The Role of Cell Metabolism in Innate Immune Memory

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Abstract
Immunological memory is classically attributed to adaptive immune responses, but recent studies have shown that challenged innate immune cells can display long-term functional changes that increase nonspecific responsiveness to subsequent infections. This phenomenon, coined trained immunity or innate immune memory, is based on the epigenetic reprogramming and the rewiring of intracellular metabolic pathways. Here, we review the different metabolic pathways that are modulated in trained immunity. Glycolysis, oxidative phosphorylation, the tricarboxylic acid cycle, amino acid, and lipid metabolism are interplaying pathways that are crucial for the establishment of innate immune memory. Unraveling this metabolic wiring allows for a better understanding of innate immune contribution to health and disease. These insights may open avenues for the development of future therapies that aim to harness or dampen the power of the innate immune response.

Introduction
Immunological memory has been classically attributed solely to the adaptive immune system. Once activated, B and T cells proliferate and confer specific long-lasting memory and thus award protection. Although invertebrates and plants lack an adaptive immune system, it is well documented that they exhibit greater resistance to subsequent infections. Specifically, systemic acquired resistance has been described in plants, whereby the first immune response confers long-term protection against a wide range of pathogens [1]. In addition, epidemiological studies point that vaccines, notably in live-attenuated forms, such as measles, polio oral vaccine, or Bacillus Calmette-Guérin (BCG), elicit nonspecific effects that reduce mortality due to protection against non-targeted diseases [2, 3]. Presently, there is a growing body of evidence in human and murine models that support the concept that the innate immune system also has features of long-term memory, whereby innate immune cells encounter an inflammatory challenge and later respond in a heightened manner to a secondary event. This phenomenon has been termed innate immune memory or trained immunity and does not rely on a particular stimulant,
Thus conferring protection against heterologous pathogens (Fig. 1) [4, 5]. Mice inoculated with the fungus *Candida albicans* became resistant to a secondary lethal infection. This protection was still observed in *Rag1*-deficient mice, demonstrating that the phenomenon is independent of T- and B-cell function [6].

Innate immune memory has been described primarily in myeloid cells, such as monocytes, macrophages, and possibly dendritic and mast cells in response to a variety of challenges. Lymphoid cells such as natural killer cells or innate lymphoid cells have also been recently shown to display long-term memory characteristics [7–9]. Monocytes or macrophages exposed to *C. albicans*, or the fungal cell wall component β-glucan, show enhanced production of proinflammatory cytokines upon an unrelated secondary stimulation such as LPS or *Mycobacterium tuberculosis* [6]. Sterile endogenous stimuli can also induce trained immunity. Oxidized low-density lipoprotein (oxLDL) induces a long-lasting proinflammatory phenotype in monocytes, which has been associated with atherosclerosis and foam cell formation [10]. However, a long-lasting protection cannot be secured by short-lived cells. Recent mouse studies have shown that BCG and β-glucan skew hematopoietic stem cells (HSCs) toward myelopoesis, generating trained monocytes and ultimately conferring partial protection against pulmonary tuberculosis and chemotherapy-induced myeloablation, respectively [11, 12]. Also, HSCs harvested from mice injected with LPS led to a decrease of mortality of the transplanted mice to *Pseudomonas aeruginosa*. This protection was still conferred by re-transplantation of the LPS exposed HSCs into a secondary host [13]. This central impact of trained immunity at the level of bone marrow progenitors was also recently explored in a human BCG vaccination study,
where it was demonstrated that the composition of HSCs did not change after vaccination. Nonetheless, BCG vaccination was shown to induce transcriptional changes in HSCs while concomitantly increasing responsiveness of innate immune cells to unrelated challenges [14]. In accordance, a cohort of BCG-vaccinated newborn infants exhibited an increase in neutrophil counts in comparison to non-vaccinated children [14]. In addition to HSCs, other stem cells may exhibit features of memory. Epithelial stem cells are progenitors of differentiated epithelial cells that form a physical barrier against external agents. Mouse models of skin sterile inflammation and fungal infection were used to show that epithelial stem cells persist after the skin returns to homeostasis and later contribute to enhanced epidermal wound healing of the previously inflamed skin [15].

Innate immune memory, in contrast to the adaptive immune memory, is not conferred neither by genomic rearrangements nor by clonal diversity and selection. Instead, the different inducers of trained immunity trigger epigenetic changes in the host cell. Trained cells harbor stable chromatin marks in gene regulatory regions that seemingly have no effect on steady-state gene expression. These chromatin adaptations modulate its accessibility to the transcriptional machinery and confer a long term increased gene transcription in response to a secondary trigger. A second hallmark of trained cells is the rewiring of metabolic pathways. Cellular metabolism has not only been correlated to cell function, but also is an important regulator of epigenetic changes. Metabolites act as signaling molecules, cofactors, and substrates that modulate the activity of chromatin modifying enzymes. This close interplay between metabolism and epigenetic reprogramming urges further exploration to understand the mechanisms underlying the different cellular metabolic pathways implicated in innate immune memory. The present review will discuss current findings that put metabolic pathways such as glycolysis, oxidative respiration, and the tricarboxylic acid (TCA) cycle at the center of the induction of trained immunity (Fig. 2). Furthermore, we will discuss future therapeutic strategies that may arise from the modulation of innate immune pathways aimed at the dampening of the trained phenotype present in proinflammatory diseases or at inducing the increased responsiveness in immunosuppressed conditions.

**Fig. 2.** Trained immunity is supported by a metabolic rewiring. Trained cells exhibit an upregulation of glycolysis, TCA, OxPHOS, glutaminolysis, and fatty acid metabolism. OxPHOS and glycolysis provide ATP, while TCA cycle intermediates modulate the function of different proteins. Succinate inhibits the degradation of the transcription factor hypoxia-inducible factor 1-α, which in turn promotes the transcription of various proinflammatory and glycolytic genes. Succinate also inhibits the enzymatic activity of histone demethylase lysine-specific demethylase 5 (KDM5), while acetyl-CoA is an acetyl donor for histone acetyltransferases. This promotes epigenetic changes that facilitate the transcription of proinflammatory genes triggered by a posterior stimulation. NADH, nicotinamide adenine dinucleotide; me, methyl; ac, acetyl; TCA, tricarboxylic acid; OxPHOS, oxidative phosphorylation; ATP, adenosine triphosphate. Figure was created with Biorender.com.
Immunometabolic Pathways Involved in Trained Immunity Glycolysis and Oxidative Respiration

Cells are extremely plastic and can adjust their metabolism according to their bioenergetic needs that change to adapt to microenvironmental changes. Glucose breakdown rapidly produces adenosine triphosphate (ATP) and pyruvate. Pyruvate can then be used in the TCA cycle or be further oxidized to lactate. Glycolysis is often increased in activated T cells [16] and proinflammatory macrophages [17]. In accordance, trained macrophages have an increased glucose consumption and release high levels of the glycolysis end-product lactate [18]. This pathway is not an efficient ATP producer (only 2 ATP molecules are produced from one molecule of glucose) but can be quickly stimulated and amplified. The enhanced glycolytic rate is enabled by the increased expression of glycolytic enzymes which is sustained through epigenetic changes [19]. Glycolysis increase is common to different training stimuli such as β-glucan, BCG, and oxLDL and has been described not only in the monocyte-macrophage populations but also in HSCs of β-glucan-exposed mice [12]. This hallmark of trained immunity is mediated by the AKT-mTOR HIF1α pathway [19, 20]. In vitro inhibition of this pathway decreases the enhanced responsiveness of trained cells. Also, myeloid cell specific HIF1α knockout mice exposed to β-glucan were more sensitive to an S. aureus infection than their trained wild-type controls [20, 21]. In addition, humans treated with metformin, a known inhibitor of mTOR, presented decreased responsiveness to restimulation after BCG ex vivo training [19].

In contrast, oxidative respiration relies on the Krebs cycle and mitochondrial machinery to generate ATP, which occurs more efficiently (in total 36 molecules of ATP from 1 molecule of glucose) but at a slower rate than glycolysis. In OXPHOS, molecules of the electron donor nicotinamide adenine dinucleotide are oxidized and donate electrons to the electron transport chain. There, electrons are sequentially relayed through the ETC complexes resulting in the reduction of oxygen and the pumping of protons across the inner membrane, generating water and a proton gradient. The shuttled protons then drive ATP synthesis generated by F0F1-ATP synthase. A pioneering study first described that β-glucan-induced memory in human monocytes leads to a metabolic rewiring with an increase of aerobic glycolysis and a dampened OXPHOS pathway [20]. However, further studies with lower doses of β-glucan or with BCG as an inducer of innate immune memory show that, concomitantly to an increase in glycolysis, these cells consume higher levels of oxygen [19, 22, 23]. This highly energetic state, where glycolytic and OXPHOS pathways are both potentiated, was further observed in human monocytes exposed to the sterile innate immune memory inducer oxLDL [24]. In addition, a recent study has described suggestive associations between common single nucleotide polymorphisms in genes related to OXPHOS and variation in the degree of responsiveness of β-glucan trained monocytes [23]. These data collectively suggest that the metabolic shift toward aerobic glycolysis and the classically described Warburg effect is not common to the establishment of innate immune memory. Instead, enhanced OXPHOS appears as a regulator of the induction of the trained phenotype.

Interestingly, the enhanced electron-hungry OXPHOS pathway might contribute to the higher NAD(P)+/NAD(P)H ratio observed in trained cells [20, 21], which may influence mitochondrial metabolism. Mitochondrial ATP production relies on nicotinamide adenine dinucleotide, which is produced by the TCA cycle and fatty acid β-oxidation. Curiously, monocytes exposed to oxLDL exhibited increased mitochondrial ROS production, which was ablated upon mTOR inhibition [21]. These findings might suggest a relevant role for mitochondria function and metabolism in the induction of trained immunity that warrants further exploration.

TCA Cycle and Amino Acid Metabolism

In addition to providing for the energetic needs of the cell, metabolic rewiring also modulates the metabolic substrates available. Different metabolites have been attributed signaling functions by inhibiting the activity of enzymes or triggering protein post-translational modifications. Specifically, the TCA cycle integrates multiple catabolic and anabolic pathways. In short, glycolysis supplies pyruvate that is metabolized into acetyl-CoA, the fuel of the TCA cycle. Acetyl-CoA can also be derived from the β-oxidation of fatty acids, while the TCA cycle intermediate citrate can be used as a precursor for lipid biosynthesis. An alternative carbon source is provided by glutaminolysis, which replenishes the TCA cycle by the conversion of glutamate into α-ketoglutarate (α-KG). Also, the oxidation of succinate to fumarate by the enzyme succinate dehydrogenase, known as complex II of the ETC, shuttles an electron to power ATP production by the OXPHOS pathway [25, 26].

The induction of innate immune memory hinges on the accumulation of TCA cycle intermediate metabolites,
which may result from the interplay between glycolysis and glutaminolysis [18]. The transcription factor HIF1α, activated in trained cells, blocks the conversion of pyruvate to acetyl-CoA by inhibiting pyruvate dehydrogenase [27]. Conversely, glutamine metabolism is increased and supplements the TCA cycle with α-KG derived from glutamate. Glutaminolysis inhibition impaired the prototypical increased responsiveness of β-glucan trained macrophages [18]. In addition to glutamine, trained cells show an enhanced consumption of the amino acid aspartate; however, its role in the induction of trained immunity remains to be clarified. Methionine consumption is also increased in β-glucan exposed monocytes [18]. It is intriguing to consider that methionine is the precursor of S-adenosylmethionine (SAM), a methyl donor critical for DNA and histone methylation which plays a central role in chromatic dynamics and gene expression patterns [28].

Metabolic rewiring is crucial for long-term gene expression and epigenetic changes in innate immune memory. Trained monocytes amass increased levels of succinate and fumarate, which are competitive inhibitors of multiple α-KG dependent dioxygenases (α-KGDD) [29]. The α-KGDD family of enzymes include histone and DNA demethylases such as the Jumonji (Jmj)-KDM5 family which actively remove lysine trimethylation, such as H3K4me3 and H3K27me3 marks [30]. The cell-permeable monomethyl fumarate was shown to increase the responsiveness of human monocytes, enrich the activating mark H3K4me3 deposition at the promoter regions of the proinflammatory genes TNFA and IL6, while also inhibiting the activity of KDM5 [18]. Succinate and fumarate also have the relevant role of stabilizing HIF1α by the inhibition of prolyl hydroxylases, which constitutionally promote the degradation of HIF1α. The increased levels of the transcription factor HIF1α also promote IL-1β transcription and upregulation of glycolysis [31]. HIF1α induces the expression of histone demethylases [32], adding another possible indirect way through which the ratios of succinate, fumarate, and α-KG could influence epigenome remodeling.

SDH, or complex II, oxidizes succinate into fumarate and shuttles an electron along the ETC. β-glucan trained monocytes show increased expression of different SDH subunits [33]. Also, genetic human studies have identified suggestive SNPs in SDH subunit coding genes associated with changes in responsiveness [23, 33]. The regulation of transcription of SDH genes has been shown to be modulated by the lysine methyltransferase Set7. In vitro pharmacological inhibition of Set7 or the use of SETD7 knockout mice led to a decrease of the β-glucan-induced upregulation of SDHB transcripts [23]. Interestingly, another modulator of SDH activity also regulates the trained immunity phenotype: itaconate is derived from the TCA cycle and produced by macrophages in proinflammatory conditions [34]. Considered as a break mechanism to the proinflammatory response, itaconate is attributed anti-inflammatory and antioxidant properties. It activates the master regulator of the antioxidant response, the transcription factor NRF2 [35], and is an inhibitor of SDH by occupying the binding pocket of succinate [36]. Notably, the in vitro β-glucan trained phenotype is hindered by concomitant exposure to the membrane permeable derivative dimethyl itaconate, possibly due to SDH inhibition [33]. This break in the TCA cycle could in turn compromise the fumarate accumulation necessary for epigenetic remodeling. Itaconate has also been described to alkylate thiol groups and interact with the ROS scavenger glutathione [37], the glycolytic enzymes lactate dehydrogenase [35], glyceraldehyde 3-phosphate dehydrogenase (GAPDH) [38], and fructose-bisphosphate aldolase [39]. Interestingly, lactate dehydrogenase and GAPDH are also possible targets for succinylation [31]. However, whether such mechanisms play a role in innate immune memory remains to be elucidated. The discovery of itaconate as a relevant immunomodulator in the context of mammalian responses is recent [34, 40, 41], and exciting new studies are exploring the intricate role of this metabolite in diverse inflammatory contexts [42–44].

**Lipid Metabolism**

Cells obtain fatty acids from de novo lipogenesis and exogenous uptake. Glycolysis, glutaminolysis, and the TCA cycle contribute to produce citrate, which in turn is converted to acetyl-CoA and gives rise to palmitate. Palmitate can be further elongated and desaturated to generate a multitude of lipid species, which can be subsequently stored as lipid droplets. These organelles contribute to the production of inflammatory mediators that regulate immune responses [45]. The stored fatty acids can be degraded to restore acetyl-CoA levels through β-oxidation. Interestingly, in vitro exogenous supplementation of acetyl-CoA induced a proinflammatory phenotype in human monocytes [46]. Acetyl-CoA serves as an acetyl donor that may favor histone acetylation and fuels the cholesterol pathway and fatty acid biosynthesis. Mevalonate, a metabolite of the cholesterol pathway, has been shown to regulate trained immunity [47]. Indeed, statins, which inhib-
it the rate-limiting enzyme that produces mevalonate, decreased the responsiveness induced by β-glucan, BCG, and oxLDL [47]. Moreover, mutations in mevalonate kinase, which lead to mevalonate accumulation, trigger an autoinflammatory clinical phenotype, the hyper immunoglobulin D syndrome. These patients show a constitutive trained immunity profile which could contribute to their clinical picture of sterile inflammation attacks [47]. Central trained immunity also relies on the mevalonate pathway as shown by the accumulation of various species of cholesterol esters in HSCs. Inhibition of the enzyme upstream of mevalonate in a mouse model of β-glucan induced innate immune memory reduced HSC expansion bias toward myelopoiesis [12]. Interestingly, systemic increase of cholesterol and inflammatory mediators also triggered innate immune memory in an atherosclerosis mouse model. Monocytic cells isolated from mice exposed to high-fat diet for a limited period showed increased responsiveness to ex vivo stimulation, even after the animals returned to a chow diet. Concomitantly the hematopoietic cell compartment proliferated and was biased toward granulocyte monocyte precursor cells, in an NLRP3 inflammasome-dependent manner [48].

Fatty acid synthesis also appears to play a role in the induction of innate immune memory. Hematopoietic progenitors of β-glucan-trained mice were enriched for lipids with shorter and more saturated acyl chains with a decrease of alkyl-ether-linked phosphatidylcholines and phosphatidylethanolamines [12]. In accordance, HSC of mice trained with LPS and exposed to a second injection of LPS exhibited increased expression of genes related to fatty acid metabolism and oxidative phosphorylation [13]. The hormone aldosterone has also been recently described to induce trained immunity in vitro and epigenetically regulate the promoter regions of genes of the fatty acid synthesis. Moreover, the increased innate immune responsiveness induced by aldosterone was ablated by inhibition of this pathway [49].

The nuclear liver X receptor (LXR) is a transcription factor that coordinates gene expression linked to cholesterol and fatty acid metabolism. Pharmacological inhibition of LXR decreased the responsiveness of BCG-trained macrophages [46]. Moreover, LXR activation alone could replicate the macrophage trained phenotype, which was also dependent on the mevalonate pathway. Other transcription factors such as sterol regulatory element binding protein and peroxisome proliferator-activated receptors are regulators of lipid metabolism; however, their involvement in the induction of innate immune memory warrants further exploration [50].

**Perspectives for Therapies**

Trained immunity is considered an evolutionary adaptation that allows for long-lasting broad-spectrum resistance to pathogens, increasing the host’s capacity to mount an immunological response. Conversely, this responsive immune state can become maladaptive and exacerbate chronic inflammatory conditions if inappropriately activated by endogenous ligands. Characterizing the specific changes that distinct innate immune cells subsets undergo in different diseases, dependent on stimuli and microenvironments, could aid in enhancing or curbing the effects of trained immunity.

Inhibition of innate immune memory could be beneficial for inflammatory conditions found in atherosclerosis [10], organ transplants [51], central nervous system inflammation [52], lupus erythematous [53], and chronic diseases associated with the Western-type lifestyle [48, 54]. On the other hand, the enhanced responsiveness of trained cells can be harnessed to improve an impaired host defense, such as in patients suffering from immunodeficiencies, immunoparalysis [55, 56], immunosenescence [57], or cancer [12]. Tumor-associated macrophages are often immunosuppressive and allow tumor progression. In this line, BCG is often used as an immunotherapy in the treatment of bladder cancer [58]. Although immune system activation allows for elimination of cancer cells, chronic inflammation may also promote neoplastic transformation and tumor progression [59]. Consequently, it is of importance to explore therapies that induce trained immunity in a durable manner but that can also be inhibited once clinical benefit is achieved. Neurodegenerative diseases are associated with chronic inflammation and may be exacerbated by peripheral infections, and possibly by trained immunity. Innate immune memory triggered on a genetic mouse model of Alzheimer by LPS led to an increase of amyloid plaque formation and associated neuritic damage [60]. Also associated with age is immunosenescence, the gradual deterioration of the immune system [61]. Therapies that combine the induction of adaptive and innate immunity could ameliorate immunosenescence. In this regard, BCG vaccination decreased the incidence of new respiratory infections in elderly individuals in a recent randomized clinical trial [62]. At the same time, BCG vaccination administration prior to other vaccines was able to boost antibody responses [63, 64]. In addition, BCG vaccination has been hypothesized as a preventive measure against the newly emerged SARS-CoV2 virus infection and multi-
ple randomized controlled trial are currently testing this potential [65].

Innate immune memory could be inhibited or potentiated at different levels, namely by engaging specific receptors or modulating cellular metabolism and epigenome. Fungal ligands rely on the dectin-1 pathway [66], the metabolite mevalonate signals through the insulin growth factor 1 receptor [47], and N-acetyl muramic acid has been shown to induce an NOD2-dependent trained immunity phenotype [67]. Also, clinical trials have shown the therapeutic value of anti-IL-1β for rheumatoid arthritis, gout, type 2 diabetes and cardiovascular disease [68, 69], possibly through the inhibition of trained immunity mechanisms. Inhibition of glycolysis and mTOR as well as the mevalonate pathway could contribute to a decrease in trained immunity, as is seen in vitro with the use of rapamycin [18, 20] and fluvastatin [47], respectively. However, epigenetic rewiring ensures the longevity of innate immune memory leading to persistence disease. As such, modulating the epigenetic landscape with inhibition of histone or DNA methyltransferases is an attractive alternative [70]. As mentioned above, different inhibitors of innate immune memory pathways have been tested in vitro, but in vivo usage encompasses new challenges. Adequate targeting to myeloid and HSCs is crucial as it decreases off-target effects and enhances bioavailability while decreasing toxicity. The use of functionalized nanoparticles could aid in targeting and in the delivery of desired compounds. In a mouse model of heart transplant, the trained phenotype of the infiltrating cells was reduced by administration of a nanoparticle that targeted myeloid cells for inhibition of mTOR [51]. In addition to pharmacological modulators, the recently developed CRISPR/dCas9 could also be exploited in the future. dCas9 blocks transcription of a targeted gene by decreasing its accessibility or can also be modified to facilitate the recruitment of transcriptional activators and thus promote the transcription of a specific gene [71]. This technology is currently under development, but it might prove a viable therapeutic option in the future, for gene expression regulation in general, and specifically in the field of innate immune responses.

**Conclusion**

Innate immune memory relies on several metabolic pathways that provide the metabolites and energy necessary for epigenetic changes. Glycolysis, oxidative phosphorylation, glutaminolysis, the TCA cycle, and cholesterol metabolism have been shown to be at the center of this enhanced responsiveness. Other mechanisms, such as mitochondrial respiration, redox, and lipid metabolism may be possible players in the induction of trained immunity and warrant future exploration. However, further technological advancements in the field of metabolomics and epigenetics are necessary to promote the identification of further relevant pathways and the interplay between them. Ultimately, innate immune memory has been attributed a contributing role for the pathogenesis of autoinflammatory and autoimmune diseases. In contrast, an impaired innate immune system might be stimulated by inducers of trained immunity and regain responsiveness. This dichotomy makes trained immunity an attractive therapeutic target. However, the development of future effective and safe therapies will first call for an increased effort toward thorough understanding of not only the molecular mechanisms regulating innate immunity but also of the contribution of non-immune and microenvironmental players.

**Conflict of Interest Statement**

M.G.N. is a scientific founder of TTxD.

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**Author Contributions**

A.V.F. wrote the first draft and J.D.A. and M.G.N. edited and reviewed it.

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Cellular Metabolism and Innate Immune Memory

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