Survivors of sepsis and other forms of critical illness frequently experience significant and disabling cognitive and affective disorders. Inflammation, ischemia, and glial cell dysfunction contribute to this persistent brain injury. In this issue of the JCI, Hippensteel et al. show that endothelial injury in animal models of sepsis or endotoxemia leads to shedding of heparan fragments from the endothelial glycocalyx. These fragments directly sequester brain-derived neurotrophic factor and impair hippocampal long-term potentiation, an electrophysiologic correlate of memory. The authors further explore the specific characteristics of heparan fragments that bind neurotrophins and the presence of these fragments in the circulation of patients who survive sepsis. This study highlights an important mechanism by which vascular injury can impair brain function.
The vasculature in sepsis: delivering poison or remedy to the brain?

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Persistent brain injury in sepsis survivors
Mortality from sepsis has fallen markedly with improvements in supportive care (1). Although over 14 million patients survive a hospitalization for sepsis each year, they are still not made whole by the life-saving care they receive (2). Rather, sepsis survivors frequently experience functional disability, along with increased risk of rehospitalization and death. Persistent brain dysfunction is a common and life-changing consequence of critical illness survivorship and results in both cognitive decline and a high incidence of affective disorders such as anxiety, depression, and posttraumatic stress (3, 4).

Multiple factors contribute to persistent brain dysfunction in sepsis survivors. Neuropathologic studies of sepsis in patients have largely been confined to those who succumb to their acute illness and have revealed that ischemia, hemorrhage, and white matter injury commonly occur at the extremes of critical illness (5). It is unclear, however, whether brain injury that results from the hypoperfusion, hypoxia, and circulatory dysfunction that underlie multiorgan dysfunction syndrome in sepsis is treatable or preventable by means other than improved supportive care of the underlying critical illness. Studies using animal models, therefore, have focused heavily on inflammatory mechanisms that extend beyond the period of critical illness and into the period of survivorship. Neuroinflammation, including activation of microglia and astrocytes, is also a prominent component of brain injury in sepsis (6, 7).

Survivors of endotoxemia, abdominal infection, or pneumonia experience multiple persistent inflammatory insults to the brain. Neutrophils and inflammatory monocytes traffic and invade the brain parenchyma (8). Expression of cytokines and alarmins in the brain persists for weeks following sepsis (9, 10). Microglia are especially susceptible to long-term reprogramming after high-dose endotoxin exposure (11), though the balance of tolerance and priming that results from persistent exposure to proinflammatory signals over time is probably governed by a network of interactions among glia and neurons (12). In addition to the activation of intrinsic neuroinflammatory responses, sepsis may result in increased blood-brain barrier permeability and entry of inflammatory signals from the circulation (13, 14).

Despite evidence pointing to multiple mechanisms of persistent brain injury in sepsis survivors, the connection from a panoply of vascular and inflammatory insults to an actual persistent, reversible neuronal dysfunction in sepsis survivors remains obscure. Cytokine signaling and direct activation of innate immune receptors play an important role in the maintenance and elimination of synapses in both normal physiology and disease (15, 16), and these same mechanisms are presumably at work in the brains of sepsis survivors. Inhibition of IL-1 signaling and microglial activation rescues impairment of the long-term potentiation (LTP) that occurs acutely after sepsis or endotoxemia (17, 18). Though cytokines act directly on neurons and glia to change their intrinsic physiology and structure, inflammation may also activate a cascade of potentially reversible dysfunctions in the processing of neurotrophins and other signals important for circuit homeostasis. For example, brain-derived neurotrophic factor (BDNF) is an agonist of the TrkB receptor kinase that is required for long-lasting facilitation of synaptic transmission in LTP. In aged animals, systemic infection has been associated with prolonged increases in IL-1β expression and reductions in hippocampal BDNF, as well as sustained but reversible impairment of LTP (19).

BDNF sequestration as a mechanism of learning and memory impairment
In this issue of the JCI, Hippensteel and colleagues demonstrate that microvascular injury in sepsis results in impairment of learning and memory by a far more
direct mechanism: sequestration of BDNF by products of endothelial glycocalyx injury (20). The endothelial glycocalyx is a meshwork of extracellular protein and carbohydrate molecules that supports endothelial function (21). Sepsis degrades the glycocalyx and leads to pathologic vascular permeability and leukocyte adhesion. Here, however, the authors found that heparan sulfate fragments, constituents of the glycocalyx itself, interfered with brain function. Circulating heparan fragments were elevated in patients with sepsis and were associated with cognitive impairment in sepsis survivors. Murine sepsis survivors demonstrated deficits in hippocampus-dependent behavior and LTP impairment. While total levels of hippocampal BDNF were not reduced after sepsis, sulfated heparan fragments functionally sequester BDNF (Figure 1). The authors were not able to specifically disrupt the interaction of heparan sulfate fragments and BDNF to restore LTP. However, supplementation of BDNF ex vivo reversed the electrophysiologic effect of endotoxemia, and direct TrkB agonism overcame the inhibitory effect of highly sulfated heparans on LTP.

**Endothelial injury products in future studies of brain injury**

The possibility that products of endothelial injury can interfere directly with brain function has several implications for future studies dealing with the prevention and treatment of brain injury associated with sepsis. The dominant model of persistent brain injury in sepsis ascribes neuronal dysfunction to a combination of circulatory, immune, and glial dysregulation. Cognitive dysfunction has been associated with vascular injury during sepsis, but this has usually been framed in terms of hemodynamic impairment or loss of barrier function (14, 22). The complex relationships among peripherally derived inflammatory mediators and leukocytes, microglia, and astrocytes in sepsis survivors that impinge on neuronal function depend on cascades of signaling pathways and changes in transcriptional programs that alter intrinsic neuronal function and expression of neurotrophins and physical-
affective disorders that cause significant disability, reduce quality of life, and contribute to morbidity and mortality. The mainstay of treating acute brain dysfunction has for many years been to alter neuromodulatory neurotransmission, mainly with typical or atypical antipsychotics. Recent landmark studies have demonstrated that this approach fails to improve brain function or alter short-term mortality (24, 25). While data on long-term outcomes are still forthcoming, the inability of antipsychotic medications to reverse or prevent delirium illustrates that simply masking behavior, such as agitation, will not restore the important homeostatic mechanisms that the brain needs to recover from acute injuries and prevent long-term dysfunction. We need a more nuanced understanding of the underlying pathophysiology of brain injury in survivors of critical illness, and here Hippiensteel and colleagues have cast new light on previously unexplored but important mechanisms.

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