Heme Deficiency Is Associated with Senescence and Causes Suppression of N-Methyl-d-aspartate Receptor Subunits Expression in Primary Cortical Neurons

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ABSTRACT

Heme is a crucial component of many pharmacological and toxicological processes, and studies have suggested that heme deficiency may play a role in cellular ageing. A model of ageing neurons was established using prolonged cultures of BALB/c mouse primary cortical neurons. Aged neurons displayed a senescent phenotype and a marked up-regulation of cathepsin-L expression. Down-regulation of the candidate neuron-specific genes for N-methyl-d-aspartate (NMDA) receptor subunits (NMDA1 and -2) and neurofilament light peptide (NF-L) were found to be characteristic of the aging process as reported in vivo (Brain Res 907:71–83, 2001; Brain Res Mol Brain Res 99:40–45, 2002). In contrast, the genes for the controlling enzymes of heme synthesis and degradation (5-aminolevulinate synthase 1 and heme oxygenase 1, respectively) were up-regulated, implying depletion of a regulatory heme pool. Inhibition of heme synthesis (by 70–80%) at different enzymic steps by succinyl acetone and N-methylprotoporphyrin IX resulted in the earlier lowered expression of NMDA1 and -2 and NF-L. Exogenous hemin added to heme-depleted cells rescued the expression of these neuron-specific genes. Culture of cortical neurons from BALB/c FecI/H9256 mutant mice demonstrating depressed heme synthesis showed premature senescence and reduced expression of NMDA1 and -2 receptor subunits and NF-L compared with wild-type cells. Our findings suggest that reduced availability of heme in neurons associated with senescence may have significant effects on synaptic function.

Heme serves as the prosthetic moiety of numerous hemoproteins. Besides acting as the key component of hemoglobin and cytochromes (including those of drug metabolism), heme regulates many processes of pharmacological importance by controlling pathways through interaction with key regulatory proteins (Ogawa et al., 2001; Taoka et al., 2002). The brain is a high consumer of oxygen, and mitochondrial cytochromes are particularly vital for maintaining normal neural metabolic function. In addition, hemoproteins are important constituents of signaling processes in the brain. Circadian clock and heme biosynthesis are reciprocally regulated, with heme acting via core clock mechanism member NPAS2 (Kaasik and Lee, 2004). Both the production of nitric oxide and some of its actions are mediated through hemoproteins, such as guanilyl cyclase (Boehning and Snyder, 2003). Likewise, the signaling function of carbon monoxide as a neurotransmitter/neuromodulator (Ingi et al., 1996; Boehning and Snyder, 2003) is utterly dependent on heme. The only known biosynthetic source of CO in the brain is as a cleavage product of heme catalyzed by heme oxygenases (HMOX) (Ingi et al., 1996). Type 1 HMOX can be markedly induced under conditions of stress, thereby increasing the requirement for substrate (Sassa and Nagai, 1996; Sassa, 2004). In nonerythroid tissue, including brain, increased requirement for intracellular heme can be detected as up-regulation of the gene Alas1 for aminolevulinate synthase 1 (ALAS1), the first step of heme synthesis, and commonly accepted as a response to a lowered regulatory heme pool (De Matteis et al., 1981; De Matteis and Ray, 1982; Sassa and Nagai, 1996). The availability of heme may also be a limiting factor in the ability of neuronal cytochrome P450 enzymes to metabolize drugs and chemicals (Meyer et al., 2002). In contrast, larger amounts of exogenous heme, produced in hemorrhage of stroke, may be toxic to neurons (Goldstein et al., 2003). Some genetic defects in heme synthesis in humans are associated with diseases with neurological symptoms and the inappropriate accumulation of heme precursors perhaps contributes to development of neuropathy (Rank et al., 1993; Lindberg et al., 1999).

On the other hand, neuropathy may be the consequence of heme deficiency, causing dysfunction of crucial signaling pathways and functioning of such hemoproteins as cyto-

ABBREVIATIONS: HMOX, heme oxygenase; ALAS, aminolevulinate synthase; NMDA, N-methyl-d-aspartate; C7, threshold cycle; NF-L, neurofilament light polypeptide; PCR, polymerase chain reaction; RT, reverse transcription; SA, succinyl acetone; NMP, N-methylprotoporphyrin IX.
chrome P450 (Lindberg et al., 1999). In the liver, many drugs and chemicals affect heme synthesis, and this may also occur in the brain (De Matteis and Ray, 1982). Furthermore, ALAS1 activity in the brain, as in the liver, declines with age (Paterniti et al., 1978; Bitar and Shapiro, 1987), and the binding of heme to amyloid β in Alzheimer’s disease may be an additional limitation on its availability (Atamna and Frey, 2004). Aging has an implication for every part of the brain, but the frontal cortex and parts of the hippocampal system seem especially affected, which has been linked to expression of synaptic N-methyl-D-aspartate (NMDA) receptor subunits ζ1 and ζ2, which are important for learning and memory (Eckles-Smith et al., 2000; Ossowska et al., 2001; Magnusson et al., 2002; Uylings and de Brabander, 2002). In experiments, proliferation of glial cells or fibroblasts. The cells were grown in a concentration of senescent cells was performed with Senescent Cells Staining Kit (Sigma-Aldrich) in accordance with manufacturer’s instructions by detection of β-galactosidase histochemically at pH 6 (Dimri et al., 1995).

RNA Extraction and Quantitative Real-Time PCR Analysis. Treated and untreated cells from different time points were collected, and total RNA was isolated by using TRI-reagent (Sigma-Aldrich). cDNA synthesis was carried out using random primer and Superscript II (Invitrogen). PCR primers were selected using the Primer Express v2.0 Software program (Applied Biosystems, Foster City, CA).

Primer sequences are shown in Table 1. Primers were designed to cross exon-exon boundaries and the concentration optimized (300–900 nM) to ensure that the efficiency of the target amplification and the efficiency of the endogenous reference amplification are approximately equal. PCR was performed using SYBR Green PCR Master Mix, primers, and 100 ng of reverse-transcribed cDNA in the PRISM 7700 Sequence Detection System (Applied Biosystems). The ther-

### Materials and Methods

#### Primary Cell Culture

Primary cortical neurons were prepared from male and female 14-day-old fetuses of the BALB/c mouse strain bred in house. The BALB/c Fech−/− mouse strain was obtained from the Jackson Laboratories (Bar Harbor, ME). The Fech−/− mutant (abbreviated here to Fech mouse) contains a point mutation in the ferrochelatase gene (Tutois et al., 1991; Davies et al., 2005). Mice were bred by homozygous mating and maintained in a negative pressure isolator at 21°C under reduced light to protect from skin lesions. Isolated brain cortex of embryos was gently dissociated to release the neurons, which were washed twice in Neurobasal medium containing 2% B-27 supplement (Sangerman et al., 2001). After attachment of the cells, the cultures were stained with a mixture of the membrane-permeable dye SYTOX (500 nM) for 5 min at 37°C. The amount of normal, necrotic (damaged/SYTOX-permeable membrane, normal nuclei) and apoptotic (impermeable membrane, condensed/fragmented nuclei) cells were scored with a fluorescence microscope and no significant difference was observed at any time between controls and any of the treatments performed in this study. Identification of senescent cells was performed with Senescent Cells Staining Kit (Sigma-Aldrich) in accordance with manufacturer’s instructions by detection of β-galactosidase histochemically at pH 6 (Dimri et al., 1995).

#### Inhibition and Measurement of Heme Synthesis.

To inhibit heme synthesis, cells were cultured in serum-free medium with 0.5 mM succinyl acetone (Sigma-Aldrich, Dorset, UK) or with 1 μM N-methylprotoporphyrin IX (Frontier Scientific Porphyrin Products, Logan, UT) continuously for the duration of the experiments. For measurement of heme synthesis, cells were incubated with 0.4 μCi of [3,5,7H]ALA (2.6 Ci/mmol; PerkinElmer, Boston, MA) for 24 h. Heme was extracted from the cells by acetonitrile and diethyl ether. The amount of radioactivity in extracted heme was measured by liquid scintillation counting as described previously (Shedlofsky et al., 1987). Total recovery of radioactivity from all fractions was the same for treated and untreated cells. Ferrochelatase activity of cortical tissue was measured as reported for lymphocytes (Rossi et al., 1988). For heme recovery experiments, heme as hemin (0.1 μM) was added to culture medium in the presence of bovine serum albumin in a 1:1 M ratio (Taketani et al., 1998).

#### Necrosis, Apoptosis, and Senescence.

Cell viability in heme depletion experiments was estimated by using SYTOX/Hoechst double-staining method. To determine the amount of necrosis or apoptosis in the neurons, the cultures were stained with a mixture of the membrane-permeable dye Hoechst-33342 (500 ng/ml) and the membrane-impermeable dye SYTOX (500 nM) for 5 min at 37°C. The amount of normal, necrotic (damaged/SYTOX-permeable membrane, normal nuclei) and apoptotic (impermeable membrane, condensed/fragmented nuclei) cells were scored with a fluorescence microscope and no significant difference was observed at any time between controls and any of the treatments performed in this study. Identification of senescent cells was performed with Senescent Cells Staining Kit (Sigma-Aldrich) in accordance with manufacturer’s instructions by detection of β-galactosidase histochemically at pH 6 (Dimri et al., 1995).
mal-cycler protocol was: stage 1, 50°C for 2 min; stage 2, 95°C for 10 min; and stage 3, 40 cycles at 95°C for 15 s and 60°C for 1 min. Each sample was run in triplicate. Quantification was performed using the comparative CT method ($\Delta\Delta CT$). Data are presented as the mean ± S.D. ($n=3–7$ for each group). Statistical significance was assessed as $P<0.05$ using one-way analysis of variance.

**Immunoblotting.** Proteins were extracted from primary neurons after 14 and 21 days of culture using lysis buffer (7 M urea, 50 mM Tris-HCl, pH 7.5, and 5 mM dithiothreitol) followed by brief sonication. SDS electrophoresis and immunoblotting were performed (Davies et al., 2005) using chemiluminescence detection (ECL; Amersham Pharmacia, Buckinghamshire, UK) and primary antibodies from the following sources: HMOX1 AND HMOX2 from Stressgen (Victoria, BC, Canada), NMDA1, neurofilament light polypeptide (NF-L), and $\alpha$-tubulin from Santa Cruz Biotechnology (Santa Cruz, CA). Results were quantified using densitometry and ImageQuant 5.2 software (GE Healthcare, Little Chalfont, Buckinghamshire, UK). Statistical significance of data was estimated using two-tailed student’s t test.

**Results**

**Senescence of Primary Cortical Neurons.** To establish the aging of primary neurons in culture over time, we examined the changes associated with senescence. After 20 days in vitro, morphological changes of the aged cells were observed, namely an adoption of enlarged shape that is characteristic for the senescent phenotype (Campisi, 2005). Electron microscopy revealed signs of swollen appearance of the cells and reduced density of neurites on day 25 (data not shown). $\beta$-Galactosidase activity detected by histochemical staining is a measure of increased residual lysosomal activity at a suboptimal pH and has been used to identify senescent human cells in culture and in vivo (Dimri et al., 1995; Kurz et al., 2000).

The proportion of the senescent cells steadily increased in neuron cultures (Figs. 1, A–C, and 2A). We also detected a significant increase in cathepsin L expression over time (Fig. 2B). Up-regulation of proteases such as cathepsin L is associated with the development of senescence phenotypes causing a disruption of tissue integrity and function (Varela et al., 2005). The proportion of cells detected showing apoptosis and necrosis was not significantly changed during this time.

**Comparison of Heme-Related and Neuron-Specific Gene Expression.** The temporal pattern of gene expression during the ageing of cultures of neurons for up to 25 days was established by real-time RT-PCR (Fig. 3) of selected genes known to be critically associated either with heme metabolism or neuronal function. Greater expression of the gene for ALAS1 ($Alas1$) reflects an increased requirement for heme, whereas induction of the $Hmox1$ gene associated with the heme catabolism enzyme HMOX1 may reflect potential increased degradation possibly as a stress response (Sassa, 2004). There was initial down-regulation of all the genes in comparison with fresh embryonic cells 1 or 2 days before differentiation of the cultures. However, between days 2 and 6, the expressions began to recover and, by day 10, were comparable with freshly isolated cells. At later stages of culture (days 21 and 25), ALAS1 expression was markedly increased. The increase in ALAS1 expression probably reflected a negative feedback from a lowered regulatory heme pool as cells aged and was compatible with a marked increase in the amount of HMOX1 mRNA (80-fold by day 25). In contrast, expression of HMOX2 showed much lower induction.

Increased expression of ALAS1 and HMOX1 was preceded by decreased expression of the neuron-specific genes for NMDA receptor subunits ($\zeta$1 and $\zeta$2 and NF-L that had recovered from plating and were induced to the greatest level.
on day 14 (Fig. 3, D–F) reflecting maximum neurite networking. For instance, an increase in NMDA1 mRNA was observed from day 2 and, by day 14, reached a significant maximum of 22-fold but declined to only 5-fold higher on day 25. Complementary changes in HMOX1, HMOX2, and NMDA2i protein expression were demonstrated by immunoblotting (Fig. 3G). Thus, lower expression of the genes associated with neurons was associated with aging of the cells and inversely related to up-regulation of the genes for ALAS1 and HMOX1. NMDA2i temporal expression pattern was similar to that of NMDA1 but with changes that were less marked.

**Heme Deficiency Results in Down-Regulation of Neuron-Specific Genes.** To test whether heme deficiency per se could effect neuronal gene expression, cells were cultured continuously with succinyl acetone (SA), a specific inhibitor of ALA dehydratase (Tschudy et al., 1981), or NMP, a specific inhibitor of ferrochelatase (De Matteis and Marks, 1996). Heme synthesis in cultures was significantly depressed by both treatments (Fig. 4A). On days 12 and 18 of culture in the presence of SA, ALAS1 expression was significantly higher than in control cells (Fig. 4, B and C). The expression of HMOX1 was similarly induced, although no difference between heme-depleted and control cells was observed in the expression of constitutive HMOX2. Increased HMOX1 protein, but not HMOX2, was confirmed by Western blotting (Fig. 4E). Inhibition of heme synthesis by NMP resulted in similar changes in ALAS1 and HMOX1 expressions (Fig. 4D). In contrast, in both heme-depleted cultures, expression of NMDA receptor subunits 1 and 2 and NF-L were significantly depressed compared with neurons not exposed to SA or NMP (Fig. 5). By day 18, when the control culture was also displaying signs of aging, expression of NF-L was detected at similar levels in treated and untreated neurons. A similar pattern but with more profound changes was detected in NMP-treated cells (Fig. 5D). Thus, disruption of heme synthesis with SA and NMP resulted in premature changes in expression of genes associated with aging of untreated cells.

**Protection by Heme.** Neurons treated with SA to depress heme synthesis were cocultured with heme at a level (0.1 μM as hemin) added in albumin thought to balance heme synthesis without inducing heme degradation (Taketani et al., 1998; Sassa, 2004). As shown in Fig. 4B, SA induced up-regulation of the genes for ALAS1 and HMOX1 and down-regulation of NMDA1 and NMDA2i. In the presence of additional heme, this was mostly reversed so that only small changes in expression of NMDA1 and NMDA2i were observed and ALAS1 and HMOX1 were little affected (Fig. 6, A–D). To compare this finding with the effect of heme deficiency on an inhibitory neurotransmitter type receptor, we investigated the expression of the GABA_A receptor. The most common receptor configuration includes the α1 subunit (Wassef et al., 2003). We did not detect marked changes in the expression of GABA_A α1 in SA-treated neurons, and additional heme did not alter expression of the gene (Fig. 6E).

**Premature Senescence of Fech Neurons.** The Fech mutant BALB/c mouse contains a point mutation in the ferrochelatase gene; the resulting enzyme exhibits less than 5% of normal ferrochelatase activity in the liver and spleen. This is associated with insufficiency of erythropoietic heme synthesis, hemolytic anemia, splenic enlargement, and marked dysfunction of liver metabolism (Tutois et al., 1991; Davies et al., 2005). Likewise, ferrochelatase activity of brain cortex from Fech mice was measured at <3% of that in wild-type BALB/c brain (Fig. 7A). In addition, when heme synthesis in cultured neurons was estimated, it was decreased by 47% compared with that of BALB/c primary cells (Fig. 7B). Fech neurons were more sensitive to the stress of plating and had less ability to survive over the long term in primary culture than wild-type BALB/c neurons. Staining for β-galactosidase activity showed earlier signs of senescence compared with control BALB/c neurons so that by day 12, 89.1 ± 3.2% (n = 5) of the Fech cells were identified as senescent (Fig. 1D), whereas this degree of senescence was not observed until much later with control BALB/c cells.

**Down-Regulation of Neuron-Specific Genes in Fech Neurons.** Heme related gene expression patterns in cultured Fech neurons (Fig. 7, C and D) showed changes similar to heme-depleted cultures caused by treatment with SA and NMP. Increased expression of ALAS1 was already detected on day 14, not later as observed with BALB/c neurons. Elevated expression of HMOX1 at the time of preparation of primary culture reflected that the Fech embryo cells were already under stress at the time of isolation. Neuron-specific genes were expressed at lower levels than control BALB/c neurons during differentiation of the culture. NMDA1 expression in Fech neurons was less than 20% of that in a control culture on days 6 and 30% on day 14. NMDA2i detected in Fech neurons was 57% and 53% on days 6 and 14 of culture, respectively, compared with the wild-type cells. Treatment with exogenous heme (0.1 μM hemin) largely rescued expression of NMDA1 in Fech neurons (Fig. 6F).
In these studies, we demonstrated that, subsequent to their differentiation, primary cortical neurons in prolonged culture develop characteristics of senescence, using a method that is unique for senescent cells (Dimri et al., 1995). In addition, the expression of genes for synaptic NMDA receptor subunits (which are important for learning and development) and for NF-L (which are associated with healthy neurite growth and networking) reached a maximum after 14 days but began to decline rapidly by day 21. Expression of NMDA receptor subunits detected at the later stages of neuron culture was less than half that identified on day 14. The decrease in NMDA1 and -2 subunit expressions seemed to be directly associated with aging of the cortical neurons after initial differentiation. The decline of expressions of NMDA receptor subunits NMDA1 and -2 (NR1 and NR2A in the rat; NMDA1 and NMDA2 in humans) in an age-dependent manner in vivo is well established and may also have effects on agonist and antagonist binding to the intact receptor (Eckles-Smith et al., 2000; Ossowska et al., 2001; Magnusson et al., 2002). Thus, the use of primary mouse cortical neurons rather than cell lines provided not only a more physiologically relevant model but also the possibility of comparing gene expression patterns in the aging process. The NMDA receptors are important in neuronal cell signaling because of their role in intracellular Ca\(^{2+}\) homeostasis, and their dysfunction can be linked to loss of neuronal integrity and death (Nicotera, 2003). The maximum expression of the gene for NF-L at day 14 reflected a demand for cytoskeleton protein during intense development of the neurites. Again, subsequent lowered expression of NF-L by day 21 may be linked with degenerative processes in older cortical neurons. In additional studies (T. Chernova and A. G. Smith, unpublished data), gene array analysis demonstrated that other synapse-linked genes may also be down-regulated in older cell cultures, whereas at the peak of differentiation, many genes were expressed to a degree similar to that in adult frontal brain cortex. On the other hand, we have found little evidence for marked changes in the expression of inhibitory receptor GABA\(_A\) subunit \(\alpha1\) over time.

![Graphs showing changes in gene expression](https://example.com/graphs)

**Fig. 3.** Changes in gene expression in prolonged culture of primary cortical neurons. Heme-related (A–C) and neuron-specific (D–F) gene expression relative to \(\beta\)-actin expression estimated by real-time RT-PCR. G, detection of HMOX1, HMOX2, NMDA1 receptor subunit, and NF-L proteins by immunoblotting in representative cultures of cortical neurons on day 21 compared with day 14. Equal amounts of protein were loaded on to the gels. All were significantly different.
In sharp contrast to genes associated with neuronal function, expression of genes associated with control of heme synthesis and catabolism (those for ALAS1 and HMOX1) were markedly induced in older cultures of neurons on day 21 and longer, by which time the proportion of senescent neurons in cultures was estimated at approximately 86%. Alas1, the gene for the rate-controlling enzyme of nonerythroid heme synthesis, is usually considered to be up-regulated in liver by repression of a negative feedback mechanism as a consequence of a depleted regulatory heme pool (Sassa and Nagai, 1996). Our findings agree with in vivo studies that this mechanism probably also operates in the brain (De Matteis et al., 1981; De Matteis and Ray, 1982). Up-regulation of Alas1 in neurons may be evidence of limitation of heme supply with aging of the cells. This was probably compounded by the simultaneous greater expression of the heme-degrading enzyme HMOX1, which is often associated with conditions of cellular stress and may be an attempt to gen-

Fig. 4. Effects of inhibitors on heme synthesis and gene expression in primary cortical neurons. Neurons were cultured in the presence of 0.5 mM SA or 1 μM NMP for 12 or 18 days. A, treated and control cells were incubated with [3H]ALA for 24 h on day 12 and then labeled heme was extracted and measured (n = 5). B and C, expression of ALAS1, HMOX1, and HMOX2 in SA-treated and control cultures on days 12 and 18, estimated by real-time RT-PCR. D, ALAS1, HMOX1, and HMOX2 expressions in NMP-treated primary cortical neurons and control cells on day 12. *, statistically different from control group (p < 0.05). E, detection of HMOX1 and HMOX2 proteins in representative cultures of succinyl acetone-treated and control neurons on day 12.

Fig. 5. Effects of inhibition of heme synthesis on NMDA1 and NMDA2 receptor subunits and NF-L expression in cortical neurons. A–C, effects of SA at days 12 and at 18 estimated by real-time RT-PCR. D, gene expression in NMP-treated neurons and control cells at day 12 (D). 1, NMDA1; 2, NMDA2; 3, NF-L. *, statistically different from untreated control group (p < 0.05).
erate the antioxidant bilirubin as well as the signaling molecule CO (Sassa, 2004). However, this mechanism must depend on the synthesis of considerable amounts of extra heme as a substrate in addition to that required for normal cellular functions, because little free heme is likely to be available for this purpose in cells (Sassa, 2004). Elevated activity of heme oxygenases may exacerbate a problem, already present, of inadequate heme supply in aging cells.

Although the changes in older culture of primary neurons were consistent with heme depletion partly associated with induction of HMOX1, and also a decline in expression of NMDA receptor subunit genes, it was not clear whether there was any link between these systems. To address this point, differentiating cultures of neurons displaying maximum expression of NMDA subunits were treated with SA or NMP to inhibit different steps in heme synthesis, thereby potentially producing heme-depleted cells (De Matteis et al., 1981; Tschudy et al., 1981; Sassa and Nagai, 1996). Treated cells did indeed show significantly depressed heme synthesis, with patterns of \( \text{Alas1} \) and \( \text{Hmox1} \) up-regulation observed in much older cultures of untreated cells. In contrast, there was decreased expression of NMDA receptor subunits and NF-L, especially with NMP. This could be considered premature aging of the primary neurons and seemed to indicate that there could be a direct relationship between heme supply and the expression of these neuron-specific genes. We were unable to exclude from this the possibility that SA or NMP might inherently affect gene expression or some other process that might influence that of the subunits of NMDA receptor. However, when heme was added back at physiologically relevant levels to SA-treated cells, expression of the \( \text{Alas1} \) and \( \text{Hmox1} \) genes was decreased and expression of those for NMDA receptor subunits was mostly restored.

The use of primary cortical neurons from \( \text{Fech} \) mice obviated the need for chemical intervention to cause a depleted...

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**Fig. 6.** Rescue of gene expression in SA-treated neurons by heme. Heme added back for 24 h to neurons after treatment with SA for 12 days eliminated up-regulation of ALAS1 (A) and HMOX1 (B). Recovery effects of added heme on NMDA receptor subunit expression in BALB/c primary neurons (C and D) and in \( \text{Fech} \) primary neurons (F) estimated by real-time RT-PCR. Expression of GABA\( _{\alpha} \)-\( \alpha1 \) subunit in primary neurons treated with SA and heme (E). * , statistically different from untreated control group (\( p < 0.05 \)).

**Fig. 7.** Heme synthesis and gene expression in primary cortical neurons from \( \text{Fech} \) mice compared with time-matched BALB/c cells. A, reduced ferrochelatase activity in brain cortex from adult mice. B, heme synthesis in cortical neurons from \( \text{Fech} \) mice compared with BALB/c mice after incubation with \( ^{3} \text{H} \)ALA for 24 h on day 12 of culture as in Fig. 3 (\( n = 3 \)). Heme related (C–E) and neuron-specific (F–H) gene expression in BALB/c and \( \text{Fech} \) neurons estimated by real-time RT-PCR on days 6 and 14 of culture. * , statistically different from control group (\( p < 0.05 \)).
heme supply (Tutoiu et al., 1991). This allowed investigation of whether 1) up-regulation of HMOX1 in a stressed environment was the primary event with subsequent up-regulation of Alas1 or 2) heme depletion per se occurred first and was compounded by induction of HMOX1 activity as cells aged. Not only was Alas1 up-regulated to a greater degree in Fech primary culture than in wild-type BALB/c cells at any time point as a result of defective heme synthesis but also neurons displayed signs of premature senescence. Modest induction of HMOX1 expression in Fech embryonic cells was possibly associated with providing defense against oxidative stress but at the same time enhancing the potential for heme depletion. It is noteworthy that after an initial increased expression compared with that in BALB/c cells, NMDA receptor subunits, and characteristics of heme deficiency may be compounded by induction of HMOX1 activity as cells aged. This allowed investigation of the relationship between neuronal function and heme levels in neurons of different age and species. Studies of neuronal degeneration in animal models have shown that heme deficiency may lead to the deterioration of neuronal cells with subsequent up-regulation of heme synthesis. This results in a net increase in heme levels and suggests that heme itself could affect expression of these neuron-specific genes. An interesting finding of different effects of heme deficiency on GABA neurotransmitter receptor subunit and glutamate receptor NMDA subunit may lead to better understanding of why heme-distorted metabolism results in certain clinical manifestations in patients. One of the possible explanations of this distinctive difference in effects for NMDA and GABA receptors could be related to regulation by different signaling pathways (Zhu et al., 2002; Kumar et al., 2005).

What could be the mechanism of the senescence of cortical neurons being affected by heme supply? Studies of human brain cell lines have shown that inhibitors of heme synthesis can activate NO synthesis and alter zinc and iron metabolism, and that cells fail to differentiate or undergo a successful cell cycle (Atamna et al., 2002). This might be due in part to a decrease in mitochondrial function, especially complex IV, as a consequence of specific disruption in the synthesis of heme a. However, hepatic mitochondrial respiratory chain activities, including complex IV (cytochrome oxidase), remained unchanged or were increased in Fech mice (Navarro et al., 2005). Observations of neurite outgrowth have indicated that heme may act by regulation of kinases concerned with structural proteins and receptors (Ishii and Maniatis, 1978). It is interesting that inhibition of heme synthesis interferes with neuron growth factor-induced outgrowth of PC-12 cells by diminishing a subset of neuron-specific genes expressed via the Ras/mitogen-activated protein kinase signaling pathway, including NF-L (Zhu et al., 2002; Sengupta et al., 2005).

In summary, long-term primary cultures of mouse cortical neurons displayed senescence, decreased expression of NMDA receptor subunits, and characteristics of heme deficiency, as have been observed in vivo for aging. Down-regulation of NMDA receptor subunit expression was potentiated by both inhibitor and mutant models of heme depletion, strongly suggesting a mechanistic link between these metabolic processes.

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