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Nitric oxide mediates glial-induced neurodegeneration in Alexander disease

Liqun Wang1, Tracy L. Hagemann2, Hermann Kalwa3, Thomas Michel3, Albee Messing2 & Mel B. Feany1

Glia play critical roles in maintaining the structure and function of the nervous system; however, the specific contribution that astroglia make to neurodegeneration in human disease states remains largely undefined. Here we use Alexander disease, a serious degenerative neurological disorder caused by astrocyte dysfunction, to identify glial-derived NO as a signalling molecule triggering astrocyte-mediated neuronal degeneration. We further find that NO acts through cGMP signalling in neurons to promote cell death. Glial cells themselves also degenerate, via the DNA damage response and p53. Our findings thus define a specific mechanism for glial-induced non-cell autonomous neuronal cell death, and identify a potential therapeutic target for reducing cellular toxicity in Alexander disease, and possibly other neurodegenerative disorders with glial dysfunction.
A strocytes are now known to influence a variety of complex nervous processes including sleep, learning and memory, and breathing. At the molecular and cellular biological levels, astrocytes play a critical role in regulating the permeability of the blood–brain barrier, responding to oxidative stress, modulating synaptic transmission and controlling synaptogenesis and synaptic pruning. Given the plausible role of many of these processes in neuronal dysfunction and death during neurodegeneration, astrocytes seem poised to be critical players in neurodegenerative disorders. Indeed, morphological changes in astrocytes are prominent in all neurological disorders characterized neuropathologically by significant cell loss or tissue damage. However, rigorously defining a role for astrocytes in neurodegenerative disorders has been challenging. First, in most disorders, neuronal pathology is quite prominent, raising the possibility that changes in glial cells are secondary phenomena. Second, it has been challenging to identify the molecular underpinnings of glial-induced neuronal death and test cell-type-specific mechanisms in vivo.

To address these limitations, we have developed a model of a glial-based neurodegenerative disorder, Alexander disease, in the genetically accessible model organism Drosophila. Alexander disease offers a unique opportunity to study glial-mediated neurodegeneration, because the primary abnormality in patients is glial: dominant missense mutations in the gene encoding glial fibrillary acidic protein (GFAP), the intermediate filament of astrocytes. Clinical presentations of this rare, but informative, disorder range from infants with prominent leukodystrophy and seizures to adults with relative preservation of myelin and a movement disorder. In all cases, the specific neuropathological manifestation of Alexander disease is the formation of eosinophilic, beaded inclusions termed Rosenthal fibres within astrocytes. Given evidence from patients and mouse models for a toxic dominant gain-of-function mechanism related to abnormal aggregation and toxicity of GFAP in Alexander disease, we have modelled the disorder by expressing disease-linked forms of human GFAP in fly glia. Our Drosophila model replicates key features of the human disorder, including formation of Rosenthal fibre-like inclusions, seizures and non-cell autonomous neurodegeneration. At the molecular level, protein misfolding, oxidative stress and dysregulated glutamate transport are critical mediators of disease pathogenesis in our fly model, correlating well with results from mice and analysis of human tissue.

An important, perhaps the most important, strength of studying human disease in Drosophila is the ability to perform unbiased forward genetic screens, which have the potential to provide hypothesis-independent insights into disease pathogenesis. Accordingly, we have performed genetic screening in our fly model of Alexander disease. Through our screen, we identified nitric oxide (NO) as a critical mediator of non-cell autonomous neurodegeneration. Further, to dissect the cell-type specificity of the effect, we develop a system to manipulate gene expression independently in neurons and glia. Using a newly developed dual-expression system, we demonstrate a key role for NO synthesized and released in glia acting through neuronal NO synthase (NOS) to promote neuronal cell death.

Results

NO signalling is upregulated in Drosophila Alexander disease. Our Drosophila model of Alexander disease is based on expression of human disease-associated mutant GFAPR79H in fly glia using the standard GAL4/UAS bipartite expression system and the glial-specific driver repo-GAL4 (refs 8,13). Our model recapitulates multiple aspects of authentic Alexander disease, including glial toxicity and secondary non-cell autonomous neurodegeneration. To identify genetic modifiers of GFAP toxicity in an unbiased pattern, we performed an unbiased forward genetic screen using a collection of 2,239 transgenic RNA interference (RNAi) lines. To monitor toxicity of GFAP to glia under screening conditions, we used the transgenic caspase reporter developed by Williams et al. By monitoring caspase activation in 10-day-old adult flies expressing GFAPR79H and RNAi targeting individual genes in glial cells, we recovered a number of candidate suppressors and enhancers of GFAP toxicity. One of these candidate suppressors was nitric oxide synthase (Nos).

Because NO represents a novel, but plausible, glial-derived neuronal cell death signalling molecule in Alexander disease, we explored NO signalling further in our Alexander disease model. We first examined levels of Nos protein (encoded by a single Nos gene in flies) in control and GFAPR79H transgenic animals using a rabbit polyclonal antibody to Drosophila Nos. A significant increase in Nos in GFAPR79H flies compared with controls was detected by western blot analysis (Fig. 1a). Double-label immunofluorescence using neuronal- and glial-specific markers confirmed the presence of Nos in glial cells (Supplementary Fig. 1a). The specificity of the antibody was confirmed on both western blot and immunostaining using the Nos allele, a deletion mutant of Nos removing the entire oxygenase domain. To probe Nos activity in our GFAPR79H transgenic flies, we performed nicotineamide adenine dinucleotide phosphate (NADPH)-diaphorase histochemistry. Consistent with Nos protein upregulation, significantly increased NADPH-diaphorase staining was present in GFAPR79H transgenic flies (Fig. 1b). In addition, with a sensitive and specific NO sensor, Cu2(FL2E)19, we observed significantly increased NO production in the brains of GFAPR79H transgenic flies compared with controls (Fig. 1c). To explore potential mechanisms responsible for upregulation of Nos in our transgenic animals, we evaluated the activity of a well-characterized reporter of STAT expression, 10XSTAT-GFP. We observed upregulation of STAT expression specifically in GFAPR79H transgenic flies (Supplementary Fig. 1b).

We then examined the effect of GFAPR79H expression on cellular pathways downstream of NO production. NO reacts with superoxide (O2-) to make peroxynitrite (ONOO-), a reactive oxidant and cytotoxic agent that can nitrate tyrosine residues of proteins; these modified proteins represent an important indicator of nitrosative stress. Western blot analysis performed with a previously characterized polyclonal anti-nitrotyrosine antibody demonstrated increased levels of nitrotyrosine-containing proteins in GFAPR79H transgenic flies compared with controls (Fig. 1d). Soluble guanylate cyclase (sGC) is an important specific target of NO-mediated nitrosylation. Nitrosylation of sGC activates the enzyme and increases cyclic guanosine monophosphate (cGMP) production. cGMP, in turn, is an important second messenger for a number of downstream targets, including cyclic nucleotide-gated channels (CNGs). Using a sensitive chemiluminescent immunoassay, we found significantly increased levels of cGMP in brains of GFAPR79H flies compared with controls (Fig. 1e).

We also investigated levels of Nos protein, nitrotyrosine-containing proteins and cGMP in transgenic flies that express wild-type human GFAP (GFAPWT) at levels equivalent to GFAPR79H, and show no evidence of toxicity or inclusion formation (Supplementary Fig. 1f, GFAPWT). We detected no significant difference in Nos protein levels, nitrotyrosine-containing proteins or cGMP levels between GFAPWT transgenic flies and control flies (Supplementary Fig. 1c–e), demonstrating specificity for disease-associated mutant GFAPR79H. However, as seen in murine models, increased expression of wild-type human GFAP does result in the
**Figure 1 | NO signalling is upregulated in GFAPR79H transgenic flies.** (a) Western blot shows increased expression of Nos in GFAPR79H transgenic flies. The blot was reprobed for actin to illustrate equivalent loading. Quantification of Nos protein levels in control and GFAPR79H transgenic flies is shown in the graph at the right. *P = 0.0078, t = 6.371, df = 3; unpaired t-test, n = 4 per genotype. (b) NADPH-diaphorase staining is increased in the brain of GFAPR79H transgenic flies. Quantification is shown in the graph at the right. *P = 0.0021, t = 5.800, df = 5; unpaired t-test, n = 6 per genotype. (c) Time-lapse imaging reveals increased nitric oxide production *in vivo* in GFAPR79H transgenic flies with a nitric oxide sensor Cu2(FL2E). Scale bar, 10 μm. *P < 0.05, t = 2.833 (6 min); 3.173 (8 min); 3.137 (10 min); two-way ANOVA, n = 7 per genotype. (d) Nitrotyrosine (N-tyr)-modified proteins are increased in GFAPR79H flies. *P = 0.0273, t = 2.902, df = 6; unpaired t-test, n = 4 per genotype. (e) Significantly increased cGMP in GFAPR79H transgenic flies as assayed by the chemiluminescent immunoassay kit. **P < 0.001, t = 6.109, df = 16; unpaired t-test, n = 9 per genotype. Flies were 20 days old in all panels. Control (Ctrl) is repo-GAL4/+ . GFAPR79H is repo-GAL4, UAS-GFAPR79H/+ .

formation of Rosenthal fibre-like inclusions (Supplementary Fig. 1f, 2 × GFAPWT). We further investigated three additional Alexander disease-associated mutant forms of GFAP, GFAPR239H, GFAPL352P and GFAPA364P. Each of these mutant forms of GFAP formed Rosenthal fibre-like inclusions in *Drosophila* glia (repo-GAL4 driver; Supplementary Fig. 1g,h). We also observed inclusion formation when human mutant GFAP was expressed in specific glial subtypes (Supplementary Fig. 1i,j), including astrocyte-like glia (alrm-GAL4 driver) and ensheathing glia (mx0709-GAL4 driver)26. However, we did not observe significant cellular toxicity with expression in either of these glial subtypes, most likely due to lower levels of transgene expression mediated by these drivers, as demonstrated by reduced β-galactosidase (β-gal) expression mediated by alrm-GAL4 compared with repo-GAL4 (Supplementary Fig. 1k).

Modulation of Nos signalling alters GFAP toxicity *in vivo*. To determine whether GFAP toxicity is influenced by Nos activity, we first manipulated the single *Drosophila* Nos gene genetically. Overexpressing Nos in glia using a UAS-Nos transgene and the glial driver repo-GAL4 markedly increased the number of TdT-mediated dUTP nick end labelling (TUNEL)-positive cells, while reducing Nos with either of the two independent transgenic RNAi lines significantly decreased the number of TUNEL-positive cells in the brains of GFAPR79H flies (Fig. 2a,e; Supplementary Fig. 2a). Increasing Nos in the absence of transgenic human GFAPR79H did not produce toxicity (Fig. 2a). Western blot analysis demonstrated marked reduction in Nos levels following RNAi-mediated gene knockdown (Supplementary Fig. 2b). To ensure that modulation of GFAP toxicity was not simply due to alterations in GFAP protein levels, we examined GFAP expression in genetically modified backgrounds and observed equivalent GFAP protein levels (Supplementary Fig. 2c). Consistent with positive TUNEL staining, the nuclei of dying cells were often condensed (43.36 ± 2.06%) and fragmented (32.82 ± 1.64%) in GFAP transgenic flies (Supplementary Fig. 2d). Significant numbers of
cells with cleaved caspase (Dcp-1) were present, and activation of hid and th (also known as Diap1), key regulators of apoptosis, was observed using transgenic reporter lines 27,28 (Supplementary Fig. 2e–g). To explore the specificity of our findings, we altered activity of the Nos pathway in flies expressing mutant human ataxin 3, the aggregation-prone polyglutamine expanded protein expressed in patients with spinocerebellar ataxia type 3, in glia 29.

In addition to cell death, our Alexander disease model flies also exhibit seizure activity 8. We therefore assessed the influence of Nos on seizure frequency in GFAPR79H transgenic flies. As with

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Nos pathway modifiers, including Nos overexpression and knockdown, did not alter the toxicity of mutant ataxin 3 as monitored by vacuolation in the brains of transgenic animals (Supplementary Fig. 2h).

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Figure 2 | Genetic and pharmacological modulation of Nos activity alters GFAPR79H toxicity. Nos was overexpressed with a UAS-Nos transgene (Nos OE) and reduced with UAS-Nos RNAi lines (Nos RNAi). GFAP represents UAS-GFAPR79H. (a) The number of TUNEL-positive cells is increased in GFAPR79H transgenic flies with overexpression of Nos (Nos OE) and decreased with reduced expression of Nos (Nos RNAi). **P<0.0001, F = 72.53, df = 29; n = 6 per genotype. Flies were 20 days old. Genotypes are indicated in the figure label. Driver: repo-GAL4. (b) Seizure frequency is increased with overexpression of Nos (Nos OE) and decreased with reduced expression of Nos (Nos RNAi). *P = 0.0264, x^2 = 4.926, df = 1; **P<0.0001, x^2 = 24.731, df = 1; n>100 per genotype. Flies were 1 day old. Genotypes are indicated in the figure label. Driver: repo-GAL4. (c) The Nos inhibitor L-NAME reduces the number of TUNEL-positive cells in GFAPR79H transgenic flies in a dose-dependent manner while D-NAME, the inactive enantiomer, shows no effect. *P<0.01, **P<0.001, F = 12.41, df = 27; n≥6 per condition. Flies were 15 days old. Genotype: repo-GAL4, UAS-GFAPR79H/+ . (d) Seizure frequency is reduced with L-NAME administration, but not with D-NAME. **P<0.0001, x^2 = 24.987, df = 1; x^2 = 24.083, df = 1; n>100 per condition. Flies were 3 days old. Genotype: repo-GAL4, UAS-GFAPR79H/+ . (e) Overexpression of Nos (Nos OE) increases, while reduced Nos expression (Nos RNAi) decreases the total number of TUNEL-positive cells (total, left), including both dying neurons (middle) and glia (right) in GFAPR79H transgenic flies. **P<0.0001, F = 13.18; 9.627; 17.06; df = 23; 23; 23; n = 6 per genotype. Flies were 20 days old. Genotypes are indicated in the figure label. Driver: repo-GAL4. + : repo-GAL4, UAS-GFAPR79H/+ . Statistical tests: one-way ANOVA with Tukey’s multiple comparison test in a, c, and e; x^2-test in b and d.
cell death, increased Nos exacerbated seizure activity, while Nos knockdown decreased seizure frequency (Fig. 2b). Thus, manipulation of Nos levels influences disease-relevant behaviour in our model.

Nos activity can also be controlled pharmacologically. 1-NAME is a well-characterized Nos inhibitor with documented activity in *Drosophila*; D-NAME is the inactive enantiomer of 1-NAME[29]. To extend our genetic results and assess Nos as a potential therapeutic target, we fed flies orally with either 1-NAME or D-NAME. We found that inhibition of Nos activity with 1-NAME decreased cell death and seizure frequency, while D-NAME did not alter cell death or the number of seizures in GFAP R79H transgenic flies (Fig. 2c,d; Supplementary Fig. 2i). 1-NAME had no effect on the levels of GFAP protein (Supplementary Fig. 2j). In our fly model of Alexander disease, expression of GFAP R79H specifically in glia triggers non-cell autonomous neurodegeneration,[8], as also occurs in human patients[31]. Since NO is a diffusible signalling molecule, we hypothesized that NO released from glia triggers neuronal cell death. To begin to test our hypothesis, we examined glial and neuronal cell death in GFAP R79H flies with altered levels of Nos. We and others have previously observed loss of endogenous cell-type-specific markers in apoptotic cells.[8,32]. We therefore introduced β-gal, expressed using *UAS-lacZ* and *repo-GALA4*, as a transgenic marker and performed double labelling with TUNEL to identify dying glia specifically and distinguish neuronal from glial cell death (Supplementary Fig. 2k). With our cell-type-specific labelling of apoptotic cells, we observed that overexpression of Nos in glia significantly promoted neuronal cell death, consistent with a non-cell autonomous effect of glial NO release (Fig. 2e, middle).

However, we also observed modestly increased glial cell death with Nos overexpression, as well as reduction in both neuronal and glial cell death with Nos knockdown. These data are consistent with effects of Nos on toxicity in both cell types. Thus, to test the specific role of NO in neurons we wanted to block the effects of glial-derived NO in neurons. Given our evidence for increased cGMP in GFAP R79H transgenic flies (Fig. 1e), we decided to focus on the NO target sGC. In *Drosophila*, genes encoding the alpha and beta subunits of sGC, *Gyc* a and *Gyc* b, have been identified and functionally characterized.[33]. To independently and simultaneously manipulate gene expression in neurons and glia, we constructed a dual-glial–neuronal expression system by combining the GAL4/UAS system and the Q system[12,34]. The Q system is a bipartite expression system, which is conceptually similar to the GAL4/UAS system but instead relies on the regulatory genes from the *Neurospora qa* gene cluster. The GAL4/UAS and Q systems do not interact and can thus be used for independent simultaneous manipulation of gene expression.[34].

Our strategy for independent manipulation of glial and neuronal gene expression is illustrated in Fig. 3a. GFAP R79H (QUAS-GFAP R79H) was expressed in glia using the Q system glial driver *ET31-Q*.[34]. These transgenic flies have frequent Rosenthal fibre-like inclusions and increased seizure frequency (Supplementary Fig. 3a–c), recapitulating the phenotypes observed when GFAP R79H is expressed in glia with *repo-GAL4* driver.[8]. Transgenic RNAi targeted to *Gyc* 99B and *Gyc* b 100B was then expressed in neurons using the *elav-GAL4* driver. β-Galactosidase was expressed in neurons via *UAS-lacZ* and *elav-GAL4* as a control for expression of an unrelated protein in neurons. Using the dual-expression system, we found that GFAP toxicity was markedly reduced when sGC expression was reduced specifically in neurons using either of the two independent RNAi lines targeting *Gyc* 99B or *Gyc* b 100B (Fig. 3b; Supplementary Fig. 3g, left). Reduction in GFAP toxicity was not due to alterations in GFAP protein levels, as determined by western blot analysis (Supplementary Fig. 3d). We verified significant reductions in messenger RNA for *Gyc* 99B and *Gyc* b 100B with expression of transgenic RNAi targeting these loci using RT–PCR (Supplementary Fig. 3e,f). Taken together, these data support our hypothesis that glial-derived NO activates neuronal sGC to mediate glia-induced non-cell autonomous neuronal cell death. As with manipulation of Nos in glia, sGC knockdown in neurons rescued both neuronal and glial cell death (Fig. 3b; Supplementary Fig. 3g, middle and right). Identification of TUNEL-positive cells as neurons or glia in Fig. 3b was based on morphology, as informed by cell-type-specific labelling experiments (Fig. 2e), given difficulties encountered in introducing an additional transgene into the complex genotypes illustrated in Fig. 3a.

iNOS signalling is upregulated in Alexander disease mice. To further investigate the role of NO signalling in Alexander disease, we probed the pathway in two well-characterized mouse models of the disorder. Mice expressing either a disease-associated point mutant (GFAP+/R236H) or wild-type human GFAP (GFAP Tg) develop clinical and pathological features reminiscent of human Alexander disease, with GFAP Tg mice exhibiting more pathology compared with mice heterozygous for the knock-in allele[6]. Compared with the single *Nos* locus in *Drosophila*, mammals have three distinct *Nos* genes with encoded proteins having different cellular localization, regulation, catalytic properties and inhibitor profiles: neuronal NOS (nNOS or NOS1), inducible NOS (iNOS or NOS2) and endothelial NOS (eNOS or NOS3).[35]. We focused our investigations first on nNOS and iNOS, because in the brain eNOS is mainly expressed in vascular endothelial cells, while nNOS and iNOS are mainly expressed in neurons and astrocytes, respectively, and have been implicated in glial and neuronal apoptosis.[35]. Using an nNOS-specific antibody, we were not able to detect differences in nNOS expression between Alexander disease model mice and controls (Supplementary Fig. 4a). In contrast, we observed significant induction of iNOS in Alexander disease model mice. Western blot analysis revealed increased iNOS expression in the cortex of GFAP Tg mice compared with age-matched littermate controls (Fig. 4a). Similarly, double-label immunofluorescence revealed increased iNOS in astrocytes (Fig. 4c; Supplementary Fig. 8), which was validated by a second antibody reagent (Supplementary Fig. 4b).

We also examined downstream components of the pathway and observed a prominent increase in nitrotyrosine-containing proteins by both western blot analysis (Fig. 4b) and immunostaining with a polyclonal anti-nitrotyrosine antibody[22–24] (Fig. 4c; Supplementary Fig. 8). To confirm and extend prior characterization[22–24], the anti-nitrotyrosine antibody was validated using nitrated BSA on immunoblot and tissue sections or by reducing nitrotyrosine to aminotyrosine on Western blot analysis (Fig. 4b). In the animals overexpressing wild-type GFAP[6,7], iNOS pathway induction, we also examined brain tissue from a widely used transgenic mouse model of tauopathy based on expression of...
human P301L mutant tau. These mice show significant astrogliosis and formation of insoluble tau-containing protein aggregates at 7 months of age. We did not observe significant upregulation of iNOS in astrocytes of tauopathy model mice (Supplementary Fig. 4g).

Oxidative stress and p53 act downstream of Nos. We have previously demonstrated a critical role for oxidative stress in mediating cell death in our Drosophila model of Alexander disease. To explore a possible link between Nos and oxidative stress in GFAPR79H transgenic flies, we expressed a sensitive oxidative stress reporter, GstD1-lacZ, in which expression of β-gal (lacZ) is controlled by the GstD1 promoter and increases in response to oxidative stress. When we performed double-label immunofluorescence for β-gal and cell-type-specific markers (repo to identify glia and elav to mark neurons), we detected activation of the reporter in glia specifically (Fig. 5a). When we overexpressed Nos together with GFAPR79H in glia using the repo-GAL4 driver, we further activated the reporter. Conversely, knockdown of Nos decreased reporter activation. Overexpressing Nos in the absence of GFAPR79H did not activate the reporter (Fig. 5b). These data are consistent with a role for oxidative stress downstream of Nos in promoting glial toxicity.

NO can promote apoptosis through DNA damage and p53 activation. To determine whether DNA damage and p53 activation are mediators of cell death in GFAPR79H flies, we first examined levels of the Drosophila DNA damage response marker phosphorylated H2Av (H2Av-pS137) and p53 using western blot analyses. Both markers were significantly increased in GFAPR79H transgenic flies compared with aged-matched control flies (Fig. 6a,c). Immunohistochemical staining also revealed robust activation of H2Av-pS137 and p53 in GFAPR79H transgenic flies (Supplementary Fig. 5a,b). Using double-label immunofluorescence, we determined that both H2Av-pS137 and p53 were increased in glial cells (Fig. 6b,d). Further, H2Av-pS137 and p53 colocalized in the same glial cells (Fig. 6e).

To examine the functional role of p53 in GFAP toxicity, we reduced p53 levels using either a transgenic RNAi line or a p53 null allele. Both manipulations significantly reduced cell death in GFAPR79H transgenic flies (Fig. 6f). Reducing p53 levels also significantly decreased seizure frequency in GFAPR79H transgenic flies (Fig. 6g), demonstrating a beneficial effect of modulating p53 at the organismal level. The reduction of GFAP toxicity was not due to alterations in GFAP levels, as determined by western blot analysis (Supplementary Fig. 5d). We confirmed decreased (p53 RNAi) or absent (p53null) p53 protein expression using

Figure 3 | Knockdown of soluble guanylyl cyclase (sGC) in neurons reduces GFAP toxicity. (a) Schematic diagram of the dual-transcriptional system. To independently manipulate gene expression in glia and neurons, the GAL4/UAS system and the Q system are used together. GFAPR79H (QUAS-GFAPR79H) is expressed in glia with the Q system glial-specific driver ET31-QF (green box) and RNAi lines (UAS-RNAi) targeting sGC are expressed in neurons with the GAL4/UAS system neuronal-specific driver elav-GAL4 (pink box). Nitric oxide synthase (Nos) produces NO from l-arginine and O2. NO diffuses to neighbouring neurons and activates sGC, which converts GTP to cGMP. By expressing RNAi directed to sGC in neurons, NO signalling is blocked. (b) Knockdown of two Drosophila sGC subunits, Gyc99B and Gyc100B, with independent RNAi lines, markedly reduces the number of TUNEL-positive cells in GFAPR79H transgenic flies (total, left). Both neuronal (middle) and glial (right) cell death are rescued by reducing expression of Nos in glia (Fig. 5b). These data are consistent with a role for oxidative stress in promoting glial toxicity.

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**P < 0.0001, compared with UAS-lacZ control, F = 15.32; 11.62; 10.15; df = 29; 29, 29, one-way ANOVA with Tukey’s multiple comparison test, n = 6 per genotype. Flies were 20 days old. Genotype: elav-GAL4; ET31-QF; QUAS-GFAPR79H/UAS-RNAi. RNAi lines are indicated in the figure label.
western blotting and a p53-specific antibody (Supplementary Fig. 5e). Reducing p53 levels using transgenic RNAi did not alter the toxicity of mutant ataxin 3, as monitored by vacuole formation (Supplementary Fig. 2d), consistent with specificity of the rescue of GFAP toxicity. To probe the relationship between NO signalling and p53, we counted the number of p53-positive cells in GFAP R79H flies with genetically altered Nos levels. Overexpression of Nos significantly increased the number of p53-positive cells, while knockdown of Nos decreased the number of p53-positive cells (Fig. 6h). These findings together suggest a role for p53-mediated cell death downstream of Nos activation.

To further test our model, we determined whether increased levels of p53 can promote death of adult glia in the absence of transgenic GFAP Tg. Since overexpression of p53 during development produces organismal lethality, we reduced developmental expression of p53 using the temperature-sensitive GAL4 inhibitor GAL80ts (ref. 42). We then observed TUNEL-positive cell death in flies expressing p53 using the glial repo-GAL4 driver (Supplementary Fig. 5c). These findings confirm the ability of p53 to promote cell death in adult glia, and taken together with our other results, support a model in which DNA damage and p53 act downstream of Nos to promote glial cell death.

We next examined DNA damage and p53 accumulation in mouse models of Alexander disease. Double-label immunofluorescence revealed upregulation of H2AX-pS139, p53 and p53-pS15 in GFAP R236H and GFAP Tg mice (arrows), but not in age-matched wild-type littermate controls (arrowheads). GFAP was used to identify astrocytes. 4,6-diamidino-2-phenylindole (DAPI) labels nuclei. Scale bar, 20 μm. See Supplementary Figs 8 and 9 for larger images.
Increased iNOS level and activated p53 in patient tissue. Having observed upregulation of the iNOS pathway and activation of p53 in *Drosophila*, as well as in mouse models of Alexander disease, we investigated these two pathways in brains from patients with Alexander disease. Western blot analysis of frontal grey matter revealed a statistically significant increase in iNOS expression in Alexander disease patients (Fig. 8a). To evaluate the specificity of our findings, we measured iNOS expression by western blot in frontal grey matter from patients with metachromatic leukodystrophy, a childhood neurological disorder with some clinical and pathological features in common with Alexander disease. In contradistinction to our findings in Alexander disease, we observed a decrease in iNOS protein levels in metachromatic leukodystrophy patients (Supplementary Fig. 4h). Consistent with our findings in experimental animal models of Alexander disease, immunohistochemical analysis and double-label immunofluorescence showed a robust and consistent increase in p53 in astrocytes of brain tissue from all Alexander disease patients examined (Fig. 8b).

Discussion

Although morphological alterations are characteristic of astrocytes (so-called reactive gliosis) in any brain condition with major parenchymal injury, defining the specific role astrocytes play in promoting neuronal pathology has been difficult. Significant neuronal pathology is present in many serious neurological disorders, including neurodegenerative disease, stroke, epilepsy and other brain syndromes. Thus, distinguishing between a primary causal or secondary reactive role for glia in these pathologies has been difficult. To define a primary role of astrocytes in neurodegeneration, we focused here on Alexander disease. Alexander disease is caused by dominant mutations in GFAP, an intermediate filament protein of astrocytes. Therefore, in Alexander disease neurodegeneration is linked directly to astrocyte dysfunction.

Identifying the signalling molecule(s) and pathways that mediate toxic effects of astrocytes on neurons has been a second major challenge in understanding the role astrocytes play in neuronal dysfunction and death. Here we have used a fly model of Alexander disease, which replicates major features of the human disorder, to perform a forward genetic screen aimed at identification of secreted signalling molecules that promote neuronal death. Our screen recovered Nos, the single fly Nos16. We provided strong evidence that NO signalling through cGMP is activated in our fly model, and using both genetics and pharmacology we demonstrated that NO signalling is critical for neuronal death. Importantly, we also showed activation of the same NO signalling pathway in astrocytes of well-characterized mouse models of Alexander disease, as well as upregulation of iNOS in post-mortem brain samples from patients with Alexander disease. Previous experiments have demonstrated that STAT3 can bind to the iNOS promoter and stimulate transcription.44 Using a well-characterized STAT reporter20, we observed upregulation of STAT in GFAP transgenic flies, suggesting that activation of the JAK/STAT pathway could contribute to upregulation of Nos in Alexander disease.

Defining the cell-type-specific effects of soluble signalling molecules in vivo has represented a further barrier in understanding the molecular mechanisms of glial-based neurodegeneration. To address cell-type specificity, we have developed a system to independently manipulate gene expression in glia and in neurons in our Alexander disease *Drosophila* model (Fig. 3a). Using the dual-expression system, we provided strong evidence that glial-derived NO activates cGMP signalling in neurons to promote neuronal cell death by reducing the NO receptor, sGC, specifically in neurons. Interestingly, however, we also found that knockdown of sGC in neurons reduces glial cell death as well as neuronal cell death (Fig. 3b). These findings are consistent with signalling, perhaps via NO, from neurons back to glia in our system. A similar amplification process based on release of
neuronal NO has been reported in a cell culture model related to experimental autoimmune encephalomyelitis. 45

Our findings raise the possibility that non-cell autonomous nervous system injury in other 'gliopathic' disorders might also be driven by glial-derived NO. Indeed, NO has been implicated in a variety of neurological diseases with prominent astrocytic pathology, including Alzheimer's disease and Parkinson's disease. 46 Genetic deletion of iNOS is neuroprotective in experimental models of these disorders, supporting a functional role for the NO pathway in mediating neuronal pathology. In the current work, we have documented activation of the DNA damage response downstream of NO production in fly and mouse models of Alexander disease. Glia in both Drosophila and mouse models showed nuclear accumulation of p53 and evidence of DNA damage, as assayed by development of pH2AX/γ-positive foci (Figs 6 and 7). We also found increased levels of p53 in brains from patients with Alexander disease, supporting the relevance of our findings in experimental models to the human disease. Interestingly, astrocytes from patients with Alzheimer's disease also show nuclear accumulation of p53 (ref. 49) and pH2AX, raising the possibility of similar pathways of glial pathogenesis in the most common human neurodegenerative disorder.

In summary, here we have used Drosophila genetics, supported by analysis of murine models, to implicate NO as a critical astrocyte-derived neuronal cell death signalling molecule. We have further provided evidence that NO promotes glial dysfunction and death through oxidative stress, DNA damage...
Figure 7 | DNA damage response markers are increased in astrocytes of Alexander disease mouse models. (a–c) DNA damage response markers H2AX-pS139 (a), p53 (b) and p53-pS15 (c) are increased in astrocytes of GFAP R236H/+ and GFAP Tg mice (arrows), but not in age-matched wild-type littermate controls (arrowheads). GFAP marks astrocytes. 4,6-Diamidino-2-phenylindole (DAPI) labels nuclei. (d) p53 and p53-pS15 colocalize with H2AX-pS139 in GFAP Tg mice (arrows). DAPI labels nuclei. (e) H2AX-pS139, p53 and p53-pS15 colocalize with CNGA4 in GFAP Tg mice (arrows). DAPI labels nuclei. Mice were 3 months old in all panels. Scale bar, 20 μm.

Figure 8 | Increased iNOS and p53 in Alexander disease patients. (a) Quantification of iNOS protein levels reveals an increase in Alexander disease patients. *P = 0.0042, t = 2.326, df = 8; unpaired t-test, n = 5 for both ctrl and AxD. (b) Double-label immunofluorescence shows activation of p53 in astrocytes of Alexander disease patient (arrows), but not in control (arrowheads). Images were taken from the grey matter in frontal cortex. GFAP labels astrocytes. 4,6-Diamidino-2-phenylindole (DAPI) marks nuclei. Scale bar, 10 μm.
and p53 activation, well-documented target pathways in astrocytes and other cell types. Importantly, we used pharmacological inhibition of NO production as a potential therapeutic target in Alexander disease, and possibly in other gliopathies as well. Indeed, abnormal signalling was a potential therapeutic target in Alexander disease, and pharmacological inhibition of NO production to support NO

Methods

**Drosophila stocks and genetics.** All fly crosses were performed at 25 °C; adults aged at 29 °C to increase transgene expression. The following stocks were obtained from Bloomington Drosophila Stock Center: repo-UAS-GAL4 (glial driver in the GALA/UAS system), G431-QF (glial driver in the Q system), elav-GAL4 (neuronal driver in the GALA/UAS system), da-GAL4, tub-P-GAL80, UAS-NosRIBa RNAi #1 (TRIP#F03220), UAS-laUZ, UAS-GcyG98 RNAi #1 (TRIP#F03176), UAS-GcyG98 RNAi #2 (TRIP#GL01038), UAS-GyGl100B RNAi #2 (TRIP#F03214), p5311-188 (p5sgm) and p53 RNAi (TRIP#F02153). UAS-GyGl100B RNAi #1 (VDRC#100706) was from the Vienna Drosophila RNAi Center. Additional stocks used include nosA115 from P. O’Farrell, hid-lacZ (hid-ta10-lacZ) from A. Bergmann, th-GFP from J. Jiang, GstD1-lacZ from D. Bohmann, alrm-GAL4 and m20709-GAL4 from M. Freeman, 10XStur92E-GFP from N. Perrimon and UAS-ATXN-7Q78 from N. Bohnert. UAS-NosRNAi #2 was generated following the second-generation protocol from the TRIP at the Harvard Medical School by cloning short hairpin targeting nos into the VALIUM20 vector. Genetic Services performed embryo injections. Oligo sequences were as follows: 5’-TGGACGATTCTTCAGATGATA-3’ (sense sequence), 5’-TATGAAATGCTGATCGAG-3’ (antisense sequence). QUAS-GFAPR79H was constructed by digesting both UAS-GFAPR79H constructs and the VALIUM20 vector with EcoRI and NotI, followed by subcloning of the GFAPR79H sequence into the pQUAST vector. BestGene performed embryo injections.

**Transgenic mice.** Five-month-old male GFAPTg (Tg73.7) mice were used for western blot and 3-month-old GFAP-Tg26465 mice and GFAPTg (Tg73.7) mice, both male and female mice in the FVB/N background, were used for immunofluorescence staining. Same-sex age-matched wild-type littermates were used as controls. All procedures were approved by the Institutional Animal Care and Use Committee of the Graduate School of the University of Wisconsin-Madison. Taupathy transgenic mice Tg(tau924)Q510 (ref. 39) were 7 months old; controls were age-matched littermates not carrying the tau transgene.

**Drug feedings.** 1-NAMe (Cayman chemical) and 1-NAME (Sigma) were dissolved in water at 5 and 10 mM (Fig. 2c,d) and then mixed with instant food. Newly eclosed flies were fed on drug-embedded food every 3 days. Samples were boiled for 10 min at 100 °C and then centrifuged at 100,000 × g. The supernatant was used for western blots. For standard western blot analysis, adult Drosophila heads were homogenized in 1 × Laemml buffer (Sigma). For murine samples, cortex was homogenized in homogenization buffer (50 mM Tris, 150 mM NaCl, pH 7.4, with protease inhibitor cocktail, Thermo Scientific) and centrifuged at 100,000g for 1 h at 4 °C. The supernatant was used for determination of cGMP concentration following the manufacturer’s instructions (aCella-cGMP immunoassay, Cell Technology). Each data point represents the mean ± s.e.m.

**NADPH-diaphorase staining.** The NADPH-diaphorase staining protocol was adapted from Ott and Elphick. Briefly, brains were dissected in PBS and fixed in freshly mixed metanol and formalin (1:1) for 2 h on ice, followed by rinsing in PBS. Samples were then incubated in a PBS solution containing 0.3% Triton X-100, 0.5 mM NADPH (Sigma) and 0.2 mM NBT (Fisher). Incubations were carried out first in a cold room and then at room temperature until a proper blue developed. Brains were rinsed with PBS and images were acquired using a SPOT camera. Six brains for each genotype were analysed. Each data point represents the mean ± s.e.m.

**NO measurement.** The NO chemical sensor FL2E (Stem Chemicals) was dissolved in dimethylsulphoxide (DMSO) and mixed with CuCl2 immediately before use to form Cu2FL2E39. Brains were dissected in PBS and added to 10 μM Cu2FL2E. Time-lapse imaging was performed for each brain on a SPOT camera with an interval of 2 min for a total of 10 min. Seven brains for each genotype were dissected and imaged. Image was used to quantify fluorescence intensity, and two-way ANOVA with the Bonferroni post test was used for statistical analysis. Each data point represents the mean ± s.e.m.

**cGMP measurement.** A single fly head was homogenized in sample buffer and then centrifuged at 600g for 15 min at 4 °C. The supernatant was used for determination of cGMP concentration following the manufacturer’s instructions (Invitrogen). Data are presented as both cell counts (Figs 2a,c and 3b) and fold changes (Supplementary Figs 2a,i and 3g) to facilitate assessment of the magnitude of the changes observed. Statistical analysis was performed using one-way analysis of variance and Tukey’s multiple comparison test. Each data point represents the mean ± s.e.m.

**Western blots.** For standard western blot analysis, adult Drosophila heads were homogenized in 1 × Laemml buffer (Sigma). For murine samples, cortex was homogenized in homogenization buffer (50 mM Tris, 150 mM NaCl, pH 7.4, with protease inhibitor cocktail, Thermo Scientific) and centrifuged at 100,000g for 1 h at 4 °C. The supernatant was used for determination of cGMP concentration following the manufacturer’s instructions (aCella-cGMP immunoassay, Cell Technology). Each data point represents the mean ± s.e.m.
Human samples. Frozen frontal cortex from five controls (mean age 10 years, range 1–28 years; 2 females and 3 males), five Alexander disease patients (mean age 13 years, range 1–27 years; 2 females and 3 males) and four metachromatic leukodystrophy patients (mean age 10 years, range 3–21 years; 2 females and 2 males) were obtained from the NICHHD Brain and Tissue Bank for Developmental Disorders at the University of Maryland, Baltimore, MD. Informed consent for tissue distribution was obtained by the NICHHD Brain and Tissue Bank. The studies were performed under IRB-approved protocol 0033P000813 from Partners Healthcare to M.B. Feany. GFAP mutations in the Alexander disease patients included R79C, R239H (2 cases), K63E and L359V. All cases had typical neuropathology of Alexander disease, including multiple Rosenthal fibres. Post-mortem intervals were comparable between cases and controls and were less than 24 h in all cases. For immunostaining analysis, tissue was thawed and fixed in 4% paraformaldehyde overnight before paraffin embedding.

Confocal microscopy. All the fluorescent images were taken as Z-stacks on a confocal microscope (a Zeiss LSM 510 or a Leica SP8 X confocal microscope at Harvard NeuroDiscovery Center Enhanced Neuroimaging Core facility or an Olympus Fluoview 1000 confocal microscope at Harvard Medical School Neurobiology Imaging facility, NINDS P30 Core Center grant #NS072030) and visualized in two-dimensional projections of Z-stacks. Control and experimental samples were imaged with the same laser setting and the same Z-stack thickness.

Real-time PCR. RNA was isolated from Drosophila head homogenates using QIAzol (Qiagen) and reverse-transcribed using the High-Capacity cDNA Reverse Transcription kit (Applied Biosystems) according to the manufacturer’s instruction. Real-time PCR was performed and monitored using SYBR Green PCR Master Mix (Applied Biosystems) in a stepOnePlus Real-Time PCR system (Applied Biosystems) according to the manufacturer’s instructions. Drosophila ribosomal protein Rpl53 was used as a control. Each data point represents the mean ± s.e.m. Primers: Gyx99B: 5′-GTGGGCAATACGCGCCGCA-3′; 5′-CCTGGCCTCGTCTCAT-3′ CT-3′. Gyf110B: 5′-AAATGTTGACGGCCGGCA-3′; 5′-CAGAATGAGCTCGACGTG-3′ Rpl53: 5′-GACCATCGCGCCAGTATC-3′; 5′-CGGGCGACGACTGTCTGT-3′.

Statistical analysis. GraphPad Prism (v5) was used to perform two-tailed unpaired Student’s t-tests (for two groups), one-way ANOVA with Tukey’s post hoc tests (for three groups or more) and two-way ANOVA with Bonferroni’s post tests (for three groups or more) and two-way ANOVA with Bonferroni’s post. P < 0.05 was considered statistically significant. No statistical methods were used to predetermine sample sizes; sample sizes were similar to those reported in prior publications. Variance in each group was not calculated, but assumed to be similar. All photocromographs shown (immunolabelling, light microscopy and confocal microscopy) were representative of at least three biological replicates. All quantitative data are presented as mean ± s.e.m. except for seizure frequency (percentage of flies with seizures). Data for all experiments were collected and processed randomly, but no formal randomization was carried out. Data collection and analysis were not performed blind given the nature of the experiments performed.


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Author contributions

L.W. and M.B.F. designed the study, analysed data and wrote the manuscript. L.W. conducted experiments. T.L.H., A.M., H.K. and T.M. provided advice and reagents, and assisted with manuscript editing. M.B.F. and A.M. obtained funding for the research.

Additional information

Supplementary Information accompanies this paper at http://www.nature.com/naturecommunications

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