Effects of lactate and carbon monoxide interactions on neuroprotection and neuropreservation

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Abstract

Lactate, historically considered a waste product of anaerobic metabolism, is a metabolite in whole-body metabolism needed for normal central nervous system (CNS) functions and a potent signaling molecule and hormone in the CNS. Neuronal activity signals normally induce its formation primarily in astrocytes and production is dependent on anaerobic and aerobic metabolisms. Functions are dependent on normal dynamic, expansive, and evolving CNS functions. Levels can change under normal physiologic conditions and with CNS pathology. A readily combusted fuel that is shuttled throughout the body, lactate is used as an energy source and is needed for CNS hemostasis, plasticity, memory, and excitability. Diffusion beyond the neuron active zone impacts activity of neurons and astrocytes in other areas of the brain. Barrierogenesis, function of the blood-brain barrier, and buffering between oxidative metabolism and glycolysis and brain metabolism are affected by lactate. Important to neuroprotection, presence or absence is associated with L-lactate and heme oxygenase/carbon monoxide (a gasotransmitter) neuroprotective systems. Effects of carbon monoxide on L-lactate affect neuroprotection – interactions of the gasotransmitter with L-lactate are important to CNS stability, which will be reviewed in this article.

Key words: astrocytes; biochemical interactions; CNS; gasotransmitters; microglia; neuropreservation; neuroprotection; oligodendrocytes

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Introduction

Lactate (discovered in 1780, considered a waste product of metabolism) is a requisite molecule for whole body metabolism and normal central nervous system (CNS) development and functioning. It is the preferred substrate for energy purposes and a precursor of lipids in oligodendrocytes, astrocytes, and neurons during the perinatal period, but is preferentially associated with ischemia of the brain. Evidence now, however, indicates that lactate is found in the CNS under physiological conditions. Neuronal activity signals induce formation primarily in astrocytes. While it is used as an energy source, it is also needed for CNS hemostasis, plasticity, memory, and excitability and is now recognized as a necessary metabolite in the coordination of whole-body metabolism as a readily combusted fuel that is shuttled throughout the body, a potent signaling molecule, and a hormone. These functions are dependent on CNS requirements for normal dynamic, expansive, and evolving CNS physiological functions. Presence or absence is also important to CNS neuroprotection and is associated with L-lactate and heme oxygenase/carbon monoxide (CO) (a gasotransmitter) neuroprotective systems. Effects of CO on L-lactate affect neuroprotection - interactions of the gasotransmitter with L-lactate are important to CNS stability, which will be reviewed in this article. The following databases were systematically searched: MEDLINE, EMBASE, and Cochrane. Databases were searched from their inception to December 31, 2020.

Sources of Lactate in the Central Nervous System

Considered a waste product of anaerobic metabolism, lactate has known beneficial roles in the CNS and its production is not limited to anaerobic metabolism. CNS concentration depends on serum lactate levels, oxygen availability, neuronal firing, degradation, and metabolic rate. Lactate serves as an important metabolic fuel and an intercellular messenger – its diffusion beyond the neuron active zone impacts activity of neurons and astrocytes in other areas of the brain. Contributing to neuronal metabolic support by transferring lactate through cytoplasmic myelinic channels and monocarboxylate transporters (MCTs), oligodendrocytes are active participants in this process – lactate is released by myelinating oligodendrocytes and then used by axons for generation of mitochondrial adenosine triphosphate. Oligodendrocytes are associated with the cell-to-cell lactate shuttle and the astrocyte-neuron lactate shuttle (lactate shuttling between cells producing lactate and cells consuming lactate is important to oligodendrocyte metabolism). These processes allow lactate to be used as a signaling molecule, energy source, and gluconeogenic precursor in the CNS. Lactate also affects barrierogenesis and function of the blood-brain barrier (BBB), which is a buffer between oxidative metabolism and glycolysis, and may be the preferred fuel for brain metabolism. Lactate shuttles within and between cells and lactate uptake across the BBB suggests it is integral to CNS homeostasis although its transport across the BBB is rate-limited and lactate production within the brain reflects the needs of the CNS (metabolism and utilization is dynamic). Astrocyte-to-neuron lactate shuttling occurs during rest, but with excitation glycolysis of glucose to lactate (exceeding rate of mitochondrial fuel oxidation) in the neurons provides for increased energy needs. In the newborn mammalian brain, elevated blood lactate is utilized...
Dephosphorylation of fructose-1,6 biphosphate: biology/metabolism and is similar between individuals.21-23

Flow of blood affects BBB cell structure – cell differentiation.29 Clearance of substances from the CNS involves perivascular efflux, metabolism, and the BBB.30,31

L-lactate is considered a volume transmitter and can travel large distances from the site of production (i.e., muscle) to a site of consumption (i.e., CNS) with monocarboxylate transporters (MCTs) mediating diffusion of lactate between extracellular space and cells. Lactate passage across the BBB by nonsaturable diffusion or via MCTs in the plasma membrane – equilibration between lactate concentration in the blood and brain results (lactate then equilibrates with pyruvate via lactate dehydrogenase) – can be restricted.32 With normal plasma lactate levels, non-saturable diffusion can be as high as 50% of lactate transport from the blood to the CNS and, with high plasma lactate levels, near instantaneous equilibration occurs. MCTs carry L-lactate (most abundant monocarboxylate in the brain) across different cell membranes including the BBB – cotransport of monocarboxylate anion and proton occurs. Four carriers (MCTs 1–4) account for bidirectional, electroneural 1:1 co-transport of protons and monocarboxylic acids (i.e., lactate). MCT2 has the highest affinity for lactate – regionalization of MCTs 1–4 has been suggested to indicate involvement in different aspects of lactate metabolism and their production varies under different conditions. Accessory proteins (i.e., embigin, basigin, neuroplastin) are necessary for their activity.32-36 Efflux by passive transport which is mediated by MCT1 which is present in both the abluminal and luminal membranes.30,37

Production of lactate in the brain

Important major metabolic pathways, catabolic and anabolic, necessary for CNS life/growth/division include gluconeogenesis/glycolysis, fatty acid β-oxidation, urea cycle, pentose phosphate pathway, oxidative phosphorylation, and the citric acid cycle. Glucose, the obligatory CNS energy substrate in mature humans, depends on glucose transport/metabolism, is mostly consumed by oligodendrocytes and astrocytes, and metabolites produced and released by astrocytes and oligodendrocytes are used by neurons as energy sources. The preferred energy substrate of the brain, lactate, is an important energy substrate for axons, is important in CNS health, and is an important source earlier in life and is now thought to be an important metabolite needed in the adult brain. Present in astrocytes, microglia, neurons, and oligodendrocytes, lactate production is via multiple pathways and its timely metabolism and elimination promotes CNS health and well-being – metabolic switching to lactate in the CNS has been linked to resistance to disease, stress, and injury. Metabolic processes in the CNS, dependent on the presence of multiple cell types and level of development/maturity, result in a pyruvate/lactate ratio that decreases with maturation and differs in regions of the brain. Astrocyte-neuron interactions, important to lactate metabolism, reflect dynamic needs and its production – glycolysis and glycogen metabolism are necessary for normal brain function, growth, and healing and can result in the production of lactate. CNS lactate production includes the following pathways: gluconeogenesis/glycolysis pathway, fatty acid β-oxidation, urea cycle, pentose phosphate pathway, oxidative phosphorylation, and citric acid cycle.4,7,38,39

Gluconeogenesis/glycolysis pathway

CNS generation of glucose from non-carbohydrate precursors (gluconeogenesis) is a multistep process – enzymes used in glycolysis may catalyze reversible reactions. There are, however, three irreversible reactions:

1) Conversion of pyruvate to phosphoenolpyruvate:
Pyruvate carboxylase: Pyruvate (cytosol) -> pyruvate (mitochondria) -> oxaloacetate (mitochondria) -> oxaloacetate (cytoplasm);
Phosphoenolpyruvate carboxykinase: Oxaloacetate (cytoplasm) -> oxaloacetate (cytosol);
Phosphoenolpyruvate carboxykinase: Pyruvate (cytosol) -> phosphoenolpyruvate;

2) Dephosphorylation of fructose-1,6 biphosphate:
Fructose 1,6-bisphosphate (cytoplasm) -> fructose-6-phosphate (cytoplasm);

3) Dephosphorylation of glucose 6-phosphate:
Glucose-6-phosphate (cytoplasm) -> glucose (cytoplasm).

Pyruvate kinase catalyzes the conversion of phosphoenolpyruvate to pyruvate. Pyruvate can then be converted to lactate via

Uptake through the BBB

The highly vascularized BBB, a protective interface between the body and the CNS, selectively allows passage of substances, including lactate, into the brain. Initially identified by Paul Ehrlich28 (injected a dye into the bloodstream of mice and noticed that the dye did not enter the brain or spinal cord), the term BBB was introduced by Lewandowsky29 after studies showed that different substances injected into the ventricles of the brain resulted in neurologic symptoms that were not seen when injected intravenously. Electron microscopy showed that the brain endothelial cells construct a barrier between blood and CNS with a single layer of brain endothelial cells in contact with neurons, astrocytes, pericytes, and vascular smooth muscle cells. BBB function depends on signaling and crosstalk of this neurovascular unit.26,27 Tight junctions (barrier to passage of ions and molecules through the paracellular pathway and to the movement of proteins and lipids between the apical and the basolateral domains of the plasma membrane) and adherent junctions (allow epithelial cells to establish polarity with different proteins and lipids in the apical and basal plasma membranes) maintain the integrity of the BBB. Although it is best known for its barrier function of restricting transport of toxic and/or harmful substances from the blood to the brain, the BBB also has a carrier function that is responsible for the removal of metabolites and transport of nutrients to the brain. Its barrier function depends on the paracellular barrier, transcellular barrier, enzymatic barrier, and efflux transporters. Passive diffusion and specific transport proteins are necessary for its carrier function.28 Flow of blood affects BBB cell structure and function, increases BBB tightness, and affects endothelial cell differentiation.29 Clearance of substances from the CNS includes perivascular efflux, metabolism, and the BBB.30,31

Pyruvate kinase catalyzes the conversion of phosphoenolpyruvate to lactate (cytoplasm) -> pyruvate (mitochondria) -> oxaloacetate (mitochondria) -> oxaloacetate (cytoplasm).
lactate dehydrogenase. Availability of enzymes varies between cells. For example, astrocytes exhibit significant 6-phosphofructo-2-kinase/fructose-2,6-bisphosphatase-3 activity, a key mechanism for regulating glycolysis and gluconeogenesis through synthesis or hydrolysis of fructose-2,6-bisphosphate. Preferential production of lactate via the gluconeogenesis/glycolysis pathway is dynamic and determined by the needs, metabolism, and health of the CNS.40-45

**Fatty acid β-oxidation pathway**

Lipids (classified into five major subcategories – fatty acids, triglycerides, sphingolipids, sterol lipids, and phospholipids) are important to normal CNS function and development and serve as bioactive molecules, energy substrates, acting as building blocks or a combination of these. Fatty acid imbalance affects CNS development and function. Fatty acids are the essential component of all of the lipids and consist of a carbon chain that ends in a carboxylic acid functional group (subclassified as short-chain fatty acids (2–4 carbons), medium-chain fatty acids (6–12 carbons), long-chain fatty acids (14–18 carbons), and very long-chain fatty acids with 18+ carbons.) If saturated, all of the carbons have hydrogen atoms and single bonds only exist between carbons whereas unsaturated fatty acids have carbon chains where a double bond has been introduced (mono- versus polyunsaturated defines whether there is more than one double bond.) Fatty acid β-oxidation results in the formation of acetyl-CoA as noted in the equation: β-oxidation of fatty acids→→→→→→→→→→Acetyl-CoA (Figure 1).

Mitochondrial β-oxidation of fatty acids results in energy—repeating sequence of four reactions catalyzed by acyl-CoA dehydrogenase, enoyl-CoA thiolase, hydroxyacyl-CoA dehydrogenase, and ketoacyl-CoA thiolase. Acetyl-CoA results with a fatty acyl molecule that is two carbons shorter –reaction cycle repeats (Figure 1). With an odd number of carbon atoms, propionyl-CoA is the final product. The acetyl-CoA enters the tricarboxylic acid cycle and pyruvate and, possibly, lactate may be generated. Propionyl-CoA is converted into succinyl-CoA (substrate for gluconeogenesis via oxaloacetate formation). Pyruvate and, possibly, lactate may be generated.46-50

Cytoplasmic fatty acyl-CoA is converted to fatty acyl carnitine by carnitine acyl transferase I, an enzyme of the inner leaflet of the outer mitochondrial membrane. Fatty acyl carnitine is then transported by an antiport in exchange for free carnitine to the inner surface of the inner mitochondrial membrane. There carnitine acyl transferase II reverses the process, producing fatty acyl-CoA and carnitine. This shuttle mechanism is required only for longer chain fatty acids as follow:

Urea cycle

While the urea cycle and the tricarboxylic acid cycle (TCA) are independent cycles, they are linked. Fumarate that is produced in the cycle is an intermediate in the TCA cycle, metabolized to malate, and then to pyruvate and lactate (Figure 2). Urea cycle - Citrulline→→→→→→argininosuccinate→→→→→→arginine and fumarate
TCA cycle - Fumarate→→→→→→malate,51-54

Pentose phosphate pathway

Taking place in the cytosol, the pentose phosphate pathway is anabolic more than catabolic, generates reducing equivalents, produces ribose 5-phosphate (needed for synthesis of nucleotides and nucleic acids) and erythrose 4-phosphate (needed for synthesis of aromatic amino acids), and is linked to glycolysis. During the oxidative phase (irreversible), glucose 6-phosphate is metabolized to ribulose 5-phosphate. The latter can be metabolized to fructose 6-phosphate during the nonoxidative phase (reversible.)
Oxidative phosphorylation
Cellular respiration depends on glycolysis (cytosol) which results in pyruvate, pyruvate transformation into acetyl-CoA (mitochondria), the citric acid cycle (acetyl-CoA is modified in the mitochondria to produce energy precursors), and oxidative phosphorylation (or electron transport-linked phosphorylation) – process where electron transport from the energy precursors from the citric acid cycle leads to the phosphorylation of adenosine diphosphate producing ATP (occurs in the mitochondria). Uncoupling between glycolysis and oxidative phosphorylation involves the partitioning between pyruvate (the primary substrate for glucose-driven oxidative phosphorylation) and lactate. In the mitochondria, succinate is reduced to fumarate by Complex II. The latter enters the Krebs cycle and pyruvate (and then lactate via lactate dehydrogenase (LDH)) can be produced. Nitric oxide oxidation also inhibits the mitochondrial respiratory chain at cytochrome c oxidase in astrocytes and the majority of glucose consumed is released as lactate.

Citric acid cycle
Oxidation of acetyl-CoA to CO₂ by the citric acid cycle is the central process in energy production. The process also results in the formation of intermediates which can be converted primarily to fatty acids, glucose derivatives, γ-aminobutyric acid (GABA), glutamate, glutamine, and aspartate. Balance between the two biochemical processes related to the cycle (anaerolysis and cataplerosis) is essential to normal CNS physiology. Pyruvate carboxylation is important for the net synthesis of glutamate, GABA and aspartate (anaerolysis) and is associated with cerebral activity - flux through pyruvate carboxylase is higher in the awake state compared to deep phenobarbital anaesthesia. However, the brain cannot fully metabolize 4- and 5-carbon molecules – they must be removed by cataplerosis, a process that may be linked to biosynthetic processes resulting in synthesis of fatty acids and amino acids and gluconeogenesis.

Glutamate seems to be the molecule connecting anaerolysis and cataplerosis in the Krebs cycle. Cytosolic malic enzyme and phosphoenolpyruvate carboxykinase in astrocytes converts malate and oxalacetate into lactate which leaves the brain (blood or periventricular system.) Efflux of lactate allows for ongoing pyruvate carboxylation.

Movement of lactate in the brain
Lactate shuttles
The lactate shuttle hypothesis describes the movement of lactate within and between cells and is based on the observation that lactate is formed and utilized continuously in diverse cells under both anaerobic and aerobic conditions. Interconversion of lactate and pyruvate occurs via lactate dehydrogenase and is produced at sites with high rates of glycolysis and glycogenolysis. Lactate plays an important role as a shuttle and can be shuttled to adjacent or remote sites and can be used as a gluconeogenic precursor or substrate for oxidation. The hypothesis also addresses the role of lactate in lipolytic control, redox signaling, and gene expression and affirms the role of lactate in cell signaling, endocrine/autocrine/paracrine link between oxidative and glycolytic metabolism, and in the delivery of gluconeogenic and oxidative substrates – L-lactate is considered the link between aerobic and glycolytic pathways (substrate for mitochondrial respiration and a product of glycolysis.) Glycolytic lactate producer cells provide lactate that is taken up by the blood and mitochondrial reticulum (oxidative lactate consumer cells.) The anion is metabolically available under aerobic conditions and is important in a complicated feedback loop – lactate production is promoted with decreased ATP levels which leads to the cells ability to promote ATP homeostasis. Redox state, concentration gradient, and/or pH gradient are known to drive these shuttles. Cell-to-cell lactate interactions are important to CNS function. There is a normal physiological function and relationship between astrocytes and neurons – the astrocyte–neuron lactate shuttle. An increase in synaptic activity or low glucose conditions result in transfer of lactate from astrocytes to neurons which sustains neuronal oxidative metabolism. Astrocytes express MCT4 (a low affinity transporter for lactate suggesting that lactate produced by glycolysis is exported.) Neurons express MCT2 (a high affinity transporter for lactate.) The hypothesis is that astrocytes produce lactate which is then taken up by neurons and oxidized for energy. There are other CNS cell-to-cell lactate shuttles as well and these include microglia and their preferential utilization of lactate under pathologic conditions, the activity of lactate shuttles in traumatic brain injury and neurodegenerative diseases and the astrocyte–microglia lactate shuttle.

Cell-to-cell, when coupled to intracellular, lactate shuttles allow for the simultaneous presence of glycolytic (i.e., glucose
to lactate) and oxidative (i.e., lactate to pyruvate to acetyl-CoA) pathways. This intracellular lactate pathway – lactate exchange and conversion into pyruvate maintaining the redox balance in the cytosol and mitochondria – is thought to exist between cytosol and mitochondria and between cytosol and peroxisomes.\textsuperscript{15,74}

**Glymphatic system**

About 60% to 68% of the brain’s total water content is within the intracellular space and 32–40% in the extracellular space (12% to 20% in the interstitial fluid (ISF), 10% in the cerebrospinal fluid (CSF), 10% in the blood.) The CSF, formed by the choroid plexuses (no BBB), depends on osmotic and hydrostatic movement of fluid, crystalloids, and colloids from plasma into stroma. The glymphatic system, integral to CNS fluid balance and removal of waste, has glial-dependent perivascular channels formed by astroglial cells, is dependent on aquaporin (AQP) 4 water channels on astrocyte endfeet, has peri-venous clearance and peri-arterial influx pathways, is connected to the lymphatic system associated with the dura, cranial nerves, and vessels which are located at skull exits, and allows continuous ISF and CSF removal of potentially harmful molecular products. Activity is increased during general anesthesia and during sleep and is decreased with stroke, diabetes mellitus, Alzheimer’s disease, and during aging.\textsuperscript{75-87} CNS lactate (higher while awake than during sleep and associated with reduction with cerebral glycolysis), a biomarker of the sleep-wake cycle, is exported as lactate by the glymphatic system – brain lactate concentration being inversely correlated with glymphatic-lymphatic clearance and resulting lactate/pyruvate ratio (and NADH/NAD redox potential) – and decreases rapidly with the onset of sleep. Elimination of lactate during sleep combined with the lack of elimination during wakefulness results in an increase of redox state with wakefulness (relative to sleep).\textsuperscript{83-85}

**Elimination of Lactate from the Central Nervous System**

Maintaining a normal lactate level in the CNS depends on its utilization \textit{versus} its accumulation. While the resting brain releases lactate (release from various brain structures differs under basal conditions), this increases incrementally with activation of the brain.\textsuperscript{86} Total lactate clearance is determined by its metabolism in the CNS, excretion via capillaries, or excretion through the CNS – clearance systems can be via CSF, ISF within the CNS, and blood.\textsuperscript{86-88} Three major models of lactate elimination via brain fluids have been hypothesized: 1) meningeal lymphatics,\textsuperscript{89} the glymphatic system,\textsuperscript{83,86} and the intramural parietal system.\textsuperscript{91,92}

The lymphatic system is a part of the vascular system. Peripherally, plasma is filtered through the vascular endothelium of capillaries in the extracellular space with over 80% being reabsorbed back into venous vessels and the remainder in the extracellular fluid or ISF. ISF is then filtered back into the lymphatics which return it into the venous system. In the CNS, the volume of brain fluid is fixed, there is BBB restriction of fluid filtration from plasma into the brain, and there is no common agreement about ISF formation in the brain. The three models of ISF formation that have been proposed include production by brain metabolism, cerebral capillary secretion of solutes, and ISF is a fraction of recycled CSF (flows from choroid plexus into subarachnoid space and then into perivascular space where CSF merges with ISF that has been generated by cerebral capillaries) – lymphatic communication with brain fluids affects the clearance of lactate as well as that of waste products, bacteria, viruses, toxins, etc.\textsuperscript{93-95} The extensive meningeal lymphatic vessel network that helps with macromolecular clearance and immune cell trafficking in the CNS also communicates with the glymphatic system, a system that allows CNS perfusion by ISF and CSF, and complements intramural, parietal clearance.\textsuperscript{96-98}

**Excess or too Little Lactate**

Maintenance of lactate levels is necessary for normal health and well-being – an excess or relatively too little can result in changes in body metabolism. Fluctuations are not usually due to low oxygen levels as hyperlactatemia can occur with normal tissue perfusion/oxygenation and relative hypolactatemia with poor tissue perfusion/oxygenation. Although lactate (the major gluconeogenic precursor, a signaling molecule, and a major energy source) production occurs normally during rest, it is increased with sepsis, pancreatitis, trauma, heart failure, etc.\textsuperscript{68,99-102} The resulting lactic acidosis, overproduction or underutilization of lactic acid with the body not being able to adjust to these changes, is associated with a buildup of acid and an imbalance of the bodies pH level. Elevated levels are used as a biomarker for risk and therapy and high levels are associated with an increased risk of death independent of organ failure and shock.\textsuperscript{103-105} Cellular metabolic reprogramming, metabolic inflexibility, and a decrease in cellular proliferation occur.\textsuperscript{106-111} Effects on the cardiovascular system can result in hyporesponsiveness of the vascular system to vasopressors and reduced cardiac contractility. NAD+/NADH ratio, production of reactive oxygen species, regulation of genes, release of vascular endothelial growth factor/interlukin-1/ transforming growth factor-β, and activation of Sirtuins are affected.\textsuperscript{101,109-112} Severe lactic acidosis has been suggested as a prerequisite of brain infarction from complete ischemia. This may be due to elevated brain glucose metabolism to lactate and its effect on astrocytes as excessive lactate is considered to be detrimental to the CNS and can affect recovery from brain trauma, anoxia, epilepsy, and ischemia.\textsuperscript{102} However, metabolism and cellular interactions determine if lactate is a waste product or a useful substrate. While acute and chronic pathologies alter brain metabolism resulting in changes in lactate concentration, lactate is a preferred fuel over glucose in brain preparations.\textsuperscript{32,112,113} When there is an increase in blood lactate in exercising humans (or when exogenous lactate is supplied), lactate is the major gluconeogenic precursor and substitutes for glucose as an energy substrate – cerebral lactate uptake increases. Improved cognitive function in brain-injured rats given intravenous lactate therapy and rapid increases of lactate MCT protein expression in rat brains following traumatic brain injury suggest that lactate metabolism in humans following traumatic brain injury is significant to recovery. The roles for lactate in normal cerebral metabolism and with CNS
pathology are being redefined. Lactate supports hepatic and renal gluconeogenesis which is important to brain metabolism, production by the body indirectly supplies glucose to the injured brain, isotopically labeled lactate is directly consumed and used by the injured human brain, and exogenous lactate infusion augments cerebral substrate supply when glycolysis is affected.\textsuperscript{115-116} Lactate changes, reflective of overall metabolic health, are affected by heme-oxygenase (and CO) and affect the health of the CNS.

**Carbon Monoxide Effects on Lactate in the Central Nervous System**

CNS extracellular lactate concentration increases with neuro-pathology as well as CNS activation - both result in significant release of lactate.\textsuperscript{117} The latter, which can be associated with excitotoxicity, may result in neuronal damage and death as excitotoxicity is a major component of CNS pathology.\textsuperscript{118-122} L-lactate (protects neurons against excitotoxicity, a regulator and metabolite in the CNS, regulates the expression of synaptic plasticity and neuroprotection genes in cortical neurons, promotes angiogenesis/immune escape/cell migration, serves as a glucose-sparing substrate, regulates cerebral blood flow, and plays an important role in learning and memory) acts as a signaling molecule in pathological states,\textsuperscript{123-125} is neuroprotective, and effects are directed at neurotransmission. CO has been found to modulate L-lactate levels in astrocytes providing evidence that the heme oxygenase and L-lactate neuroprotective systems interact.\textsuperscript{8}

The relationship between CNS L-lactate and the heme oxygenase/CO system is complex. The cytoprotective properties of the heme oxygenases are attributed to the production of CO. Endogenous production originates from heme metabolism (at least 86%) and heme-independent sources which include iron-ascorbate-catalyzed lipid peroxidation of microsomal lipids and phospholipids, photo-oxidation of organic compounds, auto- and enzymatic oxidation of phenols, and reduction of cytochrome b5. Heme oxygenase isozymes catalyze the first step of heme metabolism, and heme-oxygenase affects dilation of the vasculature to these stimuli. Apoptosis and oxidant-generating pathways are also inhibited.\textsuperscript{126,127} heme oxygenase affects dilation of the vasculature to these stimuli. Apoptosis and oxidant-generating pathways are also inhibited. CO binds to smooth muscle cell Ca\textsuperscript{2+}-activated K\textsuperscript{+} channel BKCa) channel-bound heme which leads to an increase in Ca\textsuperscript{2+} sparks-to-BKCa coupling (also binds directly to the BKCa channel at different locations) resulting in vasodilation. Inhibition of heme oxygenase affects dilation of the vasculature to these stimuli. Apoptosis and oxidant-generating pathways are also inhibited.\textsuperscript{128,129} Lactate may act as a dose-dependent regulator of cerebral microcirculation by affecting the hyperemic response, which may be related to a cytosolic redox impairment, and optimization of blood flow may interact with nitric oxide and arachidonic acid metabolites released by neuronal

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and glial cells during neural activity. As both CO and lactate affect calcium channels and NMDA, the relationship to each other, calcium channels/NMDA, and CNS local blood flow are important questions that have not been addressed. Further studies are needed.

**Glycogen synthesis and storage**

A polymer of glucose, glycogen is present in the CNS at low concentrations relative to the liver and skeletal muscle and, in developed brains in mammals under normal conditions, is located predominantly in astrocytes (has been found in embryonic neurons.) Synthesis is a multi-step process, is tightly regulated, and glucose transport across the BBB is needed. Glycogen synthase and the glycogen branching enzymes elongate glucose chains and introduce branch points. Astrocytes depleted of glycogen are not as successful at supporting neurons as are those not depleted of glycogen.

Astrocyte glycogen metabolism is important for functioning of the CNS. The substance is too large to be released and travel between cells, but studies have shown that astrocytes release lactate into the media. Modulated by a number of factors, neuronal activity signals result in astrocyte release of lactate – formed predominantly in astrocytes from glucose or glycogen and transferred from astrocytes to neurons to match the neuronal energetic needs and to provide signals that modulate neuronal functions and important to CNS health and disease. CO modulates L-lactate levels (and, therefore, glycogen metabolism) in astrocytes and is being further studied.

**Formation of neuro-glial-vascular unit and glial-vascular interface**

Composed of neurons, astrocytes, microglial cells, endothelial cells, pericytes, smooth muscle cells, and circulating blood cells, the neuro-glial-vascular unit interact to control synaptic communication, BBB function, local blood supply, neuronal development and surveillance/immune function. Models (Buxton-Want model of vascular dynamics, Hodgkin-Huxley formulation of neuronal membrane excitability, biophysical model of metabolic pathways) of the neuro-glial-vascular unit describe neural dynamics and propound that the energy inside of the neuron is dynamic and depends on neural activity – lactate in astrocytes can fuel neuronal activity – and local vasculature. The astrocyte-neuron lactate shuttle is predicted by these models – temporal dynamics of tissue lactate, tissue glucose and oxygen consumption, and of BOLD signal reported in human studies correctly predicted, transfer of lactate from astrocytes to neurons noted in response to activity, dynamics of extracellular lactate and oxygen as observed in vivo in rats correctly predicted, neuronal oxidative metabolism increased first upon activation with a subsequent delayed astrocytic glycolysis increase. This requires MCT. Nitric oxide has been shown to down-regulate MCT 1. The effect of CO on MCT’s and, therefore, lactate effects on the neuro-glial-vascular unit have not been studied.

**Molecular Regulation of pH**

Normal functioning of the CNS is affected by pH as changes affect the tightly regulated micro-circuits. Several processes regulate H+ in the CNS – intracellular H+ sequestering, CO2 diffusion, H+ buffering, membrane transport of acid/base equivalents across cell membranes, carbonic anhydrase activity, monocarboxylic acid transporters, vascular-type proton ATPase, etc. Acid extrusion as well as Na+/H+ exchange, Na+-HCO3- cotransport, and Na+-dependent Cl-/HCO3- exchange mediate pH levels. In the spinal cord, cells have two types of molecules in the cell membrane that detect pH levels with one reacting to acidic pH (acid sensing ion channel subunit 3) and the other to alkaline pH levels (PKD2L 1.) With neuronal activity, lactate is released and pH becomes more acidic. Lower pH and higher pH are inhibitory on motor activity via the secretion of somatostatin from nerve terminals of the pH-sensing central canal cells. Ion channel gating/ conductances, metabolite exchange and neuronal excitability, synaptic transmission, and intercellular communication via gap junctions are dependent on pH and subsequent information processing. CNS lactate and CO can result in changes in pH. Normalization of pH can occur at multiple levels as noted above and both lactate and CO affect several of these processes. Studies to evaluate the interaction of lactate and CO on CNS pH have not been done.

**Water transport and homeostasis**

Water transport and homeostasis in the CNS is partly dependent on cotransporters – active and passive modes have been suggested. Numbers and types of cotransporters per cell and unit water permeability vary. Water channel proteins, AQPs, are from a large family of major intrinsic proteins that are integral to water transport and homeostasis in the human brain and mediate water flux between the four water compartments in the CNS. Eight AQPs are expressed in the brain – AQP1, AQP3, AQP4, AQP5, AQP7, AQP8, AQP9, and AQP11 with AQP1 and AQP4 expressed in the highest concentrations. Orthodox AQPs only conduct water while aquaglyceroporins also have the ability of conducting other small molecules. AQP9 is involved in water flux through plasma membranes and is also permeable to monocarboxylates (i.e., lactate). AQP4, the most abundant AQP in the brain, is concentrated in the astrocytic foot processes, is important to BBB defense, neuroplasticity, removal of waste, and homeostasis of water (transports water into the brain but also moves water out of the brain in times of edema).

AQPs also conduct gas molecules. AQP1 has been associated with nitric oxide permeation when reconstituted in lipid vesicle and in vivo. Nitric oxide is produced by both neuronal nitric oxide synthase and endothelial nitric oxide synthase – AQP4 is found in the proximity of both neuronal nitric oxide synthase and endothelial nitric oxide synthase (exists in plasma membranes of end feet of astrocytes which surround endothelial cells of capillaries). AQP4 is more adapted for gas conduction when compared to AQP1. Whether AQPs can conduct gas molecules (and which gases) and whether AQP-mediated gas conduction is clinically important are not clear. Initial studies suggest that lactate and CO are probably affected by AQP’s. Further studies are needed to determine affects by AQPs on lactate and CO and interactions between gas conduction.
lactate, CO and AQPs.

**Neurotransmitter homeostasis**

There are more than 40 neurotransmitters in the human CNS – excitatory (glutamate, acetylcholine, histamine, dopamine, norepinephrine, epinephrine), inhibitory (GABA, serotonin, dopamine), neuromodulators (dopamine, serotonin, acetylcholine, histamine, norepinephrine), and neurohormones (releasing hormones, oxytocin, vasopressin). Their homeostasis, ability of a cell/system of cells in the brain to identify a change and respond with a compensatory response that restores baseline function, results as an interaction of several mechanisms including regulation of release of neurotransmitters by autoreceptors, non-synaptic production of the neurotransmitters, neurotransmitter vesicular release from the presynapse, uptake by transporters, and diffusion of the neurotransmitters. Neurotransmitter gradients exist and their maintenance is crucial for neurological processes – glia surrounding the synapse are a major effector of outcomes. Computational models have shown that specific glial configurations may be necessary, non-synaptic sources are needed to maintain extracellular concentrations of neurotransmitters, transporter and non-synaptic source densities can co-vary to provide tone on presynaptic autoreceptors, and synaptic glial configuration can be suggested for a given neurotransmitter and a given extracellular neurotransmitter concentration.\(^{155-157}\) CO (and nitric oxide) is a neurotransmitter as well. Not stored in synaptic vesicles, release of CO with depolarization results in activation of biosynthetic enzymes – heme-oxygenase 2 is activated by phosphorylation by casein kinase 2.\(^{158,159}\) The effects of lactate and CO on homeostasis of the CNS neurotransmitters continue to be studied. Lactate serves as a signal to neurons to release more noradrenaline and CO-mediated facilitation of CNS cell interactions affects neural circuits.\(^{160-162}\) The interaction between lactate and CO in affecting these responses is unknown and need to be studied.

**Ion homeostasis**

**Ca\(^{2+}\)**

CO affects Ca\(^{2+}\) signaling (necessary for synaptic transmission, gene transcription, CNS excitation, and memory storage) and may control the switch between glucose and lactate utilization during CNS activation, synaptic activity, and with neuropathology. Maintenance of normal CNS function requires regulation of calcium homeostasis – CO regulation of calcium channels (calcium activated K\(^{+}\) – distributed in both excitable and non-excitatory cells, highly sensitive to intracellular calcium concentrations and voltage, and able to decrease voltage-dependent Ca\(^{2+}\) entry through membrane hyperpolarization and serve as negative feedback regulators – and Ca\(^{2+}\) channel families) is necessary to normal CNS function. The mechanisms by which CO regulates the calcium channels are unclear, remain controversial, and require further study. Telezhkin et al.\(^{163}\) found that cysteine residue 911 in the \(\sigma_{1}\) terminal tail of human BKCa\(_{\alpha}\) subunit is important for activation by CO. Bak et al.\(^{164}\) proposed a compartmentalized Ca\(^{2+}\)-induced limitation of the malate-aspartate shuttle that defines pre- and post-synaptic compartments metabolizing glucose and glucose plus lactate in which the latter displays a positive correlation between oxidative metabolism of glucose and Ca\(^{2+}\) signaling. Lactate is essential to calcium homeostasis as well. Takata and colleagues\(^{165}\) determined that support of synaptic function by lactate utilization is associated with activation of NMDA and L-type calcium channels. Studies to determine the relationship between lactate and CO on calcium homeostasis are needed.

**Cl\(^{-}\)**

Chloride (Cl\(^{-}\)) homeostasis involves multiple complex interactions and is critical for synaptic inhibition, volume regulation, and determines effects of glycine-mediated and GABA neurotransmission.\(^{166}\) Absence/presence and function of chloride transporters and channels (differ in brain cell types) affect direction of Cl\(^{-}\) fluxes and cell physiology.

GABA, an excitatory neurotransmitter during CNS development is inhibitory in the adult. The switching of GABA from excitatory to inhibitory has been correlated with the intracellular concentration of Cl\(^{-}\) – in early development intracellular Cl\(^{-}\) is higher than extracellular concentrations, but in adults intracellular Cl\(^{-}\) is lower than extracellular Cl\(^{-}\). When GABA binds to GABA\(_{A}\) receptors, the Cl\(^{-}\) ion channel opens, Cl\(^{-}\) influxes, and postsynaptic membrane hyperpolarizes with inhibition of the Ca\(^{2+}\) influx of the postsynaptic neuron.\(^{167}\) The role switch is related to the expression of cation/chloride co-transporters NKCC1 and KCC2 and parallels the maturation of the CNS. Inhibition by GABA depends on Cl\(^{-}\) influx and its continuation on Cl\(^{-}\) extrusion from the cell (electrochemical Cl\(^{-}\) gradient does not get depleted) – in the adult CNS, KCC2 is crucial for the inhibitory effects of GABA (and glycine) in synaptic circuits.\(^{168}\) Cation/chloride co-transporters have fundamental roles in differentiation, disease, neuronal proliferation, damage, recovery, and functions associated with plasmalemmal receptor- and channel-mediated effects (i.e., synaptic plasticity). These ion transporters depend on the electrochemical force of Na\(^{+}\) and K\(^{+}\) ions which interact with the Na\(^{+}\)-K\(^{+}\) ATPase and its modulators – regulation of NKCC1 and KCC2 is by insulin-like growth factor, brain-derived neurotrophic factor, and cystic fibrosis transmembrane conductance regulator, and by phosphorylation of enzymes.\(^{169}\) Structure and mechanism of these two cation/chloride co-transporters determine electroneutral transport of sodium, chloride, and/or potassium across membranes and, thereby, impact ion absorption and secretion, maintenance of ion homeostasis, and regulation of cell volume – ion-translocation pathways, key residues for transport activity, and ion-binding sites have been defined.\(^{170,171}\) These ion transporters determine GABA actions – lactate has been associated with GABA release and CO (as an excitatory molecule) effects in the hypothalamic supraoptic nucleus are dependent on GABA.\(^{172,173}\) These relationships require further studies defining interactions between Cl\(^{-}\), GABA, CO, and lactate on health and disease of the CNS.

**K\(^{+}\)**

Four main ions fluctuate with rhythmic firing in the CNS. These include Ca\(^{2+}\), Na\(^{+}\), Cl\(^{-}\), and K\(^{+}\). Synergy between these ions results in an oscillatory activity. Astrocytes regulate neuronal firing by affecting extracellular concentrations.
Extracellular K⁺ levels in the CNS are low relative to intracellular levels and the gradient is important for synaptic communication and electrical signaling (with neuronal activity, a transient rise in extracellular levels is seen). If homeostasis is not maintained, wide-spread depolarization of CNS cells can occur and compromise normal synapses, neuronal firing, and uptake of neurotransmitters.¹⁷⁴⁻¹⁷⁷

Neuronal discharge results in the activation of the voltage-gated potassium channels with K⁺ moving to the extracellular space. The accumulating K⁺ is taken up by neighboring astrocytes and redistributed through the syncytium via the activation of gap junctions.¹⁷⁸,¹⁷⁹ In astrocytes, glycogenolysis is needed for the uptake of increased extracellular K⁺.¹⁸⁰,¹⁸¹ Kir channels (seven subfamilies with each having multiple members) and the Na⁺-K⁺ pump are required for regulation of the extracellular K⁺.¹⁷⁴ Kir channels establish and regulate the resting membrane potential of excitable cells and, in astrocytes, affect potassium buffering (and, therefore, neuronal excitability).¹⁸² In the CNS, Kir4.1 is a modulator of brain-derived neurotrophic factor. Dysfunction of astrocytic Kir4.1 channels facilitates its expression in astrocytes (activating the Ras/Raf/mitogen-activated protein kinase kinase/extracellular signal-regulated kinase pathway) and may be linked to psychiatric disorders.¹⁸³,¹⁸⁴

The relationship of CO and lactate to Kir channels and possibly brain-derived neurotrophic factor in the CNS has not been defined. CO, however, inhibits Kir channels in cardiomyocytes as does H₂S.¹⁸⁵,¹⁸⁶ The effects of gasotransmitters on Kir channels in the CNS have not been studied. Studies are needed to evaluate CO effects on Kir channels and brain-derived neurotrophic factor in the brain and the role of lactate in these metabolic processes (glycogenolysis is important).

**Organ**

**Effects on the lymphatic system**

CNS immune privilege and drainage are affected by the relationship between the CNS lymphatic system (system that allows CNS perfusion by the CSF and ISF) and meningeal lymphatics (major role in drainage of ISF, CSF, CNS-derived molecules, and immune cells from CNS and meninges to peripheral lymph nodes). Meningeal lymphatic vessels, also important in exchange of soluble contents between the CSF and ISF, are keys to removing cellular waste, solutes, and immune traffic from the CNS.⁹⁷,¹⁸⁷,¹⁸⁸ Connection between the two systems allows continuous flow of CSF that perfuses the brain to be drained to peripheral nodes.⁹⁶ Lactate is partially cleared from the CNS by these two systems. While heme oxygenase is involved in subarachnoid hemorrhage and its clearance through meningeal lymphatics,¹⁹⁰ interactions between lactate clearance, CO, and the CNS lymphatic systems have not been defined and requires study.

**Effects on the BBB**

BBB integrity is a prerequisite of normal CNS health. Its disruption, as with stroke, is associated with cerebral edema and neurologic deficiency.¹⁹⁰,¹⁹¹ This disruption has been shown to be alleviated by CO-releasing molecule-3, which reduces Evans blue leakage and brain edema, increases expression of platelet derived growth factor receptor β, Zonula occludens-1, and laminin, decreases the expression of matrix metalloprotease-9, thus protecting pericytes, tight junction proteins, and matrix proteins.¹⁹² The heme oxygenase-1 pathways are also important to the migration and differentiation of endothelial progenitor cells important to BBB integrity.¹⁹³,¹⁹⁴ Lactate also affects barriergenesis and functioning of BBB.¹⁹⁵ Interactions between CO and lactate on the BBB have not been defined and need to be studied.

**Cellular and network**

**Synaptic plasticity**

Synaptic plasticity is important for maintaining homeostasis in the body, for sharing essential information among neurons, and for regulating synaptic transmission or electrical signal transduction to neuronal networks. Upregulation or downregulation of CO affects this plasticity and, therefore, neuropathologies.¹⁹⁵,¹⁹⁶ Lactate is also a signaling molecule in synaptic plasticity – L-lactate stimulates the expression of synaptic plasticity-related genes such as Arc, c-Fos, and Zif268 through a mechanism involving NMDA receptor activity and its downstream signaling cascade extracellular signal-regulated kinase 1/2.¹²⁹,¹⁹⁷,¹⁹⁸ Inhibition of the latter increases synaptic vesicle exocytosis by increasing calcium influx through L-type calcium channels.¹⁹⁹ Downregulation of mitogen-activated protein kinase kinase/extracellular signal-regulated kinase 1/2 signaling pathways by CO has been suggested as a possible mechanism.²⁰⁰,²⁰¹ The interactions between CO and lactate and their impact on synaptic plasticity require further study.

**Synaptogenesis/maintenance/elimination**

Changes in brain structure and organization as we experience, learn, and adapt is neuroplasticity (continues throughout life). Connections within the brain are constantly becoming stronger or weaker. This intricately regulated and multi-step process is defined by neuronal synapse formation/maturation and depends on key regulators, occurs simultaneously in different brain regions and between different types of neurons (a single neuron receives thousands of synaptic inputs), and are affected by extrinsic input, neuronal activity, intrinsic signaling pathways, and sensory experience.

Changes in synaptic or neuroplasticity are associated with multiple neuropsychiatric conditions. CO is important to restoring neuroplasticity in the CNS and in structuring synaptic plasticity and is being evaluated as a neurotherapeutic.¹⁹⁴,¹⁹⁵ The biochemical changes that occur with CO is not well defined and being further evaluated. Lactate is a regulator of synaptic genes and affects synaptogenesis, maintenance, and elimination.⁷⁰,¹⁹⁴,²⁰² The relationship between CO and lactate on synaptogenesis/maintenance/elimination has not been defined.

**Neurogenesis**

The process by which new neurons are formed in the brain, neurogenesis, is crucial when an embryo is developing and continues in certain brain regions after birth and throughout our lifespan. Diversity of neurons results from regulated neurogenesis during embryonic development – neural stem cells (complex identity/functions ranging between quiescence/activation/intermediary subtypes) differentiate and become a
specialized cell type at specific times and regions in the CNS and are affected by external factors. Mitochondrial dynamics are now known to be important during this process and neural stem cell fate decisions – mitochondria-regulated energetics is linked to neuronal development.

Integral to this process, mitochondria are central to determining replication and differentiation of these precursor cells in the CNS. Transition to a more differentiated cell type is accompanied by downregulation of glycolysis and fatty acid oxidation pathways and mitochondrial biogenesis. Lactate levels affect neurogenesis as does the presence of CO. Studies are needed to further evaluate the interactions of lactate and CO on neurogenesis (and effects on neuropathologies).

Neuronal development and guidance
The human brain contains more than $1 \times 10^{11}$ neurons and each neuron connects to other neurons – axons and dendrites grow out, interact with other cells, and selectively protrude to create a functional, integrated, coordinated, working network. Initiated from the late embryonic stage and defining neuronal progenitor differentiation, neuronal development is determined by axon targeting/growth, dendrite maturation, synapse formation requiring integration of intrinsic genetic programs and the extrinsic factors, and chemoaffinity. Synaptic connections, continuously formed and eliminated, determine neural circuit plasticity.

Pruning of dendrites and axons to mature neural circuitry is necessary for normal CNS health. Within the first 2 years of life (in humans), more than half of the connections formed in utero are eliminated. Pruned neurites are important for pre-remodeling networks, may serve as pioneering neurons that can guide other neurons, eliminate initially formed extensive connections made in the earlier wiring process, eliminate excessive branches, and simplify developmental programs.

Microglia (constitute up to 10–15% of all cells in mammalian CNS) are associated with synapse elimination and influence synaptic strength – neuronal activity can directly activate microglia (detect functional state of neuron.) As the CNS matures, microglia localize in different regions throughout the brain. This is thought to be due to the microenvironment and functional differences – classical activation and alternative activation related to anti-inflammatory reactions and tissue remodeling.

Microglia monitor CNS parenchyma and affect clearance of cells in the healthy brain, neuronal excitability, neurogenesis, and synaptic activity. Phenotype can change from surveilling to pro- or anti-inflammatory in response to conditions. Upregulation of specific pathways based on stimuli can alter mitochondrial metabolism – changes in metabolism are seen with activation of microglia. CO affects microglia respiration and depends on the concentration. A CO-releasing molecule 3 affects microglia activation. Microglial metabolism (including lactate metabolism) is affected with neuroinflammation. Interaction between these two systems with pruning and neurodegenerative diseases has not been defined.

Defining the architecture of the CNS
Brain architecture continues to develop over time and is an ongoing process that begins before birth and continues into adulthood. It is comprised of billions of connections between individual neurons across different areas of the brain – simpler neural connections and skills form first followed by more complex circuits and skills – and is dependent on the interaction of genes and experiences. Whole brain volume changes throughout life – growth occurs during childhood and into adolescence, a gradual volume decrease is then seen and possible more growth or no tissue loss between around 18 and 35 years of age, followed by volume loss which increases with age. Brain reference atlases to identify the complex anatomical architecture of the brain help to link structure to function.

After birth, connections continue to be modulated based on environment and experiences with genes affecting the building of the different cellular membranes and defining the molecules and ions that are used for the functioning CNS. Synapses and microcircuits as well as neurons, local circuits, and systems/ pathways are affected. Experiences are crucial to the development of brain architecture and affect genes – genes can be turned on or off.

Ability to change the architecture of the brain in response to experiences decreases over time. Anatomical architecture and structural/functional networks affect functional interactions in the brain – changes of structural connectivity are paralleled by changes in functional connectivity and characterize brain changes in aging. Rise in brain lactate is seen with aging of the brain - the mechanism varies with different brain regions (LDH status is important.) The change architecture of the aging brain is under oxidative stress – heme oxygenase genes are induced. CO (products of reaction catalyzed by heme oxygenase) has profound effects on the CNS. The roles and interactions of lactate and CO on changing CNS architecture needs further study.

Systemic level
Effects on sleep
Lactate is a biomarker of sleeping – lactate concentrations remain higher during waking relative to sleep and with forced sleep deprivation, increase with waking, and decrease within minutes of sleep. During sleep, there is decreased ISF lactate concentration, decreased production, and increased perivascular clearance of lactate. Activation of the glymphatic system with change-of-state is associated with lowering CNS lactate during sleep. This changes the lactate/pyruvate ratio (and, therefore, the NADH/NAD redox potential). In those patients with obstructive sleep apnea, exhaled CO correlates with hypoxia during sleep. Postsleep circulating CO level is increased initially, and venous lactate is increased (improved with continuous positive airway pressure). The proinflammatory transcription factor nuclear factor kappaB (affected by lactate and CO) is also increased. Interactions between CO, lactate, and nuclear factor kappaB during sleep and in obstructive sleep apnea need further studies.

Regulation of energy balance
The brain is integral to maintaining energy balance – the hypothalamic nuclei and the brainstem, main brain areas for energy balance regulation, are affected by nonhomeostatic and
homeostatic circuits (interrelated and integrated). Neural circuits in the CNS process information regarding food consumption, food, food-related cues, and feeding and energy balance are associated with higher brain functions and degenerative processes. 240–242 5’ adenosine monophosphate-activated protein kinase (AMPK), metabolic regulator of the entire organism, senses AMP:ATP and/or adenosine diphosphate/ATP ratios and maintains energy balance by promoting ATP production by increasing the activity/expression of proteins involved in catabolism and conserving ATP by switching off biosynthetic pathways. Selective activation of hypothalamic AMPK negates hypothalamic glucose/lactate-sensing mechanisms. 243 Activated AMPK also stimulates heme oxygenase-1 gene expression and elevates the nuclear factor erythroid 2-related factor 2/heme oxygenase-1 signaling axis. 244,245 Further studies are needed to define the relationships between AMPK, lactate, and CO associated with regulation of energy balance.

**Chemosensing**

Chemosensing (generation of a response to the presence of a chemical stimulus) is a fundamental sensory function and depends on a complex combination of receptor-mediated transmembrane signals, lipid modifications, protein translocations, and differential activation/deactivation of membrane-bound and cytosolic components. Some are mediated by somatosensory nerve endings of the trigeminal, glossopharyngeal, and vagus nerves – chemicals activate chemosensory receptors via mechanisms distinct from other sensory systems and the nerve endings can trigger a range of sensations and reflexes. Overlapping sensitivity of individual ion channels to chemical agonist can result in combinatorial activation of different classes of nerve endings by a single chemical. Complex interplay between chemical agonists and chemosensory receptors allow for chemosensory mimicry by chemically distinct agents. Multiple areas of the CNS are involved in eliciting a response to a particular chemical stimulus.

Chemosensory mechanisms may be an important causative factor in some neurologic diseases, but studies to evaluate the roles of lactate and/or CO in chemosensing are lacking. However, lactate infusion is a provocative challenge for panic disorder and is thought to be related to chemosensory mechanisms. 246–248 CO effects on this are unknown. Studies to evaluate lactate and CO interactions on chemosensing are needed.

**Summary**

Lactate is no longer considered “just a waste product of metabolism.” Overall and CNS health are affected by its presence or absence and its biochemical properties. CO is now known to be critical to the production of lactate and its roles (as a fuel, hormone, and/or signaling molecule) in the CNS. The neuroprotective ability of L-lactate partially depends on the integrity of the heme-oxygenase system and, in particular, CO. Further evaluation is needed of the L-lactate/heme-oxygenase/CO interactions to better understand the effects on its application in CNS health and neuropathologies.

**Author contributions**

VLM developed idea, researched and wrote entire manuscript. The final version of manuscript has been read and approved by the author.

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