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Metabolic reprogramming: an innate cellular defense mechanism against intracellular bacteria?

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Abstract

The limited metabolic resources of a cell represent an intriguing “conflict of interest” during host-pathogen interactions, as the battle for nutrients might determine the outcome of an infection. To adapt their metabolic needs, innate immune cells such as monocytes, macrophages or dendritic cells reprogram their metabolism upon activation by microbial compounds. In turn, infection by intracellular bacteria provokes metabolic alterations of the host cell that benefit the pathogen. Here we discuss the state-of-the-art knowledge on metabolic reprogramming of host cells upon activation or infection with intracellular bacteria. The study of the host- and pathogen-driven metabolic alterations that seem to co-exist during infection is an emerging field that will define the metabolic pathways that might be targeted to combat infection.
Introduction

Cellular metabolism comprises the controlled biochemical processes that occur within a cell to maintain life. It is generally divided into anabolic processes (the build-up of macromolecules) and catabolic processes (the breakdown of macromolecules). To maintain constant cellular conditions in response to changes in the environment (homeostasis) a cell regulates its specific needs by the fine-tuning of these metabolic pathways.

During host-pathogen interactions, host and pathogen fight for the nutrients needed for cellular catabolism, as resources are generally limited [1,2]. Obtaining energy and ready-to-use basic molecules is key to build up new cellular components, to grow and reproduce, or to respond to environmental changes. Thus the result of this metabolic clash is key for the outcome of host-pathogen interactions and the infection process.

Cellular metabolism is one of the most complex and intricate cellular systems. To simplify, two main catabolic programs work to obtain energy, “glycolysis” and “oxidative phosphorylation” (OXPHOS) (Figure 1A). During glycolysis that is performed in the cytosol, cells convert glucose into pyruvate by anaerobic glycolysis and then pyruvate is fermented to lactate. Thereby 2 ATP molecules are generated per molecule of glucose. During OXPHOS the pyruvate generated by anaerobic glycolysis is routed to mitochondria where it is oxidized to CO$_2$ via the tricarboxylic acid (TCA) cycle. The released energy (stored as NADH) is used by the electron transport chain (ETC) to pump protons across the membrane. This generates an electrochemical gradient used to synthesize ATP by the mitochondrial F$_1$-F$_0$-ATPase.

Under the OXPHOS program, the cell produces 36 ATPs per molecule of glucose [3].

Interestingly, many proliferating cells (such as embryonic stem cells, cancer cells or activated T-cells) prefer the glycolytic program although this metabolic pathway is less efficient for ATP production as compared to OXPHOS. In contrast, tumour cells convert most of the absorbed glucose to lactate even under oxygen-rich conditions, a program that was termed “aerobic glycolysis” or the “Warburg effect” (Figure 1B) [4]. Thus elevated glucose uptake and lactate secretion are metabolic hallmarks of solid tumours [5,6]. Although the question why cancer cells prefer a “less efficient” metabolic pathway to grow is still a matter of debate, it seems that aerobic glycolysis coupled to fermentation is a fast process, easy to regulate, that fulfils all metabolic requirements of a proliferating cell. The Warburg program maximizes biosynthetic pathways by redirecting cytosolic glycolysis and the mitochondrial TCA cycle to the biosynthesis of nucleotides, amino acids and lipids (Figure 1B). The fast withdrawal of pyruvate and TCA intermediates in the mitochondria, which are normally
dedicated to the OXPHOS program and their redirection to biosynthetic pathways, explains
the reduction of oxygen consumption after the switch to the Warburg metabolism. Thus
Warburg metabolism seems to be induced to reprogram cellular metabolism towards
macromolecular synthesis.

Reprogramming to Warburg-like metabolism has been observed during activation of diverse
immune cells and, interestingly, also during infection of human cells by intracellular bacterial
pathogens such as *Mycobacterium tuberculosis* [7-11], *Legionella pneumophila* [12], *Brucella abortus* [13], *Chlamydia trachomatis* [14,15] or *C. pneumoniae* [16]. Although the Warburg
program in proliferating T-cells and cancer cells can be seen as program necessary for
proliferation, the induction of the Warburg program in infected cells raises intriguing
questions. Why non-proliferating immune cells, such as monocytes, dendritic cells (DCs) or
macrophages, switch to a Warburg-like metabolism during infection? What is the impact of
such metabolic changes on the pathogen that obtains its nutrients only from the infected host
cell? Is metabolic reprogramming beneficial or detrimental for intracellular bacteria? Are
these metabolic alterations driven by the host cell itself in response to bacteria, as a defence
mechanism, or are they induced by the pathogen for its own benefit?

**Non-proliferating immune cells reprogram metabolic pathways upon stimulation with
microbial compounds**

Monocytes, DCs and macrophages are differentiated, non-proliferative, innate immune cells
that, once activated by microbial stimuli, orchestrate the activation of other immune cells
through the secretion of immuno-regulatory cytokines, but they also increase glycolytic fluxes
and modulate their metabolism.

In DCs, glucose uptake is increased and accompanied by lactate production shortly after
their activation by microbial compounds that engage Toll-like receptors (TLRs) such as
bacterial lipopolysaccharide (LPS). After TLR stimulation, a signalling cascade involving
TBK1, IKKɛ and Akt promotes the association of the glycolytic enzyme hexokinase-2 (HK2)
with mitochondria increasing its enzymatic activity and therefore the glycolytic flux, however,
cellular ATP is still provided by OXPHOS [17]. At these early time points, glycolysis
supplies DCs with citrate, which fuels the synthesis of fatty acids that are required by
activated DCs to increase the size of the endoplasmic reticulum (ER) and the Golgi apparatus
[17]. Thus, the fast and early metabolic switch of DCs to a glycolytic program due to the
stimulation with microbial compounds is linked to fatty acid synthesis and the adoption of the
secretory state. At 24 h of DC activation by LPS, iNOS-derived NO inhibits cellular
OXPHOS, which almost disappears and DCs rely exclusively on glycolysis, a Warburg-like
metabolic program is installed [18].

In murine bone-marrow-derived macrophages (BMDMs), metabolic reprogramming to a
glycolytic Warburg-like program is necessary for transcription of the pro-inflammatory
cytokine IL1β. Upon LPS stimulation, BMDMs show a broken TCA cycle and intermediates,
such as succinate or citrate, accumulate [19]. Furthermore, LPS induces the up-regulation of
pyruvate kinase M2 (PKM2) dimer and tetramer levels. In its enzymatically inactive form
(dimers), increased PKM2 induces succinate accumulation [20]. PKM2 dimers also migrate to
the nucleus where they induce PKM2-mediated transcription of IL1β. In its enzymatically
active form (tetramers), LPS-induction of PKM2 also increases the rate of glycolytic fluxes,
as the reaction catalysed by PKM2 is the rate-limiting step of glycolysis. Accumulated
succinate stabilizes HIF1α, which can now translocate to the nucleus and, together with
PKM2 dimers, drive IL1β gene expression. Succinate in mitochondria is oxidized by
succinate dehydrogenase and leads to the reverse flow of the ETC, which produces
mitochondrial reactive oxygen species (mROS) that also induce HIF1α-mediated IL1β mRNA
expression [21]. Accumulation of citrate is due to the LPS-induced expression of the
mitochondrial citrate carrier, which diverts citrate from the mitochondrial TCA cycle to the
cytoplasm. Cytoplasmic citrate is then used to generate NO, ROS and prostaglandins by
BMDMs, as well as to produce the antimicrobial metabolite itaconate [22-24].

Taken together, exposure of murine macrophages to bacterial LPS reprograms their
metabolism from OXPHOS to glycolysis, rewiring TCA intermediates such as citrate and
succinate to biosynthetic pathways. As itaconate has direct antimicrobial proprieties against a
wide range of intracellular bacteria, such as M. tuberculosis, Salmonella enterica or
L. pneumophila [25], the redirection of citrate for the synthesis of itaconate represents a
prominent example of how metabolic reprogramming of activated macrophages becomes a
defence mechanism against infection by intracellular bacteria. In addition to its antibacterial
proprieties, itaconate has also immunoregulatory functions during infection due to its
inhibitory actions on the production of mROS and certain cytokines and chemokines as
shown for BMDMs infected with S. enterica serovar Typhimurium [26], and for mice
infected with M. tuberculosis [27].

Recently it was shown that human monocytes isolated from blood also shift to a Warburg-
like metabolism when stimulated with LPS, but not upon stimulation with the TLR2 ligand
Pam3Cys (P3C) or whole-pathogen lysates. Indeed, in P3C stimulated monocytes, increased glycolysis was accompanied by increased OXPHOS, which was required for retention of their phagocytic capacity and cytokine production [28]. Elevated succinate, itaconate and citrate levels were only observed in LPS-stimulated monocytes, but not in P3C-stimulated monocytes. Interestingly, exposure of human monocytes to *M. tuberculosis* whole cell lysates did not affect OXPHOS, in contrast to infection of monocytes with living *M. tuberculosis* or after exposure to LPS as OXPHOS reduction was observed [9,28]. Moreover, exposure of monocytes to *Escherichia coli* or *Staphylococcus aureus* whole-cell lysates leads to increased OXPHOS [28], suggesting that heterogeneous metabolic responses may be elicited depending on the specific pathogen and TLR activation pattern involved.

These results indicate that metabolic reprogramming of innate immune cells upon stimulation with microbial components rewrites cellular metabolism to biosynthetic pathways that drive growth of cell size and production and secretion of cytokines and antimicrobial compounds. It is critical to clarify whether these metabolic alterations are stimuli-specific (LPS vs. P3C), cell-type specific (macrophages vs. monocytes vs. DCs), host-species specific (mouse vs. human) and/or bacterial-species specific (*S. aureus* vs. *M. tuberculosis*).

**Intracellular bacteria reprogram the host metabolism to a Warburg-like metabolism**

*M. tuberculosis* [7-11], *L. pneumophila* [12], *B. abortus* [13], *C. trachomatis* [14,15], *C. pneumoniae* [16] or *Listeria monocytogenes* [29,30], were reported to modulate the metabolism of their host cells to a Warburg-like metabolism [31]. Yet only few studies addressed the metabolism of primary cells during infection thus one must be careful with the interpretation of metabolic changes as most cancer cell lines have an enhanced Warburg metabolism due to their tumour origin [31]. This has been demonstrated for *L. monocytogenes*, as high induction of glucose uptake and glycolysis was only observed in primary BMDMs, and not in the murine macrophage-like cells line J774A.1. However, intracellular bacteria replicated to five fold higher numbers within the cancer cell line than in primary cells, suggesting that the Warburg metabolism exhibited by cell lines might promote bacterial replication [29].

The changes to a Warburg-like metabolism induced by many intracellular bacteria come in different styles (Figure 3). *Chlamydia trachomatis* infection of primary human umbilical vein endothelial cells leads to a specific Warburg-like program. Cellular glucose uptake and the phosphate pentose pathway (PPP), a nucleotide biosynthetic pathway derived from the
glycolytic flux are increased [15] (Figure 3A, Chl). The infection-dependent upregulation of
the PPP supplies the pathogen with nucleotides for intracellular replication, which serve as
nutrients together with amino acids, and lipids that C. trachomatis needs to sequester from the
host cell [32]. Thus, the induction of this Warburg-like program during infection seems to
benefit the pathogen.

L. pneumophila induces a Warburg-like metabolism characterized by the upregulation of
glycolysis and reduction of OXPHOS dependent on the injection of the bacterial effector
MitF in the host cell that induces reduction of OXPHOS through fragmentation of
mitochondrial networks [12]. This metabolic reprogramming of infected cells to a Warburg-
like program appears essential for bacterial growth and might be related to the need of amino
acids, the main source of energy for growing L. pneumophila [3,33], which can be
synthesized from redirected glycolytic and/or TCA intermediates (Figure 3B, Lp).

M. tuberculosis infection induces a Warburg-like program when infecting human primary
macrophages, that is characterized by decreased OXPHOS and upregulation of glucose uptake
and glycolysis coupled with the redirection of glycolytic intermediates to the synthesis of
large lipid bodies that accumulate in the macrophage (Figure 3C, Mtb). These lipid bodies,
together with the lactate produced by glycolysis, serve as nutrients for intracellular growth of
the pathogen [34-36], suggesting that the biosynthetic role of the Warburg-like program
benefits the intracellular replication of M. tuberculosis.

Although it is not clear yet whether Salmonella enterica, induces a Warburg-like
metabolism in the host cells or not, it has been reported that S. enterica modulates the
metabolism of the cell it infects. During in vivo infection of mice with S. enterica serovar
typhimuriumOXPHOS is completely shut-off [37], while glycolysis might be induced [38].
However, the most prominent metabolic alteration induced by S. typhimurium seems the
piracy of glucose by the pathogen. In mouse BMDMs Salmonella disrupts glycolytic fluxes to
reroute glucose from host macrophages to its vacuole (Figure 3A, St) and thereby this
pathogen reduces host glucose availability [39,40]. Interestingly, inhibition of glycolytic
fluxes creates a unique metabolic defect that activates the NLRP3 inflammasome [40].

Intracellular bacteria modulate autophagy, a catabolic program and a cellular defense
system

Another important metabolic response to infection is the cellular process known as
autophagy, a self-degradative process that is key for balancing sources of energy in response
to nutrient stress and to remove intracellular pathogens in response to infection [41-43]. Interestingly, autophagy also regulates cytokine production and secretion in immune cells upon microbial stimulation [44,45].

Host-driven and pathogen-driven alterations of autophagy occur during infection with intracellular bacteria. As host-induced autophagy degrades invading pathogens, evasion of autophagy is key for pathogens such as *M. tuberculosis*, *S. typhimurium* or *L. pneumophila* that inhibit autophagy initiation upstream of autophagosome formation, or *Shigella flexneri* that evades autophagy recognition by masking the bacterial surface [46-49]. Interestingly, *L. pneumophila* encodes two secreted effectors, *RavZ* and LpSPL that inhibit autophagosome formation at two different stages [47,50]. Other intracellular bacteria such as *Anaplasma phagocytophilum*, *Yersinia pseudotuberculosis*, *Coxiella burnetii* and *Francisella tularensis* hijack autophagosomes during infection to redirect the by-products of the autophagic degradation of cellular components to their vacuoles for their own nutritional use, which finally promotes bacterial replication [51]. For instance, *A. phagocytophilum* secretes the effector *Ats-1* to induce autophagosome formation to obtain nutrients contained in them [52], showing that pathogens may activate autophagy also for their benefit.

Taken together, these results suggest that modulation of autophagy during bacterial infection is host- as well as pathogen-driven and leads to alterations of the metabolism and the immune defences of the infected cell.

**Concluding remarks**

Some metabolic host responses, such as activation of the HIF1α axis, commonly occur in macrophages and other cell types upon bacterial infection [53]. However, pathogen specific reprogramming of the cellular metabolism [28] and metabolic alterations triggered by bacterial effectors [12], which seem beneficial for the pathogen, also exist. A recurrent debate is whether the observed metabolic changes in infected cells are host-driven in response to bacterial invasion, or driven by the pathogen to benefit its intracellular survival and replication. As some metabolic alterations such as autophagy modulation or induction of Warburg-like programs, seem dependent on bacterial effectors and beneficial for the pathogen, both processes seem to happen in parallel and thus the observed metabolic phenotypes during infection are a mixture of host-driven and pathogen-driven metabolic reprogramming, reflecting the battle for resources in an infected cell.
Furthermore, some pathogen-mediated metabolic alterations during infection seem to trigger host immune responses. An example is the NLRP3 inflammasome activation following the inhibition of glycolytic fluxes at the level of GAPDH by S. enterica [40], or the binding of bacterial sugars to the glycolytic enzyme HK2, a member of the sugar kinase family that binds glucose and surprisingly triggers NLRP3 inflammasome activation when binding bacterial peptidoglycan-derived N-acetylglucosamine [54].

Although many puzzling enigmas remain to be clarified, a novel aspect may emerge from the “immune surveillance hypothesis” [55]. While this hypothesis defends that pathogen-mediated disruption of host physiology leads to immune activation, we think that an additional novel aspect should be added: specific checkpoints might exist along cellular metabolic pathways that, when altered, trigger immune responses such as inflammasome activation. We thus suggest the term “Danger-Associated Metabolic Modifications” (DAMMs) for alterations in the metabolism of host cells that trigger defence responses. OXPHOS dysfunction, altered glycolytic fluxes or piracy of metabolic intermediates induced by intracellular bacteria might be DAMMs linking pathogen-driven and host-driven metabolic reprogramming. Future work should shed light on these exciting and complex host-pathogen interactions.

**Declarations of interest:** none.

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**Comprehensive and excellent review describing how pathogenic, intracellular bacteria and immune cells mutually modulate the metabolism. Together with reference 27, a complete compendium of the state-of-the-art of the field.**


**Outstanding article that elegantly deciphers the role of PKM2 dimers and tetramers in the regulation of the Warburg-like metabolism induced by LPS in macrophages, and its consequence on infection with *Mycobacterium tuberculosis* or *Salmonella typhimurium*.


** Important article showing for the first time the heterogenous nature of metabolic responses of human primary monocytes to different TLR stimuli or whole-bacterial cell lysates.


**Outstanding paper showing the immune surveillance function of the metabolic enzyme Hexokinase.
Figure legends

**Figure 1.** General cellular metabolism (A) and the metabolic Warburg program (B). Large arrows and pathway names in bold represent increased metabolic fluxes. Anabolic pathways appear in violet. Blue electron transport chain (ETC): active optimal OXPHOS; pink ETC: reduced OXPHOS.

**Figure 2.** Metabolic reprogramming of innate immune cells upon exposure to TLR4 or TLR2 ligands. (A) In DCs, TLR4 engagement by LPS increases glycolysis through the activation of a TBK1-Akt-HK2 signalling axis. Glycolysis supplies DCs with citrate, which fuels the synthesis of fatty acids that are required by activated DCs to increase their secretory capacities. (B) In BMDM, LPS induces dimer and tetramer levels of PKM2. Increased PKM2 dimers lead to succinate accumulation and dimers also migrate to the nucleus, where they induce IL1β transcription. LPS-induction of PKM2 tetramers increases the rate of glycolysis. A broken TCA cycle leads to OXPHOS reduction and succinate and citrate accumulation. Accumulated succinate stabilizes HIF1α that can be translocated to the nucleus and drive IL1β gene expression. In mitochondria, succinate oxidation leads to mROS production and subsequent HIF1α-mediated IL1β mRNA expression. Accumulated citrate is used to generate NO, ROS and prostaglandins by BMDMs, as well as to produce the antimicrobial metabolite itaconate. (C) Human monocytes shift to a Warburg-like metabolism when stimulated with LPS, but not upon stimulation with the TLR2 ligand P3C. In P3C-stimulated monocytes, increased glycolysis is accompanied by increased OXPHOS. Elevated succinate, itaconate and citrate levels were observed in LPS-stimulated but not in P3C-stimulated monocytes.

**Figure 3.** Metabolic reprogramming of host cells infected by intracellular bacteria. (A) Salmonella enterica serovar Typhimurium (St) diverts glucose to the bacterial vacuole, while Chlamydia spp. (Chl) induce the Pentose Phosphate Pathway (PPP) to produce nucleotides for bacterial replication. (B) Legionella pneumophila (Lp) induces glycolysis, reduces OXPHOS, and might take advantage of this metabolism to acquire amino acids from the host cell. (C) Mycobacterium tuberculosis (Mt) induces glycolysis, reduces OXPHOS, and increase fatty acids synthesis to use lipid bodies and glycolytic-generated lactate as nutrients for its own growth.
A

Glycolysis

Glucose

2ATP

Pyruvate

Lactate

OXPHOS

ETC

36ATP

TCA

B

Glycolysis

Glucose

2ATP

Pyruvate

Lactate

Lipid and Nucleotide synthesis

Amino acid and Fatty acid synthesis

OXPHOS

ETC

PPP

TCA