Nucleolar Disruption and Cajal Body Disassembly are Nuclear Hallmarks of DNA Damage-Induced Neurodegeneration in Purkinje Cells

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Abstract

The Purkinje cell (PC) degeneration (pcd) phenotype results from mutation in nna1 gene and is associated with the degeneration and death of PCs during the postnatal life. Although the pcd mutation is a model of the ataxic mouse, it shares clinical and pathological characteristics of inherited human spinocerebellar ataxias. PC degeneration in pcd mice provides a useful neuronal system to study nuclear mechanisms involved in DNA damage-dependent neurodegeneration, particularly the contribution of nucleoli and Cajal bodies (CBs). Both nuclear structures are engaged in housekeeping functions for neuronal survival, the biogenesis of ribosomes and the maturation of snRNPs and snoRNPs required for pre-mRNA and pre-rRNA processing, respectively. In this study, we use ultrastructural analysis, in situ transcription assay and molecular markers for DNA damage, nucleoli and CB components to demonstrate that PC degeneration involves the progressive accumulation of nuclear DNA damage associated with disruption of nucleoli and CBs, disassembly of polyribosomes into monoribosomes, ribophagy and shut down of nucleolar and extranucleolar transcription. Microarray analysis reveals that four genes encoding repressors of nucleolar rRNA synthesis (p53, Rb, PTEN and SNF2) are upregulated in the cerebellum of pcd mice. Collectively, these data support that nucleolar and CB alterations are hallmarks of DNA damage-induced neurodegeneration.

Keywords
Cajal bodies, DNA damage, gene silencing, nucleolus, Purkinje cell degeneration, ribophagy.

INTRODUCTION

Nuclear architecture includes the organization of chromosome territories and distinct nuclear bodies, such as nucleoli, Cajal bodies (CBs), nuclear speckles of splicing factors and promyelocytic leukemia (PML)-nuclear bodies (6, 8, 14, 24, 41, 48). The nucleolus plays a fundamental role in the synthesis of rRNAs and their assembly into ribosomal particles (57, 65). Other non-ribosomal functions include the biogenesis of several categories of ribonucleoprotein particles, cell cycle control and stress response (1, 36, 51).

In mammalian cells, the nucleolus has three basic structural components: (i) fibrillar centers (FCs), which concentrate components of the rDNA transcription machinery such as the RNA polymerase I and the upstream binding factor (UBF); (ii) the dense fibrillar component (DFC), the site of transcription of rRNA genes and early rRNA processing; and (iii) the granular component (GC), where the assembly of preribosomal particles takes place [for review, see (57, 65)]. The structural configuration of the nucleolus is largely determined by its activity in ribosome biogenesis. Thus, mammalian cells that require a high level of ribosome biogenesis commonly have large nucleoli with numerous small FCs (5, 57).

Among nuclear bodies, the CB is an intriguing nuclear structure discovered by Santiago Ramón y Cajal in vertebrate neurons [for a review, see (40)]. Illustrating the importance and plurifunctionality of this structure, CBs play a fundamental role in the biogenesis, transport and recycling of small nuclear- (snRNPs) and nucleolar ribonucleoproteins (snoRNPs) involved in pre-mRNA and pre-rRNA processing, respectively [for a review, see (14, 24, 47, 48, 66)]. Molecular analysis has shown that CBs are enriched in p80 coilin, a molecular marker of this structure (4, 56), splicingosomal snRNPs, the survival motor neuron protein (SMN) and CB-specific RNA [scRNAs; (10, 14, 16, 24)]. Additionally, CBs share with the nucleolus the proteins fibrillarin, Nopp140, NAP57 and some snoRNPs (34). CBs are prominent structures in mammalian neurons and their number and size correlate with neuronal size and transcriptional activity (5, 39).
The Purkinje cell (PC) is one of the most spectacular neurons known because of its magnificent planar dendrite tree and enormous abundance of dendritic spines. Moreover, PCs are the sole efferent output of the cerebellar cortex and play a fundamental function in the cerebellar control of movement (30, 50). As a result, PC degeneration and death lead to a severe dysfunction of motor control (22, 42). At the cell nucleus level, the predominant euchromatic configuration, the presence of prominent nucleolus and CBs and the great abundance of nuclear pores correlate with a high activity in transcription and nuclear export of RNA (25, 50), which is required for the maintenance of strong synaptic activity in this cerebellar neuron. However, these cellular properties make PCs particularly vulnerable to oxidative stress and to other genotoxic agents that can induce DNA damage during lifespan and neurodegenerative diseases (9, 63).

The PC degeneration (pcd) mutant mouse is characterized by an early cerebellar ataxia. This clearly distinguishable neurological sign match with a massive loss of PCs between postnatal day 15 (P15) and P30 (42). A late degeneration of photoreceptors in the retina, mitral cells in the olfactory bulb, and a discrete population of thalamic neurons also occurs [for a review see (7, 26, 49, 70)]. The early morphological features in predegenerative PCs include alterations of the rough endoplasmic reticulum (RER) and basal accumulation of polyribosomes (42). In the pcd mutant mice only the nna1 gene is affected, which encodes an ATP/GTP-binding protein (Nna1) with a putative zinc carboxypeptidase domain that is essential for PC survival (13, 72, 73). Nna1 contains nuclear localization signals and is distributed in both the nucleus and cytoplasm, being particularly enriched in the outer mitochondrial membrane (13, 27). Although the primary function of Nna1 in nuclear physiology remains elusive, it has been suggested that it has a role in chromatin remodeling (27). Moreover, a recent proteomic analysis has revealed mitochondrial dysfunction and altered proteolytic processing of Nna1 interacting proteins in pcd mice (13).

Defects in DNA repair pathways have recently emerged as a new fundamental pathology associated with polyglutamine (polyQ) diseases, such as Huntington’s disease and several spinocerebellar ataxias, as well as other neurodegenerative disorders (9, 20, 46, 55, 58). Indeed, in a previous work (70), we demonstrated that pcd mutation induces DNA damage during the predegenerative stage in mitral cells. Similarly, our preliminary studies demonstrate the presence of DNA damage foci in degenerating PCs of pcd mice. The rapid accumulation of DNA damage in PCs of this mutant mouse between 3 and 4 weeks of age, which ultimately leads to cell death, provides a relevant neuronal system for investigating nucleolar and CB mechanisms implicated in neurodegeneration. In this way, the nucleolus is involved in the biogenesis of ribosomes, an essential housekeeping function (8, 57). Similarly, CBs are functionally linked to the nucleolus through the biogenesis and supply of snoRNPs required for pre-rRNA processing (14, 34, 48). In this study, we analyze how the accumulation of unreppaired DNA in pcd mice PCs correlates with a general inhibition of RNA synthesis, disruption of the nucleolus and CBs and severe dysfunction of ribosome biogenesis that ultimately leads to neuronal death. Therefore, the alterations of the nucleolus and CBs represent a fundamental component in PC neurodegeneration pathway.

**MATERIALS AND METHODS**

**Mouse genotyping**

C57BL/6J male mice heterozygous for the pcd1J mutated gene were purchased from Jackson Laboratories (Bar Harbor, ME, USA). They were mated with DBA/2J females without the pcd mutation. The pcd1J allele was secluded with the genetic background of the C57BL/6J strain whereas the normal allele was associated with the genetic background of the DBA/2J strain. This differential segregation of both pcd and normal alleles allowed the control and heterozygous mice to be differentiated in order to expand the colony and genetically typify the experimental and control animals from the same litters. To investigate the genotype, DNA was extracted from the tail of the mice and the PCR was performed using the primers proposed for the markers D13Mit250 and D13Mit283 in the web resource of Jackson Laboratories: (http://www.informatics.jax.org/searches/probe.cgi?38700; http://www.informatics.jax.org/searches/probe.cgi?41581). PCR products of microsatellite regions D13Mit250 and D13Mit283, which have different sizes in both the C57BL/6J and DBA/2J strains, were resolved by electrophoresis in a 3% agarose gel. The animals were kept, handled and sacrificed in accordance with the Council of the European Communities and current Spanish legislation and the experiments were approved by the Bioethical Committee of the University of Salamanca.

**Microarray analysis**

To check the gene expression changes in pcd1J mice, we compared the gene expression profiles of control and pcd mice. Three control and three 20-day-old PCD mice were used. They were decapitated and the vermis of the cerebellum was rapidly removed. Total RNA from the vermis was extracted with TRIzol, and purified using RNeasy mini-kit (Qiagen 74104, Qiagen, Valencia, CA, USA) and RNase-free DNase I digestion. Once the samples of total RNA were obtained, the integrity and pureness of the RNA were determined using the Agilent 2100 Bioanalyzer (Agilent Technologies, Palo Alto, CA, USA). The messengers (mRNA) were retrotranscribed to yield cDNA using a commercial kit from Gibco/BRL (Superscript Choice System for cDNA synthesis). Thereafter, the cDNA was put under *in vitro* transcription in the presence of biotin labeled nucleotides to generate cRNA using the IVT kit (Affymetrix, Charleroi, Belgium). cRNAs were degraded by alkaline digestion and used for hybridizations with commercial chips, but only after having been subjected to a second quality control with a biochips Affymetrix test (TestArray 3). The hybridizations were made with the GeneChip Mouse Genome 430 2.0 Array of Affymetrix. The level of signal was calculated using the Robust Microarray Analysis algorithm (29), whereas the differential expression was calculated using the significance analysis of microarrays that include an estimation of the error by means of the false discovery rate.

**Tissue preparation**

After deep anaesthesia with a mixture of xylazine (Rompum, Bayer, Kiel, Germany) and ketamine hydrochloride (Imalgene, Merial, Lyon, France), mice were perfused with heparinized saline.
for 1 minute, and fixative solution containing 4% paraformaldehyde and 0.2% saturated picric acid in 0.1 M phosphate buffer (PB), pH 7.4, for 15 minutes. After perfusion, the vermis of the cerebellum was dissected and postfixed in the same solution for 2 h at room temperature. Then, tissue blocks were washed in PB, and cryoprotected with 30% sucrose, overnight at 4°C until they sank. Thirty µm-thick sagittal sections were cut using a freezing-sliding microtome (Leica Frigomobil, Jung SM 2000, Leica, Nussloch, Germany), the slices being collected in PB and stored at −20°C in a freezing mixture containing 30% glycerol and 30% polyethylene glycol in PB, until required.

**Immunofluorescence**

Cryo-cut sections from the vermis of control and mutant mice of 20 days of age were treated with 1% NaBH4 in PB for 20 minutes at room temperature, rinsed in PB (3 × 10 minutes), and incubated for 1 h in blocking serum (5% goat serum and 0.1% Triton X-100 in PB). The primary rabbit polyclonal anti-calbindin antibody (1:7,000; Swant, Switzerland) was incubated in the same solution overnight at 4°C. Cy2-conjugated secondary anti-rabbit antibody (1:500; Jackson Laboratories) was then applied for 2 h at room temperature. Finally, the sections were washed in phosphate-buffered saline (PBS) pH 7.4 mounted, and coverslipped with antifade solution.

Squash preparations of tissue fragments from the vermis fixed by perfusion with 3.7% paraformaldehyde in PBS were processed as previously described (53). Briefly, small blocks containing the PC layer were isolated and cut into small fragments. Each tissue fragment was transferred to a drop of PBS on a siliconized slide (SuperFrostPlus, Menzel-Gläser, Germany). A coverslip was then located at the top of the slide, and the tissue was squashed by mechanical percussion with a histologic needle to dissociate neuronal perikarya. The preparation was then frozen in dry ice, and the coverslip removed by using a razor blade. By using this procedure, most PCs remained adhered to the slide. Cell samples were then sequentially processed in 96% ethanol at 4°C for 10 minutes, and PBS at 4°C. Finally, the samples were stored at 4°C until used.

To perform the immunostaining on the dissociated PCs, the samples were sequentially treated with 0.1 M glycine in PBS for 15 minutes and 0.5% Triton X-100 in PBS for 45 minutes. Then, they were incubated with the primary antibody overnight at 4°C, washed with 0.05% Tween 20 in PBS, incubated for 45 minutes in the specific secondary antibody conjugated with FITC or Texas-red (Jackson Laboratories), rinsed in PBS, and counterstained with propidium iodide (1:2,000) for 15 minutes. Double immunofluorescence experiments were also carried out. Finally, the samples were mounted with the antifade medium Vectashield (Vector, Burlingame, CA, USA). They were examined with a laser confocal microscope (Zeiss LSM 510) using a 63× plan-apochromatic objective (1.4 NA), and argon ion (488 nm) and HeNe (543 nm) lasers.

The following primary antibodies were used. goat polyclonal anti-B23 protein (Santa Cruz Laboratories, Santa Cruz, CA, USA), mouse monoclonal anti-UBF (Santa Cruz Laboratories), rabbit polyclonal serum 204.3 anti-coilin (kindly provided by Dr A.I. Lamond, University of Dundee, UK), rabbit polyclonal anti-p53 (Santa Cruz Laboratories), mouse monoclonal anti-fibrillarin (Abcam, Cambridge, MA, USA), mouse monoclonal antibody anti-H2AX (Upstate, Waltham, MA, USA), which recognizes the phosphorylated variant of histone H2AX on Ser 139, and a human autosome antibody that recognizes the Sp100 protein of PML bodies (kindly provided by Dr Marcos del Hoyo, University Hospital “Marques de Valdecilla,” Spain).

The quantitative analysis of CBs and FCs was performed in cerebellar vermis samples from wild-type (n = 3) and pcd (n = 3) mice. For the determination of the mean number of FCs per PC nucleolus, squash preparations were immunostained for UBF (FCs) and the number of UBF-positive nucleolar spots was counted on serial confocal sections of the whole nucleolus. A minimum of 50 nucleoli for either wild-type or pcd experimental groups were used. The quantification of CBs was performed in squash preparations immunolabeled with the anti-coilin antibody and counterstained with propidium iodide. CBs were counted by direct examination of the different focal planes throughout neuronal nuclei and using a 63× oil immersion objective. A minimum of 200 PCs for each wild-type and pcd experimental groups were examined. All quantitative data were analyzed by the Student t-test to compare both experimental groups. Statistical significance was established at P < 0.01.

**Run-on transcription assay in situ**

Active transcription sites were labeled by incorporation of 5′-FU into nascent RNA as previously reported (11). Briefly, under anesthesia mice were given an i.p. injection of 5′-FU (Sigma, Dorset, UK) at the doses of 5 µL/g of a stock solution of 0.4 M 5′-FU in 0.9% saline. The mice were killed 60 minutes after the injection. The animals were fixed by perfusion with 3.7% paraformaldehyde in HPEM buffer [2X HPEM: Hepes, 60 mM; Pipes, 130 mM; ethylene glycol tetraacetic acid (EGTA), 20 mM; and MgCl2·6H2O, 4 mM] containing 0.5% Triton X-100 for 10 minutes. The vermis were removed, postfixed in the same solution for 20 minutes, washed in HPEM buffer containing 0.5% Triton X-100 for 10 minutes and cut into small fragments. Mechanical PC dissociation was performed as previously described. Then, the samples were sequentially treated with proteinase K (0.25 µg/mL in buffer Tris 1 M, pH 8) for 1 minute at 25°C, 0.1 M glycine in PBS containing 1% bovine serum albumin (BSA) for 15 minutes, and 0.01% Tween 20 in PBS for 5 minutes. The incorporation of the 5′-FU into nascent RNA was detected with a mouse monoclonal (clone BU-33) anti-BrdU antibody (Sigma, Madrid, Spain), diluted 1:50 in PBS (overnight at 4°C). Then, the samples were washed with 0.01% Tween 20 in PBS, incubated for 45 minutes with an anti-mouse FITC-conjugated secondary antibody (Jackson Laboratories), washed in PBS, mounted with the antifade medium Vectashield and examined with the laser confocal microscope. Some samples were counterstained with propidium iodide or processed for a double immunofluorescence labeling.

**Electron microscopy and immunogold labeling**

Three controls and three pcd mice of 20 days of age were used in order to analyze the ultrastructure of PCs. Mice were perfused with 3% glutaraldehyde in 0.1 M PB, pH 7.4. The cerebellum was removed and the vermis was isolated. Five hundred-micrometer-thick sagittal sections were obtained using a vibratome (Leica). Then, they were rinsed in 0.1 M PB, postfixed in 2% osmium
RESULTS

Previous studies have reported that PC death in pcd mice starts at postnatal day 18 (P18) and results in a massive neuronal death at P30 (42). Thus, at P20, immunostaining for calbindin, a molecular marker of PCs, clearly demonstrated the normal organization of PCs in the cerebellar cortex of wild-type mice and the neuronal loss in pcd mice (Figure 1A, B). Given that at P20 there coexists healthy PCs with different stages of neuronal degeneration in the cerebellar cortex of pcd mice, we have selected this postnatal day to study nucleolar and CBs alterations associated with this type of neurodegeneration.

To demonstrate the induction and progressive accumulation of DNA lesions in degenerating PCs of pcd mice, we performed immunolabeling for the phosphorylated histone H2AX at Ser 139 (γH2AX), a sensor for DNA damage (21), in combination with propidium iodide, a cytochemical staining for nucleic acids. In control PCs, γH2AX immunoreactivity was not detected, whereas the nucleolus and polyribosome-rich cytoplasmic domains (Nissl substance) appeared counterstained with propidium iodide (Figure 1C–E). At early stages of PC degeneration, numerous nuclear foci of γH2AX immunoreactivity were detected throughout the nucleus. They were stained with propidium iodide, reflecting a process of chromatin compaction at sites of DNA damage (Figure 1F–H). As PC degeneration took place, the expression of γH2AX propagated to extensive genome domains, resulting in a very few large γH2AX-positive nuclear domains in addition to other microfoci of DNA damage (Figure 1I–K). The activation of the expression of γH2AX in the pcd mice cerebellum was confirmed by Western blot analysis of nuclear extracts from cerebellar vermis (Figure 1L).

PC degeneration associates with a disruption of the nucleolus and cytoplasmic protein synthesis machinery

Having established that pcd mutation induces extensive DNA damage in PCs, we next investigated how this neuronal stress affects the organization and fine structure of protein synthesis machinery, particularly the nucleolus and polyribosomes. In wild-type mice, PCs exhibited typical reticulated nucleoli composed of numerous small FCs surrounded by a shell of DFC and irregular areas of GC, in addition to the nucleolus-attached heterochromatin (20). As PC degeneration took place, the nucleolus and cytoplasmic protein synthesis machinery were disrupted in many degenerating PCs, ranging from a moderate reorganization of nucleolar components to a severe nucleolar disruption with macrosegregation of fibrillar and GCs (Figure 2B–F). Thus, at initial stages of PC degeneration, the most characteristic ultrastructural feature was the reorganization of FCs into a very few larger FCs, whereas the small FCs tended to disappear (Figure 2B). This alteration was accompanied by the formation of large nucleolus-attached heterochromatin masses, reflecting the induction of gene silencing. As PC degeneration proceeded, nucleoli exhibited partial segregation of DFC and GC as well as the formation of nucleolar cavities containing electron-dense granules of variable size, whereas FCs were rarely found (Figure 2C, D). At more advanced stages of neuronal degeneration, striking abnormalities were the macrosegregation of the DFC and GC in large separated masses and, less frequently, the formation of a very large mass of GC surrounded by small aggregates of DFC (Figure 2E, F).

As the nucleolus plays a major role in biogenesis of ribosomes and maintaining protein synthesis machinery, we analyzed the reorganization of the RER and free polyribosomes associated with the nucleolar disruption in the same cells. PCs of wild-type mice
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exhibited the typical organization of RER arrays of cisterns associated with polyribosomes, some attached to the membranes and some free interspersed among cisterns (Figure 3A), as previously reported in mature PCs (42, 50). At the initial stages of PC degeneration, the stacks of RER cisterns tended to disappear and most of neurons showed a prominent mass of densely packed free polyribosomes at the basal pole of cell soma (Figure 3B), as was reported in the initial description by Landis and Mullen (42). Interestingly, in PCs that showed macrosegregation of nucleolar components and massive heterochromatinization, indicating a severe dysfunction of nucleolar and extranucleolar transcription, free polyribosomes had been replaced by monoribosomes (Figure 3C). They were closely packed and frequently appeared sequestered into cytoplasmic compartments bounded by isolated RER cisterns and identified as fibrillar centers (FCs; white arrow) and the formation of prominent masses of nucleolus-attached heterochromatin. As PC degeneration proceeds (C,D), the nucleolus shows a segregation of DFC and GC, disappearance of FCs, and formation of nucleolar cavities containing electron dense granules of variable sizes. At final stages of neuronal degeneration (E,F), nucleoli exhibit macrosegregation of DFC and GC in large separated masses (E), or the formation of a very large mass of GC together with small aggregates of DFC (F). Scale bar: 1 μm.
autophagosomes (Figure 3C and insert). The selective autophagia of ribosomes has been reported in yeast and termed “ribophagy” (35). In conclusion, the accumulation of DNA damage in PCs of pcd mice causes disruption of nucleolus, disassembly of polyribosomes and autophagic degradation of monoribosomes.

**pcd mutation induces segregation of fibrillarin and B23, reorganization of UBF and inhibition of rRNA synthesis in PCs**

In order to correlate the ultrastructural changes of the nucleolus with the reorganization of molecular markers of nucleolar components, we performed immunofluorescence with antibodies that recognized fibrillarin, the signature protein of the DFC, B23/Nucleophosmin, a marker of the GC, and the UBF for FCs (for a review, see (8, 57)). In control PCs, fibrillarin immunostaining of the nucleolus appeared as a cluster of closely packed nucleolar spherules (Figure 4A), corresponding to the shell of DFC that surrounded the FCs detected by electron microscopy. In contrast, fibrillarin was progressively segregated into irregular areas at the periphery of the nucleolus in PCs of pcd mice (Figure 4B, C).

Immunostaining for B23 revealed a non-homogenous labeling of the nucleolar body in control PCs (Figure 4D), whereas B23 was relocalized diffusely throughout the nucleoplasm and concentrated packed free polyribosomes is observed at the basal cell body (B). As PC degeneration proceeds, free polyribosomes are replaced by densely packed monoribosomes (C, and insert in C). Cytoplasmic portions containing monoribosomes appear sequestered in autophagic vacuoles bounded by isolated RER cisternae (insert in C). Scale bars = 1 μm.
in residual masses of GC in degenerating PCs showing accumulation of DNA damage (Figure 4E, F).

To further investigate the relationships between nucleolar disruption and dysfunction of rRNA synthesis in pcd mice PCs, we analyzed the reorganization of UBF, a RNA polymerase I factor essential for the activation of rDNA transcription [for a review, see (57, 67)], and the in situ transcriptional activity. Immunostaining for UBF displayed the nucleolar distribution of FCs, as previously demonstrated in other neuronal and non-neuronal cell types (5, 60). Whereas control PCs exhibited numerous small FCs, an apparent reduction in their number was observed in neurons of pcd mice (Figure 4G–I). As the number of FCs in mammalian neuronal nucleoli positively correlated with transcriptional activity [for a review, see (5)], we estimated the number of UBF-positive intranucleolar spots (FCs) by quantitative analysis. The mean number of FCs per nucleolus significantly decreased from $11.96 \pm 2.26$, in wild-type PCs, to $5.96 \pm 1.75$, in pcd mice neurons (values are mean ± standard deviation of the mean (SD). Statistical analysis was carried out using the Student $t$-test to compare both experimental groups, $P < 0.01$). Next, we investigated changes in transcriptional activity using the incorporation of 5'-FU into nascent RNA after a pulse of 60 minutes of the intraperitoneal administration of this halogenated nucleotide (11). In control PCs, nascent RNA concentrated in the nucleolus and in numerous extranucleolar transcription foci, in addition to being diffusely distributed throughout the nucleoplasm (Figure 4J). In degenerating pcd mice PCs, the global transcription rate dropped dramatically, as was indicated by the reduction in both transcription foci and the diffuse nucleoplasmic signal of 5'-FU incorporation into nascent RNA. Interestingly, these neurons had an altered nucleolar pattern of 5'-FU labeling. It appeared as a few segregated areas of 5'-FU incorporation probably corresponding to domains of segregated DFC of the nucleolus (Figure 4K). In advanced stages of PC degeneration with a great accumulation of DNA damage no signal of 5'-FU incorporation was detected, indicating a complete silencing of gene expression (Figure 4L). In conclusion, nucleolar disruption in pcd mice PCs is another nuclear hallmark of neuronal degeneration that reflects the repression of rRNAs genes and the inhibition of ribosome biogenesis.

**Microarray analysis revealed increased expression of some genes encoding negative regulators of rDNA transcription in pcd mice**

Having established the existence of a nucleolar stress response to DNA damage with disruption of the structure and function of the nucleolus in pcd mice PCs, we next investigated whether these alterations correlate with changes in gene expression of rDNA transcription repressors. Microarray analysis of cerebellar mRNA expression revealed upregulation of certain genes encoding repressor factors of rDNA transcription in the pcd mice (Table 1). The transcripts of these genes that increase by two fold or more in pcd mice compared with wild-type mice included those encoding (i) p53 and RB, two negative regulators of RNA Pol I activity (62, 71, 75), (ii) Snf2h, a component of the nucleolar remodeling complex (NoRC) involved in the epigenetic silencing of rRNA genes (64) and (iii) PTEN, a lipid phosphatase that downregulates the PI3K pathway and blocks recruitment of the basal transcription factor SL1 to the promoter of rRNA genes (18, 76). In the case of p53, we have observed by immunofluorescence a very weak or undetectable nuclear signal and a diffuse cytoplasmic staining in control PCs (Figure 4M). In contrast, the p53 nuclear staining increased and nuclear spots of higher intensity were frequently observed in pcd mice PCs (Figure 4N). The proportion of PCs containing p53-positive spots significantly increased in the pcd mice (2.2% ± 0.1%, in the wild-type mice, vs. 54% ± 0.3%, in the pcd mice, values are mean ± SD. Statistical analysis was carried out using the Student $t$-test to compare both experimental groups $P < 0.001$). Co-staining for p53 and Sp100, a typical constituent of PML bodies (6), revealed that most p53 spots colocalized with Sp100 in PML bodies (Figure 4O). This co-localization is consistent with the role of PML bodies in the cellular stress response, which requires the recruitment and activation, through post-translational modifications, of p53 in these nuclear structures (3). Taken together, these data suggest the transcriptional activation of repressors of the rDNA transcription machinery in the cerebellum of the mutant mice. Moreover, gene expression analysis is consistent with the present observations of a breakdown of nucleolar structure and disruption of protein synthesis machinery. However, it should be noted that these microarray data referred to the whole cerebellum, not only to PC mRNAs. Therefore, the possible involvement of other cerebellar cell populations, as a secondary response to the PC degeneration, cannot be excluded.

<table>
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**Table 1. Differentially expressed transcripts of genes encoding repressors of nucleolar rRNA synthesis in vermis of pcd mutants compared with wild-type mice.**

**PC degeneration disassembles CBs and relocates coilin around segregated masses of fibrillarin**

Because CBs have been linked to the organization and function of the nucleolus, particularly in mammalian neurons [for review, see (5, 31, 53)], we investigated the reorganization of CBs that accompanied the nucleolar alterations detected in degenerating PCs in pcd mice. For this purpose, we performed double immunostaining for coilin and fibrillarin on dissociated PCs in order to visualize CBs and nucleoli, respectively. Control PCs exhibited typical CBs, free in the nucleoplasm or attached to the nucleolus, in which coilin and fibrillarin colocalized (Figure 5A–C). In contrast, the pcd mice PCs showed substantial changes in the number of CBs and in the nuclear distribution of coilin. The quantitative analysis at P20 of the mean number of CBs per nucleus revealed a significant reduction in PCs from pcd mice in comparison with wild-type mice neurons ($2.03 \pm 0.39$, in control, vs. $0.48 \pm 0.16$, in pcd mice, mean ± SD, $P < 0.01$). In PCs with initial segregation of nucleolar components, CBs with very irregular shape were found (Figure 5D, F). Notably, in nucleoli with macrosegregation of their components, coilin was redistributed as a thin shell around the large and small segregated masses of fibrillarin (Figure 5G–I). In more advanced stages of nucleolar disruption...
and fragmentation, the close association of coilin with the surface of the fibrillarin-positive fragments was preserved (Figure 5J, L). At the final stage of PC degeneration, a coilin-positive mass was frequently seen juxtaposed to a nucleolar remnant of fibrillarin at the periphery of the large heterochromatin domains (Figure 5M, O).

CB alterations and coilin relocalization in PCs of pcd mice were confirmed by immunogold electron microscopy for coilin detection. CBs in control neurons exhibited the typical configuration of round structures composed of closely packed coiled threads decorated with gold particles (Figure 6A). Interestingly, in degenerating PCs CBs frequently showed an irregular morphology with loosely arranged threads, which appeared to be in a process of disassembly (Figure 6B, C). Furthermore, small aggregates of coilin were observed at the nucleolar surface (Figure 6C). Finally, immunoelectron microscopy also revealed the presence of electron-dense bodies surrounded by a shell of amorphous material decorated with gold particles of coilin immunoreactivity (Figure 6D). These structures clearly represent the ultrastructural counterpart of fibrillarin masses with their peripheral layer of coilin detected with immunofluorescence. In conclusion, the assembly/disassembly cycle of CBs is dramatically altered in degenerating PCs of pcd mice, parallel to the disruption of nucleolar structure and function. The disassembly of coiled threads of CBs may lead to the reduction in the number of these nuclear organelles. Interestingly, the close association of coilin and fibrillarin is preserved in all stages of PC degeneration: in CBs and around the nucleolus and nucleolar fragments upon disruption of CBs.

**DISCUSSION**

PCs of the pcd mouse provide a useful neuronal system to study nuclear mechanisms involved in DNA damage-dependent neurodegeneration. A defective DNA repair has been involved in the molecular pathology of a growing list of neurodegenerative disorders [for review, see (20, 46, 58)]. In the case of PCs, the inherited spinocerebellar ataxias represent an important group of human neurodegenerative diseases with PC degeneration phenotypes, some of them caused by mutations of DNA repair genes (17, 68, 69). Although the pcd mutation is a model of ataxic mouse, it shares clinical and pathological characteristics of inherited human spinocerebellar ataxias, such as difficulties in motor coordination and degeneration and loss of PCs [for review, see (19, 72)]. A recent study on the protein-protein interaction network for human ataxias and mouse models of PC degeneration, including the pcd mouse, has revealed that many ataxia-causing proteins interact in an ataxia-based network (44). Indeed, these human and mouse neurodegenerative disorders, although with different mutated genes, share molecular pathways that explain their phenotypic similarities (44, 61). Moreover, Gene Ontology analysis has revealed that nuclear compartments are the main cellular targets of the ataxia protein network (44). Taken together, these data point to the usefulness of the pcd mutant mouse as a model of PC degeneration and clinical cerebellar ataxia.

Our results indicate that PC degeneration in pcd mice involves a progressive accumulation of DNA damage, which is associated with the disruption of nucleoli and CBs and disassembly of protein synthesis machinery that ultimately leads to neuronal cell death. Nuclear hallmarks of PC degeneration are: (i) increased accumulation of the DNA-damage signal γH2AX; (ii) disruption of the nucleolus with inhibition of rRNA synthesis and ribosome biogenesis; and (iii) disassembly and reduction of CBs. Our finding by microarray analysis that four genes encoding negative regulators of rDNA transcription (p53, Rb, PTEN and SNF2) are upregulated in the cerebellum of pcd mice suggests the activation of repressors of rRNA synthesis in PC neurodegeneration. This view is also consistent with previous Gene Ontology analysis illustrating that transcription corepressor activity is a molecular function with significantly enhanced representation in the ataxia network of protein interactions (44). Furthermore, our immunofluorescence data are consistent with the contribution of p53 to PC degeneration of pcd mice. It is well known that p53 is activated by DNA damage and can trigger DNA repair or apoptosis [for a review, see (46)]. The recruitment of p53 to PML bodies detected in pcd mice PCs may enhance p53 activity by inducing its phosphorylation and acetylation mediated by kinases (HIPK2, CK1) and the acetyl transferase CBP, two post-translational modifications that occur in PML bodies in response to various cellular stresses (3, 6).

Under physiological conditions, the γH2AX signal was not detected by immunofluorescence in PCs of the P20 wild-type mouse. In pcd mice PCs, however, DNA breaks labeled with the anti-γH2AX antibody initially appear as numerous nuclear foci that propagate and coalesce into large nuclear domains of genome as PC degeneration proceeds. This finding, and our previous demonstration of DNA breaks in mitral cells (70), clearly support that the accumulation of DNA lesions is a key component in PC neurodegeneration of the pcd mouse. Although degenerating PCs are able to trigger the γH2AX signaling response to DNA damage, the persistence and progressive increase of this signal suggest that DNA lesions are refractory to repair in the mutant animals. This interpretation is consistent with the growing evidence that a defective DNA repair is engaged in the pathogenesis of several ataxias with PC degeneration phenotype (17, 20, 68, 69).

Nucleolar alterations associated with the progressive accumulation of DNA damage in PCs include a reduction in the number of FCs, segregation of nucleolar components, inhibition of rRNA synthesis and fragmentation of the nucleolus. Regarding FCs, nucleolar domains enriched in molecular components of the rRNA transcription machinery (60, 65), previous studies in other neuronal and non-neuronal cells indicate that the number of FCs positively correlates with cell size and transcriptional activity (5, 41, 57). In this context, we envisage that the reduction of FCs detected in pcd mice PCs represents an early manifestation of the disruption of rRNA synthesis machinery. Subsequently, the segregation of DFC and GC in large separated masses is a typical morphological feature of a severe nucleolar dysfunction and it has been demonstrated in cells treated with drugs that interfere with transcription of the rDNA, such as actinomycin D and doxorubicin (12, 52, 54). Moreover, the recent observation that the inhibition of rRNA synthesis triggers apoptosis in cultured cortical neurons reflects the importance of nucleolar transcription for neuronal survival (28, 32). Similarly, our finding of the nucleoplasmic translocation of B23, a resident nucleolar chaperone involved in ribosome particle assembly and nucleocytoplasmic transport (23), has been associated with states of inhibition of rRNA synthesis in culture cell lines (74). In line with this, the existence of gene silencing in nucleolar and extranucleolar domains of the genome is clearly demonstrated by the dramatic reduction of 5′-FU incorporation into nascent RNA
detected here in degenerating PCs of pcd mice and also by the absence of detectable in situ transcriptional activity in advanced stages of PC degeneration. Collectively, these data suggest that the accumulation of unrepaired DNA induces a nucleolar stress response with severe and progressive inhibition of both rRNA synthesis and nucleolar assembly of preribosomal particles in pcd mice PCs. In support of this view, a previous study (37) has demonstrated a direct link between DNA breaks and block of rRNA synthesis in laser micro-irradiated nucleoli of wild-type mouse embryonic fibroblasts. These authors found a transient block in rRNA synthesis in irradiated nucleoli, whereas transcription was preserved in undamaged nucleoli of the same cell. Interestingly, the resumption of rRNA synthesis in irradiated cells correlated with the disappearance of γH2AX foci, suggesting that restoration of transcription is dependent on a successful repair of DNA damage (37).

What is noteworthy is the close relationship between nucleolar disruption and disassembly of protein synthesis machinery observed in pcd mice PCs. The disorganization of RER arrays and the persistent accumulation of free polyribosomes are well known ultrastructural features of PC degeneration in the pcd mouse (42). The initial disassembly of RER may contribute to the neuronal dysfunction by interfering with the vesicular transport of proteins, via the Golgi complex, to different cellular targets (2). An important additional finding in our results is the accumulation of free monoribosomes observed in PCs with macrosegregation of the nucleolar components, colin appears relocalized as a thin ring surrounding the segregated masses of fibrillarin (G–I). At the final stage of degeneration, a colin-positive mass is frequently seen juxtaposed to a nucleolar remnant of fibrillarin at the periphery of the large heterochromatin domains (M–O). Scale bar = 5 μm.

Figure 5. pcd mutation causes disassembly of Cajal bodies (CBs) and relocation of colin. Confocal microscopy images from squash preparations of Purkinje cell (PC) from wild-type (A–C) and pcd mice (D–O) co-stained for fibrillarin and colin. Control PCs exhibit typical CBs free in the nucleoplasm or attached to the nucleolus, in which both proteins colocaled (A–C). D–F. At early stages of degeneration, PCs display a segregation of the nucleolus (D) and CBs with an irregular morphology (E). In more advanced stages of PC degeneration with a severe segrega-
nucleolar components, suggesting that nucleolar disruption and disassembly of polyribosomes are linked cellular processes. In fact, the disassembly of polyribosomes into free monoribosomes lacking mRNA templates is a late cellular sign in degenerating PCs and reflects a blockade of protein synthesis. In nervous tissue, the formation of free monoribosomes correlates with a dramatic decay of 28S and 18S rRNAs (38) and has been associated with programmed cell death (33). Interestingly, in the case of PC degeneration in pcd mice, a fraction of free monoribosomes appear specifically sequestered into autophagic vesicles bounded by RER membranes, a process termed ribophagy (35). This observation suggests that autophagy-related pathways are involved in the selective degradation of ribosomes in degenerating PCs of the pcd mouse.

Nucleolus and CBs are nuclear epicenters of the RNA world and their dysfunction could have important consequences for both protein synthesis and information processing and storage in neurons (31, 59). In our experimental system of PC degeneration in pcd mice, CBs are sensible nuclear targets of the general inhibition of RNA synthesis and processing, as shown by their structural disassembly, reduction in their number and redistribution of coilin to the periphery of nucleolar fragments. Although disruption of CBs has been reported in experimental models of transcriptional inhibition [for review, see (15, 39, 45)], little information is available on the behavior of CBs in neurodegenerative disorders, with the exception of the spinal muscular atrophy, in which the reduction in the level of SMN protein alters the organization of CBs and associated SMN-containing “gems” (43), and our previous observation in mitral cells of pcd mice (70). Unlike the experimental model of ultraviolet (UV)-induced fragmentation of CBs, in which coilin relocates to nucleoplasmic microfoci (15), we have not observed nucleoplasmic accumulation of coilin in the DNA damage-induced neurodegeneration of PCs. Instead, coilin redistributes around both nucleoli and nucleolar fragments and ultimately coilin and fibrillarin form cap-like structures opposed to heterochromatin clumps. Although these two cellular models of DNA damage, UV-irradiation and pcd mutation, share a down-regulation of transcription, our results suggest that the relocalization of coilin to different nuclear targets is not simply an indirect effect of transcription inhibition. Moreover, our immunocytochemical data support a cellular mechanism of disassembly of CBs mediated by the initial loosening of coiled threads followed by their fragmentation. Given the importance of CBs for the processing of pre-mRNAs and pre-rRNAs (14, 24, 48), the disassembly and loss of CBs may reflect the response of PCs in pcd mice to a reduced demand of RNA processing, and also supports the participation of these nuclear bodies in cellular stress-response pathways.

Although the pcd mutation is a model of ataxic mouse with DNA damage, other perturbations of the ataxia protein network might cause nucleolar and CB alterations without the involvement of DNA damage, particularly those that interfere with global transcriptional activity and nuclear pre-mRNA processing. Interestingly, three main hub proteins (RBM9, A2BP1 and RBPM5) in the ataxia network, which interact with several different ataxia-causing proteins, are RNA-binding proteins involved in pre-mRNA splicing and maturation [for a review, see (44)]. However, in the case of the pcd ataxic mouse, the accumulation of DNA lesions profoundly impacts upon nuclear functions in PCs, as is indicated by the progressive transcriptional silencing. In this context, we propose that the disruption of the nucleolus and CBs are early and sensitive nuclear hallmarks of DNA damage-induced neuronal degeneration in PCs and, presumably, in other neuronal types.

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