induced and mainly found associated with both the nuclear soluble and nuclear insoluble fractions (Fig. 1 C). Thus, upon Con A treatment, Daxx is dramatically upregulated and translocated from the cytosol to the nucleus.

Figure 1. Con A activation of splenocytes induces Daxx expression. (A) Daxx localization in NBs is induced upon Con A activation. PML1/+ splenocytes, before and after Con A activation, were immunostained with anti-PML and anti-Daxx antibodies. Both Daxx and PML display a strong staining upon Con A activation. Single staining (a, b, a', and b') and superimposed images (c and c') are shown. The nuclei were visualized by DAPI staining (d and d'). Representative confocal pictures are shown. Bar, 10 μM. (B) Upregulation of Daxx mRNA expression upon Con A treatment. Northern blot analysis was performed on total RNA from PML1/+ primary splenocytes, before and after 72 h of Con A activation using a Daxx cDNA probe, which detected a single 2.6-kb band. Normalization with 28S RNA is also shown. (C) Analysis of Daxx protein levels in cytosolic (C) and nuclear (N)–soluble (SOL) and insoluble fractions (INSOL). Levels of heat shock protein 90 (Hsp90) were measured as a control of equal loading.
Daxx accumulates in the PML NBs. We next investigated whether the Daxx NBs would coincide with the PML NBs. To this end, we performed coimmunofluorescence analysis with the anti-Daxx mAb and polyclonal anti-PML antibodies (see Materials and Methods). We detected PML NBs in all cells before and after Con A activation (Fig. 1 A). However, although before Con A activation, the splenocytes displayed 4.5 ± 1.5 NBs per cell, after Con A the number of NBs increased up to twofold (9 ± 4 per cell) and were much brighter (Fig. 1 A). Coimmunofluorescence staining demonstrated a complete colocalization of the Daxx dots with PML NBs, whereas not all PML NBs contained Daxx (Fig. 1 A). Moreover, in our fractionation experiments, a significant amount of Daxx was found in the insoluble nuclear fraction (Fig. 1 C), in agreement with the notion that the PML NBs are tightly associated with the insoluble nuclear matrix (9). Thus, Daxx is localized within the NBs upon splenocyte activation.

To further confirm that Daxx and PML colocalize in the nucleus, we transiently transfected Cos-1 cells with vectors expressing Daxx-GFP and PML-BFP. We analyzed Daxx and PML localization in formaldehyde-fixed cells by confocal microscopy. Daxx was found only associated with discrete NBs that coincided with the PML-specific NBs. In the same field, we could also detect the presence of cells transfected with either PML or Daxx alone. Interestingly, in cells expressing only Daxx, its localization was both nuclear speckled and nuclear diffuse. This indeed suggested that in the presence of the overexpressed PML, the diffuse Daxx protein was recruited to the NBs (Fig. 2 A).
and PML colocalization in nuclear speckles was also observed in living cells by a phase-contrast fluorescent microscope as well as in Cos cells transiently transfected with untagged PML and Daxx, where immunofluorescence analysis using anti-PML and anti-Daxx antibodies was carried out along with 4,6-diamino-2-phenylindole (DAPI) staining (not shown).

PML and Daxx Physically Interact. To test whether Daxx and PML could physically interact, we cotransfected Cos-1 cells with untagged PML and Daxx expression vectors (pSG5-PML and pcDNA3.1-Daxx) and performed coimmunoprecipitation experiments (see Materials and Methods). Coprecipitated PML or Daxx was detected by Western blot with anti-Daxx or anti-PML antibodies, respectively (Fig. 2 B), thus indicating that PML and Daxx interact in vivo. The anti-human PML antibody we have used does not cross-react with the endogenous monkey PML. However, Cos-1 cells do express PML as revealed by Northern blot analysis (not shown).

We next investigated whether endogenous PML and Daxx would physically interact in Con A-stimulated primary splenocytes. Indeed, endogenous PML and Daxx proteins could be reciprocally coimmunoprecipitated in these cells (Fig. 2 C).

To further prove physical interaction between Daxx and PML, we used a yeast two-hybrid system. Full-length Daxx was linked to the Gal4 DNA binding domain in a pAS2-1 vector, and full-length PML was linked to the Gal4 activation domain in a pACT-2 vector (see Materials and Methods). In these assays, PML and Daxx were also

Figure 3. PML is required for the NB localization of Daxx. (A) Immunofluorescence analysis of Daxx in PML$^{-/-}$ splenocytes. Splenocytes, before and after Con A activation, were immunostained with anti-PML and anti-Daxx antibodies. Single staining (a, a', and b') and superimposed images (c and c') are shown. The nuclei were visualized by DAPI staining (d and d'). Representative confocal pictures are shown. Bar, 10 μM. (B) Western blot analysis of nuclear extract from PML$^{+/+}$ and PML$^{-/-}$ splenocytes, before and after Con A activation. Hsp90 protein levels are shown as a control of equal loading.

Figure 4. Daxx is selectively induced in B lymphocytes. PML$^{+/+}$ and PML$^{-/-}$ splenocytes were treated with Con A and at 72 h B and T subpopulations were obtained (Materials and Methods). Daxx expression levels at 0 and 72 h were measured in B and T cells by Western blot with the polyclonal anti-Daxx antibody. The differential expression of Daxx is also visible in extracts from untreated B and T cells upon longer exposure (not shown). Hsp90 was used as a normalization marker for protein input.
found to interact (Fig. 2 D). All together, these data demonstrate that Daxx can physically interact with PML in vivo, under physiological conditions.

PML Is Required for the Localization of Daxx in the NBs. To test whether PML would be required for NB localization of Daxx, we performed immunofluorescence analysis of Daxx localization in PML−/− and PML+/+ splenocytes before and/or after Con A activation (Fig. 3 A). Before Con A activation, the cytoplasmic localization of Daxx was indistinguishable between PML−/− and PML+/+ splenocytes. However, in activated PML−/− splenocytes, Daxx failed to localize in the NBs, and acquired a diffused, patchy, nuclear localization pattern (Fig. 3 A). Nevertheless, the induction of Daxx was unaffected by the absence of PML, as Daxx expression in PML−/− and PML+/+ nuclear samples was virtually indistinguishable by Western blot analysis (Fig. 3 B).

Daxx Is Differentially Upregulated in B Lymphocytes. Upon mitogenic activation of splenocytes, we noticed the presence of two distinct lymphocyte populations: cells that expressed Daxx at very high levels (Daxxh) and cells expressing low levels of Daxx (Daxxl) (not shown). Therefore, we tested whether this would be due to a differential regulation of Daxx expression in the two main splenic populations, B and T cells. B and T splenocytes were isolated after Con A stimulation by magnetic positive selection with an anti-B220 antibody and an anti-CD90 antibody, respectively (see Materials and Methods). We analyzed Daxx expression levels by Western blot in these two subpopulations. We observed that Daxx was induced in both B and T cells. However, much higher levels were detected in B cells (up to 10-fold; Fig. 4). These results were confirmed by immunofluorescence analysis (not shown).

Activation-induced Cell Death Is Profoundly Impaired in PML−/− B Lymphocytes. During Con A activation, a large number of B lymphocytes underwent programmed cell death, and after 96 h the remaining cell population was almost entirely composed by T lymphocytes (not shown). The induction of cell death coincided with the induction of Daxx and its redistribution to the NBs. Since in PML−/− splenocytes, Daxx was induced but did not localize to the NBs, we tested whether upon Con A stimulation the delocalization of Daxx would affect the induction of apoptotic

Figure 5. Daxx induction correlates with B cell apoptosis. PML+/+ and PML−/− splenocytes were treated with Con A. At 0, 24, 48, and 72 h after stimulation, B lymphocytes were isolated (see Materials and Methods) and scored for apoptosis by in situ TUNEL assay (A) or by cytometric analysis of subdiploid DNA content (B).

Figure 6. PML is required for the ability of Daxx to induce apoptosis. (A) Time course of apoptosis in PML+/+ and PML−/− keratinocytes transfected with His(6×)-Daxx or, as control, His(6×)-LacZ. Apoptotic cells were scored by TUNEL as described in B. The values are expressed as mean percentage of apoptosis ± SD (n = 3). (B) TUNEL staining of PML+/+ (top) and PML−/− (bottom) keratinocytes 24 h after transfection. Transfected cells were recognized by immunofluorescence staining with an anti-His mAb. His(6×)-positive, red; TUNEL-positive, green; DAPI, blue. White arrows point to His(6×)-positive nonapoptotic (TUNEL-negative) cells; red arrowheads point to His(6×)-positive apoptotic cells (yellow). The pictures were taken with an Olympus fluorescence microscope.
The Proapoptotic Ability of Daxx Is Abrogated in PML-negative Cells. We next investigated whether Daxx upregulation would induce apoptosis in cells that express PML, and whether its proapoptotic ability would be affected in the absence of PML. To this end, we used PML-/- and PML+/+ primary keratinocytes. These cells express PML and can be easily transfected (21, 23). Furthermore, as in splenocytes, in primary keratinocytes PML and Daxx colocalize in the NBs, whereas Daxx is delocalized from the NB in PML-/- keratinocytes (not shown). We transfected PML+/+ and PML-/- keratinocytes with a His(6×)-tagged Daxx expression vector or, as control, a His(6×)-tagged LacZ vector. After transfection, cells were harvested at 24 and 48 h and scored for apoptosis by in situ TUNEL analysis (see Materials and Methods; Fig. 6 A). Overexpression of Daxx resulted in induction of apoptosis in PML+/+ primary keratinocytes, whereas this activity was completely abrogated in the absence of PML (Fig. 6, A and B). Analysis of nuclear condensation upon DAPI staining gave comparable results (not shown). Thus, the inactivation of PML selectively impairs Daxx-induced apoptosis.

Daxx is delocalized from the NBs in APL cells. We next studied the localization of Daxx in APL cells by immunofluorescence analysis. As discussed previously, PML is delocalized from the NB in APL cells that contain the PML-RARα fusion protein due to t(15;17) translocation (5). This fusion protein is present in the APL cell line NB4 (5). Staining of NB4 cells with anti-PML antibody revealed the PML protein predominantly in nuclear microspeckles (Fig. 7). Daxx also displayed an aberrant localization pattern accumulating in microspeckles (Fig. 7), the majority of which did not appear to colocalize with the PML microspeckles (Fig. 7). Treatment of NB4 cells with RA causes relocation of PML to the NBs (5). In RA-treated NB4 cells, Daxx was relocalized to the NB, along with PML (Fig. 7). These observations further demonstrate that the normal localization pattern of Daxx requires PML and intact NBs as has been shown for other NB proteins such as Sp100 (9, 12–14).

Discussion

This study leads to the definition of a PML/NB-dependent function of Daxx that is indispensable for its proapoptotic function. Mitogenic stimuli such as Con A or LPS plus IL-4 (not shown) induce the upregulation and/or relocation of Daxx into the PML N Bs. Inactivation of PML results in the deregulation of this process and impaired apoptosis of mature splenic B lymphocytes upon mitogenic stimuli.

The impaired ability of PML-/- B lymphocytes to die provides a further rationale for the increased susceptibility of PML-/- mice to lymphomas (10). Furthermore, the identification of a critical role for PML in the induction of apoptosis by Daxx and the aberrant localization pattern of Daxx observed in APL cells provide a further basis for the resistance of the APL leukemic blasts to multiple apoptotic stimuli (11, 15). Although RA by itself induces differentiation and not apoptosis in APL cell lines such as NB4 (24; and our unpublished observations), the relocation of Daxx to the NBs upon RA treatment could render the APL cells sensitive to proapoptotic stimuli.

It has been shown that Daxx may serve as an adaptor of Fas (16), and an activator of ASK1 upon a Fas apoptotic stimulus (19). The activation of ASK1 results in the phosphorylation and activation of JNK (19). JNK activation leads to phosphorylation of c-Jun, which in turn may regulate transcription of genes relevant for apoptosis, possibly inducing the activation of caspases. This pathway can operate in parallel to the Fas-Fas-associated death domain (FADD) association, which couples Fas to procaspase 8 within the death-inducing signaling complex (DISC), leading to the cascade of caspase proteolytic activation. However, the Fas-dependent activation of JNK through Daxx does not seem to be sufficient for proper transduction of the Fas apoptotic stimulus since a Fas mutant that selec-
tively binds Daxx but not FADD is profoundly impaired in cell death induction (20). The transduction of the Fas signal to Jun does not seem to be affected in the absence of PML since Jun phosphorylation upon Fas stimulation is not impaired in PML L−/− activated splenocytes (not shown). Thus, the Daxx PML NB pathway for apoptosis may act independently of the Fas-Daxx-JNK-Jun signal transduction pathway. However, this does not exclude the possibility that Fas could act through the Daxx-PML NB pathway, supported by the fact that the ability of Daxx to potentiate Fas-induced apoptosis is completely abrogated in PML L−/− cells (not shown). It is therefore tempting to speculate that Daxx could potentiate Fas-induced apoptosis through a JNK-independent PML NB-dependent mechanism, in agreement with the marked impairment in Fas-induced apoptosis observed in PML L−/− cells and mice (11).

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