Erythropoietin Modulates Cerebral and Serum Degradation Products from Excess Calpain Activation following Prenatal Hypoxia-Ischemia

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Abstract
Preterm infants suffer central nervous system (CNS) injury from hypoxia-ischemia and inflammation – termed encephalopathy of prematurity. Mature CNS injury activates caspase and calpain proteases. Erythropoietin (EPO) limits apoptosis mediated by activated caspases, but its role in modulating calpain activation has not yet been investigated extensively following injury to the developing CNS. We hypothesized that excess calpain activation degrades developmentally regulated molecules essential for CNS circuit formation, myelination and axon integrity, including neuronal potassium-chloride co-transporter (KCC2), myelin basic protein (MBP) and phosphorylated neurofilament (pNF), respectively. Further, we predicted that post-injury EPO treatment could mitigate CNS calpain-mediated degradation. Using prenatal transient systemic hypoxia-ischemia (TSHI) in rats to mimic CNS injury from extreme preterm birth, and postnatal EPO treatment with a clinically relevant dosing regimen, we found sustained postnatal excess cortical calpain activation following prenatal TSHI, as shown by the cleavage of alpha II-spectrin (all-spectrin) into 145-kDa all-spectrin degradation products (all-SDPs) and p35 into p25. Postnatal expression of the endogenous calpain inhibitor calpastatin was also reduced following prenatal TSHI. Calpain substrate expression following TSHI, including cortical KCC2, MBP and NF, was modulated by postnatal EPO treatment. Calpain activation was reflected in serum levels of all-SDPs and KCC2 fragments, and notably, EPO treatment also modulated KCC2 fragment levels. Together, these data indicate that excess calpain activity contributes to the pathogenesis of encephalopathy of prematurity. Serum biomarkers of calpain activation may detect ongoing cerebral injury and responsiveness to EPO or similar neuroprotective strategies.

Introduction
Preterm birth alters the microenvironment in the developing central nervous system (CNS) over a sustained period of maturation before, during and after birth. The cumulative injury causes an amalgam of damage termed encephalopathy of prematurity [1]. Improved identification and refinement of therapeutic interventions depends upon a precise understanding of the mechanisms and duration of injury and potential treatment targets. For ex-
ample, the emerging neurorestorative agent erythropoietin (EPO) is currently being administered in a clinical trial for extremely preterm infants born before 28 weeks estimated gestational age without adjustment of the dosing regimen to the type or severity of the initial CNS injury, the presence of cumulative secondary insults or the individual vulnerability and response of the patient to the CNS injury (PENUT trial, NCT01378273). Serum biomarkers that reflect the CNS injury and the potential for recovery offer a promising modality to stratify patients and guide the dosing of emerging interventions.

Two major mediators of CNS injury are caspase and calpain activation [2]. Apoptosis identified by cleaved caspase-3 immunolabeling has been found in white matter lesions associated with preterm birth [3–5], and activated caspase-3 has been found in the cerebrospinal fluid of preterm infants [6]. EPO is a known CNS antiapoptotic agent [7]. Using a rat model of prenatal transient systemic hypoxia-ischemia (TSHI) that mimics encephalopathy of prematurity observed in human infants who are born extremely preterm [8], postnatal EPO administered in a clinically relevant paradigm reduced injury-induced caspase-3 activation and improved functional outcomes in adult rats [9]. However, the impact of EPO on calpain signaling in encephalopathy of prematurity has not been elucidated.

Calpains, calcium-activated proteases, have an active role in cellular homeostasis both during development and in the mature CNS by degrading proteins. Calpain activity is modulated in part by the endogenous calpain inhibitor calpastatin, and both calpain and calpastatin levels are developmentally regulated [10]. Appropriate calpain activity is essential for neurodevelopment, including learning and memory [11] and axon pruning [12]. In the CNS, calpain is quantified by measuring the degradation of full-length (250 kDa) membrane-associated cytosolic protein alpha II-spectrin (αII-spectrin) into 150- and 145-kDa spectrin degradation products (αII-SDPs) [13]. By contrast, activated caspase cleaves αII-spectrin into 150- and 120-kDa αII-SDPs [13]. In pathological neurodegenerative, demyelinating or post-traumatic processes in the mature CNS, excess calpain activity degrades molecules important for neurological function [14–17]. Previously, we have shown that prenatal TSHI induces excess NMDAR (N-methyl-D-aspartic acid receptor) activation and calcium-mediated calpain activation that limits potassium-claylde co-transporter (KCC2) expression in the hippocampal CA3 subfield, and that postnatal EPO treatment reverses excess calpain activation [18]. Here, we investigated the broader impact of excess calpain activation following prenatal HI injury.

In the present investigation, we examined the impact of calpain degradation in encephalopathy of prematurity in detail by determining the extent and time course of calpain degradation of molecules important to perinatal neurodevelopment. In addition, we investigated whether postinjury EPO treatment impacts the TSHI-induced loss of expression of molecules that reflect neurodevelopmental maturation. Specifically, we examined cortical expression levels of KCC2 oligomers, myelin basic protein (MBP) and the phosphorylated neurofilament/total neurofilament (pNF/NF) ratio following prenatal injury and postnatal EPO treatment. First, we focused on these molecules because of their importance in neuronal signaling, myelinization and axonal health during perinatal brain development, respectively. Second, these molecules were investigated because they are known calpain substrates [14–17]. Finally, we sought to determine whether serum levels reflect the cerebral changes in calpain degradation and EPO treatment. We hypothesized that prenatal HI that mimics a component of encephalopathy of prematurity will cause excess CNS calpain activation, subsequent degradation of key neurodevelopmental molecules and associated signs of elevated calpain activity in serum. Moreover, we postulated that postnatal treatment with the neuroprotective agent EPO could modulate excess CNS calpain activity, and that this effect of EPO treatment on calpain activity would be reflected in the serum. We provide novel evidence that TSHI induces a marked elevation of calpain activity in the CNS during a critical point in brain development, and that postnatal EPO treatment attenuates calpain activity and preserves downstream targets essential to normal CNS maturation. Further, we show that αII-SDP and KCC2 fragments are detected in serum following injury and may prove useful as biomarkers of CNS injury and EPO efficacy.

Materials and Methods

Prenatal TSHI

The Boston Children’s Hospital Institutional Animal Care and Use Committee approved all experimental protocols. Prenatal TSHI was performed on embryonic day 18 (E18) as previously described [8, 9]. Briefly, timed pregnant Sprague-Dawley dams were purchased from Charles River Laboratories, and laparotomy was performed under isoflurane anesthesia on E18 with uterine artery occlusion for 60 min. Shams underwent anesthesia and laparotomy without artery occlusion. Pups were born at term and remained with their dams until weaning on postnatal day 21 (P21). Rats of both sexes were used in all experiments.
EPO Modulates Calpain Activity following Prenatal HI

EPO Treatment

After birth, TSHI pups were randomly assigned to receive intraperitoneal EPO (2,000 U/kg/dose; R&D Systems) or vehicle (sterile saline) administered daily from P1 to P5. We have previously found that this dosing regimen results in sustained functional improvement in adult rats following prenatal TSHI [9, 22], and have documented that no significant knowledge was gained from a sham EPO cohort [9].

Tissue and Serum Collection

For immunohistochemistry at P2, brains were drop-fixed in 4% paraformaldehyde. After fixation, the brains were immersed in 30% sucrose and processed for immunohistochemistry as below. For immunoblotting, cortical samples were collected from at least two separate litters for each group and snap-frozen at –80°C. Specific numbers of samples analyzed in each experiment are provided. For P5 and younger, cerebral cortical samples were collected from the frontal lobes. Samples from two P11 brains were micro-dissected and pooled to provide an adequate final sample for the cortical membrane preparations for the KCC2 analyses and for the white matter MBP and NF analyses. Microdissected samples from single animals were used for P28. For serum analyses, blood was collected from each pup and centrifuged at 6,000 g for 15 min, and serum was removed and stored in aliquots at –80°C for later analysis. Care was taken to avoid repeated freeze-thaw cycles.

Immunoblotting

Western blots were performed as previously described [18, 22]. For analyses of cerebral tissue, samples were sonicated, and whole cell and membrane fractions were separated via differential centrifugation. Bradford protein assays were performed per manufacturer specifications (Bio-Rad, Hercules, Calif., USA). Proteins were separated on 4–20% Tris HCl gels, with the exception of MBP Western blots in which proteins were separated on 4–10% Bis Tris XT gels (all Bio-Rad). In the experiment to better delineate αII-SDP, 7.5% Tris HCl gels were used for optimal separation in the 120- to 150-kDa range. For cerebral tissue, 30 μg was loaded per lane and after electrophoretic separation the proteins were transferred to PVDF (polyvinylidene fluoride) membranes for incubation with antibodies. The following primary antibodies were used: αII-spectrin (Santa Cruz Biotechnology, Dallas, Tex., USA; 1:100), calpastatin (Santa Cruz Biotechnology; 1:100), KCC2 for 90-kDa N-terminal fragment (Santa Cruz Biotechnology; 1:100), KCC2 oligomers (Millipore, Billerica, Mass., USA; 1:1,000), MBP (Millipore; 1:500), NF (Millipore; 1:1,000) and pNF (SMI-312; Covance, Dedham, Mass., USA; 1:500). For loading controls, β-tubulin (Covance, Piscataway, N.J., USA; 1:200) in a buffer containing 100 mM Tris-Cl (pH 7.4), 20 mM DTT (dithiothreitol), 1% Triton X-100 and 1 mM CaCl2 for 30 min at room temperature. The reaction was terminated by the addition of 8× sample loading buffer containing 0.25 M Tris (pH 6.8), 0.2 M DTT, 8% SDS, 0.2% bromophenol blue and 20% glycerol in distilled water. Proteins were separated by electrophoresis, transferred, probed with target antibodies and visualized as above.

Statistical Analyses

For analysis of two groups (sham and TSHI) at one time point, differences were compared using two-tailed Student’s t test with unequal variance. For two groups over multiple ages and for analysis of the sham, TSHI-vehicle (TSHI-veh) and TSHI-EPO groups, two-way ANOVA was performed. Bonferroni’s post hoc correction was used for multiple comparisons. Analyses were performed using SPSS 21 (IBM, Armonk, N.Y., USA), with p < 0.05 considered significant.

Result

Cerebral Cortical Calpain Activity Is Elevated following Prenatal TSHI

A summary of the experimental design and primary end points with complete quantification is illustrated in figure 1. To determine the contribution of calpain and caspase activity to αII-spectrin degradation in our in vivo model, we performed a calpain digestion assay to identify primary αII-SDPs. Specifically, to resolve the primary αII-SDPs of 120, 145 and 150 kDa, we digested P7 sham control cortex with calpain in the presence of calcium, as previously reported [23]. In a calpain digestion assay of cortical αII-spectrin, αII-SDPs appeared in the sham P7 cortex at 145 and 150 kDa, with the 145-kDa band much more prominent than the 150-kDa band (fig. 2a). By contrast, only a faint band was present at 120 kDa, suggesting relatively minimal caspase activity. Similarly, to confirm the degradation of a second calpain target, we also exam-
ined the cleavage of known calpain substrate protein 35 (p35) into a p25 fragment [25]. As expected, in the sham P7 cortex, we found an increase in the p25 fragment in calpain-digested samples in the presence of calcium (fig. 2b). Next, we compared the pattern of αII-SDPs in a calpain digestion of the P11 sham cortex assayed alongside P11 TSHI cortex probed with αII-spectrin shows a band at 150/145 kDa that is similar to a band present in sham cortex digested with activated calpain but not present in sham cortex without calpain digestion. d P11 MBP. Calpain digestion of P11 sham white matter in the presence of calcium eliminates MBP, while P11 TSHI white matter assayed in an adjacent lane shows reduced expression. e P11 pNF/ NF. Calpain digestion of P11 sham white matter in the presence of calcium eliminates pNF and NF, while P11 TSHI white matter assayed in an adjacent lane shows reduced pNF and NF expression.

Fig. 2. Calpain digestions of cortex and white matter. a P7 αII-SDPs. Representative calpain digestion of P7 sham cortex incubated with αII-spectrin antibodies reveals a primary band at 250 kDa and a fainter band at 150 kDa. In the presence of calcium-activated μ-calpain, the αII-spectrin is digested into a faint 150-kDa, a heavy 145-kDa and a very faint 120-kDa band. b P7 p35/p25. Sham P7 cortex digested with calpain shows elimination of the p35 band, with a corresponding increase in the p25 fragment. c P11 αII-SDPs. Calpain digestion of P11 sham cortex assayed alongside P11 TSHI cortex daily from P1 to P5. Red dots (color in online version only) indicate when samples were collected from serum for quantification, while gray dots represent samples collected from frontal lobes (prior to P5) and cortex and white matter (older than P5) for quantification.

Fig. 1. Summary of experimental design and major data points. TSHI was performed on E18 via laparotomy with 60 min of uterine artery occlusion. Pups were born at term (E22). TSHI pups were randomized to receive either EPO (2,000 U/kg/dose) or vehicle.
to determine whether excess calpain activity would affect calpain targets in white matter such as MBP, pNF and NF. Thus, we performed calpain digestion assays to confirm that calpain degrades these molecules in developing white matter. Indeed, the digestion of P11 sham white matter samples with calpain in the presence of calcium degraded MBP (fig. 2d), pNF and NF (fig. 2e). These patterns were compared to representative P11 TSHI-veh white matter samples. Together, these data demonstrate that our assays could detect calpain activity and that calpain degrades molecules central to gray and white matter development, including MBP and pNF/NF. Thus, we proceeded with our investigation and quantitation of excess calpain activity in this model of prenatal TSHI.

To clarify the time course of calpain activity in the cerebral cortex following prenatal TSHI, the ratios of cleaved to full-length αII-spectrin at multiple early time points after prenatal TSHI (E18 + 6 h, E19, P0 and P2). No differences after prenatal injury were evident until P2 (n = 12–15, two-way ANOVA, p = 0.007; fig. 3a), approximately 6 days after the insult on E18, suggesting the delayed appearance of excess calpain degradation of αII-spectrin and thus the delayed onset of excess calpain activity within the CNS. To confirm that excess cerebral calpain activity was present following prenatal TSHI, we also quantified the degradation of p35 to p25 at P2 and found more p25 in the TSHI cortex (n = 10, p = 0.026; fig. 3b). Together, these findings show that excess CNS calpain activity is initially detected at P2 in the CNS in this model.

Fig. 3. Excess calpain activity is induced by prenatal TSHI and modulated by EPO treatment. a αII-spectrin ratio. Representative Western blot of αII-SDPs and ratio of cleaved to full-length cerebral αII-SDPs at 6 h (E18 + 6), 24 h (E19), 96 h (P0) and 6 days (P2) after prenatal TSHI compared to sham. A significant increase in αII-SDPs is present at P2. b P2 p35/25 ratio. The ratio of p25/p35 is increased at P2 following prenatal TSHI. c P11 αII-spectrin ratio. Following prenatal TSHI, the ratio of cleaved to full-length αII-SDP at P11 is elevated compared to sham. Postnatal treatment with EPO normalizes the ratio to sham levels (two-way ANOVA with Bonferroni correction). * p < 0.05, ** p < 0.01.

Postinjury EPO Treatment Limits Calpain Degradation

EPO treatment in vivo has been implicated in reducing calpain activity in other types of mature CNS insults [26],
but this mechanism has not previously been investigated in impaired CNS development from very preterm birth. To test whether postinjury postnatal EPO treatment alters calpain activity in this rat model of encephalopathy of prematurity, we analyzed cortical cleaved/full-length αII-SDP ratios at P11 and P28. Ratios of cleaved/full-length αII-SDPs were elevated in the P11 TSHI-veh cortex (n = 14) compared to shams (n = 22, two-way ANOVA, p = 0.002) or TSHI-EPO cortex (n = 14, p = 0.016; fig. 3c), and the differences resolved by P28 (data not shown). These data show that EPO can modulate CNS calpain activity in vivo following early injury to the developing brain.

TSHI Alters the Balance between Calpain and Calpastatin

Both calpain and the endogenous calpain inhibitor calpastatin are developmentally regulated [10], and calpain activity reflects the balance of activated calpain and calpastatin. Calpastatin immunolabeling in the cortex, subplate and white matter at P2 was reduced in TSHI brains compared to shams, with the most prominent difference evident in the subplate (fig. 4a). By contrast, contrast, immunolabeling with μ-calpain antibodies is increased after TSHI. Scale bar = 20 μm. b P7 calpastatin. Cortical calpastatin expression is reduced at P7 following prenatal TSHI compared to sham. c P11 calpastatin. Following prenatal TSHI, calpastatin expression is reduced at P11 compared to sham. Postnatal treatment with EPO restores calpastatin expression to sham levels (two-way ANOVA with Bonferroni correction). * p < 0.05.

Postinjury EPO Treatment Limits Calpain Degradation of Cortical KCC2 and White Matter MBP and NF

We hypothesized that in addition to injury-induced calpain degradation of αII-spectrin, excess CNS calpain activity caused by prenatal TSHI would also degrade other known calpain substrates, including molecules essential to neurological development and function. Calpain cleaves the neuronal potassium–chloride co-transporter KCC2 into 90-kDa N-terminal and 30-kDa C-terminal fragments that are labeled with N- and C-terminal-specific antibodies [19, 20], and calpain degradation is the primary cause of reduced KCC2 levels present in the CA3 region following prenatal TSHI [18]. Here, we examined...
cerebral cortical levels of KCC2 following prenatal TSHI and postnatal EPO treatment. Compared to shams (n = 12), P11 cortical levels of oligomeric KCC2 were decreased in prenatal TSHI-veh rats (n = 12, two-way ANOVA, p = 0.024) and improved with EPO treatment (n = 12, p < 0.001; fig. 5a). Consistent with our previous studies that showed a loss of KCC2 expression in the CA3 subfield [18] and cortical layer IV following prenatal TSHI [22], these results also suggest that postnatal EPO treatment can mitigate the loss of cerebral cortical KCC2 expression induced by prenatal TSHI.

Next, to quantify the loss of MBP and NF following prenatal TSHI, we assayed MBP levels and pNF/NF ratios, respectively. At P11 MBP levels were reduced in TSHI-veh samples (n = 12) compared to shams (n = 18, two-way ANOVA, p = 0.003) and TSHI-EPO brains (n = 20, p < 0.001; fig. 5b). At P11 samples from TSHI-veh brains (n = 26) assayed for pNF/NF also showed lower levels than shams (n = 24, p = 0.007) or TSHI-EPO brains (n = 22, p = 0.015; fig. 5c). To determine whether the loss of MBP and NF persisted in juvenile brains, we examined MBP and pNF/NF levels at P28. Similar to P11, the pNF/NF ratio at P28 was also reduced following prenatal TSHI and restored with postnatal EPO treatment. e P28 white matter MBP. Likewise, the white matter MBP levels remain lower following prenatal TSHI and normalize with EPO treatment (two-way ANOVA with Bonferroni correction). * p < 0.05, ** p < 0.01, *** p ≤ 0.001.

Fig. 5. Calpain degrades molecules essential for neurodevelopment, including KCC2, MBP and NF. a P11 cortex KCC2. At P11 KCC2 oligomer expression is reduced following prenatal TSHI, and KCC2 levels are restored following postnatal EPO treatment. b P11 white matter MBP. At P11 in the midst of myelin development, MBP expression is reduced following prenatal TSHI and normalizes following EPO treatment. c P11 white matter pNF/NF ratio. Similarly, at P11 the ratio of pNF/NF is lower following prenatal TSHI and restored following EPO treatment. d P28 white matter pNF/NF ratio. At P28 in juvenile white matter, the pNF/NF ratio at P28 is also reduced following prenatal TSHI and restored with postnatal EPO treatment. e P28 white matter MBP. Likewise, the white matter MBP levels remain lower following prenatal TSHI and normalize with EPO treatment (two-way ANOVA with Bonferroni correction). * p < 0.05, ** p < 0.01, *** p ≤ 0.001.
are reduced following prenatal TSHI for a sustained, critical period of postnatal CNS development, probably due in part to calpain degradation, and that these levels are responsive to postnatal EPO treatment.

**Serum αII-SDP Levels following Prenatal TSHI**

For selected neuromolecules, serum levels reflect CNS levels, and the presence of CNS molecules in the serum is a biomarker of brain injury. As cerebral cortical αII-SDP levels are elevated following prenatal TSHI, we investigated the presence of serum 145-kDa αII-SDPs following prenatal TSHI along the same time course. Serum 145-kDa αII-SDP levels were elevated in TSHI-veh rats (n = 10) at E19, 24 h after the E18 injury, compared to shams (n = 15, t test, p = 0.012; fig. 6a). Serum αII-SDPs remained elevated in serum at P0 following TSHI compared to shams (both: n = 11, p = 0.037; fig. 6b) and at P2 (sham: n = 15, TSHI: n = 10, p = 0.022; fig. 6c). By P7, however, the difference was no longer significant (sham: n = 11, TSHI: n = 8, data not shown). Together, these results suggest that the serum αII-SDP levels indicative of excess calpain activity rise by 24 h after the injury and remain elevated through P2, approximately 6 days after the insult.

**Serum KCC2 Degradation Product Levels**

KCC2 levels rise over a critical period of cortical circuit development [18, 27], with the predominant increase in the cerebrum in the second postnatal week [22, 28]. Because calpain degrades KCC2 into 90-kDa fragments [19, 20], we hypothesized that the KCC2 fragments may be detectable in serum as a surrogate marker of cerebral neurodevelopment and injury. Serum levels of KCC2 90-kDa degradation fragments were elevated in serum from P11 TSHI-veh rats (n = 13) compared to shams (n = 10, two-way ANOVA, p = 0.02) and were normalized in TSHI-EPO rats that had received postnatal EPO treatment (n = 11, p < 0.001; fig. 6d), mirroring the decreased oligomeric KCC2 expression found in the brain at the same time point. Together, these results suggest that serum 90-kDa KCC2 fragments may serve as a biomarker of injury to the developing cerebrum and of responsiveness to successful treatment with a neuroprotective agent.

**Discussion**

Signs of excess CNS calpain activity have been found in both preclinical [23] and clinical studies of pediatric traumatic brain injury [29]. In rat models of term birth perinatal brain injury, excess calpain activity was present for >48 h after P7 HI injury [30–32] and 5 days after P10 neonatal stroke [33]. Using an in vitro model of oxygen-glucose deprivation injury at P7, the addition of EPO inhibited calpain activity [34]. Calpain activity and its responsiveness to EPO treatment have not previously been investigated in vivo in a model that mimics encephalopathy of prematurity. Here, we found evidence of both excess serum and cerebral calpain activity in vivo over an extended postnatal period, using an established model of CNS injury from preterm birth induced in rats by TSHI on E18 [8, 35]. Serum calpain activity measured by αII-SDP fragment levels was elevated 24 h after the E18 insult, remained increased through P2, and normalized by P7.

![Fig. 6](Image) Calpain degradation products are present in serum. Serum levels of 145-kDa αII-SDP are elevated at E19 (a), P0 (b) and P2 (c, unpaired two-tailed t test). Coomassie staining was used for the loading control. d P11 serum KCC2 fragment. At P11 serum levels of the 90-kDa KCC2 calpain fragment are elevated following prenatal TSHI and normalized with postnatal EPO treatment (two-way ANOVA with Bonferroni correction). * p < 0.05, *** p ≤ 0.001.
Interestingly, prenatal TSHI induces a marked elevation of cerebral calpain activity, as shown by an increase in the ratio of cleaved to full-length αII-SDP in the cerebral cortex beginning at P2, 6 days after injury. The excess cerebral calpain activity is sustained through at least P11, 2 weeks following prenatal TSHI, and approximately equal to slightly after full term in humans. The extended period of excess cerebral calpain activity correlates with the protracted period of apparent CNS injury in encephalopathy of prematurity [36, 37]. To our knowledge, this is the first demonstration that excess calpain activity is present in a model of encephalopathy of prematurity, and that degradation products can be assayed in postnatal serum.

The pattern of CNS injury observed in preterm infants is complex and multifactorial [1]. It involves injury to both white and gray matter and encompasses injury to immature oligodendrocytes, axons, neurons and developing circuits over a sustained portion of a critical window of neurodevelopment. Developing white matter is commonly assayed biochemically through the investigation of MBP, and axonal integrity is examined via the ratio of phosphorylated to total NF levels [35]. In gray matter, the excitatory/inhibitory balance appears particularly sensitive to third-trimester injury [38]. In multiple developing cerebral cortical regions, the upregulation of membrane-bound KCC2 expression determines the intracellular chloride concentration. In a simplified view, increasing KCC2 chloride extrusion leads to the transition from a depolarizing to a hyperpolarizing response to GABA (γ-amino butyric acid) activation of GABA A and glycine receptors [27]. A loss of KCC2 expression was found in postmortem cerebral samples from human preterm infants with white matter lesions [39]. In a preclinical model, prenatal global HI injury reduced KCC2 expression in the hippocampal CA3 subfield [18], and premature loss of subplate from this injury caused KCC2 loss in cerebral cortical layer IV [22]. Here, we show that cerebral levels of known calpain substrates that are also key neurodevelopmental molecules, including KCC2, MBP and NF, are reduced for an extended postnatal period through P11, consistent with prior reports that excess calpain activity contributes to KCC2 loss in the hippocampal CA3 subfield [18]. While the reduction in MBP and NF expression following prenatal injury is likely to be multifactorial, here we show that the reduced expression of MBP and pNF/NF ratio also correlates with the excess calpain activity.

EPO is an emerging neuroprotective agent with multiple mechanisms of action that is currently being tested in the PENUT trial (NCT01378273) in infants born prior to 28 weeks gestation. Using a postnatal EPO dosing regimen that has previously produced functional improvement following prenatal TSHI [9, 22] and that is similar to the dosing regimen in the PENUT trial, we found that EPO treatment from P1 to P5 significantly modulated cerebral calpain activity at P11. Postnatal EPO treatment also restored levels of MBP and the pNF/NF ratio at P11 and P28. While EPO has previously been shown to reduce excess calpain activity in the mature CNS [26], this is the first time that EPO modulation of excess calpain activity in the developing brain in vivo has been reported.

To further elucidate the potential mechanisms of EPO neuroprotection in the last trimester, we investigated the role of excess calpain activity and the degradation of molecules essential to white and gray matter function. Unbound EPO receptors drive neural cells to apoptosis as part of the normal neurodevelopmental regulation of neuronal number [40–42], and prenatal TSHI induces a marked upregulation of EPO receptor expression in the CNS without concomitant adequate ligand expression [9, 35]. Prenatal TSHI causes excess cerebral caspase-3 activation [8], and that is mitigated by postnatal EPO treatment [9]. While excess caspase and calpain activation have been implicated in models of term birth CNS injury [30, 31, 33, 43], calpain activation or its modulation has not been previously explored in vivo in a preclinical model of encephalopathy of prematurity. Excess calpain activity is associated with a sustained reduction in the cerebral expression of molecules important for neurological development and calpain targets, including the co-transporter KCC2, MBP and NF. Notably, the reduced expression of these molecules was restored by postnatal treatment with EPO. The loss of these proteins is not solely attributable to excess calpain activity and probably varies depending on the molecule. For example, KCC2 levels are quite sensitive to calpain degradation mediated through calcium activation of NMDA receptors [18–20]. By contrast, numerous stages in the development of oligodendrocytes are impacted by prenatal TSHI, including oligodendrogenesis, survival and process extension [8, 44], and EPO treatment reverses these deficits [44], suggesting that calpain degradation of myelin is only one component of myelin deficiency of encephalopathy of prematurity. Excess calpain degradation contributes to NF loss in preclinical models of traumatic injury and multiple sclerosis, and calpastatin overexpression or calpain inhibition restores the loss [12, 17, 45–47]. Calpain inhibition has also restored neurological function in preclinical in vivo models of Alzheimer’s disease, tauopathy and Parkinson’s disease [48–50]. Importantly, calpain inhibition...
improves functional outcomes in neurodevelopmental processes, including lissencephaly [51] and spinocerebellar ataxia type 3 (Machado-Joseph disease) [52]. Together, the progress in these other disease processes and our demonstration of the EPO-induced reduction of cerebral calpain activity and the associated loss of critical CNS developmentally regulated molecules suggest that the modulation of calpain activity over a sustained period may be beneficial to long-term neurological outcomes in neonates with encephalopathy of prematurity.

The mechanism of EPO-induced reduction in calpain activity in this model of encephalopathy of prematurity is not yet known. Because activated caspases cleave calpastatin [53, 54], and here the reduced levels of cerebral calpastatin expression were restored with EPO treatment, one potential mechanism for the effect of EPO on calpain activity may be by reducing calpastatin degradation and restoring the balance of calpain and calpastatin activities. Alternatively, EPO treatment may be affecting apoptosis and subsequent calpain activation through another pathway unrelated to calpastatin. For example, D’Orsi et al. [55] showed that calpain activation occurs downstream of mitochondrial engagement during excitotoxic apoptosis. Additionally, in Alzheimer’s disease μ-calpain activates GSK-3β [56]. Because EPO signaling modulates kinase activity, including activating Akt and inhibiting GSK-3β [57], it may be impacting calpain activity by altering the balance of Akt/GSK-3β activation. Another possibility involves an imbalance of vascular endothelial growth factor signaling after injury that triggers calpain activity and is modulated by EPO treatment, similar to the mechanism suggested in a preclinical model of cerebral malaria [26]. In a nonhuman primate model of neonatal asphyxia, the addition of EPO treatment to postinjury hypothermia significantly improved chronic behavioral outcomes, which suggests that EPO can synergize with other neuroprotective strategies, perhaps through the modulation of calpain activation [58]. Clarifying the details of the mechanism is beyond the scope of this work but is being actively investigated. In addition to directly limiting calpain activity, EPO may counteract the detrimental effects of excess calpain activity that limit neurorepair. For example, after stroke in the mature CNS, excess calpain activity hinders neural stem and neuroblast proliferation and migration [59]. EPO signaling promotes multiple stages of neuronal and oligodendroglial development [44, 60] and thus may counterbalance calpain-induced suppression of repair. The pattern of sustained elevation of excess calpain activity found here is similar to the sustained elevation of caspase-3 activity previously observed in this model [8, 9] and may contribute to the sustained period of persistent lack of recovery that has been implicated in human preterm infants [36, 37].

Elevated cerebral calpain activity is reflected in serum in this model of encephalopathy of prematurity. Here we have presented three novel findings related to the serum levels. To our knowledge, this is the first time that the elevation of serum αII-SDP levels has been shown with CNS calpain activity in vivo in a clinically relevant model of encephalopathy of prematurity. This suggests the possibility that serum biomarkers could be used not only to diagnose CNS injury in the neonatal period, but also to guide the use of neuroprotective interventions such as EPO. For example, preterm infants with CNS insults are prone to cumulative injuries from additional postnatal problems such as sepsis and seizures, and serum biomarkers could guide the duration of treatment for CNS protection. The need for serum biomarkers for preterm infants, as well as infants with other perinatal CNS insults, is well recognized [61–63].

Finally, calpain-mediated KCC2 degradation products were also found in serum after prenatal TSHI and appeared responsive to EPO treatment. Cerebral KCC2 expression rises significantly in the second postnatal week [64]. Calpain cleaves KCC2 into a 90-kDa fragment [18–20], which prompted us to investigate whether serum would reflect ongoing CNS injury in the postnatal period and potential restoration with EPO treatment. Sustained excess CNS calpain activity following prenatal TSHI is likely to contribute to the prolonged period of impaired neurodevelopment observed in extremely preterm infants. Moreover, postnatal EPO treatment modulated 90-kDa KCC2 levels after prenatal injury, with restoration back to sham levels. These results suggest that KCC2 calpain fragments can be assayed in serum following perinatal CNS injury, and that they may prove useful as CNS injury biomarkers of both injury and EPO responsiveness. Given the importance of KCC2 expression to neuronal health [27], serum KCC2 fragments offer a promising line of investigation for future study. Because 90-kDa levels are currently only detectable with immunoblotting to separate fragments, additional assay development is needed prior to widespread clinical utility and is already underway. Patterns of serum biomarker trends may be more informative than a single marker [65] and provide a less invasive means to monitor the type, extent and duration of CNS injury in critically ill infants who are too ill to undergo detailed high-resolution imaging studies.
EPO Modulates Calpain Activity following Prenatal HI

References


