DNA damage in tissue-resident macrophages leads to age-related neurodegeneration

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Article

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Abstract

Neurodegenerative disorders are a growing challenge for the elderly yet their etiology remains elusive. Here, we show that persistent DNA damage in tissue-resident macrophages carrying an ERCC1-XPF DNA repair defect leads to cerebellar ataxia in mice. We find that cytoplasmic chromatin fragments accumulate in the brain microglia of progeroid and naturally aged mice stimulating a type-I Interferon (IFN-I) response and are then packaged in extracellular vesicles (EVs) leading to Purkinje cell death and neurodegeneration in \( Er\tau^{CX/-} \) animals. To reduce neuroinflammation, we developed an EV-based strategy to deliver recombinant DNase I specifically in inflamed \( Er\tau^{CX/-} \) microglia \textit{in vivo}. Our approach rapidly removes dsDNAs from the cytoplasm of microglial cells and in secreted EVs; it alleviates the IFN-I response, decreases Purkinje cell death and delays the onset of neuronal decline in \( Er\tau^{CX/-} \) animals. Thus, brain microglia causally contribute to neurodegeneration allowing for the development of promising therapeutic strategies against age-related neuroinflammation.

Introduction

To counteract DNA damage, cells rely on a series of partially overlapping DNA repair pathways to preserve their vital genetic information and faithfully transmit it to progeny \(^1\). Besides cancer, the great majority of DNA repair defects in man gives rise to a heterogeneous group of progeroid (premature aging-like) syndromes that present frequently or exclusively with mild to severe neurological symptoms \(^2\). Indeed, in view of the DNA repair defect, the high oxidative load and metabolic by-products generated during normal brain metabolism are thought to trigger irreparable DNA modifications driving neuronal cell death and the progressive degeneration of the central or peripheral nervous systems \(^3\).

Xeroderma Pigmentosum F-Excision Repair Cross Complementation group 1 (XPF-ERCC1) is a heterodimeric, structure-specific endonuclease complex required for lesion excision in nucleotide excision repair (NER) \(^4,5\) that plays an analogous role in the repair of highly cytotoxic DNA interstrand crosslinks (ICLs) \(^6\). Patients with mutations in \( XPF \) may present with mild symptoms of the NER-deficient syndrome XP or exhibit adult-onset progressive neurologic deterioration, including chorea, ataxia, cognitive deficits and brain atrophy \(^7\). Recently, a single patient was discovered with mutations in \( ERCC1 \) resulting in severe pre- and postnatal developmental defects as well as pronounced neurological alterations associated with cerebellar hypoplasia and blunted cortical gyri \(^8,9\).

We and others have recently shown that DNA damage-driven inflammation causally contributes to age-related, tissue degenerative changes \(^10\)–\(^13\). However, the relative contribution of compromised genome integrity in distinct cell types, e.g. neurons or glial cells to age-related neurodegenerative features remains unknown. Using \( Er\tau^{CX/-} \) mice with an engineered ERCC1-XPF defect only in tissue-resident macrophages, we provide evidence for a fundamental mechanism by which irreparable DNA damage leads to the accumulation of chromatin fragments in the cytoplasm of microglial cells. Using functional assays coupled to an advanced simultaneous 2-3 multiphoton analysis on acute brain slices, we find that
cytoplasmic chromatin fragments are sensed, stimulating a type-I IFN response. We find that cytoplasmic chromatin fragments are packaged in microglia-derived Er$^{CX/-}$ EVs that selectively target Purkinje cells leading to their apoptotic death and the premature onset of neurodegeneration in Er$^{CX/-}$ animals. To remove cytoplasmic dsDNAs from the cytoplasm of activated microglial cells and lessen the neuroinflammatory load in Er$^{CX/-}$ mice, we developed an extracellular vesicle (EV)-based strategy to deliver recombinant DNase 1 nuclease in vitro and in vivo, targeting CD11b$^+$ cells of the brain. We find that treatment with the EV-delivered nuclease cargo rapidly eliminates inflammatory dsDNA moieties from the cytoplasm of microglia cells and in microglia-secreted EVs. This approach alleviates the Type-I IFN response and Purkinje cell death, leading to the substantial delay in the premature onset of cerebellar ataxia in Er$^{CX/-}$ animals. Taken together, our findings highlight the prominent role of tissue-resident macrophages in age-related neurodegeneration opening new avenues for the development of novel intervention strategies against neuroinflammatory disorders.

**Results**

**Loss of ERCC1 in tissue-resident macrophages triggers progressive ataxia in mice.** How DNA damage leads to the premature onset of heterogeneous pathological features in NER patients and accompanied animal models remains an intriguing question arguing for cell type-specific responses against genotoxic threats. Tissue-resident macrophages are a heterogeneous group of immune cells that reside in distinct tissue environments and are vital for tissue homeostasis and defense against foreign pathogens or environmental challenges $^{14}$. To dissect the impact of irreparable DNA damage in tissue-resident macrophages, we intercrossed animals homozygous for the floxed Ercc1 allele (Ercc1$^{F/F}$) with mice carrying the CX3CR1-Cre transgene in an Ercc1 heterozygous background (from now on denoted as Er$^{CX/-}$ animals). CX3CR1 is a CX3C chemokine receptor 1 for fate-mapping studies of the tissue-resident monocyte and macrophage compartment $^{15}$. Confocal microscopy studies in CX3CR1-Cre crossed with the Rosa YFP transgenic animals and in Er$^{CX/-}$ animals confirmed the Cx3CR1-driven YFP expression (Figure 1A) and the absence of ERCC1 expression in Er$^{CX/-}$ tissue-resident macrophages, respectively (Figure 1B). Together, these findings indicate the normative ERCC1 expression levels in Er$^{CX/-}$ tissues or cells other than the targeted cell population. Er$^{CX/-}$ mice are born at the expected Mendelian frequency and present no developmental defects or other pathological features. At 4-months of age, however, Er$^{CX/-}$ mice manifest progressively signs of ataxia that become clearly evident in the 6-months old animals. In line, when the 6-months old wild-type mice (Er$^{F/-}$; wt.) are suspended by their tails, the animals extend and shake their hind limbs to maintain balance (Supplementary Video file 1). Instead, Er$^{CX/-}$ mice keep their hind limbs in a clasped position (Figure 1C; Supplementary Video file 2) and walk with a wide gait compared to age-matched littermate control animals. Rotarod assessment reveals that hind limb coordination deficiency is apparent in ~70% of the 6-months old Er$^{CX/-}$ animals compared to Er$^{F/-}$ littermate controls (Figure 1D). Beginning at 8-months of age, Er$^{CX/-}$ animals develop kyphosis (Figure 1E) and fine tremor to front legs. The premature onset of neurodegenerative features in Er$^{CX/-}$ animals prompted us to assess the morphological and phenotypic characteristics of CNS-resident macrophages.
We find that brain microglia in 6-months old $Er^{1C^x/-}$ animals form finger-like protrusions, a hallmark of microglia activation that involves cellular locomotion and increased antigen presentation \(^{16}\) (Figure 1F). Fluorescence-activated cell sorting (FACS) analysis of freshly isolated CD11b\(^+\) cells derived from $Er^{1C^x/-}$ brains revealed a substantial increase in cell size (Figure 1G), in the expression of MHCII and CD86 proteins as well as the number of MHCII\(^+\) CD86\(^+\) cells compared to CD11b\(^+\) cells of $Er^{1F/-}$ animals (Figure 1H and). However, when compared to LPS-treated, fully activated pro-inflammatory microglia, $Er^{1C^x/-}$ microglia appear to be in a primed, yet not fully activated state (Figure 1H and Supplementary Figure S1A). Further work reveals a comparable number of microglial cells in the 3- and 6-months old $Er^{1C^x/-}$ and $Er^{1F/-}$ brains (Figure 1I). The lack of infiltrating monocytes as assessed by the histological evaluation of 3- and 6-months old $Er^{1C^x/-}$ brains (Figure 1J), the normative CD45 expression levels in the 6-months old $Er^{1C^x/-}$ brains (Figure 1J; Supplementary Figure S1B) and the comparable number of Ly6C\(^+\) bone marrow-derived macrophages in the 6-months old $Er^{1C^x/-}$ and $Er^{1F/-}$ brains (Figure 1K) diminishes the possibility of peripheral immune cell infiltration in $Er^{1C^x/-}$ brains.

**Accumulation of cytoplasmic chromatin fragments triggers a type-I IFN response in $Er^{1C^x/-}$ microglia.** Phosphorylated histone H2A.X (\(\gamma\)-H2A.X)-containing foci accumulate at sites of DNA breaks \(^{17}\). FACS analysis in single cell suspensions of $Er^{1F/+}$ brains reveal that the great majority of \(\gamma\)-H2A.X\(^+\) cells are CD11b\(^+\)NeuN\(^-\) cells (Figure 2A; upper panel). Intriguingly, we find that the percentage of \(\gamma\)-H2A.X\(^+\)CD11b\(^+\)NeuN\(^-\) cells increases gradually from 3- to 12-months old $Er^{1F/+}$ brains indicating that microglia cells accumulate rapidly DNA damage compared to neuronal cell types with aging. Consistent with the DNA repair defect, we find a significantly higher percentage of \(\gamma\)-H2A.X\(^+\)CD11b\(^+\)NeuN\(^-\) cells in $Er^{1C^x/-}$ brains compared to littermate controls across all age groups (Figure 2A; lower panel). The phosphorylated Ataxia telangiectasia-mutated protein (pATM) is a central mediator of the DNA damage response. Confocal microscopy studies confirmed the significant increase in the number of \(\gamma\)-H2A.X\(^+\)CD11b\(^+\) and pATM\(^+\)CD11b\(^+\) cells in $Er^{1C^x/-}$ brains compared to $Er^{1F/-}$ controls (Figure 2B; as indicated). Interestingly, \(\gamma\)-H2AX and pATM accumulate in the nucleus as well as the cytoplasm of microglial cells in $Er^{1C^x/-}$ brains. Unlike in $Er^{1F/-}$ microglia cells, we also detect the presence of 4′,6-diamidino-2-phenylindole (DAPI) dense foci marking the presence of chromatin in the cytoplasm of $Er^{1C^x/-}$ freshly isolated microglial cells (Figure 2C). DNA damage triggers the release of micronuclei containing whole or fragmented chromosomes into the cytoplasm \(^{18}\). Chromatin fragments are subject to autophagic degradation \(^{19}\) or else accumulate stimulating a type-I IFN immune response \(^{20}\). The presence of chromatin fragments in the cytoplasm of cells prompts us to test whether nuclear DNA leakage in the cytoplasm of $Er^{1C^x/-}$ microglia cells triggers the activation of a type-I IFN response, known to be associated with anti-viral and immune modulating functions \(^{21}\). Evaluation of the bioactive murine type-I IFN levels by means of the B16-Blue™ IFN-\(\alpha\)/\(\beta\) cell line in the extracellular milieu (brain lavage) of the 6-months old $Er^{1C^x/-}$ and aged-matched $Er^{1F/-}$ littermate control animals reveals significantly higher type-I IFN levels in $Er^{1C^x/-}$ brains (Figure 3A). Consistently, we find an increase in the mRNA levels of several interferon signature genes (ISGs) i.e. ifn\(\beta\), ifit2, ifi207 and ifi44 in $Er^{1C^x/-}$ whole brain lysates (Figure 3B).
FACS analysis in the 6-months old Er1^Cx/− cerebella and cortices reveal an increase in pSTING (phosphorylated form of stimulator of interferon genes) protein levels known to act as a sensor of cyclic d-GMP and as an adaptor protein mediating the interferon response, once activated by DNA sensors like cGAS (Figure 3C) 22. In Er1^Cx/− microglial cells, we find that cytoplasmic DAPI+ chromatin colocalizes with the autophagy protein LC3β known to be involved in membrane trafficking and substrate delivery to lysosomes 23 and lamin B1, a nuclear lamina protein associated with chromatin domains 24 that is markedly disorganized in Er1^Cx/− microglial cells compared to controls (Figure 3D; as indicated and Supplementary Figure S2A). Consistently, we find that pATM+ DAPI+ chromatin structures accumulate in the cytoplasm of Er1^Cx/− cells and are surrounded by P62, a sequestering protein that delivers substrates to forming autophagosomes 25 (Figure 3E; as indicated and Supplementary Figure S2B). Together, these findings indicate a physiological budding response of Er1^Cx/− microglial cells to remove irreversibly damaged chromatin fragments through nucleophagy. We recently showed that type-I IFN mediates the increase of lysosomal pH to delay autophagosomal degradation, thereby favoring antigen presentation and cytokine secretion 26. In agreement, we find that the lysosomal pH of freshly isolated CD11b+ cells is significantly increased indicating that lysosomal acidity is compromised in Er1^Cx/− brain-derived microglia cells (Figure 3F). The latter may well explain the pronounced accumulation of chromatin fragments in the cytoplasm of DNA repair-deficient Er1^Cx/− microglial cells. Next, we test whether cytoplasmic chromatin fragments also accumulate with natural aging. In line, we find that DAPI-stained chromatin fragments accumulate in the cytoplasm of microglial cells derived from 24-months old naturally aged mice cells (Figure 3G). Similar to 6-months old Er1^Cx/− animals, FACS analysis revealed an increase in pSTING protein levels in naturally aged cerebella (Figure 3H and Supplementary Figure S2C).

Er1^Cx/− microglia elicit antiviral-like response that triggers Purkinje neuronal cell death. A number of animal models carrying inborn defects in DDR (e.g. Atm^−/−) and genome maintenance (e.g. Csb^m/m-Xpa^−/−) develop cerebellar ataxia associated with Purkinje cell death 27,28. However, unlike in DNA repair-deficient animal models, Purkinje cells in Er1^Cx/− mice are DNA repair-proficient and do not accumulate γ-H2AX foci in their nuclei (Figure 4A). Nonetheless, TUNEL assay and staining for activated Caspase-3 in 3- and 6-months old Er1^Cx/− brain sections reveal increased cell death in the Purkinje cell layer of cerebellum (Figure 4B; as indicated). Loss of Purkinje cells is also in line with the defective coordination of hind limbs and the fine tremor seen in Er1^Cx/− animals (Figure 1C-D). Further work reveals that myelination is unaffected in the 6-months old Er1^Cx/− mice as assessed by confocal microscopy of fluoromyelin and western blotting of MBP1 protein levels (Supplementary Figure S3A-B). FACS analysis of freshly isolated brain single cell suspensions using antibodies against Calbindin (for Purkinje cells), CD11b+ (for microglia), Annexin V (for apoptosis) and Propidium Iodide (for cell viability) reveals cell death in ~30% of Purkinje cells in the 6-months old Er1^Cx/− brains (Figure 4D). Instead, we find that Er1^Cx/− microglia show no difference in Annexin V or Propidium Iodide indicating that Er1^Cx/− brain microglia is tolerant to DNA damage (Figure 4D and Supplementary Figure S4A). We previously showed that type-I IFNs and their downstream effectors are upregulated in Er1^Cx/− brains (Figure 3A-B) at a level sufficient to
elicit a type I-IFN response in IFN-responsive cells. In Er1Cxl− cerebella, we find that type I-IFN receptor (IFNAR) protein levels are higher in the neuronal population than in microglia cells; the latter is not evident in Er1Cxl− cortices or the hippocampi (Supplementary Figure S4B). Likewise, ifna and ifnb mRNA levels are preferentially higher in the 6-months old Er1Cxl− cerebella compared to age-matched Er1Cxl− hippocampi or cortices indicating the marked sensitivity of neurons inhabiting the cerebellum to microglia-elicited type I-IFN stimuli (Supplementary Figure S5). To test whether type-I IFN-mediated stimuli affect the survival of Purkinje cells, we performed flow cytometry studies to evaluate the IFNAR protein levels in this neuronal cell type. Our analysis in the 6-months old Er1Cxl− cerebella revealed that the great majority of Purkinje cells (~63%) have significantly higher IFNAR protein levels compared to the Purkinje cell population of littermate control animals (Figure 4E-F). Further analysis revealed that out of all IFNAR+ cells in the Er1Cxl brain, Purkinje cells in the cerebellum are the only ones undergoing apoptosis (Figure 4G). Taken together, our findings indicate that ablation of ERCC1 in brain-resident macrophages is associated with loss of Purkinje cells that preferentially respond to microglia-elicited antiviral stimuli.

Er1 Cxl− microglia secrete extracellular vesicles carrying γH2AX-associated chromatin. Secreted Type I IFNs enhance innate immune responses via autocrine and paracrine mechanisms and induce expression of ISGs that trigger apoptosis in targeted cells to inhibit viral replication and spread. In this work, the presence of cytoplasmic chromatin fragments in Er1Cxl− microglia and the observed cell death of DNA repair-proficient Purkinje cells prompted us to examine whether microglia cells secrete extracellular vesicles (EVs) carrying chromatin fragments that, in turn, target IFNAR+ Purkinje cells in Er1Cxl− mice. In support, sucrose gradient ultracentrifugation and FACS analysis for CD11b revealed the increased presence of microglia-derived (CD11b+) EVs in the brain lavage of 6-months old Er1Cxl− mice compared to Er1F− littermate control animals (Figure 5A). Scanning and transmission electron microscopy in this fraction revealed that microglia-derived EVs maintain a size of ~100nm (Figure 5Bi); Er1Cxl− vesicles associate with a higher density cargo as shown by their darker center marking the presence of positively charged DNA moieties (also supported by the enriched phosphorus found by scanning electron microscopy analysis) surrounded by a double membrane (Figure 5Bii, Figure 5iii and Supplementary Figure S6). Western blot analysis confirmed the enrichment of CD11b and ALIX, known to be associated with the endosomal sorting complex required for transport (Figure 5B). FACS analysis with the DNA binding benzothiazole dye (PicoGreen) specific for dsDNA in microglia-derived CD11b+ EVs in the brain lavage of 6-months old Er1Cxl− and Er1F− animals revealed a significantly higher percentage of Er1Cxl− EVs carrying dsDNA moieties (Figure 5C). Further analysis revealed that the great majority of dsDNA moieties in Er1Cxl− EVs associate with the DNA damage marker γH2AX (Figure 5D) and nuclear LaminB1 (Figure 5E) further confirming the nuclear origin of accumulated γH2AX chromatin structures in Er1Cxl− EVs. Western blot analysis of EVs derived from Er1Cxl− and Er1F− brain lavages confirmed the enrichment of γH2AX and LaminB1, P62, a reporter of autophagic activity and β-adaptin, one of the components of the AP-2 adaptor complex that binds ataxia telangiectasia-mutated gene (ATM) to assist with the
tracking of cytoplasmic vesicles \(^{31}\) (Figure 5B). Taken together, our findings suggest that \(Er1^{Cx/−}\) microglia release EVs that are of nuclear origin carrying γH2AX-associated chromatin.

**Microglia-derived \(Er1^{Cx/−}\) EVs target IFNα-responsive Purkinje cells triggering apoptosis.** Next, we sought to examine whether Purkinje cells preferentially receive microglia-derived \(Er1^{Cx/−}\) EVs. To do so, we exposed \(Er1^{F/−}\) cultures of acute brain slices with EVs derived from 6-months old \(Er1^{Cx/−}\) and \(Er1^{F/−}\) brains that were labeled with the lipophilic green fluorescent dye PKH67. To monitor the selective uptake of microglia-derived \(Er1^{Cx/−}\) EVs by calbindin\(^{+}\) cells, we subjected \(Er1^{F/−}\) brain slices to simultaneous 2-3 multiphoton microscopy analysis allowing visualizing at least 200µm detection depths. The latter approach revealed that \(Er1^{Cx/−}\) EVs are preferentially taken up by Purkinje cells when compared to \(Er1^{F/−}\) EVs; of note, the selective targeting of \(Er1^{Cx/−}\) EVs to Purkinje cells is further propagated when \(Er1^{F/−}\) brain slices are exposed to IFNα. (Figure 6A). Further work revealed the co-localization of PicoGreen-stained dsDNA and calbindin in IFNα-treated \(Er1^{F/−}\) cerebella exposed to \(Er1^{Cx/−}\) EVs indicating that the preferential targeting of \(Er1^{Cx/−}\) EVs to Purkinje cells is followed by the release of the \(Er1^{Cx/−}\) EV dsDNA cargo in recipient cells (Figure 6B). Staining of acute brain slices with caspase-3 revealed that the exposure of IFNα-treated \(Er1^{F/−}\) brains to \(Er1^{Cx/−}\) EVs for 6 hours is sufficient to trigger Purkinje cell death (Figure 6C). Thus, microglia-derived \(Er1^{Cx/−}\) EVs preferentially target and released their dsDNA cargo to IFNα-responsive Purkinje cells leading to apoptosis.

**Intranasal delivery of DNase I-loaded EVs delays the DNA damage-driven neurodegenerative features in \(Er1^{Cx/−}\) mice.** To examine the functional contribution of accumulated γH2AX-associated chromatin fragments in the inflammatory response of \(Er1^{Cx/−}\) microglia cells, we used NIH3T3 cells to generate EVs loaded with recombinant (pH-independent) DNase I. To maximize the selective targeting of DNase I EVs to \(Er1^{Cx/−}\) microglia cells, the NIH3T3-derived EVs were also decorated with a custom anti-CD11b peptide derived from a combination of a CD63 binding sequence i.e. CRHSQMTVTSRL \(^{32}\) and the aM1-domain binding peptide CP05 i.e. RKLRSLWRR \(^{33}\). Using this approach, we find that exposure of \(Er1^{Cx/−}\) microglia to DNase I EVs efficiently removes the great majority of cytoplasmic DAPI-stained chromatin fragments and ameliorates the DNA damage-driven LaminB1 disorganization in \(Er1^{Cx/−}\) microglia nuclei (Figure 7A). Consistently, exposure of \(Er1^{Cx/−}\) microglia to DNase I EVs leads to the marked decrease of secreted IFNα levels in the media of these cells (Figure 7B). We reasoned that the NIH- and microglia-derived EVs can be fused, thereby minimizing the DNA load of \(Er1^{Cx/−}\) EVs to maximize the beneficial outcome of DNase I-loaded EVs. In agreement, FACS analysis of purified EVs from \(Er1^{Cx/−}\) brain lavages pre-stained with PicoGreen and co-incubated for 4 hours with DNase I-loaded or control EVs revealed that the microglia-derived (CD11b\(^{+}\)) \(Er1^{Cx/−}\) EVs contain substantially less dsDNA compared to corresponding controls (Figure 7C). Next, we sought to test for the in vivo efficacy of DNase I-loaded EVs in ameliorating the neurodegenerative features seen in the 6-months old \(Er1^{Cx/−}\) mice. To do so, 3-months old \(Er1^{Cx/−}\) animals were exposed to an intranasal delivery of DNase I-loaded EVs twice a week after treatment with a vasoconstrictor to prevent drainage of EVs from blood vessels into the tissues lining the nasal passages. Importantly, we find that treatment of \(Er1^{Cx/−}\) animals with DNase I-EVs leads to the substantial decrease
in Purkinje cell apoptosis (Figure 7D) and in the percentage of activated MHCI\(^+\) CD86\(^+\) Er\(\Gamma\)C\(\kappa\)-/ microglia cells (Figure 7E). Next, we monitored the motor coordination of the 3-months old Er\(\Gamma\)C\(\kappa\)-/ animals treated with DNase I-EVs by measuring the latency to fall from a rotarod apparatus. Our analysis revealed that hind limb coordination in the 3-months old Er\(\Gamma\)C\(\kappa\)-/ mice treated with DNase I-EVs improves over a period of 6 weeks when compared to age-matched Er\(\Gamma\)C\(\kappa\)-/ mice treated with control EVs (Figure 7F). Further work revealed the substantial decrease of PicoGreen-stained dsDNA (Figure 7G) in the microglia-derived (CD11b\(^+\)) EVs from Er\(\Gamma\)C\(\kappa\)-/ mice treated with DNase I-loaded EVs (Figure 7H). Thus, the intranasal delivery of DNase I-loaded EVs substantially reduces type I IFN levels and the microglia-derived dsDNA-carrying EVs, thereby delaying the premature onset of neurodegenerative features seen in Er\(\Gamma\)C\(\kappa\)-/ mice. Taken together, our findings reveal that an intranasal delivery of DNase I EVs substantially decreases Purkinje cell death delaying the premature onset of neurodegenerative features seen in Er\(\Gamma\)C\(\kappa\)-/ mice.

Discussion

Until recently, endogenous DNA damage in post-mitotic neurons was thought to be the primary cause of age-related neurodegenerative disorders seen in DNA repair-deficient patients and accompanying animal models\(^{34}\). Using progeroid animals carrying a ERCC1-XPF DNA repair defect only in tissue-resident macrophages, we show that the gradual accumulation of persistent DNA lesions in brain microglia is sufficient to trigger Purkinje cell death and cerebellar ataxia in mice. Importantly, Er\(\Gamma\)C\(\kappa\)-/ mice are born with Mendelian frequency, they grow normally, are fertile and show no visible pathological signs until adulthood. Beginning at 4-months, however, Er\(\Gamma\)C\(\kappa\)-/ mice exhibit marked signs of cerebellar ataxia associated with Purkinje cell death. The latter is unexpected as in Er\(\Gamma\)C\(\kappa\)-/ animals, Purkinje cells are proficient in DNA repair with no signs of DNA damage accumulation. Instead, we find that irreparable DNA lesions lead to the buildup of H2AX-associated chromatin fragments in the cytoplasm of NER-defective, tissue-resident Er\(\Gamma\)C\(\kappa\)-/ macrophages. Importantly, cytoplasmic H2AX-associated chromatin fragments also accumulate in the microglial cells of naturally aged animals. Nuclear dsDNAs can passively diffuse into the cytosol when the nuclear envelope breaks down in mitotically dividing microglia cells\(^{18}\). In view of the DNA repair defect in Er\(\Gamma\)C\(\kappa\)-/ animals, the latter could also reflect a physiological response of the nucleus to remove byproducts of DNA damage repair or irreversibly damaged DNA fragments. Once accumulated in the cytoplasm of microglia cells, DNA moieties trigger the activation of a type I IFN response. An important finding is that Er\(\Gamma\)C\(\kappa\)-/ microglia cells secrete EVs carrying H2AX-associated chromatin and that Er\(\Gamma\)C\(\kappa\)-/ EVs target IFN\(\alpha\)-responsive Purkinje cells in Er\(\Gamma\)C\(\kappa\)-/ cerebella. Of note, in Er\(\Gamma\)C\(\kappa\)-/ animals, Purkinje cells have significantly higher IFNAR protein levels compared to littermate controls and are the only type of neurons undergoing apoptosis. Besides active targeting, the large size and multiple branching extensions of Purkinje cells likely explains how brain microglia inadvertently affect neuronal homeostasis and why Purkinje cells are particularly prone to microglia-derived stimuli.
So far, the lack of knowledge on the causal mechanisms underlying DNA damage-driven inflammation has hindered the development of rationalized intervention strategies against degenerative diseases, especially at older ages. EVs are non-immunogenic carriers allowing their therapeutic cargo to circulate for extended periods within the body. To alleviate neuroinflammation, we developed an EV-based strategy to deliver recombinant DNase 1 nuclease in inflamed Er1CX/− microglia cells in vivo. We show that the EV-delivered nuclease cargo rapidly removes dsDNAs from the cytoplasm of microglia cells as well of microglia-derived Er1CX/− EVs, thereby maximizing the beneficial outcome of the treatment. At the cellular level, we find that the EV-based removal of cytoplasmic DNAs restores LaminB1 disorganization in the nuclear membrane of Er1CX/− microglia cells and considerably lowers the secreted IFNα levels in microglia cell culture media. In mice, the EV-based therapeutic strategy lowers the percentage of activated Er1CX/− microglia cells leading to the substantial decrease of Purkinje cell death that considerably improves motor coordination in Er1CX/− animals. Thus, as DNA damage-associated chromatin fragments accumulate over time, an EV-based therapeutic scheme is a promising strategy to combat age-related neuroinflammation and improve the outcome of neurodegenerative disorders with aging.

Materials And Methods

Animals. Animals homozygous for the floxed Ercc1 allele (Ercc1 F/F) were intercrossed with mice carrying the CX3CR1-Cre transgene in an Ercc1 heterozygous background (Er1CX/− animals). Mice lacking the CX3CR1-Cre transgene in an Ercc1 homozygous background were used as wild-type controls (Er1F/+). All animals were maintained in grouped cages in a temperature-controlled, pathogen-free animal facility (IMBB-FORTH) on a 12h light/dark cycle and were fed a normal diet (Lactamin, Stockholm, Sweden). Mice had access to water ad libitum. This work received ethical approval by independent Animal Ethical Committee at IMBB-FORTH. All relevant ethical guidelines for the work with animals were adhered to during this study.

Primary cell isolation and cell assays. For single cell isolation from whole brains or selected brain areas (cortex, cerebellum, hippocampus), brains from Er1CX/− and Er1F/+ animals were excised, washed in ice-cold full medium (DMEM containing 10% FBS, 50 µg/ml streptomycin, 50 U/ml penicillin (Sigma) and 2 mM L-glutamine (Gibco)), minced and incubated in 2 mg/ml collagenase type IV at 37°C for 45 min. Collagenase activity was halted with the addition of ice-cold medium to the resultant homogenate. After centrifugation, cells were resuspended in DMEM and further homogenized using a syringe (21G needle). Filtration through a sterile pre-moistened 40µm cell strainer was used to separate the clumped cells, meninges and tissue fragments. Following centrifugation, cells were resuspended in full medium. Primary microglial cells were isolated through CD11b+ magnetic-bead selection after density gradient centrifugation on Percoll. Briefly, Percoll density gradient was prepared in polystyrene tubes by layering 5ml of 35% Percoll solution on top of 3ml 75% Percoll solution, in which the brain cells were resuspended after the final centrifugation. 2ml of 1XPBS were loaded on top of the Percoll density gradient and microglia along with lymphocytes were separated by centrifugation at 800g for 40min at 4°C. The cell band formed between the 75 and 35% layer was harvested, cells were washed with PBS and diluted in
standard growth medium. For CD11b+ microglia cell isolation, the human and mouse CD11b (microglia) Microbeads (MACS, Miltenyi Biotec) were used. Microglia cells at a density of >50,000 cells per well were placed either on poly-L-lysine coated glass coverslips in a 24-well plate or directly in each well of 24-well plate, depending on the experiment. For the LPS treatment (50 ng/ml), microglia cells were plated on wells of a 24-well plate and incubated at 37°C for 24 hrs. For confocal microscopy, microglia cells were plated (and cultured where needed) on poly-L-lysine treated coverslips and were incubated at 37°C for 18 hrs and 3 hrs, respectively. All treatments were performed 20min after their complete attachment on culture plate/slide.

**Immunocytochemistry on primary neuronal and microglia cells.** Whole brain and brain areas (cortex, cerebellum, hippocampus) from Er1<sup>Cx<sup>-/-<sup> and Er1<sup>F<sup>/+<sup> animals were excised and neuronal and microglia cells were isolated as previously described. Once isolated, cells were placed on poly-L-lysine coated coverslips, fixed with 4% F/A for 15min maximum, RT and washed 3X with 1X PBS, for 5min, RT. Permeabilization/Blocking was performed (B1 solution:1% BSA,0,5% Triton in 1X PBS) for 1h, RT. Primary antibodies in B1 solution were added on the coverslips, O/N, at 4°C. The following day, coverslips were washed thrice (B2 solution: 0,5% Triton in 1X PBS) for 10min, RT and secondary antibodies were added, along with DAPI for 2h. Finally, coverslips were washed thrice with B2 solution for 10 min, RT and then they were put on microscope slides with 80% glycerol. Imaging was performed using SP8 confocal microscope (Leica).

**Histology, Immunohistochemistry.** Er1<sup>Cx<sup>-/-<sup> and Er1<sup>F<sup>/+<sup> animals were perfused and their brains were dissected, embedded in gelatin-sucrose, frozen and kept at -80°C. Brains were then cryosectioned (tissue sections were either used directly or stored at -20°C). For the immunohistochemistry experiments, cryosections were stained following two different protocols. According to the Digitonin protocol, cryosections were encircled using Dako – Pen and were then incubated in 1XPBS, RT for 5min. The samples were further incubated in Glycine, for 5min, RT. Three washing steps followed with 1X PBS, for 6min, RT and blocking in a solution of 0.01% Digitonin in 1XPBS, RT, for 45-60 min. The primary antibody solution was placed O/N at 4°C. Samples were immunostained with the corresponding fluorescently labeled antibodies for 2h, RT. A separate 10-min incubation was carried out in DAPI and the slides were coverslipped with 80% Glycerol. Three 6 min washes with a solution of 0.2% Triton in 1X PBS were performed in between incubations. According to the Triton-X protocol, cryosections were encircled using Dako – Pen and post-fixed in ice-cold acetone at -20°C, for 10 minutes. Three washing steps followed with 1X PBS, for 5min, at RT and blocking in a solution of 5% bovine serum albumin (BSA) in 0,5% Triton-X in 1X PBS at RT, for 1 h. Tissue sections were incubated with the primary antibody solution, O/N at 4°C. Samples were immunostained with the corresponding fluorescently labeled antibodies for 1.5h, RT. A separate 10min incubation was carried out in DAPI and the slides were coverslipped with 80% Glycerol. Three 5min washes with 1X PBS were performed in between incubations. For histological analysis of Er1<sup>F<sup>/+<sup> and Er1<sup>Cx<sup>-/-<sup> tissues, samples were fixed in 4% formaldehyde, paraffin embedded, sectioned and stained with Harris’s Hematoxylin and Eosin Y solution. The TUNEL Staining was performed on brain cryosections using the in situ cell death detection kit, Fluorescein (11684817910, Roche Diagnostics,
Mannheim, Germany), according to the manufacturer’s protocol. In brief, the sections were fixed in 4%
formaldehyde for 1 h, rinsed with PBS (5 min, 2 times) RT and permeabilized in 0,1% Triton in 0,1%
sodium citrate at 4° C, for 8 min. The slides were again rinsed with PBS (5 min, 2 times), and incubated in
50µL TUNEL reaction mixture for 1 h at 37°C, dark. The reaction was terminated by rinsing the samples
with PBS (5 min, 2 times) and the sections were sealed and detected by a light microscope. The nuclei
were stained with DAPI (1:500). A positive control was also performed using DNase I (50U/ml), MNase
(10U/ml) and proteinase K (20µg/ml).

Quantitative PCR (QPCR). Quantitative PCR was performed with a CFX Connect Real-Time PCR Detection
system device (BIORAD). The generation of specific PCR products was confirmed by melting curve
analysis. Each primer pair was tested with a logarithmic dilution of a cDNA mix to generate a linear
standard curve (crossing point (CP) plotted versus log of template concentration), which was used to
calculate the primer pair efficiency \( E = 10^{-\left(\frac{-1}{\text{slope}}\right)} \). Hypoxanthine guanine phosphoribosyltransferase1
\( (Hprt-1) \) mRNA was used as an external standard. For data analysis, the second derivative maximum
method was applied: \( \frac{E_{\text{gene of interest}}}{{\Delta CP (\text{cDNA of wt. mice} - \text{cDNA of Ercc1F/−})}} \) gene of interest \( \frac{E_{hprt−1}}{{\Delta CP (\text{cDNA wt.}}}
\text{mice} - \text{cDNA) hprt−1} \).

\( Hprt \text{ F: CCCAACATCAACAGGACTCC, } Hprt \text{ R: CGAAGTGTGGGATACAGGCC, } IFNa \text{ F: CTGCTGGCTGTGAGGACATA, } IFNa \text{ R: GGCTCTCCAGACTTCTGCTC, } IFNb \text{ F: TGAACTCCACCAGCAGACAG, } \)
\( IFNb \text{ R: AGATCTCTGCTGGACCACC, } ISG15 \text{ F: GGTTGCGTGACTAATCCAT, } ISG15 \text{ R: TGGAAAGGGTAAGACCGTCCT, } IFIT2 \text{ F: AGTACAAAGAGGAGGATCTCAGC, } IFIT2 \text{ R: AGGGCAGATGTTGCACATGG, }MX1 \text{ F: GACCATAGGCTTTGACCAAA, } MX1 \text{ R: Agacttgcctttcgggaagcc, } IFIT1 \text{ F: CCAAGTGTGTTTCAATGCTCCT, } IFIT1 \text{ R: GGAAGGGGAAGAGTAGCCGAA, } IRF1 \text{ F: GGGCTGTCAATCTCTGTTCC, } \)
\( IFI207 \text{ F: CAGGCCTCAGCTTCAGACAAC, } IFI207 \text{ R: ATTTCTGGAGGACCCTTGT, } IFI44 \text{ F: AACTGAAGCTCGCAATAATGT, } IFI44 \text{ R: GTAACACAGCAATGCTCTTGT \)

EV isolation, labelling, loading and treatments. EVs were purified using the differential ultracentrifugation
protocol. Briefly culture medium was centrifuged sequentially at 300 g, (10 min), 2000 g (10 min), and
10000 g (30 min) to remove dead cells and cell debris. Extracellular vesicles were isolated with an
ultracentrifugation at 100.000g for 2 h and were then purified using a 90–10% sucrose gradient. Purified
EVs were collected after a final ultracentrifugation at 100.000g for 2 h. All ultracentrifugations were
performed at 4°C. For PKH67 staining, EVs were incubated with PKH67 (company, 500 mL 0.2 mM) for 5
min at room temperature (RT). Labelled EVs were diluted in 500 mL 1% BSA, and then pelleted at 100,000
g, washed with 1 mL PBS to remove excess dye, re-suspended in 1 mL PBS and then pelleted at 100,000
g before final re-suspension. For the extravesicular labelling of EVs against antibodies and
fluorochromes, brain lavage-derived EVs were incubated with both of them at dark, (20 min) RT. For
intravesicular staining, exosomes were fixed with 0.01% formaldehyde (15 min) at 4°C, washed with 0,2%
saponin, 5% BSA in 1xPBS (permeabilization/blocking buffer) and finally isolated after
ultracentrifugation at 4°C, 100.000g for 2 h. EVs were re-suspended and incubated in blocking buffer for
30 min, 4°C and then the immunofluorescently-labeled primary antibody was added. EVs were incubated with the primary antibody for 45 min, 4°C and the secondary fluorescently labeled antibody was added together with DAPI. For EV loading with DNase I and peptide tagging, EVs from NIH3T3 (4x107) cells were isolated and permeabilized with 0.2% saponin for 15 min at RT. Exosomes were then incubated with 30 units of DNase I (Pulmozyme, Roche) and the chimeric peptide (3518; 1µg/1µg EVs):

CRHSQMTVTSRLRKRLRWRR at 4°C for 4 hrs. For the EV-to-EV fusion experiment, EVs were isolated as described above, tagged, loaded, labelled and incubated at 4°C for 4 hrs. EVs from NIH3T3 cells (4x107) were isolated and half of them were loaded with DNase I (Pulmozyme, Roche) and the chimeric peptide (3518, 1µgr/1µg EVs), as it was previously described, while the rest remained empty (naive EVs). Their administration was performed intranasally twice a week for 6 weeks, in 3-month-old mice and its effect on the motor coordination of Er1Cx/− mice was monitored by rotarod latency assay.

**Immunoblot analysis and antibodies.** Immunoblotting brain cells were lysed in RIPA buffer (50 mM Tris-HCl at pH 8, 150 mM NaCl, 0.5% sodium deoxycholate, 1% Nonidet P-40 and 0.1% SDS) supplemented with protease and phosphatase inhibitors (Complete EDTA Free; Roche Applied Science) and equal amounts of proteins (50µgr) were subjected to SDS-PAGE on 7%, 10% and 14% gels and then transferred to PVDF membrane (Amersham Hybond). Membranes were blocked with 5% skimmed milk or 5% BSA in TBST and then incubated with primary antibodies. For western blot analysis of EVs, EV pellets were resuspended in 5X Laemli buffer, sonicated for 5 circles and loaded in the gel. Samples were normalized using antibodies for housekeeping genes (β-tubulin). The image was resolved by ECL system (Thermo Fisher Scientific and Amersham) and detected by ImageBlot (BIORAD). Relative intensity of bands was calculated with Fiji software. Antibodies against: LC3 (C-9, WB:1:500, IF: 1:500), ERCC1 (D-10, WB:1:500, IF: 1:50), LaminB1 (ab16048, WB:1:1000), p62 (SQSTM1, MBLPM045, WB:1:5000, IF:1:10000) and goat anti-rat IgG-CFL 647 (sc-362293, IF: 1:500) were from Santa Cruz Biotechnology. γ-H2A.X (05-636, IF: 1:12000) and pATM (05–740, IF: 1:100) were from Millipore. β-Tubulin (ab6046, WB:1:1000), γ-H2A.X (ab22551, WB: 1:1000), ERCC1 (ab129267, IF: 1:150) and Calbindin (ab22551, WB: 1:1000), CD81 (10037, WB: 1:1000) and Cleaved Caspase-3 (#9661, IF-IC: 1:300, IHC-F: 1:200) were from Cell Signaling Technology. CD45 (H5A5, IF: 1:200) and MAC1 (M1/70.15.11.5.2, IF: 1:200) were from Developmental Studies Hybridoma Bank (DSHB). PKH67 Green Fluorescent Cell Linker Midi Kit (MIDI67) was from Sigma Aldrich. NeuN (26975-1-Ap, IF: 1:50-1:500) was from Proteintech. MBP1 (IF: 1:200) was from Serotec. Fluoromyelin (F34652, IF: 1:300) was from Molecular probes. β adaptin gr(PA1-1066, WB, IF: 2µg/ml), Goat anti-Mouse IgG (H+L) Cross-Adsorbed Secondary Antibody, Alexa Fluor 488 (A-11001, IF: 1:500), Goat anti-Mouse IgG (H+L) Cross-Adsorbed Secondary Antibody, Alexa Fluor 555 (A-21422, IF:1:500), Donkey anti-Rabbit IgG (H+L) Highly Cross-Adsorbed Secondary Antibody, Alexa Fluor 488 (A-21206, IF:1:500), Goat anti-Rabbit IgG (H+L) Highly Cross-Adsorbed Secondary Antibody, Alexa Fluor 555 (A-27039, IF:1:500) and DAPI (62247, IF:1:500) were from ThermoFisher/Invitrogen.
**Flow cytometry.** Cells and EVs from Er1Cx/− and Er1F/+ animals were isolated and stained with fluorochrome conjugated antibodies for 20 min at 4°C in PBS/5% FBS. Antibodies used were: anti-Ly6C (128007, clone HK1.4), anti-IFNAR (Invitrogen, clone MAR1-5A3), anti-CD11b (101212, 101208, clone M1170), anti-MHCII (107606, clone M5/114.15.2) and anti-CD86 (105026, clone GL-1). For intracellular staining, cells were permeabilized and stained using the True-Nuclear Transcription Factor Buffer Set (424401, BioLegend). Secondary antibodies used were: anti-mouse IgG, PerCP (FO114) conjugated goat F(ab)2 and Alexa Fluor 488 (A-11001). Live cells were also stained for Annexin V/PI using the FITC Annexin V Apoptosis Detection Kit (556547, BD Pharmingen). Samples were acquired on a FACS Calibur (BD Biosciences) and analyzed using the FlowJo software (Tree Star). FSC SSC scatter was gated for live cells (those on axes were excluded as cell debris or cell clusters). Positive staining was considered that of MFI more or equal to 10^1. In all cases, the same number of events (for cells or EVs) were acquired from all samples tested per experiment.

**Electron microscopy.** For electron microscopy (EM) analysis of EVs, fixed EVs were deposited on EM grids and were further fixed with glutaraldehyde. Samples were first contrasted in a solution of uranyl oxalate and then contrasted and embedded in a mixture of 4% uranyl acetate and 2% methyl cellulose. EVs were examined under JEM 100C/JEOL/Japan Transmission Electron Microscope. For scanning electron microscopy, isolated EVs were diluted in distilled water and were deposited on glass slides. EVs were examined under Scanning Electron Microscope.

**Multiphoton microscopy**

A single wavelength fs laser source was used to excite simultaneously three- and two-photon fluorescence (3p-F and 2p-F, respectively) and perform three-color multiphoton imaging microscopy in the acute brain slices. For this purpose, a custom-build multiphoton microscope ([Supplementary Figure S7A](#)) was used, based on a 1030 nm fs laser (Pharos-SP, Light Conversion, Vilnius, Lithuania), which is passing through a pair of galvanometric mirrors (6215 H, Cambridge Technology, Bedford, MA, USA) before entering into an inverted microscope (Axio Observer Z1, Carl Zeiss, Jena, Germany). The beam is then reflected by a short pass dichroic mirror (FF700-SDi01, Semrock, Rochester, NY, USA) placed at the turret box of the microscope and is focused in to the sample plane with a 20x 0.8NA objective-lens (Plan-Apochromat 20x/0.8NA, Carl Zeiss). The emitted fluorescence is collected by the same objective and is filtered by a short pass filter (FF01-680/SP, Semrock) to ensure that no laser light is reaching the detectors. Then, the beam is split by a long-pass dichroic mirror (509-FDi01, Semrock), which reflects wavelengths shorter than 509 nm and let pass wavelengths longer than 509nm. The reflected wavelengths are further filtered by a band-pass filter (FF1-458/64, Semrock), which allows passing the wavelengths in the range of 458 ± 32 nm) before reaching a detector, based on a photomultiplier tube PMT-blue (H9305-04, Hamamatsu, Hizuoka, Japan). The transmitting wavelengths longer than 509 nm, are reaching a second long-pass dichroic mirror (FF-580-FDi01, Semrock) which reflects the wavelengths shorter than 580 nm and let pass the wavelengths longer than 580 nm. The reflected wavelengths are further filtered by a band-pass filter (FF01-527/20, Semrock), which allows passing the wavelengths in the range of 527 ± 10 nm) before reaching a second detector, the PMT-green (H9305-04, Hamamatsu).
Finally, the wavelengths longer than 580 nm are passing through a band-pass filter (FF-595/31, Semrock), which allows passing the wavelengths in the range of 595 ± 15.5 nm before reaching the third PMT-red (H9305-04, Hamamatsu). In our experiments the PMT-blue detects the fluorescence emitted from the dye DAPI, while the PMT-green and the PMT-red detect the fluorescence originating from the dyes Alexa Fluor 488 (or PKH67) and Alexa Fluor 555, respectively (Supplementary Figure S7B-C). Quantification of co-localization analysis was performed using JACoP plugin in ImageJ, while the co-localization indicators presented in Figure 6 were calculated using the Manders' coefficient .

**B16 – Blue™ IFN-α/β cell line / SEAP levels detection.** B16-Blue™ IFN-α/β cells derive from the murine B16 melanoma cell line of C57BL/6 origin and allow the detection of bioactive murine type I IFNs by monitoring the activation of the JAK/STAT/ISGF3 pathway and/or IRF3 pathway. Stimulation of B16-Blue™ IFN-α/β cells with murine IFN-α or IFN-β, or type I IFN inducers, such as poly(I:C), poly(dA:dT) or 5’ppp-dsRNA delivered intracellularly, triggers the production of SEAP (Secreted embryonic alkaline phosphatase) by the activation of the IRF-inducible promoter. For the B16-Blue cell cultures, cells were transferred to a T-25 tissue culture flask containing DMEM, 10% (v/v) heat-inactivated FBS, 100 U/ml penicillin, 100 µg/ml streptomycin, 100 µg/ml Normocin and 2mM L-glutamine. No selective antibiotics were added at that point, since cells have to be passaged twice before antibiotics addition. Cells were maintained in growth media supplemented with 100 µg/ml of Zeocin. Growth medium was renewed twice a week and cells were inspected daily. Cells were passaged at a 70-80% confluency. For the detection and quantification of SEAP levels, a cell suspension of 420,000 cells/ml in growth medium was prepared. 75,000 cells in growth medium were added per well (24 well plate) along with culture media from previous experiments and the plate was incubated at 37% in 5% CO2 overnight. QUANTI-Blue solution (1ml QB reagent, 1ml QB Buffer and 98ml sterile water) was prepared the following day, from which 180µl were added in each well of a 96 well plate. Duplicates of induced B16-Blue cells’ supernatants were added, along with a positive (murine IFNα) and negative control (growth medium). The plate was incubated at 37°C for 3h. After 3h incubation, SEAP levels were detected by using a spectrophotometer (Techam) at 620-655 nm.

**Acute brain slices (SNAPSHOT method).** Brains from Er1 Cx−/− and Er1 F/++ animals were excised and sliced (400µm) using a vibratome. Acute brain slices were transferred in a 12-well plate containing fresh ACSF (artificial cerebrospinal fluid). Slices were treated with labelled or unlabeled exosomes, with or without the addition of IFNα protein (12100-1, 4.99x10^6 units/ml, 1:200, PBL assay science) and were then incubated at 37°C for 4h. A 2 min fixation followed, with the slices being transferred in a 12-well plate containing heated (80°C) PFA. The slices were rinsed with 0.1 M PBS to remove residual PFA. The plate was placed on a platform rotator. Slices were permeabilized in 1 phosphate-buffered saline (PBS) tablet, 2 ml Triton X-100 (2% v/v final) and 20 ml DMSO (20% v/v final) for a minimum of 2 hr. Non-target epitopes were blocked by incubation with blocking solution (washing solution with 10% FBS) overnight at RT. Primary antibodies along with DAPI were diluted to the required concentrations in staining solution (washing solution with 2.5% FBS) and each slice was incubated with the diluted primary antibodies in a small plastic bag made by using a Manual Impulse Sealer for 6 to 10 days at 4°C on a platform rotator or a
360° rotisserie wheel. After incubation with primary antibodies, slices were washed with permeabilizing/washing solution three to five times over the course of a day. Secondary antibodies, along with DAPI were diluted to the required concentrations in staining solution and slices were incubated with the diluted secondary antibodies in a small plastic bag for 4 to 6 days at 4°C on a platform rotator or a 360° rotisserie wheel. Fluorophores were protected from exposure to light by wrapping the bags in aluminum foil. The acute brain slices were washed once again with permeabilizing/washing solution three to five times over the course of a day and they were incubated with DAPI for 4 more hours. Acute brain slices were then rinsed three to five times in PBS and prepared for imagining. To image the tissue slices, each slice was placed on a microscope slide prepared with the slide, cover glasses, and Krazy Glue, using a transfer pipet with the tip cut off. A small drop of PBS was added on top of the brain slice before the placement of the cover glass over the brain slice. Finally, corn oil was added to each side of the microscope slide and imaging was performed with a two-photon scanning microscope.

**Rotarod assay.** To assess motor impairment, $Er1^{Cx/-}$ and $Er1^{F/+}$ animals were subjected to rotarod assay test. Briefly, mice need to keep their balance on a rotating rod by walking forward. One day before testing, mice were trained at a constant rotating mode of 5 rpm for 2 min. During testing, mice were initially placed in their lanes, with the rod rotating at 5-rpm constant speed to allow their positioning. Once all mice were able to walk forward, the acceleration test was performed, in which the rod accelerated from 5 rpm to 70 rpm in 60 sec. The time (latency) it took each mouse to fall off the rod rotating under continuous acceleration (from 5 to 70 rpm) was recorded, as well as the reason for trail end (e.g., falling, jumping). The temperature, humidity, ventilation, noise intensity and lighting intensity were controlled and maintained at levels appropriate for mice. All mice were kept in a uniform environment before and after testing to avoid anomalous results being obtained.

**Quantification and Statistical analysis.** A two-tailed t-test was used to extract the statistically significant data by means of the IBM SPSS Statistics 19 (IBM) and the R software for statistical computing ([www.r-project.org](http://www.r-project.org)). Data analysis is discussed also in the Method Details section. Experiments were repeated at least 3 times. The data exhibited normal distribution (where applicable). There was no estimation of group variation before experiments. Error bars indicate standard deviation unless stated otherwise (standard error of the mean; s.e.m.). For animal studies, each biological replicate consists of 3-5 mouse tissues or cell cultures per genotype per time point or treatment. No statistical method was used to predetermine sample size. None of the samples or animals was excluded from the experiment. The animals or the experiments were non-randomized. The investigators were not blinded to allocation during experiments and outcome assessment.

**Declarations**

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Author Contributions
KG, ET, EA, EN, GN, EG, SP, IK, DK and MS performed the experiments and/or analyzed data. GG interpreted data and wrote the manuscript. All relevant data are available from the authors.

Competing interests
The authors declare no competing interests.

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Figures
Loss of ERCC1 in tissue-resident macrophages triggers progressive ataxia in mice. (A). CX3CR1-Cre-driven Rosa-YFP expression in tissue-resident macrophages and (B). ERCC1 protein staining indicating cell type-specific ablation of ERCC1 in CX3CR1-Ercc1F/- cells (Er1CX/-) indicated by the respective arrows. (C). A photograph of a 32-week-old Er1CX/- mouse and its control littermate depicting the hind limb paralysis developed in Er1Cx/- mice. (D). A graph depicting the latency to fall (seconds on the
rotating rod) during Rotarod assessment of the motor coordination of 3-, 6-, 8 and 12-months old Er1Cx/- mice and littermate Er1F/+ controls, n=6 mice per group (E). A photograph showing the kyphosis developed in 40-week-old Er1Cx/- mice. (F). MAC1/CD11b immunofluorescent staining of microglia cells (green arrow) and DAPI staining in cerebellum (cer), cortex (ctx) and hippocampus (hip) Er1CX/- and Er1F/+ brain slices. (G). FACS analysis of freshly isolated brain microglia from an Er1F/+ and an Er1Cx/- mouse of the same age and sex. Higher SSC (black box) indicates higher granularity/vesicle content. (H). Activation status of freshly isolated microglia from Er1Cx/- mice and Er1F/+ littermates. The histogram overlays MHCII (up) and CD86 (down) expression in MHCII+CD86+ cells. The graph shows the respective MFIs from three independent experiments. (I). Graph depicting the percentage (%) of CD11b+ cells in single cell suspensions of 3-6 months old Er1CX/- and Er1F/+ mouse brains. Statistical analysis indicated no significant differences. (J). Representative image of a Hematoxylin- and Eosin-stained sagittal section of 8-months old Er1CX/- and Er1F/+ mice. (K). Graph depicting the percentage (%) of Ly6C+ cells in single cell suspensions of Er1CX/- and Er1F/+ mouse brains. Statistical analysis indicated no significant differences. The asterisk “*” indicates a p-value ≤ 0.05, two-tailed Student’s t-test.
Persistent DNA damage triggers the accumulation of cytoplasmic chromatin fragments in Er1Cx/- microglia. (A). FACS analysis for γH2AX, NeuN and MAC1/CD11b in freshly isolated brain single cell suspensions of Er1F/+ and Er1CX/- brains. The histogram depicts the representative overlay of γH2AX fluorescence intensity distribution in CD11b or NeuN cell populations in the 6-months old Er1CX/- and Er1F/+ brains. The graph depicts the percentage (%) of γH2AX+ microglia cells (CD11b) and neurons.
(NeuN) in the 3-, 6-, 8- and 12-months old Er1CX/- and Er1F/+ brains (as indicated). (B). Immunostaining for MAC1/CD11b (green), γH2AX (red) and pATM (red) on brain single cell suspensions. The graphs depict the percentage (%) of cells with cytoplasmic or nuclear γH2AX or pATM puncta (C). Immunostaining for MAC1/CD11b (green) and intensified DAPI in freshly isolated microglia cells from Er1CX/- and Er1F/+ brains; the inlay depicts the DAPI-stained chromatin in the cytoplasm of Er1CX/- microglial cells. The graph depicts the percentage (%) of cells with cytoplasmic chromatin fragments. The white line is set at 5μm scale. Error bars indicate S.E.M. among n≥ 3 replicates. The asterisk “*” indicates a p-value ≤ 0.05, two-tailed Student’s t-test.
Figure 3

Accumulation of cytoplasmic chromatin fragments triggers a type I IFN response in Er1Cx/- microglia. (A). Type I IFN bioactivity (B16 reporter assay OD fold change) in 6-months old Er1Cx/- and aged-matched Er1F/- brain lavages (n=4). (B). Quantitative PCR evaluation of the mRNA levels of interferon signature genes in the brain lysates of 6-months old Er1F/+ and Er1CX/- mice (as indicated; RFU: relative fluorescent units; n=3). (C). FACS analysis of pSTING intracellular levels in CD11b+ brain cells derived...
from the cerebella (CER) or cortices (CTX) of 6-months old Er1F/+ and Er1CX/- mice (n=3). (D). Immunostaining for LC3 and LaminB1 in freshly isolated microglial cells of 6-months old Er1CX/- and Er1F/+ brains. The graph depicts the percentage (%) of cells with reorganized LaminB1 (E). Immunostaining for pATM and p62/SQTM1 in freshly isolated microglial cells of 6-months old Er1CX/- and Er1F/+ brains. The graph depicts the percentage (%) of Er1CX/- or Er1F/+ cells with cytoplasmic DAPI+ P62+ chromatin structures. (F). FACS analysis of Lysosensor Green in freshly isolated brain microglia derived from 6-months old Er1F/+ and Er1CX/- brains. The representative histogram overlay of Lysosensor Green MFI shows the significant decrease in Er1CX/- cells. The graph depicts the percentage (%) of cells with lower Lysosensor Green MFI. (G). Immunostaining for γH2AX (green) and cytoplasmic DAPI staining of freshly isolated microglia cells derived from 24-months old naturally aged cerebella. (H). FACS analysis of pSTING intracellular levels in CD11b+ cells derived from the cerebella of 2- and 24-months naturally aged mice (n=3). The white line is set at 5μm scale. Error bars indicate S.E.M. among n≥3 replicates. The asterisk “*” indicates a p-value ≤ 0.05, two-tailed Student's t-test.
Aged microglia elicit antiviral-like response that preferentially kills Purkinje neurons. (A). Immunostaining for γH2AX (red), calbindin (green) and DAPI in brain sections derived from Er1CX/- and Er1F/+ mice (n=3 animals per group, 2 sections per mouse). (B). Representative images of Apoptotic cell death (indicated by green-colored arrowheads; TUNEL) and DAPI staining in brain sagittal sections of distinct brain areas associated with motility regulation (cerebellum, cerebral cortex) of 6-months old Er1CX/- and Er1F/+ mice.
animals. The graph depicts the number of TUNEL positive cells in the cortex and cerebellar brain sections of 3- and 6-months old mice (n=3 animals per group, 4 sections per mouse). (C). Immunostaining for apoptotic cleaved-caspase 3 (red) and DAPI in sagittal brain sections of 6-months old Er1CX/- animals and Er1F/+ littermate controls; red arrowheads indicate the caspase 3 staining in the Purkinje cell layer of Er1CX/- cerebellar cortex. (D). Representative FACS plots of Annexin V-Propidium Iodide (PI) scatter of calbindin+ cells. The graph depicts the percentage (%) Annexin V+PI+ Purkinje cells (E). A representative histogram overlay (FACS analysis) of freshly isolated single cell suspensions from Er1CX/- and Er1F/+ brains stained for IFNAR and calbindin. The graph depicts the percentage (%) of IFNAR+ Purkinje cells (n=4). (F). A representative histogram overlay of IFNAR expression in Purkinje cells derived from 6-months old Er1CX/- and Er1F/+ animals. The graph depicts the MFI of IFNAR expression in calbindin+ cells (n=4). (G). FACS representative plots of Er1CX/- freshly isolated brain single cell suspensions stained for IFNAR, calbindin, Annexin V and PI. The white line is set at 5μm scale. Error bars indicate S.E.M. among n≥3 replicates. The asterisk “*” indicates a p-value ≤ 0.05, two-tailed Student's t-test.
Figure 5

Er1CX/- microglia release EVs carrying dsDNAs associated with DNA damage. (A). FACS analysis of extracellular vesicles purified with sucrose gradient centrifugation from 6-months old Er1F/+ and Er1CX/- brain lavages stained for CD11b. (B). Scanning (i) and Transmission (ii-iii) electron microscope images of purified circulating EVs from Er1F/+ Er1CX/- brain lavages. Western blot analysis of CD11b, ALIX and Tubulin proteins levels in purified circulating EVs from Er1F/+ Er1CX/- brain lavages (n=3). (C). FACS
analysis of purified EVs from Er1F/+ and Er1CX/- brain lavages stained for CD11b and PicoGreenTM (n=3). The graph depicts the % of dsDNA carrying CD11b+ EVs. (D). FACS analysis of purified EVs from 3-, 6- and 8-months old Er1CX/- brain lavages stained for γH2AX (intravesicularly), CD11b and PicoGreenTM. The histogram overlays depict the γH2AX MFI distribution between PicoGreenTM - and PicoGreenTM + EV populations across the indicated age groups. (E). FACS analysis of purified EVs from Er1F/+ and Er1CX/- brain lavages stained for CD11b, PicoGreenTM and LaminB1. The graph depicts the % of laminB1+ EVs (n=3). (F). Western blot analysis of γH2AX, LaminB1, β-adaptin, P62 and tubulin in purified EVs from Er1F/+ and Er1CX/- brain lavages (n=3). The asterisk “*” indicates a p-value ≤ 0.05, two-tailed Student’s t-test.
Figure 6

Simultaneous 2-photon and 3-photon excited fluorescence microscopy images of Er1F/+ acute brain slices. The Er1F/+ acute brain slices are cultured in the presence of Er1F/+ and Er1CX/- EVs with (+) or without (-) rIFNα as indicated and stained for (A). PKH68TM (green) (pre-labelled EVs), Calbindin (red) and DAPI where the graph depicts the co-localization co-efficient between PKH-67 and Calbindin (n=7) (B). PicoGreenTM (pre-labelled EV dsDNA), Calbindin (red) and DAPI where the graph depicts the co-
localization co-efficient between PicoGreenTM and Calbindin (n=7) and (C). Caspase-3 (green), Calbindin (red) and DAPI where the graph depicts the co-localization co-efficient between Caspase-3 and Calbindin (n=7). The white line is set at 10μm scale. Error bars indicate S.E.M. among n≥3 replicates. The asterisk “*” indicates a p-value ≤ 0.05, two-tailed Student’s t-test.

Figure 7
Er1CX/- microglia-derived EVs preferentially target IFNα-responding Purkinje cells leading to apoptosis. (A). Immunofluorescence of LC3 (red), Lamin B1 (white) and intensified DAPI on freshly isolated microglia cells derived from Er1F/+ and Er1CX/- brains, cultured in the presence of DNase I-loaded or empty (naïve) EVs. (B). Type I IFN bioactivity in the lavage of Er1CX/- brains injected intranasally with DNase I-loaded or naïve EVs (30 Units of DNase I/administration, 12 intranasal instillations, once every 3 days) (n=4) (C). FACS analysis of purified, PicoGreenTM-labelled EVs from Er1CX/- brain lavages incubated with DNase I-loaded or naïve EVs (n=4) to assess the exosome-to-exosome fusion. (D). FACS analysis of freshly isolated cerebella from Er1CX/- brains treated with DNase I-loaded or naïve EVs stained for Calbindin, Annexin and PI (as indicated). (E). FACS analysis of freshly isolated brain single cell suspensions stained for CD11b, MHCII and CD86. The graph depicts the percentage % of MHCII+CD86+ microglial cells of 6-months Er1CX/- mice treated with DNase I-loaded or naïve EVs. (F). Graph depicting the latency ratios during Rotarod Assay (seconds on the rod before treatment/ seconds on the rod after treatment) of Er1Cx/- and littermate animals treated with DNase I-loaded or naïve EVs (no of mice / group=3; No of experiments=3). (G). FACS analysis of purified EVs from Er1F/+ and Er1CX/- brain lavages of mice treated with DNase I-loaded and naïve EVs stained for CD11b and PicoGreenTM. Representative plots depicting the elimination of dsDNA moieties upon in vivo DNase I administration. The graph depicts the percentage (%) of PicoGreenTM + EVs. Immunofluorescence of MAC1/CD11b (green), Lamin B1 (red) and intensified DAPI on freshly isolated brain single cell suspensions of Er1F/+ and Er1CX/- mice treated with DNaseI-loaded or naïve EVs. The white line is set at 5μm scale. Error bars indicate S.E.M. among n≥3 replicates. The asterisk “∗” indicates a p-value ≤ 0.05, two-tailed Student’s t-test.

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