The aldehyde hypothesis: metabolic intermediates as antimicrobial effectors

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There are many reactive intermediates found in metabolic pathways. Could these potentially toxic molecules be exploited for an organism's benefit? We propose that during certain microbial infections, the production of inherently reactive aldehydes by an infected host is a previously unappreciated innate immune defence mechanism. While there has been a significant focus on the effects of aldehydes on mammalian physiology, the idea that they might be exploited or purposefully induced to kill pathogens is new. Given that aldehydes are made as parts of metabolic programmes that accompany immune cell activation by the cytokine interferon-gamma (IFN-γ) during infections, we hypothesize that aldehydes are among the arsenal of IFN-γ-inducible effectors needed for pathogen control.

1. Introduction

All forms of life are susceptible to infection, whether it is by viruses, bacteria, fungi or parasites. Humans have evolved highly complex defence mechanisms against virtually all forms of infectious agents, and these mechanisms may be tailored to their targets. For example, while antibodies are critical for controlling many viral infections, they are often ineffective against pathogens that live within a host’s cells. Unicellular microbes, either bacterial or eukaryotic, are often captured by phagocytes such as neutrophils and macrophages. These soldiers of the immune system have numerous mechanisms to inhibit or eliminate microbial growth, including the production of pore-forming peptides, reactive oxygen and nitrogen species; the import and export of transition metals and exposure to destructive enzymes such as proteases and lipases (reviewed in [1]).

The human-specific pathogen Mycobacterium tuberculosis appears to frequently evade sterilization from infected hosts. As a consequence, and by some estimates, M. tuberculosis has infected nearly one-third of the world’s population, with most individuals controlling their infections without medical intervention. While this number has been recently debated [2], it is clear that M. tuberculosis kills about 1.7 million people annually, making it a leading cause of death by a single infectious agent [3]. Activation of macrophages by the cytokine interferon gamma (IFN-γ) is crucial for control of mycobacterial infections [4]. Historically, the importance of IFN-γ has been attributed to its ability to induce the expression of nitric oxide synthase (iNOS), and mouse studies have shown an essential role for nitric oxide (NO) in robustly controlling M. tuberculosis growth [5]. While these data provide compelling evidence of its importance, it has been challenging to demonstrate that NO has a definitive role in controlling M. tuberculosis in humans, given the paucity of humans defective for iNOS and the lack of a robust in vitro infection model using human cells. Furthermore, several animal studies have suggested a role for NO in the regulation of inflammation that is separate from its putative antimicrobial activity [6,7]. In addition to NO, copper (Cu) may also help control M. tuberculosis infections [8–11]. In vitro macrophage data show the mobilization of the...
Mycobacterium tuberculosis

2. Mycobacterium tuberculosis is highly susceptible to changes in its aldehyde metabolism

Several studies have indicated that M. tuberculosis is highly susceptible to aldehydes. Disruptions in various mycobacterial metabolic pathways can result in increased intracellular aldehydes, reducing bacterial survival under in vitro and in vivo conditions. In one study, a screen for compounds against M. tuberculosis identified a class of pyrimidine-imidazoles (PIs) that strongly inhibit growth, possibly by increasing the amount of several intracellular aldehydes [14]. Pethe et al. [14] found that PIs cause M. tuberculosis to accumulate glycerol phosphate while depleting adenosine triphosphate (ATP). Glycerol phosphate can enter the glycolytic pathway and be converted into dihydroxyacetone phosphate (DHAP), which isomerizes into glyceraldehyde-3-phosphate (GAP). Furthermore, methylglyoxal (MG), also known as pyruvaldehyde, can spontaneously form from DHAP and is known to be toxic. Interestingly, the authors of this study demonstrated that the exogenous addition of GAP, DHAP and MG to cultures of M. tuberculosis is toxic, demonstrating the direct antimicrobial activity of these aldehydes or aldehyde precursor (DHAP). Further supporting the authors' hypothesis that the accumulation of glycerol phosphate or its downstream products leads to toxicity, a genetic screen for M. tuberculosis mutants resistant to the PIs identified mutations in glpK (glycerol kinase, Rv3696c). Because GlpK phosphorylates glycerol into glycerol-3-phosphate, its disruption could potentially prevent the accumulation of GAP, DHAP and MG, thereby reducing overall aldehyde levels in bacteria grown in glycerol.

Given that several aldehydes possibly accumulate in M. tuberculosis treated with PIs, Pethe et al. [14] proposed that the PIs targeted one or more aldehyde detoxification enzymes. To test this hypothesis, the authors pulled down proteins that interact with PIs and identified four putative MG detoxification enzymes. Ectopic production of one of these proteins Rv0577 improves bacterial growth in glycerol and provides some resistance to two of the PI compounds, suggesting it has a role in aldehyde detoxification. A caveat to these studies is that the ability of the PIs to directly inhibit the activities of one or more of the identified enzymes was not tested. Interestingly, GlpK was also the focus of two recent independent studies that determined glycerol metabolism in vitro or in vivo makes M. tuberculosis more susceptible to several front-line tuberculosis (TB) drugs. The Alland group found that naturally occurring frame shift mutations in glpK appear to be selected for in M. tuberculosis clinical isolates, while the Sassetti laboratory used a transposon sequencing screen to find mutants that provided increased fitness in antibiotic-treated mice [15,16]. Both groups found defects in glycerol metabolism made bacteria more tolerant to several different antibiotics. While neither study determined how disruption of glpK protected M. tuberculosis, Sassetti and co-workers [7] proposed that MG or other products of glycerol metabolism could increase antibiotic efficacy. It is tempting to speculate that aldehydes like MG or GAP could help potentiate the activity of one or more drugs against M. tuberculosis and other microbes.

In addition to interference with glycerol metabolism, several other laboratories noted that disruption of certain reactions around the tricarboxylic acid cycle or glyoxylate shunt result in the accumulation of aldehydes in M. tuberculosis. In a study by Puckett et al., deletion of the malate synthase gene (Rv1837c) resulted in the accumulation of glyoxylate, making bacteria highly attenuated for in vitro and in vivo growth [17]. Similarly, a study by the Nathan laboratory determined that disruption of 2-hydroxy-3-oxoadipate synthase (HOAS, Rv1248c), the E1 component of α-ketoglutarate dehydrogenase, results in the accumulation of succinate semialdehyde and glyoxylate [18]. A mutant defective in HOAS is highly sensitive to NO, although the connection between HOAS and NO sensitivity is unclear. These studies aligned well with a report from the Darwin laboratory; work in the Nathan laboratory previously determined that M. tuberculosis mutants defective in proteasome activity are highly sensitive to NO, although it was unknown how protein degradation was linked to this phenotype [19]. Remarkably, a genetic suppressor screen found that a single proteasome substrate called Log (lonely guy, Rv1205) is responsible for the NO sensitivity of proteasomal mutants. Log catalyses the final step in the production of signalling molecules known as cytokinins, which can be broken down into adenine and various aldehydes. Indeed, cytokinin-associated aldehydes are sufficient to sensitize wild-type M. tuberculosis to NO [19]. Interestingly, the Darwin laboratory also found that a mutation in glpK suppresses the NO-sensitive phenotype of a proteasome mutant, further supporting a link between aldehyde accumulation and NO sensitivity [19]. Collectively, these data strongly suggest M. tuberculosis must tightly regulate its intracellular aldehyde levels to prevent toxicity from aldehydes alone, as well as mitigate sensitization of the bacteria to NO and possibly other chemicals by the aldehydes. However, more research is required to clearly establish mechanistic links between intracellular aldehyde accumulation and bacterial fitness.

3. Phagocytes produce a variety of aldehydes

There are abundant data that show macrophages and neutrophils produce copious amounts of aldehydes. For example, Heinecke and co-workers [20] showed that activated neutrophils can oxidize almost any amino acid into an aldehyde. This process requires myeloperoxidase, which is released from granules in neutrophils during infections, as well as by hydrogen peroxide (H2O2) produced by an oxidative
burst in phagocytes. Reactive oxygen species (ROS) such as H$_2$O$_2$ can also oxidize lipids into highly toxic lipid aldehydes including 4-hydroxynonenal (4-HNE) and malondialdehyde (MDA).

The activation of macrophages and other cell types to produce ROS and mobilize other antimicrobial effects requires IFN-$\gamma$, which is one of the most important cytokines needed for the control of $M$. tuberculosis [4]. The downstream effectors of IFN-$\gamma$ signalling needed to control bacterial growth are, however, incompletely understood. Recently, the Stanley and other laboratories have shown that IFN-$\gamma$ strongly induces aerobic glycolysis during $M$. tuberculosis infections of mouse or human macrophages [21–23]. Aerobic glycolysis, also known as the Warburg effect, is a pathway by which cells rely exclusively on glycolysis for the production of ATP, even in the presence of oxygen. Several laboratories have reported the observation that a ‘Warburg-like’ effect is induced in M1 macrophages in various infection settings and that this metabolic programme supports host control of infections with $M$. tuberculosis, Legionella pneumophila and Streptococcus pneumoniae [22,24,25]. Furthermore, the IFN-$\gamma$ responsive transcription factor HIF-1$\alpha$ requires aerobic glycolysis for its stabilization in macrophages during immune activation. The fact that HIF-1$\alpha$ is essential for control of $M$. tuberculosis as well as other pathogens, strongly suggests that aerobic glycolysis is an integral component of a host-protective immunometabolic programme [21,22,25].

Relevantly, the ability of Salmonella Typhimurium and Brucella abortus to establish chronic infection in M2 macrophages, which rely on oxidative metabolism rather than aerobic glycolysis, has been linked to excess glucose levels. It has also been proposed that the induction of aerobic glycolysis is exploited by various pathogens for their benefit, for example, by providing nutrients like lactate or lipids that benefit bacterial growth (reviewed in [26]). However, given that glycolytic intermediates or side-products are aldehydes in mammals just as they are in bacteria, we propose that the induction of glycolysis is a host-beneficial adaptation with respect to $M$. tuberculosis and possibly other infections.

4. The aldehyde hypothesis

Based on data from the Stanley laboratory, we took a closer look at glycolysis as a pathway that generates aldehydes to contribute to pathogen control downstream of IFN-$\gamma$ activation. Estimates of GAP and other glycolytic intermediate concentrations in cells range from 15 $\mu$M to 1.5 mM, depending on the metabolic state of the cell [27,28]. Relevantly, $M$. tuberculosis-infected macrophages release up to 1.6 mM MG into culture supernatants [29]. At the time of this study, the authors hypothesized that the mycobacteria produce MG to kill macrophages; however, it is more likely that macrophages produce MG in response to infection. Indeed, even sterile immune activation with lipopolysaccharide results in high levels of MG production in macrophages [30–33].

We also hypothesize that the decreased detoxification of aldehydes could benefit a host during infections. A substantial proportion of the world’s population of Asian descent harbour a specific loss-of-function mutation in aldehyde dehydrogenase 2 (ALDH2), which greatly reduces its activity. The mutant allele is known as ALDH2*2 (rs671), which has a codon 487 glutamate to lysine substitution. Hetero- or homozygous carriers of ALDH2*2 who drink alcohol accumulate acetaldehyde, the toxicity of which results in a characteristic facial flushing response. Given that the ALDH2*2 allele incurs deleterious effects even in the absence of alcohol consumption, it appears that there was a strong selective pressure that maintained this mutation in humans. Shin and co-workers [34] performed an association analysis among a cohort of Korean men with various disease states and the ALDH2*2 allele. Strikingly, the only significant correlation they observed was individuals with TB were less likely to have the mutant allele. Much like the mutation that causes sickle cell anaemia protects against malaria in Africa, it is possible that the proliferation of the ALDH2*2 allele served an analogous function against $M$. tuberculosis infections in Asia.

The authors of this study presumed a healthier lifestyle caused by decreased alcohol consumption in individuals with inactive ALDH2 contributed to more resistance to acquiring TB. While alcohol abuse is linked to poor TB outcomes, the precise reasons are unknown and may have either to do with immune fitness, antibiotic regime compliance or a combination of both [35]. Given that it is estimated the ALDH2*2 allele arose approximately 3000 yr ago [36], it is impossible to know if alcohol consumption played a role in TB epidemiology at that time. Perhaps another compelling argument for a protective effect of aldehydes conferred by the ALDH2*2 allele is the appearance of the so-called hypervirulent ‘Beijing’ or ‘W’ $M$. tuberculosis strains. This lineage appears to have established in China between 2000 and 7000 yr ago [37,38]. While the mechanisms (or definitions) of bacterial hypervirulence are somewhat unclear, these strains have resulted in several outbreaks outside of China, suggesting that they are more transmissible or virulent. We speculate that these strains evolved to be more transmissible within a population with higher intrinsic resistance to $M$. tuberculosis, either due to increased resistance to aldehydes or by reducing the induction of aldehyde-generating pathways by infected cells.

In addition to their anti-pathogen activity, aldehydes are also reactive with and damaging to self. Thus, humans have evolved to make numerous enzymes that convert aldehydes into less harmful molecules. The glyoxalase system in humans, consisting of the enzymes glyoxalase 1 (GLO1), glyoxalase 2 (GLO2) and reduced glutathione, detoxifies MG into D-lactate and is present in all mammalian cells. Modulation of MG levels by GLO1 is thought to influence the pathogenesis of numerous diseases including diabetes, atherosclerosis and neurodegenerative disorders, among others [39]. MG levels are high during sepsis, a hyper-inflammatory state usually caused by bacterial infection [32]. During inflammation, high levels of ROS are produced. ROS can directly suppress GLO1 function [39], and because ROS detoxification also requires glutathione, it can compete with the glyoxalase system. It is therefore possible that during infection, the production of ROS could lead not only to the reactive aldehydes 4-HNE and MDA but also to increased levels of MG. It would be interesting to test whether genetic or pharmacological inhibition of GLO1 could lead to enhanced control of bacterial infection.

5. A tuberculosis drug in hand?

Disulfiram (DSF) or antabuse was developed over 50 yr ago to treat alcohol abuse and works by inhibiting ALDH2,
with the hypothesis that these genes encode detoxifying products of this lipid aldehyde encountered in animal tissues. Two enzymes, RhaA1 and RhaA2, were identified to have detoxification activity in vitro. Heterologous expression of these genes in Bacillus subtilis, a non-pathogenic bacterial species, provides some protection against 4-HNE in vitro. While this study showed L. monocytogenes infection induces 4-HNE in infected mouse tissues, the rhaA genes do not provide a fitness advantage to the parental strain in mice. It is possible that additional or different gene products are needed for 4-HNE resistance or that deletion of rhaA genes results in the induction of compensatory pathways to mitigate aldehyde toxicity in vivo. Another reason why no difference was observed could be that given that L. monocytogenes is generally not pathogenic to immunocompetent hosts and can be readily cleared, 4-HNE-sensitive mutants might not necessarily show fitness defects in immunocompetent infection models.

While the mechanism(s) of aldehyde toxicity have not been determined in any of the above studies, it is possible that aldehydes target numerous pathways, collectively inactivating bacterial growth or survival in vivo. This non-specific, ‘death by a thousand cuts’ feature of aldehydes may make it an ideal weapon against invading microbes.

7. Going forward

We propose a model whereby mammalian macrophages exploit aldehydes produced during inflammation to target invading pathogens (figure 1). How could this model be tested, given that testing the contributions of specific host cell effectors in inhibiting bacterial growth can be challenging? In some cases, proposed effectors are essential for normal cellular processes in the host. Other challenges manifest in the numerous limitations of cell culture infection models. For example, in vitro cell culture models of M. tuberculosis infection are generally limited to 6–10 d, after which mimicking the ALDH2*2 mutation in humans [40]. More recently DSF was identified in a screen to repurpose existing clinical drugs for use in TB patients. The treatment of M. tuberculosis-infected mice with DSF results in a significant reduction in bacterial burden, comparable to treatment with the antibiotic rifampin [41]. Because the screen that identified DSF activity against M. tuberculosis was performed in the absence of mammalian cells, the authors of this study did not assume that the inhibition of ALDH2 was linked to the anti-tubercular activity of DSF in mice. In an effort to understand the mechanism of action of DSF, Wolschendorf and co-workers [42] found that in acidic or Cu-rich conditions, DSF breaks down into diethylthiocarbamate (DETC), which complexes with Cu ions. DETC-Cu complexes can enter M. tuberculosis bacilli, resulting in bacterial death. Thus, it is presumed that DSF directly kills M. tuberculosis in vivo in a Cu-dependent manner. By contrast, DSF does not use Cu to inactivate ALDH2 activity and instead is likely to catalyse the formation of a stable intramolecular disulfide bond between two active site cysteines in ALDH2 [43].

We do not know if DSF is readily converted to DETC in vivo or if there is enough bioavailable Cu available to complex with DETC if formed. These observations thus question a Cu-dependent anti-mycobacterial activity of DSF in vivo. We therefore hypothesize the inhibition of ALDH2 and concomitant aldehyde accumulation from various metabolic pathways including inflammation contributes to the killing of M. tuberculosis bacilli in mice treated with DSF. This activity could be in addition to a direct Cu-dependent killing mechanism on bacteria. The prospect of DSF having both a direct anti-bacterial effect and a ‘host-enabling’ mechanism of action could be a strong justification to move DSF into the clinics for the treatment of TB.
host cells die. Given that M. tuberculosis infections can last for years and even decades, it is difficult to envision a week-long experiment modelling the reality of natural infection. Additionally, in vitro monolucute fails to reproduce the complex cellular interplay encountered in vivo (i.e. the presence of T cells is critical for the robust control of mycobacterial infections). While these and other reasons have made it challenging to quantify the relative contributions of different effectors, mouse infection models along with bacterial mutants with variable aldehyde susceptibility should make the aldehyde hypothesis testable.

Data accessibility. This article has no additional data.

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