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ABOUT COVER

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Effect of a fever in viral infections — the ‘Goldilocks’ phenomenon?

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Abstract

Acute infections, including those due to Coronaviridae and other viruses, often stimulate a febrile response. A mild fever appears to improve outcome; it appears to diminish viral replication by several mechanisms, including virion entry into host cells and genome transcription, and improving host defence mechanisms against the pathogen. However, a fever may also damage host cellular and tissue function and increase metabolic demands. At temperatures at the lower end of the febrile range, the benefit of the fever appears to outweigh the detrimental effects. However, at higher temperatures, the outcome worsens, suggesting that the disadvantages of fever on the host predominate. A non-infective fever is associated with a worse outcome at lower temperatures, suggesting that hyperthermia carries less benefit in the absence of infection. This review discusses the risks and benefits of a fever on the host response, focusing on the effects of a fever on viral replication and host response, and the detrimental effect on the host.

Key Words: Fever; Virus; Coronavirus; Infection; Body temperature regulation; Hyperthermia

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Core Tip: Acute infections, including those due to coronavirus and other viruses, often stimulate a febrile response. A mild fever appears to improve outcome; it appears to diminish viral replication by several mechanisms, and improve host defence mechanisms against the pathogen. At higher temperatures, the outcome worsens, suggesting that the risks of fever on the host outweigh the benefit. A non-infective fever is associated with a worse outcome at lower temperatures. This paper discusses

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why a mild fever may be better than no, or very high fever.

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INTRODUCTION

The presence of a fever in infection and its potential benefit has been known for over two millennia, with the Greek philosopher Parmenides suggesting: “Give me the power to produce fever and I’ll cure all disease”^[1].

In humans, normal core temperature varies between around 36.5 °C and 37.8 °C^[2]. A raised temperature may be due to processes when the hypothalamus has a new regulatory set point causing the temperature to rise (for example, infection or inflammation), often called *fever*. Other incidences of a raised temperature may be due to heat gain in excess of heat loss (for example, in heat waves, after exertion or exercise, or illicit drug-taking), despite no change in the regulatory set point, a process often defined as *hyperthermia*^[3]. Definitions of different degrees of fever also vary between different sources, but one consensus statement defines a fever in humans as equal to or greater than 38.3 °C^[4], that is, 0.5 °C higher than the physiological range. Similarly, there is not a consensus on the definition of extreme hyperthermia, but there are significant effects above around 40 °C, suggesting that this temperature or higher poses particular risks.

A fever has numerous effects on the virus and on the human host. Whether a fever is useful or should be treated remains unclear. This review summarises the benefit and disadvantages of fever in acute viral infection, including what is known about pyrexia in coronavirus infection.

OBSERVATIONS FROM NATURE

Fever is a common immunological response to bacterial and viral infections, and is conserved across the animal kingdom, suggesting that it might provide an evolutionary benefit^[5].

A mildly raised temperature in patients with infection in the first 24 h following admission to the intensive care unit (ICU) is associated with a better outcome compared with normothermia or hyperthermia above 40 °C^[6,7]. In a study of elderly patients with pneumonia, survival was significantly lower in patients who lacked fever (29%) when compared with patients who developed a febrile response (4%)^[8]. A temperature greater than 38.2 °C is protective against invasive fungal infections in critically ill patients^[9].

In contrast with a fever in response to sepsis, a non-pyrogenic fever does not appear to have any benefit. A temperature of 37.5 °C or greater at any point during an ICU admission trends towards a worse outcome, and becomes significant at temperatures greater than 38.5 °C^[7].

Similarly, there is an increase in mortality for non-infectious neurological conditions (traumatic brain injury or stroke) in ICU at temperatures above 38.0 °C within the first 24 h, compared with normothermia; in patients with a brain infection, no significant increase in mortality is seen compared with normothermia until the temperature reaches above 40 °C^[10].

However, fever may come at a cost: Fever during an ICU admission is associated with greater organ dysfunction, prolonged hospitalisation, and prolonged periods of mechanical ventilation^[11,12]. A randomised controlled trial found that the return to normothermia from 38.5 °C in critically ill, shocked patients resulted in significant vasopressor reduction by 50% at 12 h, improved early mortality outcomes, and higher rates of shock reversal^[13].

In the treatment of acute febrile illness, antipyretics are often recommended. However, there is little evidence of benefit^[14], and evidence that they may prolong acute varicella zoster infection without any amelioration of symptoms^[15,16], prolong the

duration of acute rhinovirus illness^[17], and may worsen mortality^[18]. In a recent systematic review, antipyretics suggested an increased mortality when used in influenza infections in animals^[19].

In addition to humans, other warm-blooded animals^[20], cold-blooded animals^[21,22] and indeed plants^[23] raise their temperature when acutely infected with a pathogen, and in animals at least^[20-22], survival is often improved.

EFFECT OF FEVER ON VIRUS

Viruses, from the Latin word *virus*, meaning *poison*, are tiny ubiquitous micro-organisms, usually a few hundred nanometers in size. A virion contains a single nucleic acid (RNA or DNA) core surrounded by a protein coat, and requires the cellular processes of animals, plants, and bacteria in order to replicate.

Viruses are often classified according to the Baltimore system^[24], named after the professor of biology and Nobel laureate David Baltimore. Seven Baltimore groups are described, categorised by whether the viral genome is made of deoxyribonucleic acid (DNA) or ribonucleic acid (RNA), whether the genome is single- (ss) or double-stranded (ds), whether the sense of a single-stranded RNA genome is positive or negative, and whether reverse transcriptase (RT) is involved in replication (see Table 1). All of the viruses below may cause fever in acute infection, although the degree of fever generation depends on the virulence of the micro-organism and host response to the infection.

Coronaviruses are a group of related single-stranded, positive-sense large RNA viruses that cause diseases in mammals and birds. In humans, these viruses cause respiratory tract infections that can range from mild (for example, the common cold), to lethal [for example, severe acute respiratory syndrome (SARS), Middle East respiratory syndrome (MERS), and SARS coronavirus 2 (SARS-CoV-2)]. They have characteristic club-shaped spikes that project from their surface, similar to the solar corona, from which their name derives.

That a febrile response to acute viraemia may be beneficial to host survival and inhibitory to viral replication and shedding has been known since at least the 1970s^[25-27]; more recent work has improved the understanding of the mechanisms involved.

Basic viral replication

Viruses, including the coronaviruses, are acellular and cannot replicate without incorporating themselves into a host cell and relying on their host cells to provide the basic machinery to allow replication.

The first stage in the viral life-cycle is entry of the virion into the host cell. This often requires interaction with a specific receptor on the host cell surface, allowing fusion of the host cell and viral membranes, or endocytosis. In the case of SARS-CoV-2, this host cell receptor has been identified as the angiotensin-converting enzyme-2 (ACE2) protein^[28]. Alternative mechanisms for some viruses to enter the host cells are suggested; for example, the African swine fever virus (ASFV) may enter host cells *via* micropinocytosis (non-receptor-mediated entry, involving non-selective endocytosis of fluid and solutes)^[29]. ASFV infection in pigs often results in high fever in hosts.

Once inside the host cell, the viral genome can be transcribed and replicated. The viral capsid is first removed and degraded by viral enzymes or host enzymes, which releases the viral genomic nucleic acid. The method of replication of the viral genome varies between DNA and RNA viruses and between the structure of the genome. The process culminates in the synthesis of new viral proteins and genome, which are then assembled into new virions and released from the host cell.

The method of viral release from the host cell depends on the virion structure. Enveloped viruses are often released by budding, acquiring a phospholipid envelope from the host cell. Nonenveloped viruses exit an infected cell by lysis, resulting in destruction of the host cell.

The effect of pyrexia on viral replication

The presence of a pyrexia appears to affect various stages of the replication process. Temperature appears to affect entry of the virion into cells *via* a number of mechanisms. Temperature has a direct effect on the fluidity of cell membranes, with early work suggesting that the increase in fluidity with increasing temperatures is a consequence of virion entry into the cell^[30], with variable effects on viral entry. For example, membrane fluidity plays a key role in the accumulation of binding sites for

Table 1 Examples of fever-causing viruses, categorised according to the Baltimore classification

Class	Type	Examples of virus	Examples of disease in humans
I	dsDNA virus	Adenoviruses	Bronchitis, pneumonia, gastroenteritis
		Herpesviruses	Herpes, chickenpox
II	ssDNA virus	Parvoviruses	'Slapped cheek syndrome', arthritis
III	dsRNA virus	Reoviruses	Gastroenteritis
IV	ssRNA virus	Coronaviruses	COVID-19, SARS, MERS
		Togaviruses	German measles, encephalitis
V	ssRNA virus	Orthomyxoviruses	Influenza
		Rhabdoviruses	Rabies
VI	ssRNA-RT virus	Retroviruses	AIDS, role in cancer
VII	dsDNA-RT virus	Hepadnaviruses	Hepatitis

ss: Single-stranded; ds: Double-stranded; RT: Reverse transcriptase; COVID-19: Coronavirus disease 2019; SARS: Severe acute respiratory syndrome; MERS: Middle East respiratory syndrome; AIDS: Acquired immunodeficiency syndrome.

human immunodeficiency virus 1 (HIV-1) cell entry; increases in temperature enhance HIV-1 adsorption and infectivity^[31]. In contrast, increased membrane fluidity may be inhibitory to hepatitis C virion entry^[32]. Temperature-dependent increases in fluidity appear to stimulate T-cell activation^[33], suggesting enhanced viral detection and clearance. Whether the entry of SARS-CoV-2 into cells is affected by temperature-induced changes in host cell membrane fluidity remains unclear.

The influenza virus is internalised *via* receptor-mediated endocytosis, which requires an acidic environment for the endosome to trigger viral and endosomal membrane fusion^[34,35]. High temperatures appear to increase endosomal pH, adversely affecting influenza virus entry and intracellular transportation^[36].

Once inside the host cell, the efficacy of viral genome transcription and replication may be modulated by changes in temperature. For coronaviruses, as with other RNA viruses, replication is performed by an RNA-dependent RNA polymerase^[37]. Features of this enzyme are remarkably conserved across RNA viruses^[38], including in the recently described SARS-CoV-2 RNA polymerase^[39], and thus temperature-dependent effects on its activity in other viruses may be analogous. Temperatures greater than 41 °C have been demonstrated to destabilise viral RNA polymerase in highly pathogenic avian influenza (HPAI), reducing its replicative ability^[40]. While the viral messenger RNA transcription may be maintained at higher temperatures, genome replication is markedly reduced, due to reduced association of the polymerase with positive-sense RNA templates^[41]. A raised temperature also appears to cause a reduction in the cytokine interleukin (IL)-6 levels after human cell infection with influenza virus^[36]; IL-6 is required for a number of cytokine-mediated processes involved in viral replication^[42].

However, certain influenza A strains may adapt to different host temperatures; reassortment of the RNA polymerase subunits has been shown to confer increased thermal stability^[43,44], and this ability may allow the virus to cross species barriers.

EFFECT OF FEVER ON HOST IMMUNITY

The human immune system protects against invasive pathogens and is composed of innate and adaptive components. The innate system provides nonspecific defence mechanism (physical barriers and stimulation of inflammation and recruitment of white blood cells, complement and cytokines). The adaptive component is specific to a particular pathogen, mediated by antibodies and complement proteins. Almost every step of both components appears to be stimulated by generation of a fever^[21].

Innate immunity

Neutrophils are the first immune cell populations to be recruited to a site of infection, but their precise role has not yet been determined^[45]. They appear to display a number

of antiviral mechanisms, including phagocytosis and destruction of infected cells, blockage of viral genome replication, and production of antiviral cytokines^[45]. Neutrophil production^[46] and activity^[21] are increased by the presence of a fever, and a fever appears to improve attraction and migration of neutrophils to the site of infection^[47].

Similarly, fever stimulates the cytotoxicity of natural killer (NK) cells and increases migration to the site of infection^[21]. While the precise role is again undefined, NK cells are likely to contribute to cytolytic killing of virus-infected cells, production of pro-inflammatory cytokines, and stimulation and recruitment of adaptive immune cells^[48]. Elevated temperatures inhibit expression of MHC class I proteins and increase heat shock protein 70 (HSP70) production by tumour cells, both of which enhance NK cytotoxicity. Whether a similar response is seen in virus-infected cells is unclear^[49].

Dendritic cells (DC) serve as antigen-presenting cells (APCs), by processing antigen material and presenting it on the cell surface to components of the adaptive immune system. Febrile temperatures have a variety of actions on DCs, including enhancing phagocytic activity, and increasing production of interferon-alpha (IFN α). Interferon-alpha appears to have a number of effects in acute viral infection, many inhibiting viral replication^[50]. Expression of MHC class I and II molecules and co-stimulatory proteins, for example CD80 and CD86, is increased, enhancing T- and B-lymphocyte activity^[21]. Migration of dendritic cells is also enhanced by a febrile temperature, with increased migration of epidermal DCs to the lymphatic system^[51].

Adaptive immunity

Lymphocyte trafficking is vital to ensure lymphocytes migrate to the required site of inflammation and through lymphoid tissue; it allows interaction between dendritic cells and lymphocytes, and ensures that there is exposure to antigen-presenting cells. Febrile temperatures enhance a number of mechanisms involved with effective migration, including binding of adhesion molecules and stimulation of endothelial trafficking molecules^[21]. T-lymphocyte stimulation and cytotoxicity is enhanced by febrile temperatures^[52,53].

Heat shock proteins

Heat shock proteins (HSPs) are a family of proteins that are produced by cells in response to thermal stress and other stressors including exposure to cold, UV light toxins and hypoxia, and during wound healing or tissue remodelling. HSPs are one of the most phylogenetically conserved classes of proteins and are found in virtually all living organisms. They have a wide range of critical roles in maintaining cellular homeostasis and in protecting cells from stressful conditions, reflecting in large part their ability to facilitate folding of nascent protein or refolding of denatured protein.

In the context of viral infections, HSPs are stimulated both by viral-induced fever and by viral infection itself^[54]. They have effects on both the pathogen and host; the exact interaction is still not well understood, and it is still debated whether the outcome may benefit the host or the pathogen. They have been variously described as having an enhancing^[55-59] or deleterious^[54,60] effect on the mechanisms of viral replication or activation. In particular, HSP70 has been implicated as a chaperone for the replication complex in influenza A infection^[61], although conversely may inhibit the nuclear export of newly synthesised influenza ribonucleoprotein complexes and thus viral morphogenesis^[60]. In fruit flies, the absence of heat shock transcription factor results in higher mortality in viral infection, suggesting that heat shock proteins may confer a benefit to survival in viral infection^[62]. However, other organisms, such as salmonella, rely on stimulation of HSPs to enhance infectivity^[63], suggesting the variable and pleiotropic effects of HSPs in viral replication and host defence.

There is little understanding currently of the interaction between heat shock proteins and the Coronaviridae family, but recent work has suggested that the SARS-CoV uses host cellular endoplasmic reticulum as a site for the synthesising and processing of its viral proteins, and that it can induce the unfolded protein response (UPR)^[64]. The UPR is a ubiquitous cellular stress response related to endoplasmic reticulum (ER) stress, which is activated in response to an accumulation of unfolded or misfolded proteins in the lumen of the ER. The UPR aims to restore normal function of the cell by a number of mechanisms, including halting protein translation, degrading misfolded proteins, and activating signaling pathways involved in protein folding. If restoration of function is not achieved, apoptosis may be triggered^[65]. HSP90 appears to improve stability of associated intracellular protein kinases^[66], with HSP90 inhibition accelerating cell lysis^[67]. There is therefore a potential role for HSP90 inhibitors, known to have chemotoxic effects, as cytotoxic agents against coronavirus-infected cells^[68].

Effect on pharmacology

A further effect of the febrile response is the effect on the efficacy of various medications. One study of 17 different antibiotics on 432 strains of bacteria at temperatures between 35 and 41.5 °C showed a progressive increase in antimicrobial activity as the temperature increased^[69]. Little is known about the effect of temperature on the activity of antiviral pharmaceutical agents.

DETRIMENTAL EFFECTS OF HYPERTHERMIA ON THE HOST

If there were no alternative benefit, conventional preconception would suggest that any departure from physiological homeostasis would be deleterious to an organism. Thermal dysregulation is no exception to this, and there appear to be injurious consequences observed in hyperthermia in humans. The deleterious effects of hyperthermia have been described similarly in a variety of causes, suggesting that it is the presence of a raised temperature, rather than the cause, that is primarily important. While much of the evidence comes from non-infective heat injury, biological plausibility suggests that similar effects also occur in infective fevers.

Mechanisms of injury of hyperthermia

The mechanism of injury in hyperthermia, whether due to an infective or non-infective cause, appears to result from both direct heat stress on molecular, cellular and tissue-level structures (see [Figure 1](#)), and from local and systemic cytokine-driven responses. Pyrogenic cytokines such as IFN γ , IL-1b and IL-6, which are closely associated with the viral fever pathway^[70], have been observed in mechanisms responsible for capillary leak, myocardial suppression, and systemic inflammation^[71]. IL-1 and IL-6 knockout mice tend to manifest an attenuated febrile response^[72,73]. Longer duration and higher magnitude of IL-6 peaks in heatstroke patients are associated with poor outcome irrespective of maximum core temperature^[74], and are proportional to symptoms and the degree of fever in influenza^[70]. However, the pleiotropic nature of these cytokines makes it difficult to accurately model a causal relationship in disease.

Effects on cellular mechanisms

Hyperthermia results in disruption of transmembrane transport mechanisms, affecting ionic flux and gradient balance^[75]. Impaired protein and nucleic acid synthesis is also observed^[75], which may continue beyond re-establishment of normothermia^[76]. At temperatures of 40 °C, nuclear processes are interrupted by damage to nuclear scaffolding structures^[77]. Direct cellular death is observed above 41 °C, and the increase in rate of cell death rises significantly beyond this point^[75,77].

Effects on barrier dysfunction

Temperatures in excess of around 39-40 °C are associated with an increase in enteric epithelial permeability^[78,79], and the translocation of enteric bacteria and endotoxins, improved when subsequently treated with antibiotics against enteric bacteria^[80-82]. This is supported by evidence of bacterial infections and elevated circulating procalcitonin levels, a sensitive biomarker of sepsis, in more than 50% of heatstroke patients^[83].

Neuroprotection conferred by the highly selective permeability of the blood-brain barrier (BBB) is increasingly lost at temperatures of greater than 38-39 °C with loss of structural integrity^[84]. The changes in endothelial permeability of the BBB occur at similar temperatures to the endotoxaemic process postulated in the gastrointestinal (GI) tract, and this association may contribute to neurological impairment^[85]. A prospective observational study of ICU patients describes intestinal permeability as a predictor of development of multiple organ dysfunction syndrome (MODS)^[86].

Poor endothelial barrier function in hyperthermic states is also associated with cytokine-mediated extravasation of neutrophils and macromolecules from lung vasculature which, compounded with increased oxygen toxicity, aggravates acute lung injury in critical illness at temperatures above around 39 °C^[47,87,88].

Organ- and system-specific dysfunction

In addition to epithelial barrier dysfunction in the GI tract as described above, hyperpyrexia is associated with disruption of cellular structural mechanisms and exacerbation of oxidative stress in splanchnic tissue as a consequence of increased free radical production at the tissue level^[89]. Intestinal ischaemia is observed, with circulatory diversion away from splanchnic tissues^[78].

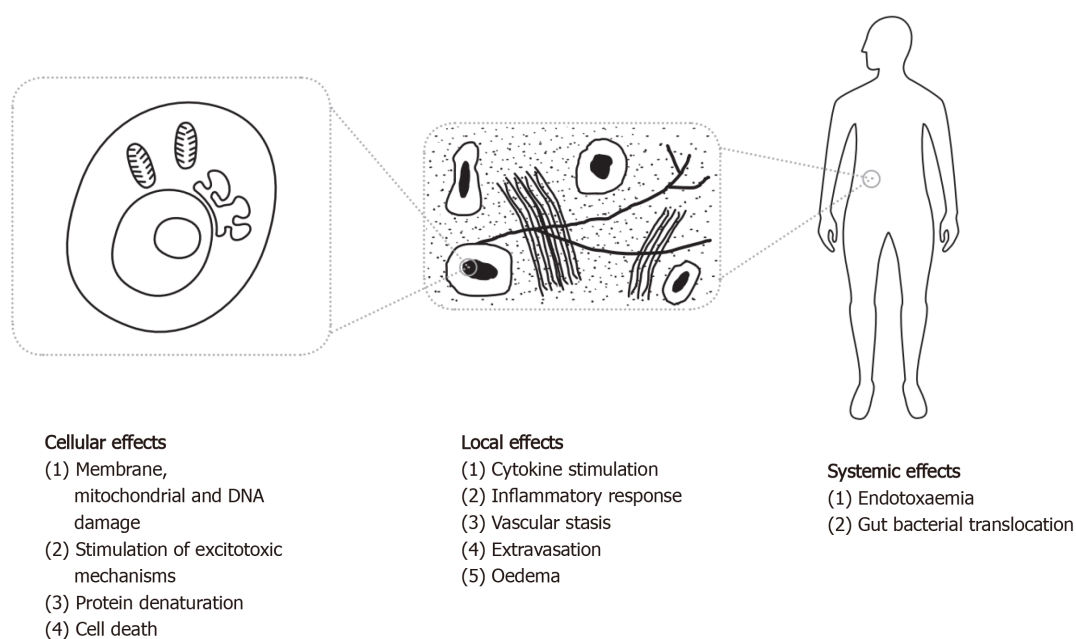


Figure 1 Diagrammatic representation of the mechanisms of damage from hyperthermia. Citation: Walter EJ, Hanna-Jumma S, Carraretto M, Forni L. The pathophysiological basis and consequences of fever. *Crit Care* 2016; 20: 200. Copyright© The Authors 2016. Published by Springer Nature Group.

Neurological cytotoxicity and dysregulation are well documented in heat stress. There is evidence of higher cortical dysfunction, which may be chronic or permanent^[85,90], and neuroradiological studies suggest a number of discrete pathologies, including oedema, ischaemia, haemorrhage, and encephalitic and accelerated degenerative changes^[91,92]. Many parts of the central nervous system are vulnerable^[91,92], but the Purkinje neurons are particularly susceptible^[85,93]. Mammalian studies suggest that this particular thermosensitivity is related to the preferential expression of heme oxygenase-1 in the Purkinje cells^[94].

Intravascular and haematological homeostasis is also disrupted by hyperthermia. Animal models demonstrate vasoplegia, venostasis and capillary hyperpermeability at sustained temperatures of greater than 40 °C^[95]. There is evidence of coagulopathy in more than 40% of cases, compounded by impaired platelet aggregation above 38 °C^[96,97]. Hepatic dysfunction and hepatocellular structural damage is often observed alongside coagulopathic states in hyperthermia^[98-100].

Acute kidney injury, sometimes sufficiently severe to require renal replacement therapy, has been described across heat stress conditions of various aetiologies^[96,101,102], the magnitude of which correlates with mortality^[101]. Similar to dysfunction in other organs, renal capillary bed dilatation, vascular stasis, and interstitial haemorrhage are seen^[95]. This may be further exacerbated by stimulation of the renin-aldosterone-angiotensin system which in turn reduces renovascular flow at temperatures as low as 38 °C^[103].

Metabolic cost of pyrexia

The association between pyrexia and hypermetabolism is well established; however, there are little data on the effect on patient outcome. A calorimetric study on 204 critically ill patients found fever to be the most significant single determinant of hypermetabolism of those observed, above even the presence of SIRS^[104]. Another study determined that cooling patients from 39.5 °C to 37 °C resulted in a reduction in oxygen uptake, carbon dioxide production, cardiac output, and oxygen extraction fraction of approximately 20%^[105]. A one-degree increase above 37 °C results in a 10% increase in metabolic rate^[106]. The ability of the organism to cope with this increased demand is likely to depend on the metabolic reserve of an individual patient, but may be significant.

CONCLUSION

Current epidemiological and biological evidence suggests that viral replication is

reduced and that human host defence mechanisms are enhanced at degrees of mild fever, compared with normothermia. However, at higher degrees of fever, (around 39-40 °C), mortality again increases, suggesting that any benefit at mildly elevated temperatures is outweighed by the damage the fever causes on the host. This 'U'-shaped curve, demonstrating improved outcomes at mild febrile temperatures, does not appear to be replicated in non-pathogenic states, suggesting that the damage that the fever exerts on the host is not matched by any perceived benefit, and that this permissive pyrexia may only be true with an infective aetiology.

What is less clear is the effect that modulating the temperature has on clinical outcomes, or the optimal temperature. The latter is likely to be impacted by the physiological reserve of the patient and their ability to tolerate the increase in metabolic demands that a fever requires.

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