Neuroserpin polymers cause oxidative stress in a neuronal model of the dementia FENIB

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A b s t r a c t

The serpinopathies are human pathologies caused by mutations that promote polymerisation and intracellular deposition of proteins of the serpin superfamily, leading to a poorly understood cell toxicity. The dementia FENIB is caused by polymerisation of the neuronal serpin neuroserpin (NS) within the endoplasmic reticulum (ER) of neurons. With the aim of understanding the toxicity due to intracellular accumulation of neuroserpin polymers, we have generated transgenic neural progenitor cell (NPC) cultures from mouse foetal cerebral cortex, stably expressing the control protein GFP (green fluorescent protein), or human wild type, G392E or delta NS. We have characterised these cell lines in the proliferative state and after differentiation to neurons. Our results show that G392E NS formed polymers that were mostly retained within the ER, while wild type NS was correctly secreted as a monomeric protein into the culture medium. Delta NS was absent at steady state due to its rapid degradation, but it was easily detected upon proteasomal block. Looking at their intracellular distribution, wild type NS was found in partial co-localisation with ER and Golgi markers, while G392E NS was localised within the ER only. Furthermore, polymers of NS were detected by ELISA and immunofluorescence in neurons expressing the mutant but not the wild type protein. We used control GFP and G392E NPCs differentiated to neurons to investigate which cellular pathways were modulated by intracellular polymers by performing RNA sequencing. We identified 747 genes with a significant upregulation (623) or downregulation (124) in G392E NS-expressing cells, and we focused our attention on several genes involved in the defence against oxidative stress that were up-regulated in cells expressing G392E NS (Aldh1a1, Apoe, Gpx1, Gstm1, Prdx6, Scara3, Sod2). Inhibition of intracellular anti-oxidants by specific pharmacological reagents uncovered the damaging effects of NS polymers. Our results support a role for oxidative stress in the cellular toxicity underlying the neurodegenerative dementia FENIB.

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1. Introduction

The serpinopathies are protein conformational diseases characterised by the polymerisation and intracellular deposition of mutant variants of the serpins (serine protease inhibitors), within the endoplasmic reticulum (ER) of the cells that synthesise the protein. This common pathological mechanism results in liver disease upon polymeric mutations in alpha-1 antitrypsin (A1AT), and a rare but fatal neurodegenerative dementia caused by mutations in neuroserpin (NS) [reviewed in Roussel et al., 2011]. The latter is an autosomal dominant condition known as FENIB (familial encephalopathy with neuroserpin inclusion bodies) (Davis et al., 1999). Described in patients carrying one of six different mutations in the NS gene: S49P, S52R, S49P, S52R, ...
factor κ in vitro cells can be isolated from several regions of the mouse foetal brain, expression of NS has not been developed to date. Mouse neural progenitor cell lines have the in vivo dividing neurons through a well-defined protocol (Conti et al., 2015), precluding a decrease in locomotor activity, with decreasing mobility correlating to increased polymer content in the brain (Miranda et al., 2008). Despite these results, the mechanism of toxicity of NS polymers in cell models of disease has been elusive so far. Accumulation of NS polymers within the ER fails to induce a classical unfolded protein response (UPR), contrarily to a truncated version of NS (delta NS) lacking the last third of the aminoacidic sequence, which does not fold properly, does not polymerise and activates the UPR (Davies et al., 2009). Instead, NS polymers activate the ER overload response through a poorly understood signalling pathway that depends on Ca²⁺ and leads to the activation of nuclear factor eB (NF-eB) (Davies et al., 2009). Nevertheless, three different cell model systems transiently transfected COS-7 cells, stable inducible PC12 cells and stable inducible HeLa cells, failed to show clear signs of cell malfunction and death upon NS polymer accumulation (Miranda et al., 2008). However, the mechanism of toxicity of NS polymers in cell models of disease is not well understood, and its relevance for the toxicity of NS polymers in FENIB is not clear. The relevance of this pathway was confirmed by an increase in apoptosis of cells expressing G392E NS upon pharmacological inhibition of their anti-oxidant defences.

2. Material and methods

2.1. Reagents and antibodies

Unless stated otherwise, reagents, buffers, culture media and serum for cell cultures were purchased from Sigma-Aldrich (Milan, Italy). Custom-made rabbit polyclonal anti-NS antibody and rabbit polyclonal anti-GAPDH antibody were from Abcam (Cambridge, UK). The mouse monoclonal anti-NS antibodies were made in-house as reported before (Miranda et al., 2008). Anti-KDEL was from Enzo Life Sciences (through 3VChimica S.r.l., Italy), anti-GM130 from BD Transduction Laboratories and anti-catalase from Merek Millipore (both through SIAL S.r.l., Italy). Goat polyclonal anti-rabbit-HRP (horseradish peroxidase) and rabbit anti-mouse-HRP are from Sigma-Aldrich (Milan, Italy). Goat anti-mouse IgG-Alexa Fluor® 488 and -Alexa Fluor® 594, and goat anti-rabbit IgG-Alexa Fluor® 594 were from Life Technologies (Milan, Italy).

2.2. Generation and culture of neural progenitor cell cultures with stable neuroserpin expression

The open reading frame of human NS was amplified by PCR and subcloned into the pTP6 expression vector. This vector contains the CAGG promoter followed by the NS gene, an internal ribosome entry site (IRES) and the gene for puromycin resistance, allowing selection for simultaneous expression of both genes. NPCs derived from the mouse cerebral cortex at embryonic day 13.5 and protocols to culture them in vitro were previously described (Soldati et al., 2012). Briefly, NPCs were maintained in expansion medium consisting of basal medium [DMEM/F12 (LifeTechnologies, Milan, Italy), 1% penicillin/streptomycin, 0.1 M l-glutamine (LifeTechnologies, Milan, Italy), 1 mM Hepes, 7.5% NaHCO₃, 0.6% glucose] supplemented with 20 ng/ml of human recombinant epidermal growth factor (EGF; R&D System, through Aurogene, Rome, Italy), 10 ng/ml of human recombinant basic fibroblast growth factor (bFGF; R&D System, through Aurogene, Rome, Italy), and 1% N2 supplement (LifeTechnologies, Milan, Italy). NPCs were routinely cultured at 37 °C in a 5% CO₂ atmosphere within flasks previously coated with 10 μg/ml poly-ornithine (for 4 h) and 5 μg/ml laminin (for 12 h), seeding them at a cell density of 0.5–1 × 10⁶ cells/cm² andpassaging them every 3–5 days using Accutase. NPCs expanded for up to 20 passages in vitro since their original derivation were used for this work. Early passage NPCs were transfected with pTP6-NS in three different versions: wild type, G392E and delta NS. The transfection was performed with Amaza Nucleofector.
device and mouse Neural Stem Cell Nucleofector kit (Lonza, Cologne AG, Germany) according to manufacturer instructions. Two days after transfection, NPCs with stable integration of the pTP6-NS transgene were selected with puromycin (1.6 μg/ml), giving rise to cell lines consisting in mixed populations of cells with random stable insertion of the transfected plasmid DNA. A NPC line expressing pTP6-hrGFP (Vallier et al., 2004)(Pratt et al., 2000) was used as a negative control for non-specific effects due to the overexpression approach. For neuronal differentiation, NPCs were plated in expansion medium (20,000 cells/cm²) and 24 h later this was replaced with neuronal differentiation medium [basal medium containing 10 ng/ml of bFGF, 1% N2, 2% B27, and 0.5 μM DAPT (Tocris Bioscience, Bristol, UK)]. After 6 days this medium was partially replaced with fresh medium. After 6 days under these conditions, NPCs differentiated into a mixed culture of neurons (around 60–70%) and glial or residual progenitor cells [as reported in Sinno et al., 2013].

2.3. SDS and non-denaturating PAGE and western blot analysis

Each cell pellet from proliferative cells was obtained from a T25 flask adding 200 μl of lysis buffer (150 mM NaCl, 50 mM Tris-Cl, pH 7.5, 1% v/v Nonident P-40, plus 1 mM protease inhibitor cocktail). Differentiated cells were lysed as for proliferative cells but adding 50 μl of complete lysis buffer directly into the flask (T25) and using a cell scraper (Corning Inc., through Sigma-Aldrich, Milan, Italy) for cell detachment and lysis. The soluble fraction was collected by centrifugation at 17,000 g, 4 °C for 15 min, and proteins in the insoluble pellet were extracted by heating at 95 °C in lysis buffer containing 10% v/v beta-mercaptoethanol and 4% w/v SDS. 50 μg of total protein from each lysate for proliferative cells and 65 μl (corresponding to one flask) for differentiated cells, and the equivalent volume of each culture medium, were mixed with loading buffer without or with beta-mercaptoethanol and SDS as above and analysed by non-denaturing 7.5% w/v acrylamide PAGE (Miranda et al., 2008) or 10% w/v acrylamide SDS-PAGE (Miranda et al., 2008) as indicated. For enzymatic digestions, 20 μg of protein and the equivalent volume of culture medium were incubated with 1000 U of endoglycosidase H (endoH) for 3 h at 37 °C. Proteins were then separated by SDS-PAGE and analysed by western blot as described above. The HRP signal was developed using the LiteAblot PLUS and TURBO extra sensitive chemoluminescent substrates (Euroclone, Milan, Italy) and exposed to film or visualised on a ChemiDoc system (BioRad, Italy).

2.4. Sandwich ELISA

Quantification of NS was performed by sandwich ELISA with anti-NS antibodies made in house as described previously (Miranda et al., 2008). Briefly, 96-well plates (Costar 3590, Corning Inc., New York, US) were coated with antigen-purified rabbit polyclonal anti-NS antibody (2 μg/ml), washed (0.9% w/v NaCl, 0.05% v/v Tween20) and blocked with blocking buffer (PBS, 0.25% w/v bovine serum albumin, 0.05% v/v Tween20, 0.025% w/v sodium azide). Standards (recombinant purified monomeric or polymeric NS) and samples were diluted in blocking buffer and incubated for 2 h. After washing, wells were incubated with either a pool of monoclonal antibodies (1A10 and 10BB, 0.5 μg/ml each) or with an anti-NS polymer monoclonal antibody (7C6, 1 μg/ml). Rabbit anti-mouse IgG–HRP labeled antibody was used for detection with TMB substrate solution, and HRP activity was measured in a GloMax plate reader (Promega) at 450 nm. The concentration of NS in each sample was calculated against the appropriate standard curve (monomeric protein for WT NS, polymers for G392E NS), normalised to the total protein concentration of the sample (Bradford assay), and graphed in arbitrary units (AU).

2.5. Immunofluorescence staining and confocal microscopy

Cells for immunocytochemistry were grown on coverslips pre-treat- ed with poly-ornithine and laminin, in 24-well plates. Proliferating cells (around 60% confluence) or cells differentiated for six-days in vitro were washed, fixed in ice-cold 4% (w/v) paraformaldehyde for 30 min at room temperature and washed again. Cells were then treated with permeabilisation solution (0.5% saponin, 1% BSA, 0.5% EDTA pH 8 in PBS) for 10 min, then incubated for 30 min in blocking buffer (PBS plus 0.1% saponin, 0.5% BSA, 2% normal goat serum, 5 mM glycine, 0.1% sodium azide), and immunostained with anti-NS rabbit polyclonal antibody or the anti-NS polymer monoclonal antibody 7C6, anti-KDEL and anti-GM130 antibodies, and the corresponding secondary antibod- ies (goat anti-mouse IgG-Alexa Fluor® 488 and -Alexa Fluor® 594, and goat anti-rabbit IgG-Alexa Fluor® 594). Nuclear DNA was counter- stained with DRAQ®5 (Abcam, Cambridge, UK). Coverslips were mounted with FluorSave (Calbiochem, VWR International, Milan, Italy) plus 2% DABCO. Imaging was performed on a Zeiss 780 confocal microscope.

2.6. MTT assay

The culture medium was replaced by a solution of MTT [3-(4,5-di- methylthiazol-2-yl)-2,5-diphenyltetrazolium bromide, 0.5 mg/ml] in culture medium. After 3 h of incubation at 37 °C, this solution was re- moved and the formed formazan was solubilized in 250 μl of isopropanol plus 0.1% v/v HCl and 0.1% v/v NP-40. The optical density of the blue formazan was measured in a GloMax plate reader (Promega) at 560 nm.

2.7. RNA extraction and real-time RT-PCR

Proliferating NPCs plated in 6-well plates were lysed in 300 μl/well of RLT buffer (Qiagen RNeasy Mini Kit). Total RNA was purified out using Qiagen RNeasy Mini kit and eluted in 50 μl of RNase-free water. RNA was quantified using a Thermo Scientific NanoDrop 2000 spectrophotometer. NPC cultures differentiated for 6 days in T25 flasks were lysed in 400 μl of RLT buffer/flask; 20 μl of RNeasy-free water were used for RNA elution. For real-time reverse-transcriptase polymerase chain reaction (real-time RT-PCR), RNA was reverse-transcribed using the Qiagen QuantiTect reverse transcription kit and amplified on a Rotor-Gene Q (Qiagen), using Qiagen Quantifast SYBR Green PCR kit. Primers for real-time RT-PCR were designed using Primer3 (http://bioinfo.ut.ee/primer3/) and purchased from Sigma-Aldrich Co (Milan, Italy). Primer sequences used for this study are listed in Supplementary Table 2. Melt curve analysis was performed at the end of each real-time RT-PCR run. All primer pairs used for this study showed a single amplification peak, which was absent in the corresponding no-template control reaction, indicating primer specificity. Relative gene expression levels in different samples were determined with the built-in comparative quantitation method using Ecfial or Rpl19 as reference genes, with similar results. Statistical analysis of experimental data was performed with Microsoft Excel software.

2.8. RNA-sequencing

Total RNA was extracted and quantified as described above. RNA-Seq library preparation and sequencing was performed by the IGA Technology Services (Udine, Italy), using the Illumina-TruSeq RNA Sample Preparation Kit V2 (Illumina, San Diego, USA) according to manufacturer’s instructions. The final libraries for single-read sequencing of 50 base pairs were carried out on an Illumina HiSeq2000. Each sample produced about 20 millions of reads. Read quality was evaluated using FastQC (version 0.11.2, Babraham Institute, Cambridge, UK) tool. Reads were mapped to the mouse Ensembl GRCm38 build reference gene- nome using Tophat (version 2.0.12) (Trapnell et al., 2012) using default settings plus the “—no-novel-juncs” option. Gene structure annotations corresponding to the Ensembl annotation release 75 were used to build a transcriptome index and provided to Tophat during the alignment step using the “—G” parameter. The same gene annotations were used.
to quantify the gene-level read counts using HTSeq-count (version 0.6.1) script. Subsequently differential gene expression analyses were performed using Bioconductor (Gentleman et al., 2004) R (version 3.2.2) (R Core Team, 2015) package DESeq2 (version 1.4.5) (Love et al., 2014). Genes significantly up- or down-regulated in NS-expressing samples (q-value <0.1) were clustered by enrichment pathway analysis using Bioconductor R packages ClusterProfiler (Yu et al., 2012) and Pathview (Luo and Brouwer, 2013), Gene Ontology Database (Ashburner et al., 2000) and Reactome Pathway Database (Croft et al., 2014). All RNA-seq raw and processed data files were deposited at NCBI in the Gene Expression Omnibus (GEO) repository with accession number GSE96096.

2.9. Neurotoxicity experiments

NPCs were cultured in differentiation medium for 6 days in 24-well plates. At day 5 of differentiation, cells were incubated with 100 μM of diethyl maleate (DEM) or 500 μM of 3-amino-1,2,4 triazole (ATZ) for 1 h, or with 100 μM H2O2 for 30 min, at 37 °C, then cultured in fresh differentiation medium for 24 h at 37 °C. For DEM plus H2O2 treatment, cells were incubated always at 37 °C, first with 100 μM DEM for 1 h, then with fresh differentiation medium for 18 h, next with 100 μM H2O2 for 30 min, and finally with fresh differentiation medium for 6 h, for a total of 24 h. At the end of the treatments, the cells were washed in PBS and fixed in 4% (v/v) paraformaldehyde for 15 min, stained for cell counting with 3 μg/ml of the nuclear staining DAPI, and analysed with a 40× objective, counting at least 150 cells distributed in 15 areas per sample.

2.10. TUNEL assay

TUNEL staining was performed by using the in situ Cell Death Detection Kit, Fluorescein (Roche, 11,684,795,910). Briefly, cells were cultured for 6 days in differentiation medium and were pre-treated or not with 100 μM of DEM. Twenty-four hours after DEM treatment, cells were fixed in 4% paraformaldehyde for 15 min at RT. Cells were then washed with PBS for three times. Cell permeabilisation and staining were performed following the manufacturer’s instructions. Briefly, cells were permeabilised in a 0.1% Triton X-100 in 0.1% sodium citrate solution for 2 min and then washed again in PBS. Finally, cells were incubated with labelling solution at 37 °C for 1 h.

3. Results

3.1. Expression and handling of wild type and G392E neuroserpin in proliferating NPCs

We aimed to create a neuronal model to study the cellular phenotype of the dementia FENIB using non-transformed cell cultures. We cloned the wild type (WT), G392E and delta variants of human NS in the bicistronic expression vector pTP6, which is transcribed into a single mRNA encoding for both the gene of interest and the antibiotic resistance gene (puromycin). Previously described NPCs (Soldati et al., 2012) derived from E13.5 mouse cerebral cortex were transfected with the plasmids containing the different versions of NS. Three stable NPC-NS lines were established by selection with puromycin. The intracellular levels of G392E NS were always detectable only in cell lysates after proteosomal blocking with the MG132 inhibitor, as expected for this truncated version of NS (Fig. 1B, upper panel). No signal for NS was detected in the samples from GFP expressing cells used as negative control. The intracellular accumulation of NS polymers was confirmed in lysates of G392E NS cells by sandwich ELISA using several monoclonal antibodies against NS, including the anti-NS polymer 7C6 monoclonal antibody, as described before (Miranda et al., 2008) (Fig. 1C).

Taken together, these results show that WT NS is correctly processed and secreted in proliferative NPCs, while G392E NS forms polymers that accumulate within the ER, as seen above in proliferative conditions (Fig. 1B, upper panel). Delta NS was recognisable in cell lysates only after proteosomal blocking with the MG132 inhibitor (Fig. 2A, upper panel). The results from non-denaturing PAGE and western blot analysis showed that WT NS was found as a monomeric protein, mostly in the culture medium, while G392E NS was present in the polymeric conformation, with most of the signal corresponding to the cell lysate (Fig. 2A, lower panels). Quantification of intracellular NS by sandwich ELISA showed that the amount of G392E NS in differentiated NPCs was lower when compared to proliferative conditions, but the intracellular levels of G392E NS were always higher than those of WT NS, and the presence of NS polymers was confirmed with the 7C6 monoclonal antibody in cell lysates of G392E cells (Fig. 2B), as seen above for proliferating cells.

We next assessed the intracellular localisation and secretion of WT and G392E NS in differentiated NPCs by treating cell lysates and culture medium with the deglycosylating enzyme endoglycosidase H (endoH), which only removes N-glycans that have
not been modified in the Golgi apparatus. We found that intracellular WT and G392E NS were sensitive to endoH, supporting their pre-Golgi nature, while WT NS in the culture medium was endoH resistant, in agreement with its correct Golgi processing and secretion (Fig. 2C). The double band observed in cell lysates of G392E NS cells collapsed to a single one after endoH treatment, supporting that the difference in migration was due to different glycosylation patterns, as described before (Moriconi et al., 2015).

The intracellular distribution of NS in differentiated NPC cultures was investigated by immunofluorescence. WT and G392E NS were broadly distributed within the cell soma and along the neurites, with G392E NS showing a punctuated appearance (Fig. 2D). No NS signal was found in differentiated GFP and delta NS expressing cells. When using the anti-NS polymer monoclonal antibody 7C6, only the cells expressing G392E NS were positive, with a clearly punctuated pattern reminiscent of NS inclusions (Fig. 2D, lower panels). We looked into the trafficking of WT and G392E NS by comparing their distribution with that of the ER localisation signal (KDEL) and the Golgi resident protein GM130. WT NS was found in partial co-localisation with both the ER and Golgi markers, indicative of correct synthesis and transport of the protein, while G392E NS co-localised almost completely with KDEL and showed no co-
localisation with GM130 in the Golgi, in agreement with its accumulation within the ER (Fig. 2E).

These results confirm that the behaviour of WT and the polymerogenic variant G392E NS seen in previous models of FENIB is recapitulated in our novel model of differentiated progenitor cells from mouse foetal cerebral cortex.

3.3. The expression of WT NS is toxic during differentiation of NPCs

In order to look at the effects of NS expression on cellular viability, we first performed MTT assays in our cell lines during differentiation in vitro for six days. As shown in Fig. 3A, we compared the vitality of cells at time 0 (proliferative conditions), and 3 and 6 days after starting their differentiation. We found that MTT activity in WT NS cells was strongly decreased after 3 days, reaching around 50% of activity at day 6. The expression of GFP, delta or G392E NS did not affect MTT activity at any of the time points analysed. We next analysed the percentage of neuronal cells generated after differentiation by immunofluorescence detection of the specific neuronal protein βIII-tubulin, a microtubule protein almost exclusive of mature neurons (Fig. 3B). In control cells expressing GFP and in cells expressing G392E NS, the differentiation protocol induced a neuronal phenotype in over 60% of the cells, while overexpression of WT NS lead to a reduced percentage of neurons of about 40% of the surviving cells. These results suggest that WT NS overexpression impairs neuron generation in our cell culture system.

3.4. Cells overexpressing G392E NS show higher levels of several genes involved in the anti-oxidant defence

Since no evident cell toxicity was found in neurons upon accumulation of G392E NS polymers, we decided to perform an unbiased RNA sequencing (RNAseq) analysis to compare the entire transcriptome of differentiated NPCs expressing GFP or G392E NS. We decided to use the GFP expressing cells rather than the WT NS expressing cells as reference sample, due to the toxic phenotype observed for the latter. In the principal component analysis of the RNAseq results the three
independent samples of GFP or G392E NS differentiated cells showed good reproducibility and tended, respectively, to cluster together, while samples from GFP and G392E NS cells were well separated due to their very different RNA profiles (Fig. 4A). Indeed, the RNAseq analysis resulted in the identification of 1960 genes whose expression was significantly modulated in G392E NS cells in comparison to the GFP samples, with 623 genes overexpressed at least 2-fold (log2 > 1) and 124 genes underexpressed at least 2-fold (log2 < −1) (Fig. 4B). These genes were selected for further analysis by Gene Ontology (GO) Reactome and Reactome Pathway Database, which highlighted the alteration of several cellular pathways potentially due to the presence of NS polymers within the ER (Fig. S1 and S2). Among them we found several pathways involved in lipid and carbohydrate metabolism and biological oxidation (Fig. S1). Since oxidative stress plays a major role in the aetiology of various neurodegenerative disorders, we were interested to see that genes overexpressed in differentiated G392E NS cells included a few that, according to our GO analysis and previous publications (Supplementary Table 1, (Han et al., 1998)(Brown et al., 2013)), and they were all significantly up-regulated in G293E NS cells (Supplementary Table 1, Fig. 4C).

To confirm that up-regulation of these genes was specifically due to the presence of polymerogenic G392E NS, we analysed their expression by real-time RT-PCR in differentiated NPCs expressing WT NS or delta NS, in addition to cultures expressing GFP or G392E NS (Fig. 5). In this analysis, we included three additional genes involved in oxidative stress, namely Gpx1 (coding for glutathione peroxidase 1), Sod2 (coding for superoxide dismutase 2) and Cat (coding for catalase), which showed a trend towards up-regulation in G392E NS cells in our RNA sequencing dataset, although not statistically significant due to high variability in the three replicates used for RNA sequencing analysis (Supplementary Table 1, Fig. 4C, Fig. S3 and data not shown). Our results confirmed that mRNA levels of these genes were significantly higher in cells expressing G392E NS when compared to control cells expressing GFP, with the exception of catalase (Fig. 5 and Fig. S3). Moreover, there were no significant differences between GFP cells and cells expressing WT or delta NS, except for Aldh1B1 that showed a moderate increase in WT cells (Fig. 5B).

These results suggest that NS polymers activate cellular anti-oxidant defences in differentiated NPCs by up-regulation of several genes involved in oxidative stress.

3.5. Differentiated G392E NS cells are resistant to an oxidative insult but show enhanced toxicity upon inhibition of the anti-oxidant defences

In order to investigate oxidative stress in our differentiated NPC cultures, we first decided to assess their sensitivity to an exogenous oxidative insult such as H2O2. We measured cell survival after exposure to
100 μM H$_2$O$_2$ and found that G392E NS cells, and to a lesser extent WT NS cells, were more resistant to the oxidative insult when compared to naïve (non-transgenic), GFP and delta NS cells (Fig. 6A), suggesting that cells expressing WT and specially G392E NS are better adapted to cope with oxidative stress.

We next decided to investigate the effect of pharmacological inhibition of the cellular defences against oxidative stress. Glutathione (GSH) is an important antioxidant metabolite that prevents cell damage caused by the presence of reactive oxygen species (ROS), lipid oxidases and free radicals, and is involved in the mechanism...
of action of several ROS-detoxifying enzymes [reviewed in Schulz et al., 2000]. Cellular levels of GSH can be depleted by treating the cells with diethyl maleate (DEM) (Mitchell et al., 1983). Catalases are strong cellular scavengers of H₂O₂ [reviewed in Dringen et al., 2005], and can be inhibited with 3-amino-1,2,4 triazole (ATZ) (Margoliash et al., 1960). Since mRNA and protein levels of catalase were similar in all differentiated NPC lines (Fig. S3, panels B and D), the inhibitory effect of ATZ should be similar for all of them. The biological effects of catalase and GSH inhibitors were examined in differentiated NPCs expressing WT, G392E or delta NS, using GFP and naïve NPCs as negative controls, by culturing the cells without or with 100 μM DEM or 500 μM ATZ for 1 h and quantifying cell survival 24 h later by counting the surviving cells stained with DAPI. We found that the pharmacological inhibition of these antioxidant defences reduced the viability of G392E NS expressing cells when compared to GFP NPCs (Fig. 6B). These data were further confirmed by counting the number of cells with condensed chromatin, an indication of apoptosis, after GSH depletion and catalase inhibition. As shown in Fig. 6C, in differentiated cultures expressing G293E NS the impairment of these antioxidant defences reduced the viability of G392E NS expressing cells when compared to GFP NPCs (Fig. 6B). These data were further confirmed by counting the number of cells with condensed chromatin, an indication of apoptosis, after GSH depletion and catalase inhibition. As shown in Fig. 6C, in differentiated cultures expressing G293E NS the impairment of these antioxidant defences caused a phenotype of chromatin condensation, which corresponded to positive staining with TUNEL (Fig. 6C, panels d, h, i). It also led to positivity to an anti-cleaved caspase-3 antibody in cells containing G392E NS polymers (Fig. 6C, panel k), confirming the apoptotic nature of this nuclear phenotype. The quantification of condensed nuclei (Fig. 6D) showed a high number of apoptotic nuclei in differentiated cells expressing G392E NS treated with DEM (100 μM) or ATZ (500 μM).

After having found that G392E NS cells showed a toxic phenotype when the antioxidant defences were blocked by DEM, we tested their response under this condition to the additional stress caused by treating them with H₂O₂. Differentiated cells were pre-incubated with DEM as described in the methods section and then further exposed to a mild H₂O₂ treatment (100 μM H₂O₂ for 30 min). As shown in Fig. 6E we found that, after treatment with DEM, the addition of H₂O₂ was more toxic to cells overexpressing G392E NS than GFP or WT NS cells, indicative of a higher sensitivity of differentiated G392E NS cells to a pro-oxidant insult when the antioxidant defences are impaired.

4. Discussion

The neurodegenerative dementia FENIB is a rare condition caused by aberrant polymerisation of NS within the ER. Studies conducted in two different animal models of FENIB, namely flies (Miranda et al., 2008) and mice (Galliciotti et al., 2007) overexpressing mutant human NS, have demonstrated that the presence of polymers is toxic to the organism, but the exact mechanism of this toxicity is still poorly understood. This is mainly due to the absence of a toxic phenotype in the cell culture models of the disease available so far (Miranda et al., 2004)(Miranda et al., 2008)(Kroeger et al., 2009)(Davies et al., 2009)(Roussel et al., 2013).
Fig. 6. Polymers of G392E NS cause oxidative stress in differentiated NPCs. A. NPCs were differentiated for 6 days and each cell line was incubated in the absence or presence of H2O2 (100 µM) for 30 min. Cell survival was quantified 24 h later by counting DAPI-positive cells. Results are expressed as the percentage of surviving neurons compared with control cultures (non treated) or with either the glutathione depletor DEM (100 µM) or the catalase inhibitor ATZ (500 µM). Neuronal survival was quantified 24 h later by counting DAPI-positive stained cells. Results are expressed as the percentage of surviving neurons compared to control cultures, and are the mean ±/− SEM of three independent repeats, t-test: **p ≤ 0.01; ***p ≤ 0.005. C. Differentiated NPCs expressing GFP or G392E NS were treated or not with 100 µM DEM and stained with DAPI (a–d), TUNEL assay (e–i) or anti-cleaved caspase-3 antibody (green) and anti-NS polymers (7C6 antibody, red) (j–k), to assess the nature of nuclei showing chromatin condensation. Arrows: nuclei with condensed chromatin.

D. Differentiated cells were incubated for 1 h with either 100 µM DEM or 500 µM ATZ and their nuclear morphology was analysed 24 h later. The toxic effect of DEM and ATZ was quantified by counting DAPI-stained nuclei with condensed chromatin. Results are expressed as the ratio between treated and non-treated cells, and data are expressed as the mean ±/− SEM of three independent repeats, t-test: *p ≤ 0.05; **p ≤ 0.01. E. Differentiated NPCs were pre-incubated for 1 h without or with 100 µM DEM and then further incubated for 30 min with 100 µM H2O2 in the presence or absence of DEM. Neuronal survival was quantified 24 h later by counting DAPI-stained cells. Results are expressed as the percentage of surviving neurons compared with control non-treated cultures, and data are the mean ±/− SEM of three independent repeats, t-test: **p ≤ 0.01. In panels A, B, D and E, 15 different fields (to a total of 150 to 350 cells) were counted per condition in each independent repeat.
Our first aim in the present work was to create a novel cell model system closer to the conditions found in FENIB. Postmortem analysis of brains from FENIB patients has shown the presence of abundant inclusion bodies in some brain regions, mainly in the cerebral cortex and the substantia nigra (Davis et al., 1999) (Hagen et al., 2011). We thus chose to use NPCs derived from the developing mouse cerebral cortex to establish cell lines overexpressing NS and a selection marker from the same promoter, so NS expression could be maintained stably by culturing the cells in the presence of puromycin. Although these cultures consisted in mixed populations with stable random integration of the transgenes, the use of a bicistronic plasmid led to similar NS expression levels by real-time RT-PCR and homogeneous presence of GFP and NS by immunofluorescence, indicating low variability in our NPC cultures.

Since FENIB is characterised by the presence of inclusion bodies in mature neurons, we differentiated our cells in vitro using previously described protocols driving efficient neuronal differentiation of NPCs (Sinning et al., 2013). This led to a postmitotic population with approximately 60% of neurons, as assessed by morphology and expression of ßIII tubulin, and 40% of glial-like cells, as previously described (Sinning et al., 2013). Our results show that, although protein levels of NS were lower in differentiated cells, NS was handled in the same way both in proliferating and differentiated NPCs, with correct secretion of the wild type variant as a mature glycoprotein and accumulation of G392E NS polymers within the ER, thus reproducing the trafficking defects characteristic of FENIB as described in previous cell models of disease (Miranda et al., 2008) (Roussel et al., 2013). Our results showed that NS expression could be maintained stably by culturing the cells in the presence of puromycin. Although these cultures consisted in mixed populations with stable random integration of the transgenes, the use of a bicistronic plasmid led to similar NS expression levels by real-time RT-PCR and homogeneous presence of GFP and NS by immunofluorescence, indicating low variability in our NPC cultures.

During the first three days of the differentiation protocol, we observed a strong decline in cultures of NPCs expressing WT NS. This could be due to a toxic effect of the overexpression of functionally active neuroserpin in this system. A similar toxicity was previously observed when generating a D. melanogaster line overexpressing functional WT NS in the fly brain (D. Crowther, personal communication). In contrast, cells expressing G392E NS did not show toxicity during routine culturing or differentiation, so we analysed them by global mRNA sequencing in comparison to control GFP cells. We found 747 genes whose expression levels were altered by at least two-fold, and decided to further investigate those involved in oxidative stress and lipid oxidation according to GO analysis and previously published studies. It has been shown that expression of the polymorphic Z variant of A1AT in mouse liver leads to upregulation of genes involved in redox regulation, namely carbonyl reductase 3 (CBR3), glutathione S transferase alpha 1 + 2 (GST A 1 + 2 ) and glutathione S transferase M3 (GST M3), and this correlated with higher ROS levels and oxidative damage in liver tissue from aged mice transgenic for Z A1AT (Marcus et al., 2012). The involvement of oxidative stress and lipid oxidation in FENIB has not been investigated so far. Our RNA sequencing analysis and real-time RT-PCR data confirmed that several genes of interest were upregulated in G392E NS cells, but not in cells expressing WT NS or the UPR eliciting mutant delta NS, except for a moderate upregulation of Aldh1b1 in WT NS cells. The most upregulated gene in G392E NS cells in comparison with GFP, WT NS and delta NS cells was Scav3, which has been shown to scavenge ROS (Han et al., 1998) and to be protective against cell death induced by oxidative stress in myeloma cells (Brown et al., 2013). Additional genes involved in the defence against oxidative stress and lipid oxidation were also significantly and specifically upregulated in G392E NS cells: Prdx6 codes for an antioxidant enzyme whose levels increase in brains of patients suffering from Alzheimer’s disease (Power et al., 2008) and Parkinson’s disease or dementia with Lewy bodies (Power et al., 2002); Aldh1 family genes code for important enzymes that prevent the accumulation of toxic aldehydes derived from lipid oxidation during oxidative stress (Makia et al., 2011); the protein products of Gsr family genes are protective against oxidative stress damage, particularly through detoxification of lipid oxidation products [reviewed in Sharma et al., 2004], and upregulation of GST M3 was detected upon expression of a polymorphic variant of A1AT in mouse liver (Marcus et al., 2012); studies in knock-out mice lacking the lipid transporter apolipoprotein E have shown increased lipid peroxidation in the cortex (Montine et al., 1999). By performing real-time RT-PCR analysis with a higher number of samples than those used for the RNA sequencing analysis, we also found that mRNA levels of Sod2 and Gpx1, two important genes involved in detoxification of ROS [reviewed in Gandhi and Abramov, 2012; Cobb and Cole, 2015], were higher in G392E NS cells. The fact that G392E NS cells show a higher expression of these genes when compared to GFP, WT and delta NS cells supports a specific response in G392E NS cells to the accumulation of NS polymers within the ER, separate from other effects due to overexpression of a heterologous protein or a misfolding protein able to activate the UPR. It has been reported that polymers of mutant Z A1AT reduce the mobility of proteins in the ER lumen, rendering the ER more susceptible to stress (Ordoñez et al., 2013). It is possible that the presence of NS polymers alters the redox balance in the ER by a similar mechanism, although further studies are needed to support this hypothesis.

Based on these results, we hypothesised that polymers of G392E NS caused low levels of oxidative stress in our differentiated NPCs, triggering a protective response involving increased expression of proteins with anti-oxidant activities. Blocking the main anti-oxidant defences by pharmacological depletion of GSH and inhibition of catalase showed that G392E NS cells became more susceptible to oxidative stress than the other cell lines, with increased cell death through apoptosis. These results support the involvement of oxidative stress and lipid oxidation in the progression of the neurodegenerative dementia FENIB, as already demonstrated for more common forms of neurodegeneration as Alzheimer’s and Parkinson’s diseases [reviewed in Gandhi and Abramov, 2012; Cobb and Cole, 2015]. Furthermore, it is known that anti-oxidative defences decrease with age, which may help to understand why FENIB mutations like G392E and G392R cause higher number of inclusion bodies in the brain at an early age: the young brain, coping better with oxidative stress, would be more resistant to polymer accumulation and thus need a higher burden of polymers to develop the disease, while lower amounts of inclusion bodies in older brains suffice to start neurodegeneration with slower polymer-forming mutant variants of NS, like the S49P (Davis et al., 1999).

A recent report has proposed a protective role for extracellular WT NS against exogenous H2O2, through an increase in phosphorylation of AKT (Cheng et al., 2016). The study does not clarify how this effect is exerted and whether it depends on the inhibitory properties of NS, but it could provide an explanation for our observation that NPCs overexpressing WT NS are more resistant than control cells to addition of H2O2, although not as much as G392E NS expressing cells. The fact that NPCs overexpressing WT NS show similar expression levels to those seen in control cells for all except one of the anti-oxidant genes studied here by real-time RT-PCR, indicates that the signalling activated by WT and G392E NS in NPCs are probably diverse: an extracellular effect for WT NS, which is efficiently secreted as a monomeric protein, and an intracellular response to the intracellular accumulation of polymers in the case of G392E NS.

In conclusion, we show here for the first time that expression of a strongly polymorphic variant of NS that causes severe dementia FENIB leads to upregulation of several anti-oxidant genes and to apoptotic neural cell death upon impairment of the oxidative stress defences, suggesting that oxidative stress and lipid oxidation may contribute to neurodegeneration in FENIB.
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Conflict of interest statement

The authors declare no conflicts of interest.

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