Infection-induced Inflammation and Cerebral Injury in Preterm Infants

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Summary

Preterm birth and infectious diseases are the most common causes of neonatal and early childhood deaths worldwide. The rates of preterm birth have increased over recent decades and currently account for 11% of all births globally. Preterm infants are at significant risk of severe infection in early life and throughout childhood. Bacteraemia and/or inflammation during the neonatal period in preterm infants is associated with adverse outcomes, including death, chronic lung disease and neurodevelopmental impairment. Recent studies suggest that bacteraemia may trigger cerebral injury even without penetration of viable bacteria into the central nervous system. Here we review available evidence that supports the concept of a strong association between bacteraemia, inflammation and cerebral injury in preterm infants, with an emphasis on the underlying biological mechanisms, clinical correlates and translational opportunities.
Introduction

Globally, more than 15 million infants are born preterm (<37 completed weeks of gestation) each year, and over one million die. The rates of preterm birth have increased over the last several decades and affect approximately 11% of all pregnancies. Preterm birth now is the most common cause of neonatal mortality and will likely surpass pneumonia as the leading cause of death in early childhood by 2015. Importantly, a significant proportion of the survivors of preterm birth suffer long-term neurological disabilities and evidence suggests that exposure to neonatal infection is a major contributor to cerebral injury in this population. Current treatment strategies for neonatal infections, however, largely focus on optimal antimicrobial activity without specifically targeting infection-induced inflammation. Accordingly, novel therapeutic approaches aimed at modulation of infection-related inflammatory responses may improve long-term outcomes. We conducted a literature search focused on chorioamnionitis, bacteraemia, sepsis, and necrotising enterocolitis in order to summarise the current state of the art with respect to mechanisms and potential mitigating agents for inflammation-induced preterm cerebral injury. Inflammation and cerebral injury caused by viral infections are beyond the scope of this review.

The burden of exposure to perinatal inflammation in preterm infants

The incidence, morbidity and mortality of neonatal infection

In developed countries, approximately 1% of all live births are affected by neonatal infections. Worldwide, infections account for two thirds of the 7.6 million annual deaths in children less than 5 years of age. The neonatal period carries the highest lifetime risk of serious infections, with an estimated 400,000 newborn deaths annually. Neonatal infections disproportionately (~80%) occur in the minority of infants born preterm (8-12%), who have a several times higher risk of invasive bacterial infection than term infants. Depending on gestational age at birth, 25-60% of extremely preterm infants (<28 week gestation) develop at least one invasive bacterial infection during their birth-related hospital admission and recurrent neonatal infections are common. Importantly, the heightened vulnerability to serious infection persists into later childhood and the infection-related morbidity and mortality is not limited to extremely preterm infants, but also affects the much larger proportion of moderate and late preterm infants.

Globally, the majority of moderate and late preterm births occur in resource-poor settings, where data are less easily collected and consequently less robust, but where the incidence of invasive infection is likely to be substantially higher than that reported for high-resource settings. The surviving preterm infants in resource-poor settings are likely to be moderately preterm but of low birth weight, further increasing the risk for neonatal and childhood infection and infection-related mortality.

The burden of exposure to perinatal inflammation

Chorioamnionitis (inflammation of the placental chorionic disc, extraplacental membranes, cord and/or amniotic fluid) affects 2-5% of all births, is intrinsically linked to premature rupture of membranes, spontaneous onset of preterm labour and is an important risk factor for early-onset neonatal infection. Large retrospective cohort studies demonstrate a strong...
inverse relationship between gestational age, birth weight and incidence of histologically
diagnosed chorioamnionitis, which is present in approximately 65% of placentae at 23 to 24
weeks of gestational age, 30% of placentae at 29 weeks gestational age, and 2-14% at
term.\textsuperscript{14,15} The clinical diagnosis of chorioamnionitis is unreliable and therefore studies
without placental histology are likely to underestimate significantly the true incidence of
chorioamnionitis and its biological effects.\textsuperscript{16} Chorioamnionitis is frequently caused by
fastidious organisms that are not readily cultured with routine microbiological techniques.
However, culture-independent methods, such as detection of conserved bacterial 16sRNA
by polymerase-chain reaction, have demonstrated the presence of microorganisms in
placental tissues and/or amniotic fluid in the majority of histologically confirmed
chorioamnionitis.\textsuperscript{17} Failure to recover fastidious organisms may also explain association of
histological chorioamnionitis with adverse short- and long-term neonatal outcomes when
routine placental bacterial culture is sterile.\textsuperscript{18} Thus even culture-negative, asymptomatic
chorioamnionitis that does not result in early-onset neonatal infection may lead to persistent
activation of the inflammatory response and have profound and pervasive effects by altering
maturation of the neonatal immune system and longer term infection risk.\textsuperscript{19,20} Emerging
data suggest that exposure to chorioamnionitis not only leads to increased neonatal
morbidity, but also may have long-term effects on immune-related outcomes such as an
increased risk of childhood asthma.\textsuperscript{21,22}

\textbf{The impact of infection and inflammation on cerebral injury and
neurodevelopment in preterm infants}

\textbf{Human data}

The potential link between perinatal inflammation, neonatal sepsis and cerebral injury was
first noted over 30 years ago, when both autopsy data and subsequently cranial ultrasound
studies showed an increased risk of periventricular leukomalacia in infants exposed to
maternal infection or neonatal sepsis.\textsuperscript{23,24} The association between maternal infection,
chorioamnionitis and a several-fold increased risk of cerebral palsy was not limited to high-
risk preterm infants, but was also observed in term infants.\textsuperscript{25-27}

The commonest lesion associated with inflammation in the preterm infant is white matter
injury, which is characterised by focal cystic periventricular leukomalacia and/or diffuse
necrosis. White matter injury is defined by loss of immature preoligodendrocytes, which
would normally mature to ensheath axons with myelin, but which are particularly
susceptible to oxidative stress and inflammation.\textsuperscript{28,29} Further mechanisms of injury involve
inhibition of neuronal precursor cell proliferation and activation of astroglia.\textsuperscript{30,31}

Over the last decade, observational studies have provided more detail on the association
between neonatal sepsis and adverse long-term neurological and neurocognitive
outcomes. Data from a large US Neonatal Network demonstrates that any form of neonatal
infection, including clinical infection, culture-proven sepsis, meningitis with or without
sepsis, and necrotising enterocolitis with or without sepsis, is associated with poor growth and
increased risk of neurodevelopmental impairment.\textsuperscript{32,33} Similarly, several studies of preterm
infants in Europe and Canada demonstrate an association between late-onset sepsis and

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adverse neurodevelopmental outcomes in childhood, with repeated infections and Gram-negative pathogens conferring the highest risk. These findings are not limited to extremely preterm infants; in a Brazilian study of moderately preterm infants, neonatal sepsis is also strongly and independently associated with increased risk of cerebral injury. Finally, in a US cohort, late-onset sepsis is independently associated with neurodevelopmental impairment in preterm infants with necrotising enterocolitis. Sophisticated magnetic resonance imaging (MRI) protocols allow increasingly detailed analyses of cerebral injury that may not be detected by cranial ultrasound examination, which is accurate for cystic periventricular leukomalacia, but has limited sensitivity for diffuse white matter injury. MRI analyses predict long-term neurological outcomes. In preterm infants who have had neonatal sepsis there is an increased risk for white matter injury on MRI and of motor impairment on clinical examination. In addition, recurrent neonatal culture-positive infection (without meningitis) is associated with a significantly greater risk of progressive white matter injury. Furthermore, late-onset sepsis in preterm infants, both microbiologically proven and clinically diagnosed, is associated with acute alteration of cerebral function, indicated by acute changes in electrographic activity and burst suppression pattern. Gram-negative neonatal sepsis has a significantly higher mortality than sepsis caused by the most commonly isolated group of Gram-positive organisms, coagulase-negative staphylococci (which accounts for ~50-75%). However, the association between sepsis and cerebral injury appears to be largely independent of the bacterial species involved, indicating that a detrimental final common pathway can be activated by diverse initial host-microbe interactions. Neonatal clinical sepsis (i.e., signs of infection with negative microbial cultures) is a risk factor for preterm infant white matter injury in univariate analysis, whereas culture-positive infections (predominantly sepsis, but also cases of urinary tract infection and pneumonia in the absence of meningitis), are also a significant risk factor for white matter injury after adjustment for common confounders. Challenges in interpreting studies of the association between infection and brain injury include i) cohort variability, predominantly consisting of retrospective studies and clinical trials subject to selection bias, non-uniform definitions of ii) chorioamnionitis (clinical versus histological) and of iii) neonatal sepsis and iv) wide variations in clinical management (Table 1). Importantly, there is no universally accepted gold standard for diagnosing neonatal infection and hence definitions commonly include variable combinations of the following: a) positive culture from sterile site, predominantly blood, cerebrospinal fluid and urine, b) clinical signs such as respiratory distress, apneas, temperature instability, feed intolerance etc., which are both non-specific and insensitive), c) elevated inflammatory markers such as C-reactive protein whose use and cut-off values are variable, and d) intention-to-treat duration of antibiotic therapy which is highly variable between units. Furthermore various methods are used to quantify cerebral injury in neonates, including postmortem examination of the brain, cranial ultrasound, MRI and standardised clinical evaluations of neurodevelopmental outcome. Despite these methodological shortcomings, there is robust evidence for an heightened risk of cerebral injury and adverse effects on neurodevelopmental outcomes following a variety of perinatal

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inflammatory exposures, including maternal infection, maternal and fetal chorioamnionitis, and early and late neonatal infection.

Humoral mediators, including pro-inflammatory cytokines such as interleukin (IL)-1, IL-6 and chemokines such as CXCL-8 (formerly named IL-8) as well as tumour necrosis factor-α (TNF), type I and II interferons and reactive oxygen species are likely to be key mediators in the pathogenesis of cerebral injury. Levels of these cytokines/chemokines are elevated in amniotic fluid, cord blood, cerebrospinal fluid and cerebral tissue of infants diagnosed with inflammation-related white matter lesions, predominantly periventricular leukomalacia. Detrimental neurotoxic effects are not only induced by direct host-microbe interaction, but may also be generated by exposure to perinatal inflammation; activation of fetal/neonatal immune cells triggered by bacterial products that activate pattern recognition receptors and/or maternal pro-inflammatory mediators that cross the placenta.

For example, exposure to intrauterine inflammation results in a fetal inflammatory response, characterised by activation of CD45RO+ T-cells and elevated levels of pro-inflammatory cytokines, which are associated with cerebral injury on MRI. Therefore, an active fetal response rather than exclusively passive transfer of maternal mediators may be a key pathogenic mechanism for cerebral damage in preterm infants. Furthermore, high plasma levels of pro-inflammatory cytokines in preterm infants with sepsis or necrotising enterocolitis are associated with increased risk of ultrasound-detected white matter injury. However, although both blood and cerebrospinal fluid cytokine levels are associated with white matter injury, plasma cytokine levels may not reflect local cytokine production in the brain and imbalance of pro- and anti-inflammatory mediators may be at least as important as absolute levels of individual cytokines.

In the newborn, cytokines can be released from activated immune cells, mainly monocytes, macrophages and T-cells. Activation can occur systemically in the blood compartment or at sites of infection; in the brain, this is predominantly by resident microglia or activated macrophages infiltrating the brain, and both mechanisms may be active simultaneously. In addition, there is increasing appreciation of the immunological implications of brain immaturity in the preterm infant, particularly with respect to the immaturity of central nervous system immune cell regulation, which may render the preterm brain exquisitely vulnerable to damage by poorly controlled and pervasive inflammation. These data also support the observation that direct bacterial infections of the neonatal brain, such as meningitis, only cause a small proportion of cerebral injuries and that significant neonatal brain injury can occur without entry of bacteria into the cerebrospinal fluid.

In the following section we review recent data from in vitro and animal models that have significantly advanced the understanding of mechanisms underlying the relationship between bacteraemia, inflammation and cerebral injury in preterm infants.

**Experimental models**

The vulnerability of the newborn brain to infection/inflammation was first described by Gilles in 1976 when systemic administration of bacterial lipopolysaccharide (LPS), that activates cells via Toll-like receptor 4 (TLR4), caused leukencephalopathy in neonatal kittens, but not in mature cats. Furthermore, the newborn feline brain is particularly
vulnerable to the effects of LPS, whereas other organs were comparatively resistant. In fetal sheep, systemic intrauterine LPS exposure leads to activation of microglia, the resident macrophage-like cells of the central nervous system and to loss of neurofilament and myelin basic protein, changes that are associated with white matter injury, specifically periventricular leukomalacia.\textsuperscript{66-68} Furthermore, these changes are associated with acute increase in blood levels of S100B protein, a marker of cerebral injury.\textsuperscript{69} In mice, microglia and neurons express TLR1-9 and TLR2, -3, -4 and -8, respectively, and stimulation of quiescent microglia with TLR agonists initiates rapid upregulation of cytokines and chemokines, indicating their importance in mediating responses to infectious organisms.\textsuperscript{70,71} In addition, microglia are the principal central nervous system cell population responsive to peripherally administered TLR agonists. Microglia-mediated neuronal injury was critically dependent on intact expression of TLR2 and 4, respectively, highlighting the biological relevance of the TLR pathways in the central nervous system.\textsuperscript{72-75} There are no studies specifically investigating leukocyte entry into the immature brain after peripheral administration of TLR agonists. However data from adult animals suggest active transport of some cytokines across the blood brain barrier or production by endothelial cells with subsequent release into adjacent cerebral tissue(Figure 1).\textsuperscript{76,77}

Arterial hypotension is common in both neonatal sepsis and experimental models, especially in Gram-negative infection and necrotising enterocolitis, and via cerebral ischemia and reperfusion can potentiate the risk of white matter injury.\textsuperscript{78} However, cerebral damage can be induced by systemic inflammation in the absence of systemic cardiovascular impairment and in the human preterm infant the relationship between infection, hypotension and white matter damage is often inadequately documented and thus incompletely understood.\textsuperscript{79}

Most neonatal bacterial infections are caused by Gram-positive organisms that do not express LPS and predominantly signal via TLR2 and other pattern recognition receptors. \textit{In vitro} experiments and \textit{in vivo} studies show that administration of TLR2 agonists (Pam3CSK4 and FSL1) as well as inactivated whole bacteria result in inhibition of neural progenitor cell proliferation and consequently in perinatal brain injury (Table 2).\textsuperscript{80-82} Cell wall preparations and secreted factors of one of the principal early-onset sepsis pathogens and most common cause of neonatal meningitis, Group B streptococcus (GBS), induce neuronal cell death \textit{in vitro}, which is dependent on the presence of microglia expressing TLR2 and the TLR-adapter protein MyD88.\textsuperscript{83} In addition, GBS-mediated activation induces microglial apoptosis via caspase-8, a potentially autoregulatory mechanism limiting ongoing innate immune activation and inflammatory damage in the brain.\textsuperscript{84}

Direct bacterial cytotoxicity and activation of the local host response with production of various inflammatory mediators, such as cytokines, prostaglandins and reactive oxygen species are the main detrimental mechanisms of bacterial infection of the newborn central nervous system.\textsuperscript{51,52,85} In contrast, it is largely unknown how cerebral injury is mediated in the absence of meningitis. Systemic inflammation can exert rapid negative impact on cerebral function that precedes peripheral organ dysfunction and can occur without bacterial invasion of the central nervous system.\textsuperscript{46,86} This is presumably augmented in part by blood-brain barrier dysfunction during infection, as exposure to bacterial cell wall components
directly increases blood-brain barrier permeability. Additional potential mechanisms, such as direct transfer of bacterial components or inflammatory mediators across endothelial cells, remain incompletely characterised (Figure 1). Neuronal apoptosis occurs within hours of pneumococcal bacteremia or systemic challenge with pneumococcal cell wall preparations in mice, and importantly, this effect does not require binding of bacterial components to the endothelium and is independent of bacteria or leukocytes entering the cerebrospinal fluid. In newborn marsupial opossums, in whom the majority of brain maturation occurs postnatally (akin to very preterm infants) LPS administration, especially repeated exposure, results in significant and sustained increases in blood brain barrier permeability, microglial activation and white matter injury. In several animal models, intraperitoneal LPS injection induces activation of microglia, intracerebral expression of TLR2, IL-1 and IL-6 and decreased hippocampal neurogenesis and cerebral damage can occur within hours of pneumococcal bacteraemia without meningitis. The pathogenic mechanisms appear to be at least partially independent of direct cytokine-mediated inflammation as anti-TNF antibodies do not reduce neuronal injury. However, overexpression of IL-10 is protective (Figure 1). Preterm infants not only show gestational age-dependent impairment of inflammatory responses, but importantly, have profoundly reduced capacity to produce anti-inflammatory cytokines, such as IL-10. The quality and quantity of anti-inflammatory responses of central nervous system cells to systemic inflammation are not defined, particularly in human newborn infants.

Exposure to bacterial pattern recognition receptor agonists increases vulnerability of the preterm brain

Bacterial infection increases the vulnerability of the preterm brain to non-inflammatory insults: in human infants the combination of maternal infection and asphyxia amplifies the risk of cerebral palsy. Administration of LPS to rat fetuses or newborn rat pups induces cerebral expression of CD14 and TLR4 and sensitises the immature brain to subsequent hypoxic-ischaemic injury. In mice pre-treatment with LPS converts a sub-threshold hypoxic insult to a critical one. Furthermore, the potentiation of cerebral injury induced by LPS exposure is dependent on intact expression of MyD88. MyD88-deficient animals displayed significantly reduced activation of nuclear factor kappa-B, inflammatory cytokines and chemokines and reduced white and gray matter injury. Deletion of the TNF gene cluster in mice results in lack of LPS-induced activation of microglia and endothelial cells and abolishes the sensitisation to hypoxic-ischaemic insult following LPS-exposure. Cerebral sensitisation after LPS exposure of fetal mice is pervasive even if the postnatal hypoxic-ischemic insult occurs as late as day 70. Of note, in rats, these sequential insults resulted in anatomical grey and white matter damage as well as behavioural and motor deficits similar to those observed in human infants.

The association between exposure to bacterial components, hypoxic-ischaemic insult and cerebral injury is not uniform, but appears to be sensitive to timing and dosage of the insults, context and, importantly, the maturity of the animal. LPS pre-treatment in adult mice actually reduced loss of cerebral tissue upon hypoxic-ischaemic insult, indicating that in mature animals LPS may have a protective preconditioning effect. In neonatal rats, the sensitisation afforded by LPS is time-dependent: LPS given either 6 hours or 72 hours prior

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to hypoxic-ischaemic insult results in increased cerebral injury, whereas injury size is markedly reduced with a 24 hour interval.\textsuperscript{108} In addition, increasing brain maturity (postnatal age) and the concurrent rise in cerebral expression of TLR4 is critical for the protective preconditioning effects of LPS.\textsuperscript{109}

### Detection of cerebral injury in preterm infants with bacteraemia

The relationship between bacteraemia and cerebral injury is complicated by the multiple pathways that may contribute to an increased risk of brain injury in preterm infants. In addition, clinical outcomes associated with bacteraemia, such as requirement for indwelling plastic devices (e.g., respiratory and feeding tubes, intravascular catheters) and total parenteral nutrition may themselves be related to alteration of cerebral development.

Intraventricular haemorrhage is the most common form of cerebral injury in very preterm infants currently detected with conventional neuroimaging, such as cranial ultrasound. Preterm infants with early-onset bacteraemia have a significantly elevated risk (up to 20\%) for high grade intraventricular haemorrhage (Figure 2A).\textsuperscript{110} Cerebellar haemorrhage may occur in up to 20\% of preterm infants, but the incidence is often underestimated by cranial ultrasound (Figure 2B). In preterm infants, the association between neonatal sepsis and increased risk for cerebellar haemorrhage further supports the concept of bacteraemia and exacerbation of neonatal brain injury.\textsuperscript{111}

The incidence of periventricular leukomalacia and white matter injury varies with the method of detection. Preliminary studies suggest that MRI is more sensitive for white matter injury than either computed tomography or ultrasound.\textsuperscript{112,113} Human MRI studies document a strong association between infection and white matter lesions in the preterm brain.\textsuperscript{44,45} Abnormalities have also been reported on white matter microstructural integrity, measured by white matter diffusion, supporting an adverse impact of infection or inflammation on the cerebral white matter.\textsuperscript{48} White matter injury is particularly severe in the setting of necrotising enterocolitis, presumably relating to the distinct and long-lasting systemic inflammation inherent to this disease, and may be further aggravated by associated systemic hypotension (Figure 3).\textsuperscript{114}

Finally, MRI can define alterations in cerebral development including reductions in cerebral growth both globally and regionally, as well as alterations in cerebral biochemistry (spectroscopy). Bacteremia is associated with alterations in brain biochemistry and reductions in cerebral growth, consistent with the known adverse effect of neonatal infection on somatic growth.\textsuperscript{32,48,115} An understanding of alterations in the sequence of normal cerebral development, combined with or independent of cerebral injury, will provide a better appreciation of the impact of bacteraemia and associated complications on the central nervous system.

In summary, neuroimaging has given a clear delineation of the breadth of impact of neonatal bacteremia from exacerbation of brain injury to impairments in brain development.
The incidence of preterm birth has risen over the past thirty years and interventions aimed at reducing this trend have largely been disappointing. Furthermore, a significant proportion of infants born preterm will have been exposed to inflammation from well before birth, and attempts at ameliorating the postnatal consequences will be challenging. Therefore, new strategies for the prevention and treatment of perinatal inflammation and neonatal sepsis are urgently needed. Interventions that reduce infection and inflammation-induced cerebral injury are of particular relevance in the context of extreme prematurity, as the detrimental effects of neonatal infection on white matter injury and long-term neurological outcomes appear intimately related to the release and circulation of pro-inflammatory bacteria-derived molecules that induce systemic inflammation. Consequently, antibiotic therapy alone does not ameliorate the risk of white matter injury, neurodevelopmental impairment and cerebral palsy associated with neonatal sepsis. Based on emerging evidence discussed above, novel protective interventions might include those targeting free-radical generation or accumulation, anti-apoptotic agents, and anti-inflammatory agents and compounds targeted at blunting the host inflammatory response to microbial products, e.g. bacterial lipopeptides and lipoteichoic acids (TLR2), LPS (TLR4), peptidogycans (nucleotide oligomerization domain 1, NOD1) and others.

Corticosteroids are potent anti-inflammatory agents and dexamethasone in particular has long been used in preterm infants, primarily for the treatment and prevention of chronic lung disease. Both in experimental models of perinatal inflammation and in chorioamnionitis-exposed human infants, corticosteroids can modulate inflammation and ameliorate related lung disease. However, dexamethasone has highly significant adverse long-term neurodevelopmental consequences, including an increased risk of cerebral palsy, precluding its universal use as an anti-inflammatory agent. Hydrocortisone may be equally effective for chronic lung disease without the detrimental neurodevelopmental outcomes associated with dexamethasone, but also has acute adverse effects, such as gastrointestinal perforation. Importantly, in addition to the significant side effects, postnatal steroids have not been evaluated systematically as immunomodulators in preterm infant infection, and thus their effects on survival and long-term outcomes in this context remain unknown.

The history of neonatology includes serious, unexpected long-term side effects of well-intended interventions with apparent short-term benefits. Any new intervention will need to be evaluated carefully with an emphasis on safety prior to being advocated for routine use, particularly at this critical period of immunological development.

N-acetylcysteine

The free radical scavenging/anti-oxidant agent N-acetylcysteine readily crosses the placenta and is considered safe during pregnancy and in the preterm neonate. N-acetylcysteine may prevent LPS-induced degeneration of oligodendrocyte progenitors and hypomyelination in the developing rat brain, an effect associated with attenuation of the intracerebral inflammatory reaction, including reduced levels of TNF, IL-1, and expression...
of inducible nitric oxide synthase.\textsuperscript{124} In mice, N-acetylcysteine administration to the pregnant dam attenuates the maternal and fetal pro-inflammatory response to intrauterine LPS administration, resulting in fewer preterm births and reduced neonatal white matter injury.\textsuperscript{125,126} In addition, N-acetylcysteine provides substantial neuroprotection against brain injury caused by the combination of LPS exposure and hypoxic-ischaemic insult in neonatal rats, suggesting that N-acetylcysteine has potential therapeutic value, especially considering the protective effects of antenatal administration to the pregnant animal with chorioamnionitis – a scenario where postnatal intervention may be of limited benefit.\textsuperscript{127} However, there are some concerns that N-acetylcysteine may compromise fetal cardiovascular stability and to date, no large clinical trials have evaluated systemic N-acetylcysteine administration in newborn infants - or their mothers - prior to delivery.\textsuperscript{128}

**Erythropoietin**

Erythropoietin, previously appreciated exclusively as a kidney-derived haemopoietic growth factor, is neuroprotective in a range of experimental models of cerebral injury. The protective mechanisms are not completely understood, but may include anti-oxidative, anti-apoptotic as well as significant anti-inflammatory action, both systemically and in the brain.\textsuperscript{129,130} Importantly, erythropoietin crosses the blood-brain barrier and interacts with cerebral erythropoietin receptors that are expressed from early in gestation.\textsuperscript{131,132} Clinical trials demonstrate that repeated administration of recombinant human erythropoietin is well tolerated and results in plasma levels that are neuroprotective in animal studies. Furthermore, pilot data suggest that erythropoietin improves neurodevelopmental outcomes in infants with hypoxic-ischaemic encephalopathy and is associated with superior developmental outcomes at 10-13 years of age in extremely preterm infants with intraventricular hemorrhage treated prophylactically with erythropoietin for anaemia of prematurity.\textsuperscript{133-135} However, recent meta-analyses confirm that early routine administration of erythropoietin increases the risk of retinopathy of prematurity in extremely preterm infants. This is of concern particularly in the context of late-onset sepsis, which also is an independent risk factor for the development of retinopathy.\textsuperscript{136,137} Large-scale clinical trials to evaluate erythropoietin for neuroprotection in preterm infants and in term infants with hypoxic-ischaemic encephalopathy are ongoing.

**Pentoxifylline**

Pentoxifylline is a synthetic xanthine-derived phosphodiesterase inhibitor that raises cellular concentrations of cyclic adenosine monophosphate thereby inhibiting production of inflammatory mediators such as TNF. Pentoxifylline has beneficial effects in models of neonatal inflammatory conditions, including sepsis and necrotising enterocolitis.\textsuperscript{138,139} Pilot studies of pentoxifylline as adjunct therapy for neonatal sepsis show that it is safe and well-tolerated with a favourable side-effect profile, and meta-analysis of the available data (a total of 227 patients from four studies) conclude that pentoxifylline reduced sepsis-related mortality.\textsuperscript{140,141} However, although pentoxifylline has also been implicated as a potential adjunct agent for treatment of hypoxic-ischaemic encephalopathy based on animal studies, there are no neonatal clinical data to date. Given its anti-inflammatory properties and promising clinical profile, pentoxifylline warrants evaluation in larger clinical trials as
adjunct therapy in neonatal sepsis, including assessment of long-term neurodevelopmental outcomes.

**Minocycline**

The tetracycline antibiotic minocycline has shown promising anti-apoptotic, anti-oxidant and anti-inflammatory effects in animal models, especially inhibiting microglial activation after cerebral insults, as well as protective effects on blood-brain barrier integrity in systemic inflammation. However, the use of tetracyclines in neonates and infants is contentious, particularly because of concerns of disruption of normal formation of bone and tooth enamel. Furthermore, while there is experimental evidence in animal models for neuroprotective effects of minocycline, even when administered after the cerebral insult, there is a lack of supportive data from human clinical trials to date.

**Immununological interventions**

Our understanding of neonatal and infant immune function and its maturation in early childhood remains incomplete. Over the past decade, a number of immune interventions aimed at preventing or improving the outcome of neonatal sepsis, such as colony-stimulating growth factors and intravenous immunoglobulin, have been unsuccessful. The mechanisms underpinning the unique susceptibility of preterm infants to invasive infections are the subject of ongoing research efforts.

Antimicrobial proteins and peptides (APPs), a group of naturally occurring molecules expressed in leukocytes and on mucosal epithelial cells possess anti-infective and immunomodulatory properties and have shown promise in animal and/or early human clinical studies. These cationic molecules kill microbes, but unlike conventional antibiotics are also able to bind microbial components and reduce their inflammatory activity by preventing their interaction with bacterial products. For example, administration of an endotoxin-neutralising recombinant 21 kDa fragment of bactericidal/permeability-increasing protein (rBPI21) in addition to a fluoroquinolone antibiotic improves survival in mice exposed to lethal radiation and reduces systemic inflammation and morbidity when given with conventional antibiotics to children with meningococcal sepsis. In human preterm infants, oral supplementation with lactoferrin, an iron-binding glycoprotein with anti-infective activities, reduces the incidence of bacterial and fungal late-onset sepsis and large international clinical trials further evaluating this approach are ongoing. Systemic administration of the synthetic immune defense regulator peptide 1018, a derivative of the human cathelicidin LL-37, resulted in marked mitigation of LPS and hypoxia-ischemia-induced cerebral injury in mice.

Overall, accumulating evidence indicates that bacterial infection triggers inflammatory pathways that damage the preterm brain even in the absence of direct bacterial entry to the central nervous system. Given the high global rate of preterm birth, the frequency of preterm brain injury, and its long term morbidities, translational research directed at defining the underlying mechanisms and adjunctive therapies is urgently needed to provide novel approaches to mitigate severe long-term neurodevelopmental consequences for this highly susceptible population.
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Search strategy and selection criteria

References for this review were identified through searches of PubMed for articles published from January, 1973, to February 2014, by use of terms “sepsis”, “bacteremia”, “necrotising enterocolitis”, “chorioamnionitis”, “newborn”, “preterm infant”, “brain injury”, “long-term outcome”, “neurodevelopment” and “intervention”. Articles resulting from these searches and relevant references cited in those articles were reviewed.
Figure 1.
This figure summarizes known and hypothetical pathways of bacteraemia-induced neuronal damage: (1) bacterial products on or shed from bacteria in the bloodstream activate endothelial pattern recognition receptors such as TLRs triggering release of inflammatory mediators into the CNS; (2) Leak of bacterial products such as lipopolysaccharide (LPS) and bacterial lipopeptide (BLP) across the blood–brain barrier that activate microglia to release inflammatory mediators; (3) Entry of leukocytes into the CNS; and (4) direct diffusion of cytokines/chemokines from the peripheral circulation across the blood–brain barrier. Hypothetical mechanisms for which there has not yet been published evidence are marked with a question mark (“?”).
Figure 2.
Intracranial haemorrhage in a preterm with bacteraemia. A preterm infant born at 24 weeks gestation following onset of maternal fever grew *E. coli* from blood cultures obtained at 45 minutes of age. He had septic shock with coagulopathy, thrombocytopenia and hypotension prompting inotropic support. His day 2 cranial ultrasound revealed left grade III and right grade IV intraventricular haemorrhage (A) with severe cerebellar haemorrhage (B). Note that areas of echodensity (brightness) indicate haemorrhage.
Figure 3.
White matter injury in a preterm infant with necrotizing enterocolitis. A preterm infant was born at 24 weeks gestation with germinal matrix hemorrhage developed severe necrotising enterocolitis at 6 weeks of age requiring surgery. (A) ultrasound 4 weeks post- surgery revealed small cystic white matter echolucencies (arrow). (B) An MRI scan 2 weeks later at 36 weeks post-menstrual age demonstrated extensive white matter injury with periventricular gliosis and extensive encephaloclastic changes.
Table 1

Key animal studies linking Bacteria-induced Inflammation with Cerebral Injury.

<table>
<thead>
<tr>
<th>Author/year</th>
<th>Animal species</th>
<th>Age at intervention</th>
<th>Intervention</th>
<th>Clinical/anatomical outcomes</th>
<th>Biomarkers</th>
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<td>Mallard et al., 2003</td>
<td>Fetal sheep</td>
<td>Days 93-96 of gestation</td>
<td>Intravenous <em>E. coli</em> LPS</td>
<td>Focal inflammatory infiltrates and cystic lesions in periventricular white matter, microglial activation, astrocyte damage, loss of oligodendrocytes.</td>
<td>None</td>
<td>150</td>
</tr>
<tr>
<td>Svedin et al., 2005</td>
<td>Fetal sheep</td>
<td>Median days 89 or 121 of gestation</td>
<td>Intravenous <em>E. coli</em> LPS</td>
<td>Microglial activation and loss of neurofilament indicative of white matter injury.</td>
<td>None</td>
<td>67</td>
</tr>
<tr>
<td>Garnier et al., 2006</td>
<td>Fetal sheep</td>
<td>Median day 107 of gestation</td>
<td>Intravenous <em>E. coli</em> LPS</td>
<td>Inflammatory infiltrates and cystic lesions in periventricular white matter.</td>
<td>Increase S100B blood levels</td>
<td>69</td>
</tr>
<tr>
<td>Stolp et al., 2007</td>
<td>Opossums</td>
<td>Postnatal day 35</td>
<td>Intraperitoneal <em>E. coli</em> LPS</td>
<td>Single LPS injection induced short-lasting blood brain barrier dysfunction. Repeated LPS exposure resulted in more profound and prolonged blood brain barrier impairment.</td>
<td>None</td>
<td>90</td>
</tr>
<tr>
<td>Orihuela et al., 2006</td>
<td>Mice</td>
<td>4-5 weeks old</td>
<td>Intranasal or intravenous live or heat-killed whole <em>S. pneumoniae</em> or purified cell wall preparation</td>
<td>Both whole bacteria and cell wall preparations induce hippocampal neuronal injury. This was partially mitigated in TLR2−/−, NOD2−/− and IL10-overexpressing mice.</td>
<td>None</td>
<td>69</td>
</tr>
<tr>
<td>Gavilanes et al., 2009</td>
<td>Fetal sheep</td>
<td>Day 111 or 123 of gestation</td>
<td>Intraperitoneal <em>E. coli</em> LPS with 2d or 14d survival</td>
<td>Fetuses with long-term survival displayed apoptosis, microglial activation and astrogliosis. Loss of mature oligodendrocytes and neurons were decreased in some regions of the brain.</td>
<td>None</td>
<td>68</td>
</tr>
<tr>
<td>Du et al., 2011</td>
<td>Newborn mice</td>
<td>Postnatal days 3-11</td>
<td>Once daily intraperitoneal TLR2 agonist (Pam3CSK4)</td>
<td>Decreased volume of gray and white matter and activation of microglia.</td>
<td>Elevated levels of IL1, IL6, CCL2 in brain homogenates after 1st injection</td>
<td>81</td>
</tr>
<tr>
<td>Dean et al., 2011</td>
<td>Fetal sheep</td>
<td>Day 102 of gestation</td>
<td>Single intravenous <em>E. coli</em> LPS injection</td>
<td>Reduced grey and white matter volume, including loss of oligodendrocytes and cortical neurons. Loss of normal maturation of EEG.</td>
<td>None</td>
<td>66</td>
</tr>
<tr>
<td>Keogh et al., 2012</td>
<td>Fetal sheep</td>
<td>Days 103-108 of gestation</td>
<td>Intravenous <em>E. coli</em> LPS infusion</td>
<td>Loss of EEG maturation, increased cerebral inflammation and caspase 3 positive cells in white matter, but no loss of oligodendrocytes and cortical neurons</td>
<td>Transient rise in plasma cortisol and IL-6</td>
<td>79</td>
</tr>
</tbody>
</table>
### Table 2

Clinical Studies Linking Bacterial Infection and Cerebral Injury in Preterm Infants.

<table>
<thead>
<tr>
<th>Author</th>
<th>Number of infants</th>
<th>Study population</th>
<th>Definition of infection</th>
<th>Central nervous system involvement</th>
<th>Key outcomes</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stoll et al., 2004</td>
<td>N=6093</td>
<td>Birth weight 401-1500g</td>
<td>Positive blood culture plus ≥5 days of antibiotics. Clinical infection: culture negative, but ≥5 days of antibiotics Positive (cerebrospinal fluid, CSF) culture plus ≥5 days of antibiotics</td>
<td>192 cases of meningitis</td>
<td>Any postnatal infection associated with poor growth and neurodevelopmental impairment. Strongest effect for sepsis/NEC</td>
<td>32</td>
</tr>
<tr>
<td>Glass et al., 2008</td>
<td>N=133</td>
<td>&lt;34 weeks</td>
<td>Intention-to-treat plus positive culture from blood, endotracheal tube, urine, skin lesion, or CSF (plus suggestive CSF findings)</td>
<td>Cases of meningitis excluded from analysis Due to small numbers, meningitis was not significantly associated with adverse outcomes</td>
<td>Recurrent postnatal infections are associated with increased risk of progressive white matter injury</td>
<td>45</td>
</tr>
<tr>
<td>Shah et al., 2008</td>
<td>N=192</td>
<td>&lt;30 weeks</td>
<td>Positive blood culture plus abnormal I/T ratio, CRP or platelet count plus ≥5 days of antibiotics Meningitis: CSF contains &gt;20 cells/mm$^3$ plus therapeutic course of antibiotics</td>
<td>3 cases of culture-negative meningitis; 2 in proven sepsis group, 1 in clinical sepsis group</td>
<td>Postnatal infection and/or NEC associated with white matter injury increased risk of motor impairment at 2 years of age</td>
<td>44</td>
</tr>
<tr>
<td>Chau et al., 2009</td>
<td>N=96</td>
<td>24-32 weeks</td>
<td>Any positive culture from blood, CSF, urine; in tracheal aspirate if associated with &gt;4 leukocytes and clinical pneumonia</td>
<td>No case of meningitis mentioned</td>
<td>Postnatal infection, but not histological chorioamnionitis is associated with white matter injury</td>
<td>78</td>
</tr>
<tr>
<td>Bassler et al., 2009</td>
<td>N=944</td>
<td>Birth weight 500-999g</td>
<td>Sepsis: positive blood culture; Meningitis: positive CSF culture, no further information given</td>
<td>22 cases of meningitis. Meningitis with or without sepsis showed strongest association with adverse outcome</td>
<td>Any postnatal infection or NEC increased the risk of late death or survival with neurosensory impairment</td>
<td>33</td>
</tr>
<tr>
<td>Martin et al., 2010</td>
<td>N=1155</td>
<td>23-27 weeks</td>
<td>Positive blood culture (taken weekly as routine part of ELGAN study)</td>
<td>Meningitis not mentioned</td>
<td>Increased risk of impaired neurodevelopment at 24 months of age in infants with surgical NEC, esp. when accompanied by sepsis</td>
<td>41</td>
</tr>
<tr>
<td>Helderman et al., 2010</td>
<td>N=108</td>
<td>24-26.6 weeks</td>
<td>Positive blood culture plus clinical signs plus ≥5 days of antibiotics. Culture-negative infection: negative culture plus clinical signs plus ≥5 days of antibiotics</td>
<td>No cases of meningitis</td>
<td>Culture-positive and clinical sepsis are associated with acute encephalopathy, but normal rate of brain maturation</td>
<td>46</td>
</tr>
<tr>
<td>Silveira et al., 2011</td>
<td>N=88</td>
<td>Birth weight 500-1500g</td>
<td>Clinical signs plus positive culture from blood or CSF</td>
<td>No case of meningitis</td>
<td>Postnatal sepsis associated with increased risk of periventricular</td>
<td>40</td>
</tr>
<tr>
<td>Author</td>
<td>Number of infants</td>
<td>Study population</td>
<td>Definition of infection</td>
<td>Central nervous system involvement</td>
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<tr>
<td>Chau et al., 2012</td>
<td>N=117</td>
<td>24-32 weeks</td>
<td>Any positive culture from blood, urine or CSF; in tracheal aspirate if associated with &gt;4 leukocytes and clinical pneumonia</td>
<td>4 cases of meningitis. Results unchanged when meningitis excluded from analysis</td>
<td>Postnatal infections (proven and clinical) are associated with abnormalities in metabolic and structural brain development</td>
<td>48</td>
</tr>
</tbody>
</table>

leukomalacia