Hypoxic Ischemic Encephalopathy (HIE) is the result of severe anoxic brain injury during the neonatal period and causes life-long morbidity and premature mortality. Currently, therapeutic hypothermia immediately after birth is the standard of care for clinically relevant HIE. However, therapeutic hypothermia alone does not provide complete neuroprotection and there is an urgent need for adjunctive therapies. Ischemic conditioning is an adaptive process of endogenous protection in which small doses of sub-lethal ischemia can provide a protection against a lethal ischemic event. Remote Ischemic Post-conditioning (RIPC), a form of ischemic conditioning, is highly translatable for HIE diagnosed immediately after birth as the conditioned ischemic stimulus is applied at the limb after the lethal ischemic episode. A number of studies in neonatal rats have demonstrated that RIPC is effective at reducing injury in focal cerebral ischemia models and improves neurological outcomes. In this review, we focus on the available data on HIE and its current treatment, models in HIE studies, ischemic conditioning/RIPC and its mechanism. We discuss in particular the effect of RIPC on neonatal brain with HIE. We postulate that combining RIPC with standard therapeutic hypothermia can be an attractive therapeutic approach for HIE.

Keywords: hypoxic ischemic encephalopathy, remote ischemic post-conditioning, hypothermia
Therapeutic hypothermia or total body cooling in treating hypoxic ischemic encephalopathy (HIE) is effective in many randomized controlled trials (Gluckman, et al., 2005; Shankaran, et al., 2005; Azzopardi, et al., 2009; Simbruner, Mittal, Rohlmann, & Muche, 2010; Zhou, et al., 2010; Jacobs, et al., 2011). It is the standard of care for infants with HIE (Higgins, et al., 2011). However, as it does not provide complete protection, the search for adjunctive therapies continues. Remote limb ischemic post-conditioning (RIPC) is the simple, inexpensive, and safe use of repetitive inflation-deflation procedure of a blood pressure (BP) cuff on the arm or leg to protect distant organs such as the brain, heart, and kidney from ischemic injury. Many preclinical studies demonstrated that RIPC is effective at reducing cerebral, myocardial or retinal injuries resulting from ischemia. In this review, we conducted a literature search to evaluate the rationale for the use of RIPC as an adjuvant treatment for hypoxic ischemic encephalopathy to standard therapeutic hypothermia.

Hypoxic Ischemic Encephalopathy
Hypoxic-ischemia (HI) is a term that is used to describe the constellation of complex physiological, cellular, and molecular changes that are induced by lack of oxygen supply to the brain (Liu, Li, & Gu, 2007; Busl & Greer., 2010). Hypoxic ischemic encephalopathy (HIE) is the result of severe anoxic brain injury during the neonatal period and accompanies life-long morbidity, including cerebral palsy, and premature mortality (Volpe, 2001). The incidence of HIE is between 1 and 8 per 1000 live births in developed countries with rates as high as 26 per 1000 live births in the developing world (Kurinczuk, White-Koning, & Badawi, 2010). Its pathophysiology involves oxidative stress, mitochondrial energy production failure, glutaminergic excitotoxicity and cell death (Marcelino, et al., 2015). The clinical symptoms of HIE include acute symptoms such as seizures, alteration of consciousness, weak breathing, poor muscle tons or metabolic derangement as well as chronic conditions such as cerebral palsy, mental retardation, learning disabilities, and epilepsy (Vannucci & Perlman, 1997; Vannucci R., 1997; Shah & Perlman, 2009).

Current treatment of HIE
As the consequences of HIE are extensive and cause significant challenges to the affected individual, their family, and society, many potential treatments have been studied (Vannucci & Perlman, 1997; Vannucci R., 1997; Johnston, Fatemi, Wilson, & Northington, 2011). Therapeutic hypothermia immediately after birth is the standard of care for clinically relevant HIE and appears to have a temporal relationship with outcome (Edwards, et al., 2010). Many studies have shown that therapeutic hypothermia inhibits key steps in the excito-oxidative cascade including energy failure and increases lactic acid, glutamate, and nitric oxide concentrations in the brain (Thoresen, et al., 1995; Thoresen, et al., 1997; Amess, et al., 1997). Based on the information from many studies and workshops presented by the Eunice Kennedy Shriver National Institute of Child Health and Human Development (NICHD), the current standard of care for HIE is the whole body cooling to a core temperature of 33.5°C for 72 hours, starting within 6 hours of birth (Higgins, et al., 2011). Initiation of therapeutic hypothermia within 6 hours after birth in neonates with moderate to severe HIE leads to significant improvement in neurocognitive outcomes and reduced mortality (Shankaran, et al., 2005; Azzopardi, et al., 2009; Gluckman, et al., 2005; Jacobs, et al., 2011; Tagin, Woolcott, Vincer, Whyte, & Stinson, 2012). Surprisingly, longer duration or lower core body temperature does not confer additional benefit and may pose some risk especially in those infants with low birth weight (Higgins, et al., 2011; Thoresen, Bägenholm, Löberg, Apricena, & Kjellmer, 1996; Shankaran, et al., 2012; Shankaran, et al., 2014). These findings are similar to animal models of HIE, wherein the optimal temperature for the P7 rat is likely between 32 °C and 33.5 °C, with no significant additional neuroprotection for rat pups cooled below 33.5 °C (Bona, Hagberg, Lobeg, Bagenholm, & Thoresen, 1998; Hobbs, et al., 2008; Wood, et al., 2016). Similar to human studies, intermediate to long duration (5 hours) of therapeutic hypothermia is more effective than 3 hours in rat pups with HI (Hooke, Chakkarakani, Liu, Hobbs, & Thoresen, 2009). However, there was no difference in neuroprotection in rat pups receiving therapeutic hypothermia from 5 hours to 10 hours (Sabir, Scull-Brown, Liu, & Thoresen, 2012).

Therapeutic hypothermia alone does not provide complete neuroprotection. The high prevalence of neurologic morbidity and mortality, even after therapeutic hypothermia, persists with death or severe neurological complications resulting in 40-50% of neonates who underwent therapeutic hypothermia (Edwards, et al., 2010). Due to the high morbidity and mortality, the development of new interventions for patients with HIE is imperative (Tagin, Woolcott, Vincer, Whyte, & Stinson, 2012; Shankaran, et al., 2014; Perlman, et al., 2010). The most likely next advance in care for these infants is in the form of an adjuvant therapy to therapeutic hypothermia (Peliowski-Davidovich, 2012).

Models of hypoxic ischemic encephalopathy
Many models of intrauterine, perinatal, and neonatal hypoxic ischemia have been reported (Northington, 2006). Maternal hypoxia or umbilical cord occlusion in fetal sheep produces a robust hypoxia-ischemia phenotype, but the high cost of lambs and large body size make this model impractical for widespread use (Gleason, Hamm, & Jones, 1990; Harris, Helou, Gleason, Traystman, & Koehler, 2001; Gonzalez, Hunter, Bennet, Power, & AJ., 2005; Lotgering, et al., 2004). The development of rodent models has provided mechanistic insight into the pathophysiology and clinical implications of neonatal HIE. The HIE model in neonatal brain was first validated by Vannucci and colleagues. Based on the Levine preparation in adult, it consists of unilateral common carotid artery ligation followed by systemic hypoxia produced by the inhalation of 8% oxygen –balance nitrogen. This model has been the most used for HIE (Rice, 1981;Vannucci & Vannucci, 1997). Importantly, studies showed that the brain of postnatal day 7 rat is phenotypically similar to an early third trimester human fetus (Clancy, 2001). Neuroprotection in less mature rats requires a more extensive hypoxic insult to reproduce the same histological findings observed at postnatal day 7 (Sheldon, Chua, & Ferriero, 1996). Vannucci’s model demonstrates a reproducible brain damage model in a spontaneously breathing rat with low mortality. No seizures or cardiopulmonary complications were noted for at least the first 50 hours of survival (Rice, 1981). Hypothermia is also efficacious in animal models of HIE including the Vannucci model (Young, Olenginski, Yagel, & Towfighi, 1983; Bona, Hagberg, Lobeg, Bagenholm, & Thoresen, 1998; Mishima, et al., 2004). Brief periods of moderate hypothermia to 32°C after injury delay cell death in 7-day-old rat pups by as much as a week, after which brain atrophy occurs (Trescher, Ishiwa, & Johnston, 1997).

Ischemic Conditioning
Ischemic conditioning is an adaptive process of endogenous protection in which small doses of sub-lethal ischemia protect the organism against a lethal ischemic event. Ischemic conditioning can be performed at a distance or remote from the ischemic target organ (Murry, Jennings, & Reimer, 1986; Gho, Schoemaker, van den Doel, Duncker, & Verduw, 1996; Schmidt, et al., 2007). Similarly, tumor growth and progression has been noted to utilize these processes. Intermittent hypoxia
has been reported within the tumors and is thought to increase tumors’ resistance to major ischemia and anti-cancer therapy (Muscari, et al., 2013). Depending on the temporal relation of the sub-lethal ischemia to the lethal ischemic event, the conditioned stimulus can be applied prior to ischemia and vessel occlusion (pre-conditioning), during ischemia and before reperfusion (per-conditioning), or after the lethal ischemic episode and through reperfusion (post-conditioning). A study found that combining limb remote per- and post- conditioning significantly reduced cerebral ischemia/reperfusion injury (Ren, et al., 2015).

There have been several studies that focus on the neurochemical basis for the neuroprotective role of these ischemic conditioning regimens. Similar pathways and molecules play a role in these conditionings but the roles and timing can be different (Alkan, 2009). Several studies focused on the differences of the conditioning on stimulated the model of pre-conditioning. There are two windows of protection: an early and late window. The mediators for the early window include phosphorylation, transporter regulation, interfering RNA, metabolic pathways such as nitric oxide (NO), increased glucose metabolism, inflammatory responses, and angiogenesis. For the late window, they include regulation of gene, regulator of programmed cell death (stimulate cell survival mechanisms and down regulated apoptotic pathways), receptor modulator like NMDA receptor activation, antioxidant capacity, and suppression of immune system (Alkan T., 2009; Dornbus & Ding, 2012; Dornbos, et al., 2013; Heyman, et al., 2016; Liu, et al., 2016; Li, et al., 2017). In addition, adaptation of glial cells after pre-conditioning including early differentiation of astrocytes may play a role in neuroprotection as well (Sen, et al., 2011).

Apart from remote conditioning with a repetitive inflation-deflation procedure of a blood pressure (BP) cuff, some studies had utilized isoflurane for post-conditioning. Isoflurane is a volatile anesthetic agent that can mimic ischemic-conditioning in both adult and neonatal brain (Kapinya, et al., 2002; Xiong, et al., 2003; Zhao, Peng, Li, Xu, & Zuo, 2007). A recent study shows that isoflurane post-conditioning can improve survival and neurological outcomes in an adult rat model of cardiac arrest (Zhang, Wu, Yu, & Liu, 2017).

The induction of neuroprotective effect is likely region-specific. A hypoxic pre-conditioning study shows a metabolic adaptation in hippocampus, parietal cortex and cerebellum of the brains of neonatal rats but not the striatum (Marcelino, et al., 2015). Another study in piglet shows reduction of cell death in the periventricular white matter, internal capsule and corpus callosum (Ezzati, et al., 2016). These coincide with the result from a study in adult rats that confirms the effect of hypoxic pre-conditioning including change in gene expression in the frontal cortex but not in the caudate, putamen and thalamus (Omata, et al., 2006).

**Neonatal HIE and Post-conditioning**

Several studies have shown that immature neonatal brain responds to hypoxic ischemic event differently from adult brain. Infants exposed to hypoxic ischemia tend to present with neonatal encephalopathy leading to difficulty in maintaining respiration, depression of tone and reflexes decreased consciousness, feeding issue and seizures (Nelson & Leviton, 1991; Hassell, Ezzati, Alonso-Alconada, Hausenloy, & Robetson, 2015). Hypoxic ischemic injury usually causes a prominent oxidative stress environment and accumulation of reactive species (Alkan, Goren, Vatansever, & Sarandol, 2008; Fatemi, Wilson, & Johnston, 2009; Penna, et al., 2014). The brain of newborns especially hippocampal cells are more vulnerable to oxidative stress and free radical oxidative damage than the brain of adult leading to the described detrimental consequences (Drunalini-Perera, et al., 2014). In acute hypoxic ischemic event, after the falling of Na+ / K+ ATP dependent pump, glutamate overflows and cytoplasmic Ca2+ concentration arises. Ca2+ triggers many downstream toxic cascades and generates high levels of reactive oxygen species. A study showed that Edaravone, a free radical scavenger could reduce the brain damage from hypoxic ischemic injury in neonatal rats (Ikeda, Xia, Kaneko, Sameshima, & Ikenoue, 2002). Hypoxic ischemia also causes changes in the regulation of genes, transcription factors and molecules relating to cell death signaling and immune responses (Guglielmotto, et al., 2009; Sameshima & Ikenoue, 2013). Mitochondrial membrane permeabilization was noted to play a major role in cell death. In adult, the mechanism of this permeabilization transition in hypoxic ischemic event is predominantly secondary to the cyclophilin D-dependent opening while Bax-dependent mitochondrial permeabilization is the key mechanism in immature neonatal brain. Bax-inhibiting peptide (BIP) was found to be associated with a reduction of Bax activation, mitochondrial permeabilization and downstream caspase activation (Wang, et al., 2010).

Post-conditioning is appropriate for HIE diagnosed immediately after birth. Remote ischemic post-conditioning (RIPC) has been well established. It is a simple, inexpensive, and safe use of repetitive inflation-deflation procedure of a blood pressure (BP) cuff on the arm or leg to protect distant organs such as the brain, heart, and retina from ischemic injury (Schmidt, et al., 2007; Gho, Schoemaker, van den Doel, Duncker, & Verdouw, 1996; Hess, Hoda, & Bhatia, 2013; Liu, Sha, & Cho, 2013; Zhang, et al., 2014; Yamaguchi, et al., 2015). RIPC provides protection in the adult and neonatal brain against subsequent lethal ischemia. (Zhou, et al., 2011; Hess, Hoda, & Bhatia, 2013). A randomized crossover study of healthy individuals showed no difference in blood flow, tissue perfusion, concentration of nitrite and platelet mitochondrial function between ischemic conditioning at thigh or arm. Thus, localization does not seem to affect the cyto-protection property of ischemic conditioning (Dezfulian, et al., 2017).

**Effects of RIPC**

Remote ischemic post-conditioning can reduce infarct size in both neonatal and adult brain. In neonatal brain, a study in rats shows that RIPC alone can significantly reduce the infarct volume after the hypoxic-ischemic brain injury with possibilities of involving opioid receptor/ AKT pathway (Zhou, et al., 2011). For adult brain, RIPC can significantly reduce infarct size in both animal and human models (Ren, et al., 2012; Zhao, Ren, Chen, & Shen, 2012; Liu, et al., 2014). RIPC reduced the infarct volume at 48 hours after hypoxic ischemia. It also significantly improved the long-term neurological functional outcomes. (Zhou, et al., 2011). However, RIPC shows no long-term improvement in brain weight (Zhou, et al., 2011; Drunalini-Perera, et al., 2014).

An application of RIPC after the neonatal hypoxic ischemic injury can improve long-term sensory, motor (coordination and strength) and cognitive impairments. These cognitive impairments include short-term memory and spatial learning and memory (Drunalini-Perera, et al., 2014). Isoflurane post-conditioning might have minimal effect on motor function but could significantly alleviate the spatial learning and memory impairments (Xu, et al., 2016). Another study also showed that physical training for 2 weeks after a hypoxic-ischemic insult in neonatal brain could reduce brain damage and decrease learning and memory impairments (Chen & Jiang, 2010).

Many studies have shown that post-conditioning should be done within 6-8 hours after hypoxic ischemic event for favorable outcome. This is likely the therapeutic window and post-conditioning might be able to reverse neuronal apoptosis (Hassell, Ezzati, Alonso-Alconada, Hausenloy, & Robetson,
However, a study shows that RIPC up to 24 hours after hypoxic ischemic event could provide a long-term neuroprotection and 3-day consecutive treatment could possibly give greater benefit (Drunalini-Perera, et al., 2014).

**Mechanism of RIPC in neonates**

The mechanisms of neonatal neuroprotection from RIPC is not well-understood but seems to involve nerve activation and limb ischemia that trigger humoral protective factors that circulate and provide protective effect throughout the body (Hassell, Ezzati, Alonso-Alconada, Hausenloy, & Robetson, 2015). Its neuroprotection can be via the transferring of protective factors such as opioids, receptor stimulation, mitochondrial permeability pore, KATP, pro-survival kinase (phosphatidylinositol-3-kinase [PI3K]/AKT, ERK pathways) (Zhou, et al., 2011; Xu, et al., 2016). This leads to increased cerebral blood flow, decrease in inflammation and upregulation of pro-survival signaling cascades. Plasma nitrite concentration seems to play an important role in increasing the blood flow. Although, there has been no study that specifically assessed neonatal NO, an adult human study shows that, after the first ischemic conditioning, plasma nitrite concentration peaks and remains elevated along with the ongoing conditioning (Dezfulian, et al., 2017). This elevated nitrite involves in cyto-protection from hypoxic/re-oxygenation injuries, mainly secondary to its reduction to NO by deoxygenated heme proteins (Kamga Pride, et al., 2014).

Apart from plasma nitrite, phosphor calmodulin-dependent protein kinase II (Phosphor-CaMKII) and Brain derived neurotrophic factor (BDNF) may play a role in neuroprotection of RIPC as a 2-week physical training after a hypoxic-ischemic insult in neonatal brain increased their expressions (Chen & Jiang, 2010). In addition, isoflurane post-conditioning could induce neuroprotection after hypoxic ischemic event partly by mediating GluR2 subunit of AMPA receptor in neonatal rats (Xu, et al., 2016).

There is a lack of studies in gene regulation after RIPC in neonatal brain. A hypoxic ischemia study in newborn piglet shows changes in gene expression of KATP channel and endothelin A receptor in the white matter (Ezzati, et al., 2016). For adult brain, a study in rats shows that local cerebral blood flow and increases in angiogenesis in the ischemic brain correlate with upregulated expressions of Notch 1 and Notch intracellular domain (NICD) surrounding the ischemic area (Ren, et al., 2016). Another adult rat study shows involvement of alteration of the expression of hypoxia-inducible factor (HIF)-related genes including vascular endothelial growth factor (VEGF) and erythropoietin (EPO) after remote ischemic pre-conditioning (Heyman, et al., 2016).

**Post-conditioning may have significant role in alleviating the surge in of oxidative stress in vulnerable neonatal brain after hypoxic ischemic injury. Post-conditioning in adult rat shows the effect of modulating reactive oxygen species. It exerts this via upregulation of anti-oxidative stress proteins such as nuclear factor erythroid 2-related factor 2 (Nrf2) and heme oxygenase -1 (HO-1) (Zhang, et al., 2014).**

Additionaly, post-conditioning may alter glial cell responses. In a newborn piglet study, applying post-conditioning resulted in reduced microglial activation in corpus callosum, which attenuated the inflammatory reaction. Post-conditioning may also improve mitochondrial metabolism leading to an increase in surviving oligodendrocytes in corpus callosum as well as in the periventricular white matter (Ezzati, et al., 2016).

With evidences from adult studies, ischemic conditioning might also involve many other molecular mechanism including neuroglobin, glial fibrillary acidic protein, lipid peroxidation level, lactic dehydrogenase, creatine kinase activities, JAK-STAT pathway, Protein kinase C pathway, antioxidant enzyme

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as well as other mRNA expressions (Alkan, 2009; Zhan, et al., 2010; Wang, et al., 2010; Liu, et al., 2012; Zhang, et al., 2014; Liu Y., Li, Li, & Zou, 2015).

Conclusion
HIE is the result of severe anoxic brain injury during the neonatal period and accompanies life-long morbidity, including cerebral palsy, and premature mortality. Therapeutic hypothermia immediately after birth is the standard of care for HIE. However, this hypothermia alone does not provide complete neuroprotection. Even after therapeutic hypothermia, high prevalence of neurologic morbidity and mortality persists. Due to that, an adjuvant therapy for patents with HIE is imperative.

Ischemic conditioning is an adaptive process of endogenous protection in which small doses of sub-lethal ischemia protect the organism against a lethal ischemic event. Post-conditioning is appropriate for HIE diagnosed immediately after birth as the conditioned stimulus is applied after the lethal ischemic episode. Remote ischemic post-conditioning (RIPC) is well established and is a safe use of repetitive inflation-deflation procedure of a BP cuff on the limb to protect distant organs from ischemic injury. A number of studies in neonatal rats have demonstrated that RIPC is effective at reducing injury in focal cerebral ischemia models. Combining this RIPC with standard therapeutic hypothermia to treat HIE is an attractive therapeutic approach.

RIPC alone can provide long-term neuroprotective effect. It is easy to use and can be started as late as 24 hours after the initial hypoxic ischemic injury in a rat model. However, beyond 24 hours, it is still unclear how long we can wait to start RIPC. Consecutive treatments may be beneficial as shown in many studies but the optimal regimen including the duration and dose is still inconclusive. Combining this RIPC with standard therapeutic hypothermia to treat HIE has the potential to be an effective synergistic treatment. However, it needs further studies to elucidate the effectiveness of this combination and their interactions as well as its feasibility and safety. In addition, another important question is whether this combination will offer the similar protection and improve neurological outcomes when we extrapolate it to human studies. Table 1 summarizes the current available data and the issues that require further attention.

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References


