Lessons From Heat Stroke for Understanding Myalgic Encephalomyelitis/Chronic Fatigue Syndrome

Dominic Stanculescu 1, Nuno Sepúlveda 2,3, Chin Leong Lim 4 and Jonas Bergquist 5,6*

1 Independent Researcher, Sint Martens Latem, Belgium, 2 CEAUL—Centro de Estatística e Aplicações da Universidade de Lisboa, Lisbon, Portugal, 3 Department of Mathematics and Information Science, Warsaw University of Technology, Warsaw, Poland, 4 Lee Kong Chian School of Medicine, Nanyang Technological University, Singapore, Singapore, 5 Analytical Chemistry and Neurochemistry, Department of Chemistry—BMC, Uppsala University, Uppsala, Sweden, 6 The ME/CFS Collaborative Research Center at Uppsala University, Uppsala, Sweden

We here provide an overview of the pathophysiological mechanisms during heat stroke and describe similar mechanisms found in myalgic encephalomyelitis/chronic fatigue syndrome (ME/CFS). Both conditions are characterized by disturbed homeostasis in which inflammatory pathways play a central role. Splanchnic vasoconstriction, increased gut permeability, gut-related endotoxemia, systemic inflammatory response, central nervous system dysfunction, blood coagulation disorder, endothelial-cell injury, and mitochondrial dysfunction underlie heat stroke. These mechanisms have also been documented in ME/CFS. Moreover, initial transcriptomic studies suggest that similar gene expressions are altered in both heat stroke and ME/CFS. Finally, some predisposing factors for heat stroke, such as pre-existing inflammation or infection, overlap with those for ME/CFS. Notwithstanding important differences - and despite heat stroke being an acute condition - the overlaps between heat stroke and ME/CFS suggest common pathways in the physiological responses to very different forms of stressors, which are manifested in different clinical outcomes. The human studies and animal models of heat stroke provide an explanation for the self-perpetuation of homeostatic imbalance centered around intestinal wall injury, which could also inform the understanding of ME/CFS. Moreover, the studies of novel therapeutics for heat stroke might provide new avenues for the treatment of ME/CFS. Future research should be conducted to investigate the similarities between heat stroke and ME/CFS to help identify the potential treatments for ME/CFS.

Keywords: post-viral fatigue, myalgic encephalomyelitis/chronic fatigue syndrome, gut permeability, systemic inflammatory response syndrome (SIRS), heat stroke, endotoxemia, ME/CFS, splanchnic vasoconstriction

INTRODUCTION

Heat stroke results from a failure of the body to maintain its normal core temperature when exposed to a high environmental temperature (i.e., “classic heat stroke”) or from strenuous exercise (i.e., “exertional heat stroke”) (1). There is no universally accepted definition of heat stroke. Based on the pathophysiology of heat stroke, Bouchama and Knochel (1) proposed to define heat stroke as “a form of hyperthermia associated with a systemic inflammatory response leading to a syndrome of multorgan dysfunction in which encephalopathy predominates.” Heat stroke is often clinically defined as “a core body temperature that rises above 40°C and that is accompanied by hot, dry skin
and central nervous system abnormalities such as delirium, convulsions, or coma” (1, 2). However, given the occurrence of multiple organ dysfunction in heat stroke victims, the Japanese Association for Acute Medicine (JAAM) recommended heatstroke clinical criteria to include “the presence of renal and hepatic complications and disseminated intravascular coagulation (DIC)” (3). The JAAM criteria do not include temperature “because elderly people often do not present with high temperature even when they are already suffering from heat stroke” (3). Heat stroke can lead to organ damage or death, and the hospital mortality rate in classic heat stroke ranges from 10% to 65%, whereas that of exertional heat stroke is 3% to 5% (4). An average of 702 heat-related deaths occurred in the United States annually during 2004–2018 (5) (we assume representing mostly cases of classic heat stroke). “Prompt recognition and effective body cooling will in most cases rapidly reverse heat-induced organ dysfunction. However, body cooling alone may not suffice to effect a full recovery, and prompt administration of adjuvant treatments may be critical for survival” (4). Several novel treatments are currently being explored for heat stroke, notably to address gut permeability and to reduce systemic inflammatory markers (4).

Myalgic encephalomyelitis/chronic fatigue syndrome (ME/CFS) is a debilitating multi-system disease of unclear etiology (6–10). The most common peri-onset events reported by patients are infection-related episodes, stressful incidents, and exposure to environmental toxins (11). “Impaired function, post-exertional malaise, and unrefreshing sleep” are considered to be the core symptoms (12, 13). Post-exertional malaise refers to an exacerbation of some or all of an individual’s ME/CFS symptoms after physical or cognitive exertion, or orthostatic stress that leads to a reduction in functional ability. The severity of the symptoms varies, where “very severely affected patients experience profound weakness, almost constant pain, severe limitations to physical and mental activity, sensory hypersensitivity (light, touch, sound, smell, and certain foods), and hypersensitivity to medications” (14, 15). The illness can be completely incapacitating (16) and “at least one-quarter of ME/CFS patients are house- or bedbound at some point in their lives” (12). Patients with milder symptoms experience a significant reduction in their previous level of functioning (8). The illness progresses over time, shifting from an early hypermetabolic state to a hypometabolic state with low energy production (17). ME/CFS is often dismissed by physicians because standard urine and blood laboratory tests produce normal results, thereby multiplying patients’ suffering (18). Deaths from ME/CFS are rare (19), however, the prevalence of the disease is estimated to be at least 1.5 million cases in the United States alone (20). There are currently no effective treatments for ME/CFS (21–25).

In the following sections, we provide an overview of key pathophysiological mechanisms in heat stroke, such as splanchnic vasconstriction, increased gut permeability, gut-related endotoxemia, systemic inflammatory response, central nervous system dysfunction, blood coagulation disorder, endothelial-cell injury, and mitochondrial dysfunction - and relate these mechanisms to similar findings from ME/CFS research. In addition, we also describe the results of initial heat stress gene transcriptome studies and the parallel findings from ME/CFS studies. Finally, we summarize key predisposing factors for both illnesses.

The central thesis of this paper is that the lessons from the human studies and animal models of heat stroke can contribute to the understanding and treatment of ME/CFS. Although this thesis might seem unlikely given the important differences in the onset and clinical outcomes between these two conditions, it is derived from a broader hypothesis that supports the notion of common pathways in the physiological responses to very different forms of stressors and disease manifestations (26–30). The paper serves as an element of a wider enquiry into commonalities in the pathology of illnesses induced by physical, infectious, and/or emotional stressors including heat stroke, ME/CFS, prolonged critical illness (chronic intensive care unit [ICU]), post intensive care syndrome (PICS), cancer-related fatigue, post-viral fatigue, post-acute coronavirus disease 2019 [COVID-19] syndrome (PACS), and long-COVID-19. The paper is in line with a history of research comparing ME/CFS with other conditions (including fibromyalgia, multiple sclerosis, and rheumatoid arthritis) to derive lessons for understanding ME/CFS.

**PATHOPHYSIOLOGICAL MECHANISMS**

Since the 1990s, human and animal model studies have shown that endotoxemia of intestinal origin (i.e., the presence of endotoxins leaked from the intestines into circulation) may play an important role in the pathophysiological mechanisms of heat stroke (1, 2, 31–33). This evidence culminated in the dual pathway model of heat stroke, which proposes that heat stroke can be triggered by two sequential but independent pathways (31, 34): (i) the heat toxicity pathway, referring to cytotoxicity resulting directly from hyperthermia (e.g., protein denaturation, organ damage, and interruption of critical cellular processes because of heat), and (ii) the endotoxemia pathway (or heat sepsis pathway), involving splanchnic vasconstriction, gut hyper-permeability (“leaky gut”), and endotoxemia culminating in systemic inflammation (or sepsis) (Figure 1). Whereas heat toxicity was traditionally considered the primary determinant of morbidity and mortality in heat stroke, it is now increasingly recognized that this pathway occurs only at temperatures above 42°C. Conversely, the endotoxemia pathway can provoke heat stroke at temperatures between 40 and 42°C (31, 33). In other words, the endotoxemia pathway precedes and acts largely independently of the heat toxicity pathway in the pathology.

**Abbreviations:** CMV, Cytomegalovirus; DIC, disseminated intravascular coagulation; EBV, Epstein-Barr virus; HSPs, heat shock proteins; IBS, irritable bowel syndrome; ICU, intensive care unit; IL, interleukin; JAAM, Japanese association for acute medicine; LPS, lipopolysaccharides; ME/CFS, myalgic encephalomyelitis/chronic fatigue syndrome; O&NS, oxidative and nitrosative stress; PACE, Post-acute COVID-19 syndrome; PICS, Post-intensive care syndrome; POTS, postural orthostatic tachycardia syndrome; ROS, reactive oxygen species; SIRS, systemic inflammatory response syndrome; TNF, tumor necrosis factor; TLR, toll-like receptor.
of heat stroke (31). Moreover, the endotoxemia pathway is increasingly considered as the lead driver of severe organ damage and the main cause of death in patients suffering from heat stroke (35).

In the following paragraphs, we provide an overview of the mechanisms of the modern paradigm of heat stroke with a focus on the endotoxemia pathway. Moreover, we describe similar findings from ME/CFS. Our focus on the similarities between the two conditions does not intend to negate the disease-specific mechanisms. Instead, it intends to highlight possible lessons from heat stroke for understanding ME/CFS.

**Splanchnic Vasoconstriction and Hypoxia In Heat Stroke**

The primary response of the cardiovascular system to an increase in the ambient heat or heat from exertion is an increase in the blood flow to the skin (i.e., cutaneous vasodilation) to dissipate the excess heat. The modern paradigm of heat stroke suggests that the maintenance of a constant blood flow implies/requires a reduction in blood flow to internal organs (i.e., splanchnic vasoconstriction) (1, 32, 36–38). However, splanchnic hypoperfusion is not unique to heat stroke and can lead to critical illnesses (i.e., severe injury or infection) (39, 40), and may thus be a response to severe stress in general. In any case, the resulting hypoxia has important consequences on the gut and brain during heat stroke (as shown below) and promotes liver damage. In addition, the strain on the cardiovascular system often results in syncope and orthostatic intolerance in heat stroke (36).

**In ME/CFS**

Akin to heat stroke, several new hypotheses of ME/CFS suggest that vasoconstriction and hypoxia are central to its pathology—respectively focused on the consequence of β2 adrenergic and M3 acetylcholine receptor autoantibodies (41, 42), a potential reduction of ACE2 expression (43), and an abnormal immune response with a concomitant impact on endothelial function (44). In addition, postural orthostatic tachycardia syndrome (POTS) is a common comorbidity of ME/CFS (45), which is a complex disorder of the vascular system also associated with gastrointestinal dysfunctions (46, 47). Impaired blood pressure variability and arrhythmia have been observed in the patients with ME/CFS (48, 49).

**In summary,** vasoconstriction and hypoxia would appear to be common to both heat stroke in at least a subset of patients with ME/CFS—despite the difference in onset events of these two conditions. Research from critical illnesses (beyond heat stroke) suggests that splanchnic vasoconstriction is a general response to stressors. Vasoconstriction and hypoxia have far-reaching implications for organs and tissues.

**Gut Permeability and Endotoxemia In Heat Stroke**

Animal models of heat stroke have shown that ischemia resulting from reduced blood flow to the gut causes mucosal damage and a weakening of the tight junctions in the gut—likely via the action of oxidative and nitrosative stress (O&NS) (1, 4, 31–33, 35–38, 50, 51)—as well ATP depletion and local acidosis (50) induced by hypoxia. The result is a leaky gut characterized by a reduction of intestinal immunity and increased permeability of the intestines (1). This condition promotes gram-negative bacteria (i.e., lipopolysaccharides, LPS) that are normally contained in the gut to enter blood circulation (1, 4, 31–33, 35–38, 50, 52). Studies have shown that in the case of heat stroke, the liver is unable to effectively clear endotoxin (31), notably because of poor circulation in the venous system that returns blood from the digestive tract and spleen to the liver (37, 50). The resulting endotoxemia is assumed to trigger a systemic inflammatory response syndrome (SIRS) in heat stroke (as shown below) (4, 31–33, 35, 36, 53). Furthermore, findings from critical illness suggest that additional forms of intestinal injury play a role in driving critical illness, including the erosion of the mucus barrier, a shift in the composition and virulence of intestinal microbes, and the inability of the host epithelium to regulate its proliferative and apoptotic response (39, 54). These elements are thought to contribute to a vicious cycle of inflammation and intestinal injury. Moreover, the reduction in gut motility, the formation of toxic gut-derived lymph, and a decrease in the secretion of gastrointestinal hormones could contribute to multi-organ dysfunction in critical illness (55, 56).

Critically, normalization of the core temperature during heat stroke through cooling does not necessarily imply a reversion to normal gut permeability or the end of endotoxemia (1). Some researchers have thus suggested and tried to repair gut function to treat heat stroke (38). This includes the administration of nutritional supplements, such as glutamine and bovine colostrum with some promising results (33, 50, 57, 58). Additionally, the use of prebiotics and probiotics is being explored as an avenue for treatment (38, 50). Others have had success in limiting intestinal injury with the use of 4-phenylbutyrate to interrupt the endoplasmic reticulum stress-mediated apoptosis pathway of intestinal cells (59). Similarly, others have studied the use of xanthine oxidase inhibitor (allopurinol) to protect the integrity of tight cell-to-cell junctions in the gut of rat models (51). The inhibition of endotoxemia by pre-treatment with LPS-specific antibodies and corticosteroids to counteract LPS was shown to protect animals from lethal heat stress (31, 60). Readers are referred to the reviews for details on the results of human and animal studies (35, 38, 50, 61).

**In ME/CFS**

Echoing the endotoxemia model of heat stroke, ME/CFS researchers have hypothesized that the translocation of gut microbes or toxins from the gastrointestinal tract and the resulting inflammation is central to ME/CFS (62–66). Studies have found gut hyper-permeability (63) and gut inflammation (67) in ME/CFS as well as increased levels of the antibodies to LPS (IgM and IgA) in serum of patients with ME/CFS (the prevalence of patients with CFS with abnormally increased IgM and IgA levels: 40% and 66.7%) (68). Moreover, researchers have induced fatigue in mice through the administration of LPS (69). Finally, studies have shown that the normalization of leaky gut in ME/CFS by administration of natural anti-inflammatory and antioxidative substances is accompanied by a
clinical improvement (70). Readers are referred to the reviews for a detailed discussion on the implication of gut permeability on the immune system and mitochondrial function in ME/CFS (7, 71).

It is worth noting that leaky gut is distinct from irritable bowel syndrome (IBS), although the former is believed to contribute to the latter (72). Leaky gut is related to changes in the permeability of the tight junctions in the gut barrier, which can be observed in the absence of symptoms (73). In research, changes in the gut barrier have been measured in urine using sugar probes of different molecular sizes, and in the circulation, using intestinal fatty acid binding protein (iFAB), and claudine-3 (74, 75). iFAB is an intracellular protein in the intestine wall and supports the transportation of fatty acid across the gut barrier. Claudine-3 is located between the tight junctions and functions as a “glue” to preserve the integrity of the gut barrier. Leaky gut is associated with diseases beyond the gut, such as diabetes (76) and autoimmune diseases (77, 78). Conversely, IBS is related to gut dysbiosis and is primarily associated with gastrointestinal discomfort (79). Whereas IBS is widely recognized as a clinical entity by the medical profession, leaky gut is largely considered the domain of alternative medicine (80).

In summary, gut hyper-permeability and endotoxemia would appear to be common to both heat stroke and at least a subset of patients with ME/CFS patients—despite the difference in onset events of these two conditions. These mechanisms have far-reaching implications on various physiological systems, including inflammation (as shown below). The results of human and animal experiments to repair gut function and mitigate endotoxemia in heat stroke might inform and provide new therapeutic avenues for ME/CFS.

**Systemic Inflammatory Response In Heat Stroke**

Bouchama and Knochel suggest that the inflammatory response during the initial “acute-phase of heat stress” is mediated primarily by interleukin-6 (IL-6) and other pro-inflammatory cytokines, which serve to protect against tissue injury and to promote repair. Indeed, IL-6 and other cytokines induced by exposure to heat stress appear to mediate fever, leukocytosis, increased synthesis of acute-phase proteins, muscle catabolism, stimulation of the hypothalamic–pituitary–adrenal axis, and the activation of leukocytes and endothelial cells (1). However, an exaggeration of the acute-phase response can precipitate a maladaptive progression from heat stress to heat stroke. Specifically, leaky gut (as described above) hastens local inflammation toward the “exaggerated” systemic inflammatory response that underpins heat stroke (1). The field of exercise immunology has demonstrated that LPS leaking from the gut activates toll-like receptor (TLR) 4 on immune and intestinal epithelial cell membranes. This activation results in the production of pro-inflammatory cytokines, e.g., IL-6, IL-1β, and tumor necrosis factor (TNF)-α (31, 33, 35, 50). Moreover, the animal models of heat stroke have shown that, when LPS is present, NLRP3 (NOD-, LRR-, and pyrin domain-containing protein 3) inflammasomes are activated and...
drive neuroinflammation (81). A vicious cycle centered around intestinal injury ensues in heat stroke involving the release of nitric oxide, the leakage of gut-related endotoxins, and increased production of inflammatory cytokines (e.g., TNF-α, IL-1β, and IFN-γ) (35). Anti-inflammatory cytokines (IL-10, IL-1 receptor antagonist (IL-1ra), and TNF receptors p55 and p75) are also elevated, but to a lesser degree (2). The mode of action of cytokines in a heat stroke remains to be fully understood (33, 36). However, the increase in inflammatory cytokine levels is associated with inflammation-associated injury and refractory immunosuppression (1, 36) and can induce fever when transported to the brain (1, 82). Mouse models indicate that glia cells are a likely source of cytokine and chemokine production (36, 83).

Critically, normalization of the core temperature does not necessarily inhibit the systemic inflammatory response in heat stroke (1). Researchers have thus suggested and tried the modulation of the inflammatory response as a form of heat stroke treatment and prophylaxis, including with the administration of IL-1 receptor antagonists (81, 84), corticosteroids (60, 85), activated protein C (86), and the inhibition of nuclear factor-κB activity (1). Others have used oleoyl ethanolamide to inhibit LPS-induced TLR-4 stimulation of the immune response (50, 87). Blood purification therapy (53) and bone marrow-derived mononuclear cell therapy (88) to reduce pro-inflammatory cytokines have been suggested or tested in clinical trials. Other clinical trials have tested the efficacy of antioxidants and anti-inflammatory pathways, such as quercetin (89). One study found that parental administration of ascorbate attenuated heat stroke–induced systemic inflammation in mice (90). Readers are referred to the reviews for details on the results of human and animal heat stroke studies (35, 50).

In ME/CFS
Akin to heat stroke, researchers have found that chronic inflammation also underlies ME/CFS (91–95), including inflammation of the brain—hence the name myalgic encephalomyelitis (96, 97). Moreover, similarly to heat stroke, O&NS has been documented in ME/CFS (98) and the cytokine profile of patients with ME/CFS differs from healthy controls (99, 100). Researchers have also linked NLRP3 inflammasomes with inflammation in ME/CFS (101). Some researchers have theorized that the complex pathways between these elements (O&NS, cytokines, and inflammation) underpin the metabolic dysfunction characteristic of the disease (102), including by hindering mitochondrial (103) and hypothalamic function (104, 105). Significantly, researchers suggest “the possibility of a vicious cycle of immune-disregulation and gut dysbiosis accompanied by poor physiological gut function” perpetuating a disease state in ME/CFS (7)—as is the case for heat stroke. Readers are referred to the numerous studies and reviews for a detailed discussion on the relationship between redox imbalance, cytokines, and inflammatory pathways in ME/CFS (7, 101, 106, 107).

In summary, inflammation associated with cytokines and O&NS is central to both heat stroke and ME/CFS—despite the difference in onset events of these two conditions. Moreover, a vicious inflammatory cycle involving altered gut-blood barrier function (leaky gut) may be a central element of both heat stroke and some patients with ME/CFS. The results of human and animal experiments to modulate the inflammatory response in heat stroke might inform and provide new therapeutic avenues for ME/CFS.

Central Nervous System Dysfunction In Heat Stroke
Delirium, convulsions, and coma are common symptoms in the clinical presentation of heat stroke. The mechanisms of central nervous system injury in heat stroke are not well understood but are likely due to the combination of ischemia, hypoxia, thermolysis (Tc > 42°C), and hemorrhage (108–110). The animal models of heat stroke identified ischemia resulting from decreased cerebral flow and increased intracranial pressure as causing extensive cerebral neuronal injury (36). Moreover, a decrease in brain blood supply and an increase in intracranial pressure are followed by a compromised blood–brain barrier and an increase in circulatory pyrogens (37). In other words, analogous to the consequences of hypo-perfusion in the gut, reduced blood flow to the brain can lead to hypoxia, inflammation, and hyper-permeability within the brain, facilitating protein and pathogen leakage from the systemic circulation into the brain (32, 36, 37, 111). The severity of this mechanism in heat stroke can lead to brain injury.

In ME/CFS
Central nervous system dysfunctions are present in ME/CFS, albeit in a different form. Brain fog and hypersensitivity to noise and light are common symptoms in patients with ME/CFS (8, 13). However, the mechanisms in ME/CFS are arguably even less well understood than in heat stroke. Brain imaging studies revealed significant neuroinflammation (112–114) and white matter abnormalities (115), which are confirmed by autopsies (116). As mentioned above, a new unifying hypothesis of ME/CFS suggests that vasocostriction and hypoxia in muscle and brain underpin ME/CFS, including cognitive impairments (41, 42). Studies have shown reduced cerebral blood flow during tilt tests (117). Akin to heat stroke, some patients with ME/CFS present signs of increased intracranial pressure (118, 119), consistent with altered blood flow. Alternatively, others have hypothesized that ME/CFS is promoted by gut microbes or toxin translocation from the gastrointestinal tract into other tissues, including the brain (62). The central nervous system is generally an immune-privileged site (i.e., protected from the peripheral immune system) (120), but pathogens that reach the brain can cause devastating inflammation (121).

In summary, central nervous system dysfunctions are central to both heat stroke and ME/CFS, albeit in different forms and to different degrees. Neuroinflammation, decreased blood flow to the brain and increased intracranial pressure might be common to both heat stroke and some ME/CFS patients. The animal models of heat stroke may inform the understanding of central nervous system dysfunction in ME/CFS.
**Coagulation Disorders and Endothelial-Cell Injury**

**In Heat Stroke**
Both the endotoxemia and the heat toxicity pathways are believed to contribute to coagulopathy dysfunction and endothelial-cell injury in heat stroke. First, intestine-derived endotoxins and cytokines activate vascular endothelial cells and coagulation factors to amplify the coagulation cascade (32, 35), which triggers vascular endothelial cell injury (122, 123). Moreover, the increase in core temperature directly leads to vascular endothelium damage (2), which in turn drives coagulation (32, 36). Bouchama and Knochel (1) describe a complex interplay among the cytotoxic effect of the heat and the inflammatory and coagulation responses of the host. Together these mechanisms result in a positive feedback loop, causing occlusion of the arterioles and capillaries (i.e., microvascular thrombosis) (32, 53, 90, 122). The vascular endothelial cell injury in heat stroke has been substantiated by evidence of differential expression of exosome lncRNAs and miRNAs (122).

Critically, normalization of the core temperature does not necessarily deactivate the coagulation process (1). Researchers have thus suggested anticoagulation therapy (53) with some initial positive results (124). Others have found that dexmedetomidine protects endothelial glycocalyx and increases the survival of rats during heat stroke (125). Readers are referred to the reviews for details on the results of human and animal studies (35, 38, 50).

**In ME/CFS**
Akin to heat stroke, hyper-coagulation was also suggested to be a contributing factor in ME/CFS (126–128), but the findings have not been consistent across the studies (129). The lesser importance or absence of coagulation dysfunctions in ME/CFS compared with heat stroke may be explained by the fact that the “heat toxicity pathway,” one of the drivers of the “positive feedback loop” driving coagulopathy dysfunction in heat stroke, is absent in ME/CFS.

Studies have shown that endothelial function is impaired in ME/CFS, both in large vessels and in microcirculation (130, 131). As in heat stroke, these findings were substantiated by indirect evidence related to the expression of miRNAs relevant to endothelial function (132). A new hypothesis for ME/CFS suggests that endothelial senescence underpins ME/CFS by disrupting the intestinal and blood-brain barriers (62).

In summary, coagulopathy dysfunction and endothelial-cell injury may be common to both heat stroke and at least a subset of patients with ME/CFS, despite the difference in onset events of these two conditions. The results of human and animal experiments to address coagulation disorders and endothelial-cell injury in heat stroke might inform and provide new therapeutic avenues for ME/CFS.

**Mitochondrial Dysfunction**

**In Heat Stroke**
There are only few studies that assess mitochondrial function in heat stroke. ATP depletion (36) as well mitochondrial swelling and vacuolization in the epithelial cells of the gut have been observed in the mice models of heat stroke (52). Moreover, animal studies have shown that apoptosis of vascular endothelial cells in heat stroke was the deleterious consequence of reactive oxygen species (ROS)-induced p53 translocation into mitochondria (133). Finally, it was shown in vitro that higher temperatures can directly affect the viscosity of mitochondria (134).

**In ME/CFS**
Akin to heat stroke, the immuno-inflammatory pathways associated with O&NS are thought to be one of the possible mechanisms driving mitochondrial dysfunction in ME/CFS (103, 107). [Alternatively, mitochondrial dysfunction could be caused by viral reactivation in ME/CFS (135, 136)]. Impaired energy production (97, 137) and reduced mitochondrial activity (138–140) are central to the pathology of ME/CFS (7). Researchers have observed mitochondrial fragmentation (135) and impaired pyruvate dehydrogenase function as well as lactate degradation deficiencies in ME/CFS (141). Moreover, irregularities in the metabolites of patients with ME/CFS (141, 142) suggest that patients experience a hypometabolic or “dauer” state (143) maintained by mitochondria; an evolutionary adaptation to protect the cells and hosts from harm (c.f., “cell damage response”) (30, 144, 145). Readers are referred to the reviews for a detailed discussion on the relationship between the immune system and mitochondrial dysfunction in ME/CFS (7, 71).

In summary, mitochondrial dysfunction may be common to both heat stroke and ME/CFS, despite the difference in onset events of these two conditions. Mitochondrial dysfunction can be driven by immuno-inflammatory and O&NS pathways. The animal models of heat stroke may inform the understanding of mitochondrial dysfunction in some cases of ME/CFS.

**Altered Gene Transcription**

**In Heat Stroke**
A human transcriptome study performed by Bouchama et al. using peripheral blood mononuclear cells (PBMCs) of healthy volunteers exposed to short but extreme environmental heat treatment found “downregulation of most of the genes involved in the respiratory chain […] with a predicted decrease in ATP production and increase in oxidative stress.” The findings suggested that the “shift to glycolysis as a source of energy” and “inhibition of mitochondrial respiration” were sustained at least an hour after the end of the heat treatment. In addition, there was a “downregulation of ubiquitin conjugation and proteasome genes with a predicted decrease of protein refolding, mono- and poly-ubiquitination, and antigen presentation” as well as “decreased expression of the major histocompatibility I and II complexes genes, suggesting reduced antigen presentation” that persisted at least an hour after the end of treatment. To be clear, the volunteers did not experience heat stroke (only heat stress), but nonetheless, the study showed an adaptive response of mitochondria to heat stress - namely, altered protein synthesis, and reduced expression of genes related to immune function - to persist at least an hour after the heat stress ended. The authors...
could not explain why cellular stress response to heat would involve a “switch to a less efficient form of metabolism” (146).

In ME/CFS
Transcriptome studies have generally found altered gene expression in patients with ME/CFS (147–154). Researchers, for example, identified changes in the expression of genes in peripheral immune cells related to inflammation and mitochondrial function (153) as well as to oxidative stress (149) and immune function (149, 150, 154). Others have found specific microRNA expression patterns associated with specific symptom severities (155). Readers are referred to the reviews for a detailed discussion on alterations in gene expression in ME/CFS (95).

In summary, initial gene transcriptome studies would appear to suggest overlaps in alterations of gene expressions between the two conditions. Specifically, alterations in genes related to mitochondrial and immune function may be common to both heat stroke and some patients with ME/CFS. Future transcriptome studies of heat stroke may inform the understanding of ME/CFS.

Predisposing Factors
In Heat Stroke
Young children, the elderly, and patients with chronic diseases are most susceptible to classic heat stroke. Further intrinsic risk factors for classic heat stroke include underlying illness, cardiovascular insufficiency, concurrent infections, and an immune-compromised state (36).

However, the predisposing factors for exertional heat stroke—which affects young and apparently healthy and active individuals—are less clear. Similar to classic heat stroke, researchers have suggested that prior bouts of illness, infection or inflammation, preexisting barrier dysfunction, and a compromised immune system may increase the risk of exertional heat stroke (50, 156, 157). Specifically, prolonged stress exposure and/or suboptimal dietary intake - which suppress the immune system and alter the microbiome of the host, thus depleting its immune and epithelial barrier functions - are thought to be the determining factors in the susceptibility to exertional heat stroke (31, 50). Accordingly, some researchers have suggested that stress and lifestyle factors set the stage for a single event that promotes LPS translocation (such as exertion) to overwhelm the suppressed immune system (31). In the central circulation, LPS can be bound to LPS-specific antibodies, monocytes, or high-density lipoprotein and transported to the liver for removal by the reticuloendothelial system (31, 34). Immune suppression compromises LPS removal by monocytes and LPS-specific antibodies, leading to an increase in the concentration of LPS in the circulation (34, 158) and thus susceptibility to endotoxemia.

In terms of prior infection, a hypothesis derived from exercise immunology research suggests that the immune system is suppressed temporarily immediately post-intense exercise, which provides a window for dormant viruses to be reactivated to cause an infection (c.f. “open window theory”) (159). More specifically, a group of researchers has shown that the increased incidence of illness in response to excessive exercise is not due to immunosuppression per se, but rather to an altered focus of the immune function: an upregulation of humoral immunity and suppression of cell-mediated immunity (160).

Regarding gender bias, a review of 22 studies found that the overall risk for heat illnesses was 2.64 times higher in men than in women, but the authors suggest this could relate to sex-associated behavioral differences rather than physiology (161). Indeed, women are disadvantaged in terms of thermoregulation because of the smaller body surface area (BSA) to mass ratio, a higher percentage of body fat, and lower maximal oxygen consumption (VO_{2max}) (158). A study showed that greater heat intolerance in women relative to men is largely explained by VO_{2max} (162). A review of 24 studies of gender differences in the armed forces suggest that being a female was associated with a greater risk of exertional heat illnesses, but men experienced a slightly higher incidence of exertional heat stroke (163) — perhaps also because of behavioral differences.

Failure to increase the expression of heat shock proteins (HSPs) has been implicated in the pathogenesis of heat stroke (146). HSPs play a central role in the maintenance of endothelial and epithelial barrier function (164, 165). Conditions associated with a low level of expression of HSPs, for instance, aging, lack of acclimatization to heat, and certain genetic polymorphisms, may favor the progression from heat stress to heat stroke (1). Initial studies have shown that selective pharmacologic induction of the expression of HSPs (e.g., with alpha-lipoic acid) can protect cells from apoptosis and inflammatory responses induced by heat stroke (166).

Finally, Bouchama and Knochel suggest that “genetic factors may determine the susceptibility to heat stroke; candidate susceptibility genes include those that encode cytokines, coagulation proteins, and heat-shock proteins involved in the adaptation to heat stress” (1). Specifically, TLR4 polymorphisms may be a genetic factor predisposing to mortality during the systemic inflammatory response to heat stroke (36). However, the importance of genetic factors as a risk for heat stroke remains unresolved.

In ME/CFS
Akin to exertional heat stroke, the predisposing factors for ME/CFS are not clear. Certainly, existing inflammation or infection is likely a risk factor (167). Echoing the “open window theory” of exercise immunology, it has long been suggested that ME/CFS results from bacterial or viral reactivation in the context of a suppressed immune system (168, 169), particularly the reactivation of Epstein-Barr virus (EBV) and cytomegalovirus (CMV) (170, 171). Cumulative life stress is considered a predisposing factor for ME/CFS (172), which is also thought to alter gut permeability and thus immune function (173). As in heat stroke, the microbiome disturbances and dysbiosis are associated with ME/CFS pathology (147, 174–179). Gender is a risk factor for ME/CFS: the female-to-male ratio ranges from 2:1 to 6:1 (12).

Moreover, the studies have shown that ME/CFS is characterized by impaired HSPs production (180) which, combined with O2&NS and low-grade inflammation, could explain muscle dysfunction and exercise intolerance (181, 182). Akin to heat stroke, it has been suggested (but not yet trialed) to
incorporate the upregulation of HSP into future treatments for ME/CFS (183).

Finally, research suggests there may be a genetic element in the pathogenesis of ME/CFS, such as polymorphism in genes related to immunomodulatory response (184, 185), hormone action (184, 186), and metabolic kynurenine pathways (187). However, genetic associations are not consistent (188) and studies have had quality-control issues (188, 189).

In summary, there are important similarities in predisposing factors for heat stroke and ME/CFS. Pre-existing inflammation or infection may make individuals susceptible to the conditions. Viral reactivation may also play a role in both conditions. Both are associated with immune suppression. Parallel hypotheses suggest that cumulative stress and associated degradation of microbiome health could set the stage for increased susceptibility for both conditions. There is a female predominance in ME/CFS; physiological differences may make women intrinsically more susceptible to exertional heat illnesses than men. In addition, insufficient expression of HSPs has been identified as a factor in both conditions. Finally, genetics are thought to be a risk factor for both conditions, but studies are inconclusive.

**DISCUSSION**

We suggest that the research on heat stroke might provide important lessons for understanding and treating ME/CFS. First, the findings from heat stroke confirm the possibility of systemic inflammation that is not associated with an exogenous pathogen (but rather with endogenous toxins). Heat stroke is primarily an inflammatory response following an external stressor (i.e., an increase in ambient temperature or over-exertion during exercise). This insight is likely important for the broader acceptance of ME/CFS as a disease that does not necessarily result from an exogenous pathogen.

Second, the findings from heat stroke describe a sequence and causality between physiological disturbances in various body systems, including the cardiovascular, gastro-intestinal, immune, and central nervous systems. The sequential pathway from splanchnic vasoconstriction to hypoxia, redox imbalance, and gut hyper-permeability (or intestinal injury) has been consistently documented in both classic and exertional heat stroke. The resulting endotoxemia, cytokine production, systemic inflammation, and decreased cerebral flow are now understood to underlie mitochondrial, central nervous system, and multi-organ dysfunction in heat stroke. This knowledge of the sequence and causality across systems is likely relevant for understanding ME/CFS because of the overlaps in anomalies across the two illnesses.

Third, heat stroke provides a model for disease perpetuation. The heat stroke literature describes a “vicious cycle” centered around intestinal injury involving the release of nitric oxide, the leakage of endotoxins, and the increased production of inflammatory cytokines. This model may also be relevant for understanding the perpetuation of disease state in ME/CFS, as these elements have also been documented in ME/CFS. However, the prevalence of endotoxemia in ME/CFS requires further study to estimate for what share of patients with ME/CFS a similar “vicious cycle” might apply.

Fourth, heat stroke research—combined with the findings from critical illness in general—suggests that intestinal injury resulting from splanchnic hypo-perfusion is a common stress response. This insight could help to understand how a variety of triggers [i.e., physical, infectious, and/or emotional stressor(s)] can result in homeostatic imbalance by initiating intestinal injury. Incidentally, intestinal permeability and microbial translocation are hypothesized to contribute to the pathology in multiple sclerosis (190), rheumatoid arthritis (191), fibromyalgia (192), and Hashimoto's thyroiditis (193)—diseases for which previous studies have described shared pathophysiological mechanisms with ME/CFS (194–198). The endotoxemia and inflammatory pathways described in these diseases echo the patterns observed in heat stroke. A comparative assessment of the overlaps (and differences) between these illnesses would surely provide an important contribution toward resolving these diseases.

Finally, the results of therapeutic studies for heat stroke in animals and people may indicate promising avenues for the treatment of ME/CFS. Heat stroke researchers have tested the use of pharmacological agents and supplements to repair the immune and barrier function of the gut, mitigate endotoxemia, and modulate the inflammatory response. By building on these findings, researchers may identify therapeutic avenues with potential for ME/CFS. The treatment of heat stroke is incomplete and remains an active area of research; a collaboration between ME/CFS and heat stroke researchers would likely benefit both fields.

The findings from heat stroke may provide lessons for understanding both the early and late stages of ME/CFS. Much of heat stroke literature describes the mechanisms occurring within the first 72 h of illness following heat stress. Specifically, adaptive mechanisms intended to deal with stressors are “exaggerated” in certain individuals, thereby overwhelming the homeostasis of the body. Similar exaggerated adaptive responses to external stressors and overwhelmed homeostasis may explain the early stage of ME/CFS. Conversely, the detailed findings from heat stroke regarding the mechanisms that prevent a reversion to a normal homeostatic state even once the stressor is removed (c.f. endotoxemia, O&NS, and pro-inflammatory cytokines) may help to understand the late stages of ME/CFS in at least a subset of patients with ME/CFS.

The existence of overlaps between heat stroke and ME/CFS might initially appear unlikely because heat stroke is a critical illness requiring intensive care, whereas ME/CFS is a chronic illness that debilitates patients for months and years. Moreover, the lack of a disease-specific biomarker and heterogeneity in the clinical symptoms of patients with ME/CFS (resulting in misclassification/misdiagnosis) - combined with the small sample size of many existing studies, pose a challenge in the assessment of overlaps between the two conditions. Yet our suggestion of overlaps between ME/CFS and heat stroke is in line with earlier hypotheses of common pathways in the physiological responses to very different forms of stressors (26–30). A difference in the severity of the vasoconstriction and “endotoxemia pathway” (i.e., intestinal injury) between the two illnesses might explain the difference in the critical/chronic natures of the illnesses. Whereas in
heat stroke the vasoconstriction and “endotoxemia pathway” are so severe that they lead to brain injury and “heat sepsis,” the same mechanisms may be less severe but underpin brain fog and chronic inflammation in some patients with ME/CFS. Speculatively, at least a subgroup of patients with ME/CFS might be thought of experiencing a series of mild exertional heat strokes or a milder but chronic form of the pathological mechanisms witnessed in exertional heat stroke. Accordingly, post-exertional malaise - a key symptom of ME/CFS according to Canadian Consensus Criteria - could be the manifestation of accentuation of these mechanisms following exertion. Additional research is required to assess the relative severity of the pathophysiologial mechanisms described above within heat stroke and ME/CFS, and identify a subgroup of patients where this hypothesis and the above lessons may be most applicable.

Heat stroke is not known to provoke ME/CFS. However, the patients surviving life threatening illnesses can develop PICS, i.e., “the cognitive, psychiatric, and/or physical disability after treatment in ICUs,” including ICU-acquired weakness (199, 200). PICS resembles ME/CFS and remains unresolved. It has been suggested that the persistence of endocrine disturbances observed during critical illness could also explain PICS (201). The similarities and differences between PICS and ME/CFS require further study. Auto-immunity—an important factor in ME/CFS pathology (91, 93, 197)—remains largely unexplored in heat stroke. However, auto-immunity has been associated with inflammatory cytokines and leaky gut (77, 78), which are present in heat stroke.

Neuroendocrine functions during heat stroke remain largely unexplored (31). However, it would seem plausible that the biphasic neuroendocrine pattern found in other critical illnesses might also exist in heat stroke. Specifically, an acute phase (in the first few hours or days) characterized by increased release of pituitary hormones, followed in some critical care patients by a prolonged phase (after a few days) characterized by the suppression of the release of pituitary hormones (202, 203). We have previously described the relevance of the mechanisms during the prolonged phase for ME/CFS (28)—notably the suppression of the pituitary gland’s pulsatile secretion of tropic hormones (204), and a “vicious circle” between inflammation, O&NS, and low thyroid hormone function (205, 206). A compilation of the relevant findings from heat stroke described above (i.e., on the vascular, gastro-intestinal, immune, and nervous systems) and the relevant findings from critical illnesses described previously (i.e., neuroendocrine dysfunctions) might provide important insights into the mechanisms that prevent recovery in ME/CFS (28).

Finally, the lessons-learned from heat stroke for ME/CFS may also be relevant for the aftermath of the COVID-19 pandemic. Many patients with COVID-19 continue to experience a variety of debilitating symptoms despite successfully clearing the virus—termed “post COVID-19 syndrome” or “long COVID-19” — that resemble ME/CFS (207–215). Moreover, the patients with ME/CFS and COVID-19 recovery patients share “molecular signatures” (216), which suggest overlaps in immune and metabolic dysregulation (101).

CONCLUSION

There may be overlaps in the pathophysiological mechanisms in heat stroke and ME/CFS, including splanchnic vasoconstriction, increased gut permeability, gut-related endotoxemia, systemic inflammatory response, central nervous system dysfunction, blood coagulation disorder, endothelial-cell injury, and mitochondrial dysfunction. Moreover, the pathophysiological mechanisms involved may be self-perpetuating in both conditions. Indeed, normalizing body temperature does not necessarily reverse the multiorgan dysfunction in heat stroke; similarly, ME/CFS persists even after the onset event (i.e., physical, infectious, and/or emotional stressor) has ended. A vicious cycle centered around intestinal injury involving the release of nitric oxide, the leakage of endotoxins, and increased production of inflammatory cytokines are central to heat stroke and have also been observed in some patients with ME/CFS. Moreover, initial gene transcriptome studies show that the alterations in gene expression related to mitochondrial and immune function may be common to both conditions. Finally, predisposing factors for heat stroke and ME/CFS appear to partly overlap, including prior inflammation and the altered gut microbiome. Genetics may contribute to the predisposition of individuals in both illnesses, but the existing evidence is inconclusive. Neuroendocrine functions during heat stroke remain largely unexplored. The lack of a disease-specific biomarker and heterogeneity in the clinical symptoms of patients with ME/CFS—combined with the small sample size of many existing studies—pose a challenge in the assessment of overlaps between these illnesses.

A full description of therapeutic trials for heat stroke and their relevance for ME/CFS is beyond the scope of the paper. However, therapeutic approaches to repair the immune and barrier function of the gut, mitigate endotoxemia, modulate the inflammatory response, and induce the expression of HSPs have been suggested or trialed for both conditions.

We suggest that lessons from the human studies and animal models of heat stroke might contribute to the understanding of ME/CFS. Specifically, the human studies and animal models of heat stroke confirm the possibility of systemic inflammation that is not associated with an exogenous pathogen. Moreover, they describe a sequence and causality between physiological disturbances in various body systems and provide an explanation for the self-perpetuation of the disease state that might also inform the understanding of ME/CFS. These studies—combined with the studies from critical illness in general—could help to understand how a variety of triggers (i.e., physical, infectious, and/or emotional stressor) might lead to homeostatic imbalance by initiating intestinal hyper-permeability (or intestinal injury). Moreover, given the similarities described above, an exhaustive analysis of the treatments already tried for heat stroke might help identify which ones could be redeployed to treat ME/CFS. Additional research is required to identify a subgroup of patients with ME/CFS where these lessons may be most applicable.

The existence of possible overlapping pathophysiological mechanisms between heat stroke and ME/CFS could further provide support for a broader hypothesis that the maladaptive mechanisms that prevent recovery in ME/CFS may be common...
to conditions induced by physical, infectious, and/or emotional stressors, such as heat stroke, prolonged critical illness (chronic ICU), PICS, cancer-related fatigue, post-viral fatigue, PACS, and long-COVID-19. We thus suggest that collaborative efforts should be sought among the researcher community across these conditions to better understand these conditions and identify treatments mitigating the functional disability that they induce.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article-supplementary material, further inquiries can be directed to the corresponding author/s.

REFERENCES

1. Bouchama A, Knochel JP. Heat stroke. N Engl J Med. (2002) 346:1978–88. doi: 10.1056/NEJMoa011089
2. Lugo-Amador NM, Rothenhaus T, Moyer P. Heat-related illness. Emerg Med Clin North Am. (2004) 22:315–27. viii. doi: 10.1016/j.emc.2004.01.004
3. Kondo Y, Hifumi T, Shimazaki J, Oda Y, Shiraishi SI, Hayashida K, et al. Comparison between the bouchama and Japanese association for acute medicine heatstroke criteria with regard to the diagnosis and prediction of mortality of heatstroke patients: a multicenter observational study. Int J Environ Res Public Health. (2019) 16:18. doi: 10.3390/ijerph160183433
4. Epstein Y, Yanovich R. Heatstroke. N Engl J Med. (2019) 380:2449–59. doi: 10.1056/NEJMra1810762
5. Vaidyanathan A, Mallay J, Schramm P, Saha S. Heat-related deaths—United States, 2004–2018. MMWR Morb Mortal Wkly Rep. (2020) 69:729–34. doi: 10.15585/mmwr.mm6924a1
6. Komaroff AL. Advances in understanding the pathophysiology of chronic fatigue syndrome. JAMA. (2019) 322:499–500. doi: 10.1001/jama.2019.8312
7. Missalidis D, Annesley SJ, Fisher PR. Pathological mechanisms underlying myalgic encephalomyelitis/chronic fatigue syndrome. Diagnostics. (2019) 9:3. doi: 10.3390/diagnostics9030080
8. Nacul L, Authier FJ, Scheibenbogen C, Lorruso L, Helland IB, Martin JA, et al. European network on myalgic encephalomyelitis/chronic fatigue syndrome (EUROMENE): expert consensus on the diagnosis, service provision, and care of people with ME/CFS in Europe. Medicina. (2021) 57:510. doi: 10.3390/medicina57050510
9. Cortes Rivera M, Mastronardi C, Silva-Aldana CT, Arcos-Burgos M, Lidbury BA. Myalgic encephalomyelitis/chronic fatigue syndrome: a comprehensive review. Diagnostics. (2019) 9:3. doi: 10.3390/diagnostics9030091
10. Domingues TD, Grabsowska AD, Lee JS, Ameijeiras-Alonso J, Westermeier F, Scheibenbogen C, et al. Herpesviruses serology distinguishes different subgroups of patients from the United Kingdom myalgic encephalomyelitis/chronic fatigue syndrome biobank. Front Med. (2021) 8:686736. doi: 10.3389/fmed.2021.686736
11. Chu L, Valencia IJ, Garvert DW, Montoya JG. Onset patterns and course of myalgic encephalomyelitis/chronic fatigue syndrome. Front Pediatr. (2019) 7:12. doi: 10.3389/fped.2019.00012
12. Institute of Medicine. Beyond Myalgic Encephalitis/Chronic Fatigue Syndrome: Redefining an Illness. Washington, DC: The National Academies Press (2015).
13. Open Medicine Foundation. Symptoms of ME/CFS. (2020). Available online at: https://www.omf.ngo/symptoms-mecfs (accessed March 27, 2021).
14. Centers for Disease Control and Prevention. Clinical Care of Patients with ME/CFS—Severely Affected Patients. (2019). Available online at: https://www.cdc.gov/me-cfs/healthcare-providers/clinical-care-patients-mefs/severely-affected-patients.html [accessed March 27, 2021].

AUTHOR CONTRIBUTIONS

DS wrote the first draft of the manuscript. All authors contributed to manuscript revision, read, and approved the submitted version.

FUNDING

Open Medicine Foundation (JB) is acknowledged for support. NS acknowledges funding from the Polish National Agency for Academic Exchange (Reference Grant: PPN/ULM/2020/1/00069/U/00001) and FCT—Fundaçao para a Ciência e Tecnologia, Portugal (Ref. Grant: UIDB/0006/2020).

15. Williams LR, Isaaco-Mariash C. Three cases of severe me/cfs in adults. Healthcare. (2021) 9:2. doi: 10.3390/healthcare9020215
16. Dafoe W. Extremely severe ME/CFS—a personal account. Healthcare. (2021) 9:504. doi: 10.3390/healthcare9050504
17. Nacul L, O’Boyle S, Palla L, Nacul FE, Mudie K, Kingdon CC, et al. How myalgic encephalomyelitis/chronic fatigue syndrome (ME/CFS) progresses: the natural history of ME/CFS. Front Neurol. (2020) 11:826. doi: 10.3389/fneur.2020.00826
18. Komaroff AL. Myalgic encephalomyelitis/chronic fatigue syndrome: when suffering is multiplied. Healthcare. (2021) 9:919. doi: 10.3390/healthcare9070919
19. McMamien SL, Devendorf AR, Brown AA, Moore BC, Moore JH, Jason LA. Mortality in patients with myalgic encephalomyelitis and chronic fatigue syndrome. Fatigue. (2016) 4:195–207. doi: 10.1080/21641846.2016.1236588
20. Jason LA, Mirin AA. Updating the national academy of medicine ME/CFS prevalence and economic impact figures to account for population growth and inflation. Fatigue: Biomed Health Behav. (2021) 21:1–5. doi: 10.1080/21641846.2021.1878716
21. Toogood PL, Clauw DJ, Phadke S, Hoffman D. Myalgic encephalomyelitis/chronic fatigue syndrome (ME/CFS): where will the drugs come from? Pharmacol Res. (2021):105465. doi: 10.1016/j.phrs.2021.105465
22. Richman S, Morris MC, Broderick G, Craddock TJA, Klimas NG, Fletcher MA. Pharmaceutical interventions in chronic fatigue syndrome: a literature-based commentary. Clin Therapeut. (2019) 41:798–805. doi: 10.1016/j.clinthera.2019.02.011
23. Kim DY, Lee JS, Park SY, Kim SJ, Son CG. Systematic review of randomized controlled trials for chronic fatigue syndrome/myalgic encephalomyelitis (CFEOM/ME). J Transl Med. (2020) 18:7. doi: 10.1186/s12916-019-02196-9
24. Castro-Marrero J, Sáez-Francàs N, Santillo D, Alegre J. Treatment and management of chronic fatigue syndrome/myalgic encephalomyelitis: all roads lead to Rome. Br J Pharmacol. (2017) 174:345–69. doi: 10.1111/bph.13702
25. Rowe PG, Underhill RA, Friedman KJ, Gurwitt A, Medow MS, Schwartz MS, et al. Myalgic encephalomyelitis/chronic fatigue syndrome diagnosis and management in young people: a primer. Front Pediatr. (2017) 5:121. doi: 10.3389/fped.2017.00121
26. Arnett SV, Clark IA. Inflammatory fatigue and sickness behaviour–lessons for the diagnosis and management of chronic fatigue syndrome. J Affect Disord. (2012) 141:130–42. doi: 10.1016/j.jad.2012.04.004
27. Craddock TJ, Fritsch P, Rice MA, Jr., del Rosario RM, Miller DB, et al. A role for homeostatic drive in the perpetuation of complex chronic illness: Gulf War Illness and chronic fatigue syndrome. PLoS ONE. (2014) 9:e84839. doi: 10.1371/journal.pone.0084839
28. Stanculescu D, Larsson L, Bergquist J. Hypothetic: mechanisms that prevent recovery in prolonged ICU patients also underlie myalgic encephalomyelitis/chronic fatigue syndrome (ME/CFS). Front Med. (2021) 8:41. doi: 10.3390/fmed.2021.628029
1. Armstrong LE, Lee EC, Armstrong EM. Interactions of gut microbiota, endotoxemia, immune function, and diet in exertional heatstroke. *J Sports Med.* (2018) 57:25475. doi: 10.1155/2018/5725475

2. Hall DM, Buettner GR, Oberley LW, Xu L, Matthews RD, Gisolfi CV. Mechanisms of circulatory and intestinal barrier dysfunction during whole body hyperthermia. *Ann J Physiol Heart Circ Physiol.* (2001) 280:H509–21. doi: 10.1152/ajpheart.2001.280.2.H509

3. Lambert GP. Role of gastrointestinal permeability in exertional heatstroke. *Exerc Sci Sport Rev.* (2004) 32:185–90. doi: 10.1097/00003667-200410000-00011

4. Hifumi T, Kondo Y, Shimizu K, Miyake Y. Heat stroke. *Intensiv Care.* (2018) 6:1–8. doi: 10.1186/s40680-018-0298-4

5. Mittal R, Coopersmith CM. Redefining the gut as the motor of critical illness. *Trends Mol Med.* (2014) 20:214–23. doi: 10.1016/j.molmed.2013.08.004

6. Deane A, Chapman MJ, Fraser RJ, Horowitz M. Bench-to-bedside review: the gut as an endocrine organ in the critically ill. *Crit Care.* (2010) 14:228. doi: 10.1186/cc9309

7. Martinez EE, Fasano A, Mehta NM. Gastrointestinal function in critical illness—a complex interplay between the nervous and enteroinnervational systems. *Pediatr Med.* (2020) 3:74. doi: 10.21037/pm-20-74

8. Soares AD, Costa KA, Wanner SP, Santos RG, Fernandes SO, Martins FS, et al. Dietary glutamine prevents the loss of intestinal barrier function and attenuates the increase in core body temperature induced by acute heat exposure. *Br J Nutr.* (2014) 112:1601–10. doi: 10.1017/S0007114514026086

9. Singleton KD, Wismehemy PE. Oral glutamine enhances heat shock protein expression and improves survival following hyperthermia. *Shock.* (2006) 25:295–9. doi: 10.1097/01.shk.0000196548.10634.02

10. Cao Y, Liu Z, Xiao W, Gu Z, Xiao G, Yuan F, et al. 4-Phenybutrylate prevents endoplasmic reticulum stress-mediated apoptosis induced by heatstroke in the intestines of mice. *Shock.* (2020) 54:102–9. doi: 10.1097/SHK.00000000000001419

11. Liu CC, Chien CH, Lin MT. Glucorticoids reduce interleukin-1 concentration and result in neuroproprotective effects in rat heatstroke. *J Physiol.* (2000) 527:333–43. doi: 10.1111/j.1469-7939.2000.01-033x.x

12. King MA, Rollo I, Baker LR. Nutritional considerations to counteract gastrointestinal permeability during exertional heat stress. *J Appl Physiol.* (1985) 130:1754–65. doi: 10.1152/japplphysiol.00072.2021

13. Sfera A, Osorio C, Zapata Martin Del Campo CM, Pereida S, Maurer S, Maldonado JC, et al. Endothelial senescence and chronic fatigue syndrome, a COVID-19 based hypothesis. *Front Cell Neurosci.* (2021) 15:673217. doi: 10.3389/fncel.2021.673217

14. Maes M, Coucke F, Leunis JC. Normalization of the increased translocation of endotoxin from gram negative enterobacteria (leaky gut) is accompanied by a remission of chronic fatigue syndrome. *Neuro Endocrinol Lett.* (2008) 29:73–94.

15. Giloteaux L, Goodrich JK, Walters WA, Levine SM, Ley RE, Hanson MR. Reduced diversity and altered composition of the gut microbiome in individuals with myalgic encephalomyelitis/chronic fatigue syndrome. *Microbiome.* (2016) 4:30. doi: 10.1186/s40168-016-0171-4

16. Morris G, Berk M, Carvalho AF, Caso JR, Sanz Y, Maes M. The role of microbiota and intestinal permeability in the pathophysiology of autoimmune and neuroimmune processes with an emphasis on inflammatory bowel disease type 1 diabetes and chronic fatigue syndrome. *Curr Pharm Des.* (2016) 22:6058–75. doi: 10.2174/1381622622666691094118222

17. Morris G, Maes M, Berk M, Puri BK. Myalgic encephalomyelitis or chronic fatigue syndrome: how could the illness develop? *Metabolic Brain Disease.* (2019) 34:385–415. doi: 10.1007/s11011-019-0388-6

18. Lakhan SE, Kirchgeissner A. Gut inflammation in chronic fatigue syndrome. *Nutr Metab.* (2010) 7:79. doi: 10.1186/1743-7075-7-79

19. Maes M, Mihaylova I, Leunis JC. Increased serum IgA and IgM against LPS of enterobacteria in chronic fatigue syndrome (CFS): indication for the involvement of gram-negative enterobacteria in the etiology of CFS and for the presence of an increased gut-intestinal permeability. *J Affect Disord.* (2007) 99:237–40. doi: 10.1016/j.jad.2006.08.021

20. Zhang ZT, Du XM, Ma XJ, Zong Y, Chen JK, Yu CL, et al. Activation of the NLRP3 inflammasome in lipopolysaccharide-induced mouse fatigue
and its relevance to chronic fatigue syndrome. *J Neuroinflammation.* (2016) 13:71. doi: 10.1186/s12974-016-0539-1

70. Maes M, Leunis JC. Normalization of leaky gut in chronic fatigue syndrome (CFS) is accompanied by a clinical improvement: effects of age, duration of illness and the translocation of LPS from gram-negative bacteria. *Neuro Endocrinol Lett.* (2008) 29:902–10.

71. Anderson G, Maes M. Mitochondria and immunity in chronic fatigue syndrome. *Prog Neuro-psychopharmacol Biol Psychiatry.* (2020) 103:109976. doi: 10.1016/j.pnpbp.2020.109976

72. Camilleri M, Gorman H. Intestinal permeability and irritable bowel syndrome. *Neurogastroenterol Motil.* (2007) 19:545–52. doi: 10.1111/j.1365-2982.2007.00925.x

73. Arrieta MC, Bistritz L, Meddings JB. Alterations in intestinal permeability. *Gut.* (2006) 55:1512–20. doi: 10.1136/gut.2005.085373

74. Shing CM, Peake JM, Lim CL, Briske D, Walsh NP, Fortes MB, et al. Effects of probiotics supplementation on gastrointestinal permeability, inflammation and exercise performance in the heat. *Eur J Appl Physiol.* (2014) 114:93–103. doi: 10.1007/s00421-013-2748-y

75. Derix JP, Loyer MD, Heineman E, Buurman WA. Non-invasive markers of gut wall integrity in health and disease. *World J Gastroenterol.* (2010) 16:5272–9. doi: 10.3748/wjg.v16.i42.5272

76. Sapone A, de Magistris L, Pietzak M, Clemente MG, Tripathi A, Cucca F, et al. Zonulin upregulation is associated with increased gut permeability in subjects with type 1 diabetes and their relatives. *Diabetes.* (2006) 55:1443–9. doi: 10.2337/diabetes.55.6.1443

77. Pasano A, Leaky gut and autoimmune diseases. *Clin Rev Allergy Immunol.* (2012) 42:71–8. doi: 10.1007/s12016-011-8291-x

78. Mu Q, Kirby J, Reilly CM, Luo XM. Leaky Gut As a Danger signal in Autoimmune Diseases. *Front Immunol.* (2017) 8:598. doi: 10.3389/fimmu.2017.00598

79. Collins SM. A role for the gut microbiota in IBS. *Nat Rev Gastroenterol Hepatol.* (2014) 11:497–505. doi: 10.1038/nrgastro.2014.40

80. Service NH. Leaky gut syndrome (2018). Available online at: https://www. heron.nhs.uk/publish/PublicationDetails.aspx?m=1&formatid=24760 (accessed November 1, 2021).

81. Zhang ZT, Gu XL, Zhao X, He X, Shi HW, Zhang K, et al. NLRP3 activation enhances anti-inflammatory effects on LPS-induced THP-1 cells by enhancing PPAR γ signaling. *Am J Physiol Regul Integr Comp Physiol.* (2015) 309:R960–8. doi: 10.1152/ajpregu.00291.2015

82. Pedersen BK, Febbraio M. Muscle-derived interleukin-6—a possible link and antiinflammatory mechanisms. *Chem Biol Interact.* (2017) 265:47–54. doi: 10.1016/j.cbi.2017.01.006

83. Chang CY, Chen JY, Chen SH, Cheng TJ, Lin MT, Hu ML. Therapeutic treatment with ascorbate rescues mice from heat stroke-induced death by attenuating systemic inflammatory response and hypothermic neuronal damage. *Free Radic Biol Med.* (2016) 93:84–93. doi: 10.1016/j.freeradbiomed.2015.12.017

84. Bland J, Gottfried CG, Eliafouri A, Rizwan M, Rosén A. Infection elicits autoimmunity and myalgic encephalomyelitis/chronic fatigue syndrome: an explanatory model. *Front Immunol.* (2018) 9:229. doi: 10.3389/fimmu.2018.00229

85. Myhill S. Diagnosis and Treatment of Chronic Fatigue Syndrome and Myalgic Encephalitis, 2nd ed.: Its Mitochondria, Not Hypochondria. White River Junction, VT: Chelsea Green Publishing (2018).

86. Morris G, Berk M, Galecki P, Maes M. The emerging role of autoimmunity in myalgic encephalomyelitis/chronic fatigue syndrome (ME/CFS). *Mol Neurobiol.* (2014) 49:741–56. doi: 10.1007/s12033-013-8553-0

87. Komaroff AL. Inflammation correlates with symptoms in chronic fatigue syndrome. *Proc Natl Acad Sci U S A.* (2017) 114:8914–6. doi: 10.1073/pnas.1712475114

88. Sweetman E, Noble A, Edgar C, Mackay A, Hellwell A, Vallings R, et al. Current research provides insight into the biological basis and diagnostic potential for myalgic encephalomyelitis/chronic fatigue syndrome (ME/CFS). *Diagnósticos.* (2019) 9:73. doi: 10.3390/diagnostics9030073

89. Glassford IA. The neuroinflammatory etiopathology of myalgic encephalomyelitis/chronic fatigue syndrome (ME/CFS). *Front Physiol.* (2017) 8:88. doi: 10.3389/fphys.2017.00088

90. Tomas C, Newton J. Metabolic abnormalities in chronic fatigue syndrome/myalgic encephalomyelitis: a mini-review. *Biochem Soc Trans.* (2018) 46:547–53. doi: 10.1042/BST20170503

91. Maes M, Kubera M, Uytterhoeven M, Vrydaghs N, Bosmans E. Increased plasma peroxides as a marker of oxidative stress in myalgic encephalomyelitis/chronic fatigue syndrome (ME/CFS). *Med Sci Monit.* (2011) 17:A4. doi: 10.25270/msm.981699

92. Montoya JG, Holmes TH, Anderson JN, Maecker HT, Rosenberg-Hasson Y, Valencia JJ, et al. Cytokine signature associated with disease severity in chronic fatigue syndrome patients. *Proc Natl Acad Sci USA.* (2017) 114:E1750–E8. doi: 10.1073/pnas.1705159114

93. Hornig M, Montoya JG, Klimas NG, Levine S, Felsenstein D, Bateman L, et al. Distinct plasma immune signatures in ME/CFS are present early in the course of illness. *Sci Adv.* (2015) 1:e1400121. doi: 10.1126/sciadv.1400121

94. Paul BD, Lemle MD, Komaroff AL, Snyder SH. Redox imbalance links COVID-19 and myalgic encephalomyelitis/chronic fatigue syndrome. *Proc Natl Acad Sci U S A.* (2021) 118:34. doi: 10.1073/pnas.2024358118

95. Armstrong CW, McGregor NR, Lewis DP, Butt HL, Gooley PR. Metabolic profiling reveals anomalous energy metabolism and oxidative stress pathways in chronic fatigue syndrome patients. *Metabolomics.* (2015) 11:1626–39. doi: 10.1007/s11306-015-0816-5

96. Morris G, Maes M. Mitochondrial dysfunctions in myalgic encephalomyelitis/chronic fatigue syndrome explained by activated immuno-inflammatory, oxidative and nitrosative stress pathways. *Metab Brain Dis.* (2014) 29:19–36. doi: 10.1007/s11011-013-9435-x

97. Hatzigeokaki E, Adamaki M, Tsilioni I, Dimitriadis G, Theoharides TC. Myalgic encephalomyelitis/chronic fatigue syndrome–metabolic disease or disturbed homeostasis due to focal inflammation in the hypothalamus? *J Pharmacol Exp Ther.* (2018) 367:155–67. doi: 10.1124/jpet.117.250845

98. Morris G, Anderson G, Maes M. Hypothalamic-Pituitary-Adrenal Hypofunction in myalgic encephalomyelitis (me)/chronic fatigue syndrome (cfs) as a consequence of activated immune-inflammatory and oxidative and nitrosative pathways. *Mol Neurobiol.* (2017) 54:6806–19. doi: 10.1007/s12035-016-0170-2

99. Pall M. “The NO/ONOO-cycle mechanism as the cause of chronic fatigue syndrome/myalgia encephalomyelitis.” In: Svoboda E, Zelenjak K, editors. *Chronic Fatigue Syndrome: Symptoms, Causes and Prevention, London: Nova Publishers* (2009).

100. Morris G, Maes M. Oxidative and nitrosative stress and immune-inflammatory pathways in patients with myalgic encephalomyelitis
(me)/chronic fatigue syndrome (CFS). *Carr Neuropharmacol.* (2014) 12:168–85. doi: 10.2174/1570159X1666131120224653

108. Shibot S, Coll R, Giltat T, Sohar E. Heatstroke: its clinical picture and mechanism in 36 cases. *Q J Med.* (1967) 36:525–48.

109. Chao TC, Sinniah R, Pakiam JE. Acute heat stroke deaths. *Pathology.* (1981) 13:145–56. doi:10.1111/j.1365-2559.1981.tb00637.x

110. Lin MT. Pathogenesis of an experimental heatstroke model. *Clin Exp Pharmacol Physiol.* (1999) 26:826–7. doi:10.1046/j.1440-1681.1999.03137.x

111. Yamaguchi T, Shimizu K, Kokubu Y, Nishijima M, Takeda S, Ogura H, et al. Effect of heat stress on blood-brain barrier integrity in iPS cell-derived microvascular endothelial cell models. *PLoS ONE.* (2019) 14:e0222113. doi:10.1371/journal.pone.0222113

112. Nakatomi Y, Kuratsune H, Watanabe Y. Neuroinflammation in the brain of patients with myalgic encephalomyelitis/chronic fatigue syndrome. *Brain Nerve.* (2018) 70:19–25. doi:10.1177/1360171117729495

113. Nakatomi Y, Mizuno K, Ishii A, Wada Y, Tanaka M, Tazawa S, et al. Neuroinflammation in patients with chronic fatigue syndrome/myalgic encephalomyelitis: an (1)H/(13)C-(R)-PK11195 PET study. *J Nucl Med.* (2014) 55:945–50. doi:10.2967/jnm.113.131045

114. Mueller C, Lin JC, Sheriff S, Maudsley AA, Younger JW. Evidence of widespread metabolite abnormalities in Myalgic encephalomyelitis/chronic fatigue syndrome: assessment with whole-brain magnetic resonance spectroscopy. *Brain Imaging Behav.* (2020) 14:562–72. doi:10.1007/s11682-018-0029-4

115. Zeineh MM, Kang J, Atlas SW, Raman MM, Reis AL, Norris JL, et al. Right arcuate fasciculus abnormality in chronic fatigue syndrome. *Radiology.* (2017) 274:517–26. doi:10.1148/radiol.14114079

116. Ferrero K, Silver M, Cocchetto A, Masliah E, Langford D. CNS findings in chronic fatigue syndrome and a neuropathological case report. *J Invest Med.* (2017) 65:974–83. doi:10.1136/jim-2016-000390

117. van Campen CMC, Verheugen FWA, Rowe PC, Visser FC. Cerebral blood flow is reduced in ME/CFS during head-up tilt testing even in the absence of hypotension or tachycardia: a quantitative, controlled study. *J Pain.* (2018) 19:655–71. doi:10.1007/s80783-018-0070-8

118. Chen HS, Tong HS, Zhao Y, Hong CY, Bin JP, Su L. Differential expression in patients with myalgic encephalomyelitis: an (1)C-(R)-PK11195 PET study. *J Nucl Med.* (2014) 55:945–50. doi:10.2967/jnm.113.131045

119. Muller C, Lin JC, Sheriff S, Maudsley AA, Younger JW. Evidence of widespread metabolite abnormalities in Myalgic encephalomyelitis/chronic fatigue syndrome: assessment with whole-brain magnetic resonance spectroscopy. *Brain Imaging Behav.* (2020) 14:562–72. doi:10.1007/s11682-018-0029-4

120. Zeineh MM, Kang J, Atlas SW, Raman MM, Reis AL, Norris JL, et al. Right arcuate fasciculus abnormality in chronic fatigue syndrome. *Radiology.* (2017) 274:517–26. doi:10.1148/radiol.14114079

121. Ferrero K, Silver M, Cocchetto A, Masliah E, Langford D. CNS findings in chronic fatigue syndrome and a neuropathological case report. *J Invest Med.* (2017) 65:974–83. doi:10.1136/jim-2016-000390

122. van Campen CMC, Verheugen FWA, Rowe PC, Visser FC. Cerebral blood flow is reduced in ME/CFS during head-up tilt testing even in the absence of hypotension or tachycardia: a quantitative, controlled study. *J Pain.* (2018) 19:655–71. doi:10.1007/s80783-018-0070-8

123. Bouchama A, Roberts G, Al Mohanna F, El-Sayed R, Lach B, Chollet-Martin S, et al. Inflammatory, hemostatic, and clinical changes in a baboon experimental model for heatstroke. *J Intensive Care.* (2015) 3:579–86. doi:10.1186/s40560-015-0039-3

124. Kobayashi K, Mimuro S, Sato T, Kobayashi A, Kawashima S, Makino H, et al. Dexamethasone preserves the endothelial glycolysis and improves survival in a rat heatstroke model. *J Anesth.* (2013) 28:808–15. doi:10.1007/s00540-012-2589-7

125. Berg D, Berg LH, Couvras J, Harrison H. Chronic fatigue syndrome and/or fibromyalgia as a variation of antiphospholipid antibody syndrome: an explanatory model and approach to laboratory diagnosis.
145. Naviaux RK. Metabolic features and regulation of the healing cycle: A new model for chronic disease pathogenesis and treatment. *Mitochondrion*. (2019) 46:278–97. doi: 10.1016/j.mito.2018.08.001

146. Bouchama A, Aziz MA, Mahri SA, Gabere MN, Diamy MA, Mohammad S, et al. A model of exposure to extreme environmental heat uncovers the human transcriptome to heat stress. *Sci Rep.* (2017) 7:9429. doi: 10.1038/s41598-017-09819-5

147. Kitami T, Fukuda S, Kato T, Yamaguti K, Nakatomi Y, Yamano E, et al. Deep phenotyping of myalgic encephalomyelitis/chronic fatigue syndrome in Japanese population. *Sci Rep.* (2020) 10:19933. doi: 10.1038/s41598-020-77105-y

148. Frampton D, Kerr J, Kellam P. Assessment of a 44 gene classifier for the evaluation of chronic fatigue syndrome from peripheral blood mononuclear cell gene expression. *PLoS One*. (2011) 6:e18672. doi: 10.1371/journal.pone.0018672

149. Gow JW, Hagan S, Herzyk P, Cannon C, Behan PO, Chaudhuri A. A gene signature for post-infectious chronic fatigue syndrome. *BMJ Med Genomics*. (2009) 2:238. doi: 10.1186/1755-8794-2-238

150. Vernon SD, Unger ER, Dimulescu IM, Rajeevan M, Reeves WC. Utility of the blood for gene expression profiling and biomarker discovery in chronic fatigue syndrome. *Dis Markers*. (2002) 18:193–9. doi: 10.1155/2002/892374

151. Nguyen CB, Alsoe L, Lindvall JM, Sulheim D, Winger A, et al. Whole blood gene expression in adolescent chronic fatigue syndrome: an exploratory cross-sectional study suggesting altered B cell differentiation and altered expression of antiviral response genes. *J Transl Med.* (2017) 15:102. doi: 10.1186/s12967-017-1210-0

152. Kerr JR, Petty R, Burke B, Gough J, Fear D, Sinclair LJ, et al. Gene expression subtypes in patients with chronic fatigue syndrome/myalgic encephalomyelitis. *J Infect Dis.* (2008) 197:171–84. doi: 10.1086/533453

153. Sweetman E, Ryan M, Edgar C, MacKay A, Vallings R, Tate W. Changes in the transcriptome of circulating immune cells of a New Zealand cohort with myalgic encephalomyelitis/chronic fatigue syndrome. *Int J Immunopharmacol*. (2019) 33:2057384118820402. doi: 10.1016/j.ijip.2018.08.001

154. Saiki T, Kawai T, Morita K, Ohta M, Saito T, Rokutan K, et al. Identification of marker genes for differential diagnosis of chronic fatigue syndrome. *Mol Med.* (2008) 14:599–607. doi: 10.2119/2007-0059-saiki

155. Nepotchatykh E, Elremaly W, Caraus I, Godbout C, Leveau C, Chalder L, et al. Profile of circulating microRNAs in myalgic encephalomyelitis and their relation to symptom severity, and disease pathophysiology. *Scientific Reports*. (2020) 10:19620. doi: 10.1038/s41598-020-76438-y

156. Epstein Y. Heat intolerance: predisposing factor or residual injury? *Med Sci Sports Exerc.* (1990) 22:29–35. doi: 10.1249/00005768-199002000-00006

157. Armstrong LE, De Luca JP, Hubbard RW. Time course of recovery and heat uncovers the human transcriptome to heat stress. *Alpha-Lipoic acid protects cardiomyocytes against heat stroke-induced apoptosis and inflammatory responses associated with the induction of hsp70 and activation of autophagy. Mediators Inflamm.* (2019) 2019:8187529. doi: 10.1155/2019/8187529

158. Epstein Y. Heat intolerance: predisposing factor or residual injury? *Med Sci Sports Exerc.* (1990) 22:29–35. doi: 10.1249/00005768-199002000-00007

159. Shen HH, Tseng YS, Kuo NC, Kung CW, Amin S, Lam KK, et al. Alpha-Lipoic acid protects cardiomyocytes against heat stroke-induced apoptosis and inflammatory responses associated with the induction of hsp70 and activation of autophagy. *Mediators Inflamm.* (2019) 2019:8187529. doi: 10.1155/2019/8187529

160. Lakier Smith L. Overtraining, excessive exercise, and altered immunity: is muscle fatigue syndrome combines increased exercise-induced oxidative stress and reduced cytokine and Hsp responses. *J Transl Med.* (2020) 18:246. doi: 10.1186/s12967-020-02419-4

161. Gifford RM, Todisco T, Stacey M, Fujisawa T, Allerhand M, Woods DR, et al. Profile of circulating microRNAs in myalgic encephalomyelitis/chronic fatigue syndrome. *Med Sci Sports Exerc.* (1990) 22:29–35. doi: 10.1249/00005768-199002000-00007

162. Sherry JW, Wettenhall RE, Scanlon D, Gooley PR, Lewis DP, McGregor N, et al. Increased d-lactic acid intestinal bacteria in patients with chronic fatigue syndrome. *In Vivo*. (2009) 23:621–8.

163. Tsai SY, Chen HJ, Lio CF, Kuo CF, Kao AC, Wang WS, et al. Increased risk of chronic fatigue syndrome in patients with inflammatory bowel disease: a population-based retrospective cohort study. *J Transl Med.* (2019) 17:55. doi: 10.1186/s12967-019-1797-3

164. Safadi JM, Quinton AMG, Lennox BR, Burnet PWJ, Minichino A. Gut dysbiosis in severe mental illness and chronic fatigue: a novel trans-diagnostic construct? a systematic review and meta-analysis. *Mol Psychiatry.* (2021) 21:1032: doi: 10.1038/s41380-021-01032-1

165. Ghali A, Richa P, Lacout C, Gury A, Beucher AB, Homedan C, et al. Epidemiological and clinical factors associated with post-exertional malaise severity in patients with myalgic encephalomyelitis/chronic fatigue syndrome. *J Transl Med.* (2017) 15:102. doi: 10.1186/s12967-017-09819-5

166. Armstrong LE, De Luca JP, Hubbard RW. Time course of recovery and heat uncovers the human transcriptome to heat stress. *Alpha-Lipoic acid protects cardiomyocytes against heat stroke-induced apoptosis and inflammatory responses associated with the induction of hsp70 and activation of autophagy. Mediators Inflamm.* (2019) 2019:8187529. doi: 10.1155/2019/8187529

167. Lacerda EM, Geraghty K, Kingdon CC, Palla L, Nacul L. A logistic regression analysis of risk factors in ME/CFS pathogenesis. *BMC Neurol.* (2019) 19:275. doi: 10.1186/s12883-019-1648-2

168. Buchwald D, Cheney PR, Peterson DL, Henry B, Wormsley SB, Geiger A, et al. A chronic illness characterized by fatigue, neurologic and immunologic disorders, and active human herpesvirus type 6 infection. *Ann Intern Med.* (1992) 116:103–13. doi: 10.7326/0003-4819-116-2-103

169. Ablashi DV. Viral studies of chronic fatigue syndrome. *Clin Infect Dis.* (1994) 18:S130–3. doi: 10.1093/clinids/18.Supplement_1.S130

170. Gifford RM, Todisco T, Stacey M, Fujisawa T, Allerhand M, Woods DR, et al. Heat shock proteins and heat adaptation of the whole organism. *J Appl Physiol.* (1985) 53:1413–7. doi: 10.1152/jappl.1985.53.3.1413
dysfunction in myalgic encephalomyelitis/chronic fatigue syndrome: a pilot study. Front Pediatr. (2019) 7:206. doi: 10.3389/fped.2019.00206
185. Carlo-Stella N, Badulli C, De Silvestri A, Bazzichi L, Martinetti M, Lorusso L, et al. A first study of cytokine genomic polymorphisms in CFS: positive association of TNF-857 and IFN gamma 874 rare alleles. Clin Exp Rheumatol. (2006) 24:179–82. doi: 10.1094/CEP-2006-07-0085
186. Vangeel E, Van Den Eede E, Hompes T, Izzii R, Del Favero J, Moorkens G, et al. chronic fatigue syndrome and dna hypomethylation of the glucocorticoid receptor gene promoter If region: associations with hpa axis hypofunction and childhood trauma. Psychosom Med. (2015) 77:853–62. doi: 10.1016/j.psymp.2015.07.002
187. Kashi AA, Davis RW, Phair RD. The IDO metabolic trap hypothesis for the etiology of ME/CFS. Diagnostics. (2019) 9:82. doi: 10.3390/diagnostics9030082
188. Dibble JJ, McGrath SJ, Ponting CP. Genetic risk factors of ME/CFS: a critical review. Hum Mol Genet. (2020) 29[R1]:R117–R24. doi: 10.1093/hmg/ddaa169
189. Grabowska AD, Lacerda EM, Nacul L, Sepulveda N. Review of the quality control checks performed by current genome-wide and targeted-genome association studies on myalgic encephalomyelitis/chronic fatigue syndrome. Front Pediatr. (2020) 8:293. doi: 10.3389/fped.2020.00293
190. Kirby TO, Ochoa-Reparaz J. The gut microbiome in multiple sclerosis: a potential therapeutic avenue. Med Sci. (2018) 6:3. doi: 10.3390/medsciences3020009
191. Tanjea V. Arthritis susceptibility and the gut microbiome. FEBS Lett. (2014) 588:4244–9. doi: 10.1016/j.fbslet.2014.05.034
192. Erdrich S, Hawrelak JA, Myers SP, Harnett JE. Determining the FEBS Lett. (2020)
193. Küçükemre Aydin B, Yildiz M, Akgün A, Topal N, Adal E, Önal H. Determining the FEBS Lett. (2020)
194. Morris G, Maes M. Myalgic encephalomyelitis/chronic fatigue syndrome and encephalomyelitis disseminated/multiple sclerosis show remarkable levels of similarity in phenomenology and neuroimmune characteristics. BMC Med. (2013) 11:205. doi: 10.1186/1741-7015-11-205
195. Maes M, Twisk FNM, Kubera M, Ringel K. Evidence for inflammation and activation of cell-mediated immunity in myalgic encephalomyelitis/chronic fatigue syndrome (ME/CFS): increased interleukin-1 tumor necrosis factor α, PMN-elastase, lysosome and neopterin. J Affect Disord. (2012) 136:933–9. doi: 10.1016/j.jad.2011.09.004
196. Natelson BH, Myalgic encephalomyelitis/chronic fatigue syndrome and fibromyalgia: definitions, similarities, and differences. Clin Ther. (2019) 41:612–8. doi: 10.1016/j.clinthera.2018.12.016
197. Sotzny F, Blanco J, Capelli E, Castro-Marrero J, Steiner S, Murovska M, et al. Myalgic encephalomyelitis/chronic fatigue syndrome—evidence for an autoimmune disease. Autoimmunity Reviews. (2018) 17:601–9. doi: 10.1016/j.autrev.2018.01.009
198. Castro-Marrero J, Faro M, Aliste L, Sáez-Francá N, Calvo N, Martinez-Martinez A, et al. Comorbidity in chronic fatigue syndrome/myalgic encephalomyelitis: a nationwide population-based cohort study. Psychosomatics. (2017) 58:533–43. doi: 10.1016/j pysym.2017. 04.010
199. Rawal G, Yadav S, Kumar R. Post-intensive care syndrome: an overview. J Transl Med. (2017) 5:90–2. doi: 10.1186/s12967-016-0016
200. Smith S, Rahman O. “Post intensive care syndrome [Updated 2020 Jun 25],” In: StatPearls [Internet]. Treasure Island (FL): StatPearls Publishing. (2020).
201. Van Aerde N, Van Dyck L, Vanhorebeck I, Van den Berge G. “Endocrinopathy of the critically ill,” In: Preiser J-C, Herridge M, Azozay E, editors. Post-Intensive Care Syndrome. Cham: Springer International Publishing. (2020), 125–43.
202. Van den Berghe GH. Acute and prolonged critical illness are two distinct neuroendocrine paradigms. Verh Akad Geneeskd Belg. (1998) 60:487–518; discussion —20. doi: 10.1210/jc.83.6.1827
203. Vanhorebeek I, Van den Berge G. The neuroendocrine response to critical illness is a dynamic process. Crit Care Clin. (2006) 22:1–15. doi: 10.1016/j.ccc.2005.09.004
204. Van den Berge G. On the neuroendocrinopathy of critical illness: perspectives for feeding and novel treatments. Am J Respir Crit Care Med. (2016) 194:1337–48. doi: 10.1164/rccm.201607-1516CI
205. Manzini A, Di Segni C, Raimondo S, Olivieri G, Silvestrini A, Meucci E, et al. Thyroid hormones, oxidative stress, and inflammation. Mediators Inflamm. (2016) 2016:675154. doi: 10.1155/2016/675154
206. Chatzitomaris A, Hoermann R, Midgley JE, Hering S, Urban A, Dietrich B, et al. Thyroid allostatic–adaptive responses of thyrotropic feedback control to conditions of strain, stress, and developmental programming. Front Endocrinol. (2017) 8:163. doi: 10.3389/fendo.2017.00163
207. Greenhalgh T, Knight M, ACourt C, Buxton M, Husain L. Management of post-acute covid-19 in primary care. BMJ. (2020) 370:m3026. doi: 10.1136/bmj.m3026
208. Dani M, Dirksen A, Taraborrelli P, Torocastro M, Panagopoulos D, Sutton R, et al. Autonomic dysfunction in “long COVID”: rationale, physiology and management strategies. Clin Med. (2020) 20:7301. doi: 10.7861/crmed.2020-0896
209. Huang C, Huang L, Wang Y, Li X, Ren L, Gu X, et al. 6-month consequences of COVID-19 in patients discharged from hospital: a cohort study. The Lancet. (2021) 397:220–32. doi: 10.1016/S0140-6736(20)32656-8
210. Townsend L, Dyer AH, Jones K, Dunne I, Mooney A, Gaffney F, et al. Persistent fatigue following SARS-CoV-2 infection is common and independent of severity of initial infection. PLOS ONE. (2020) 15:e0240784. doi: 10.1371/journal.pone.0240784
211. Komaroff AL, Bateman L. Will COVID-19 lead to myalgic encephalomyelitis/chronic fatigue syndrome? Front Med. (2021) 7 (1132). doi: 10.3389/fmed.2020.606824
212. Wildtving T, Holt N. The neurological symptoms of COVID-19: a systematic overview of systematic reviews, comparison with other neurological conditions and implications for healthcare services. Therapeutic Adv Chronic Dis. (2021) 12:2046022320976979. doi: 10.1177/2046022320976979
213. Proal AD, VanElzakker MB. Long COVID or post-acute sequelae of Covid-19 (pasc): an overview of biological factors that may contribute to persistent symptoms. Front Microbiol. (2021) 12:698169. doi: 10.3389/fmicb.2021.698169
214. Mackay A. A paradigm for post-Covid-19 fatigue syndrome analogous to ME/CFS. Front Neurol. (2021) 12:701419. doi: 10.3389/fneur.2021.701419
215. Komaroff AL, Lipkin WI. Insights from myalgic encephalomyelitis/chronic fatigue syndrome may help unravel the pathogenesis of postacute COVID-19 syndrome. Trends Mol Med. (2021) 27:895–906. doi: 10.1016/j.molmed.2021.06.002
216. Camella PH, Gonzalez-Koolwa E, Kosoy R, Charney AW, Peredejordi JF, Chandrasekar S, et al. A molecular network approach reveals shared cellular and molecular signatures between chronic fatigue syndrome and other fatigueing illnesses. [Preprint] medRxiv. (2021). Available online at: https://www.medrxiv.org/content/10.1101/2021.01.29.21250755v1 (accessed October 5, 2021).