Clinical vignette: An 8-year-old boy presents to the pediatric ICU after two days of cough with increasing secretions. The patient is progressing to respiratory failure and requires noninvasive mechanical ventilation. His past medical history is remarkable for premature birth at 25 and 6/7 weeks gestational age, cerebral palsy, developmental delay, epilepsy, and gastrostomy tube dependence. His chest x-ray is remarkable for multifocal opacities that are consistent with atelectasis. A complete blood count reveals a wbc count of 9.2 with a normal differential, Hg of 11.7, and platelet count of 276,000. A respiratory viral panel from a nasal swab returns positive for rhinovirus. Additional patient history from the parents uncovers that he has been hospitalized three times over the course of the past 2 years with a similar presentation.
Spare hypoxia, spoil the child?

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Current knowledge

The patient in this scenario illustrates a common and increasingly frequent clinical scenario in pediatric hospitals and wards. The most recent published admissions data from The Pediatric Health Information System database show that children with significant chronic medical conditions accounted for 19.2% of hospitalized patients, but a remarkable 48.9% of hospital days and 53.2% of hospital charges (1). Within this group, the most common primary chronic diagnosis was cerebral palsy.

The increasing survival of premature infants, particularly those at the extremes of prematurity and low birth weight, has resulted in a marked increase in the number of premature infants surviving into childhood. Unfortunately, many individuals born prematurely suffer chronic neurologic impairments (2). Recent data indicate that approximately 10% to 15% of extremely premature infants will go on to exhibit cerebral palsy, with an incidence that is inversely related to gestational age (3). Up to 50% will later exhibit cognitive or behavioral deficits (4). High-grade intraventricular hemorrhage (IVH) is a rare but important adverse event that affects this patient population; however, the most common radiologic and neuropathologic findings correlate these functional deficits to periventricular leukomalacia (PVL) and its accompanying neuronal/axonal abnormalities. PVL most broadly refers to a pattern of diffuse cerebral white-matter injury, with specific areas of necrosis and loss of cellular elements within the deep periventricular white matter (5). In the classic PVL description, these necrotic areas were initially quite large and subsequently evolved over time into macrocystic lesions (Figure 1). Thankfully, this type of PVL presentation has become more of a historical note, with recent imaging studies showing that severe PVL has a modern incidence of less than 5% (6, 7). In modern practice, it is far more common for areas of necrosis to be microscopic and progress to foci of glial scarring (so-called microcystic PVL). Moreover, the diffuse white-matter abnormalities are characterized by gliosis, microgliosis, and altered numbers and maturation of cells of the oligodendrocyte lineage (8). Imaging studies indicate that 50% or more of very low birth weight (VLBW) infants present with manifestations that are consistent with PVL (5).

PVL pathogenesis is multifactorial and incompletely understood. Human, animal, and in vitro data suggest that upstream physiologic derangements converge to cause the death and/or maturational arrest of oligodendrocyte precursors (preOLs), ultimately leading to the characteristic abnormalities of white-matter myelination. PreOLs are vulnerable to reactive oxygen and nitrogen species, excitatory molecules, and inflammatory mediators at developmentally specific and temporally restricted periods, helping to account for the highest incidence of PVL within infants born within a specific window of early prematurity (2, 8). The upstream events most commonly implicated in triggering these downstream sequelae are hypoxia/ischemia and inflammation. Interestingly, the mechanism whereby ischemia leads to PVL remains elusive. It is generally attributed to peculiarities of the arterial architecture serving the periventricular white matter coupled with immature autoregulation of cerebral blood flow (8–10). While the anatomy of these specific distal arterial fields is fairly well described, the idea that this predisposes the periventricular deep white matter to vascular insufficiency is merely inferred (9–11). Other regions perfused by end arteries without rich vascular anastomoses are not as prone to ischemic injury. Additionally, this theory is complicated by the lack of an animal model that recapitulates the focal changes of clinical PVL. Exposure to global hypoxia and/or ischemia in rodent and large animal models certainly results in widespread white-matter injury with frank infarction (12). In other models, to achieve more selective involvement of white matter requires adherence to an extremely narrow window of hypoxic exposure before gray matter injury ensues (2).

Research advances

In this issue, Licht, Dor-Wollman, and colleagues report on their generation of a novel murine model that recapitulates specific temporal and regional phenotypes analogous to those of human PVL (13). Spe-
poral window of vulnerability. Licht, Dor-Wollman, et al. note a lack of vessels in the affected regions of their model that corresponds to the striking paucity of arteries in histologic sections of infants with PVL (14). This model is also consistent with data from histologic sections of human fetal and postnatal brain specimens that show region-specific waves of VEGF immunoreactivity (15). Licht, Dor-Wollman, and colleagues speculate that the biology of PVL parallels that of retinopathy of prematurity (ROP), wherein hyperoxia exposure suppresses VEGF (via repression of HIF), which induces apoptosis of nascent vascular endothelial cells (16). Interestingly, a recent publication by Yuen et al. shows that in postnatal murine development, preOLs play a role in vascularization of white matter. Hypoxic HIF stabilization in these precursors directly upregulates transcription of the WNT ligands 7a and 7b, which then act in an autocrine manner to delay oligodendrocyte differentiation and myelination as well as in a paracrine fashion to stimulate angiogenesis in the corpus callosum (17). This postnatal white-matter vasculogenesis is predominantly at the capillary level (18) and distinct from the early angiogenesis that characterizes the VEGF-dependent window proposed by Licht, Dor-Wollman, et al. However, postnatal vasculogenesis in white matter does substantiate an ongoing role for HIF-mediated oxygen sensitivity in the coordination and direction of cerebral vascular development in particular as well as developmental vascularization in general (19).

Recommendations
It is now clear from the collective publication of the Neonatal Oxygenation Prospective...
Meta-analysis Collaboration (NEOPROM) studies of high versus low oxygen saturation targets that management aimed to minimize hypoxia exposure in premature infants can reduce the incidence of ROP (20–22). It is less clear what effect lower target saturations may have on neurodevelopmental outcomes, however. While neither the Canadian Oxygen Trial (COT) nor the Study to Understand Prognoses and Preferences for Outcomes and Risks of Treatments (SUPPORT) trial demonstrated a difference in the primary outcome of death or severe neurodevelopmental disability at 18 to 24 months, meta-analysis of the five trials found increased mortality in the low-saturation group (22). The low-saturation groups also had an increased incidence of necrotizing enterocolitis (NEC), a trend in the meta-analysis that was not significant in any of the individual trials. There was no substantial difference in other neurologic end points evaluated; however, these specific outcome measures were not uniform and generally encapsulated only the most blatant injuries, such as high-grade IVH. The findings by Licht, Dor-Wollman, et al. highlight the need for attention not just to IVH but also to a wider range of brain injuries in premature infants that potentially stem from exposure to hyperoxic conditions.

Perhaps most importantly, the insights from the paper by Licht, Dor-Wollman, and colleagues emphasize a potentially limited time frame of highest developmental risk. This model suggests that there is a defined transition point to VEGF independence and, by inference, decreased hyperoxia vulnerability, with respect to periventricular vascularization. Infants enrolled in the NEOPROM studies randomized to high- or low-saturation targets were maintained at their oxygen targets through 36-weeks gestational age, but this may far outlive the therapeutic benefit of relative hypoxia. At 29- to 32-weeks gestational age, when the incidence of NEC is reaching its peak (23), there may be no further neurovascular benefit from this strategy. Thus, it may be more appropriate to consider a tiered approach to oxygen saturation guidelines in premature infants that vary with their gestational age. Infants between 24- and 28-weeks gestational age, for example, could be targeted to lower \( O_2 \) saturations, with gradual increases to normal pediatric values after 28-weeks gestational age. Such an approach would therefore enable premature infants to reside in a physiologically hypoxic environment during the highly oxygen-sensitive developmental windows regulating cerebral and retinal vascularization, while preventing their exposure to potentially pathological hypoxia during later gestational ages at which diseases such as NEC are more likely to occur. The publication by Licht, Dor-Wollman, et al. therefore underscores the importance of developing more nuanced approaches to the use of oxygen therapy in the intensive care nursery than currently practiced.

Proper development requires precisely coordinated interactions between the embryo/fetus and its environment. A better understanding of the fundamental environmental mechanisms that drive mammalian development in utero will therefore help neonatologists devise more physiologically relevant treatment modalities to improve the care of premature neonates ex utero. Importantly, there is a growing urgency for increased basic research in this area as technological improvements in our abilities to sustain the lives of increasingly premature and small-for-gestational-age infants outstrip our abilities to ensure a meaningful quality of life for this incredibly at-risk population.

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