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**effect of metformin on stem cells: Molecular mechanism and clinical prospect**

Jiang LL *et al*. Effect of metformin on stem cells

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**Abstract**

Metformin is a first-line medication for type II diabetes. Numerous studies have shown that metformin not only has hypoglycemic effects, but also modulates many physiological and pathological processes ranging from aging and cancer to fracture healing. During these different physiological activities and pathological changes, stem cells usually play a core role. Thus, many studies have investigated the effects of metformin on stem cells. Metformin affects cell differentiation and has promising applications in stem cell medicine. It exerts anti-aging effects and can be applied to gerontology and regenerative medicine. The potential anti-cancer stem cell effect of metformin indicates that it can be an adjuvant therapy for cancers. Furthermore, metformin has beneficial effects against many other diseases including cardiovascular and autoimmune diseases. In this review, we summarize the effects of metformin on stem cells and provide an overview of its molecular mechanisms and clinical prospects.

**Key Words:** Metformin; Stem cells; Differentiation; Anti-aging; Anti-cancer; Mechanism

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**Core Tip:** The effect of metformin on stem cells is quickly gaining attention, because metformin modulates various physiological activities and pathological changes *via* targeting stem cells. Emerging studies suggest that metformin has broad prospects in the fields of stem cell medicine, gerontology, regenerative medicine, and cancer therapy, *etc.* In this review, we summarize the effects of metformin on stem cells and provide an overview of its molecular mechanisms and clinical prospects.

**INTRODUCTION**

Metformin (N,N′-dimethyl metformin), which is widely used in patients with type 2 diabetes, exerts hypoglycemic effects mainly by inhibiting absorption of glucose in the gut, suppressing gluconeogenesis and glycogen synthesis, and facilitating the uptake and utilization of glucose, and sensitivity to insulin of peripheral tissues[1]. It is widely accepted that metformin reduces diabetic risk factors such as obesity and improves diabetic complications such as cardiovascular disease, peripheral neuropathy, and higher fracture risk[2-5].

In recent years, studies have shown that metformin modulates many physiological and pathological processes ranging from aging and cancer to fracture healing[1,6-8]. In 2005, Evans *et al*[9] found that metformin reduces the morbidity of malignant tumors in patients with type 2 diabetes, attracting attention to explore the connection between metformin and cancer[9]. In 2013, Cabreir’s research on the anti-aging effect of metformin was published in the journal Cell. He reported that metformin increases the lifespan of *Caenorhabditis elegans* cocultured with *Escherichia coli* by altering microbial folate and methionine metabolism, demonstrating the anti-aging effect and mechanism of metformin[10]. These studies suggest that metformin has regulatory effects on various physiological activities and pathological changes. Studies have shown that stem cells play a curial role in these processes. Therefore, many scientists have studied the effect of metformin on stem cells in recent years.

Previous studies have demonstrated that metformin affects stem cell differentiation, enhances their immunomodulatory properties, and exerts anti-aging, anti-oxidative, and anti-inflammatory effects in stem cells[11-16]. This review focuses on the multiple effects of metformin on stem cells, its molecular mechanisms, and clinical prospects.

**Effect of metformin on differentiation of stem cells**

Cell differentiation refers to the process through which cells from the same source gradually produce cell groups with different morphological structures and functional characteristics. It is the basis of ontogeny that is conductive to improve the efficiency of various physiological functions. Thus, a large number of studies on stem cell differentiation have been reported. Studies have shown that metformin affects the differentiation of stem cells and progenitor cells[11,17,18]. We have summarized these effects and their molecular mechanisms (Table 1).

***Osteogenic differentiation***

Bone is a complex tissue containing several cell types, which is continuously undergoing a process of self-renewal and repair, termed bone remodeling. Many studies have indicated that metformin promotes osteogenic differentiation of stem cells and osteogenic progenitor cells. The promotive effects manifest as increased cell proliferation, cell migration, alkaline phosphatase activity, mineral deposition, and upregulated expression of osteoblast marker genes, including osteopontin (OPN), osteocalcin, and runt-related transcription factor 2 (Runx2), during osteogenic cell differentiation[8,11,19].

Metformin promotes osteogenic differentiation mainly through the adenosine 5′-monophosphate-activated protein kinase (AMPK) signaling and Runx2-related signaling pathways[20-27]. Metformin is an AMPK activator similar to 5-aminoimidazole-4-carboxamide ribonucleotide[28]. Its primary site of action is direct inhibition of complex 1 of the respiratory chain, which decreases production of ATP, leading to an increase of the AMP/ATP ratio and then activated AMPK[29]. Sedlinsky *et al*[21] submitted bone marrow progenitor cells (BMPCs) to 15 d osteoblastic induction in the presence or absence of metformin and/or compound C (an inhibitor of AMPK activation). As a result, metformin increased the P-AMPK/total AMPK ratio and production of type 1 collagen (a marker of osteoblastic differentiation) in BMPCs, whereas compound C inhibited these increases, demonstrating that metformin promoted osteoblastic differentiation of BMPCs through AMPK activation[21]. Similarly, Wang *et al*[23] treated induced pluripotent stem cell (iPSCs) with metformin, demonstrating the same effect *via* the liver kinase B1 (LKB1)/AMPK signaling pathway. LKB1 is a common upstream molecule of AMPK kinase. Inhibiting its activity markedly reverses metformin-induced AMPK activation and Runx2 expression[23]. In addition, metformin exerts a similar effect on MC3T3-E1 cells through the AMPK/growth factor independence-1 (Gfi1)/OPN axis. AMPK activation downregulates the transcriptional repressor Gfi1 and disassociates it from the OPN promoter, ultimately upregulating OPN[24]. Furthermore, metformin may promote osteoblastic differentiation through decreased acetyl coenzyme carboxylase activity and lipogenic enzyme expression induced by AMPK activation. These decreases contribute to inhibited adipogenesis and break the balance between osteogenic and adipogenic differentiation[30].

Regulation of the Runx2-related signaling pathway by metformin is the second mechanism to promote osteogenic differentiation. Runx2 promotes mesenchymal stem cells (MSCs) to differentiate into preosteoblasts and inhibits adipogenic and chondrogenic differentiation[31]. Marofi *et al*[25] treated human bone marrow stromal cells (hBMSCs) with metformin and found that metformin promoted osteogenic differentiation through the Twist1/Runx2 signaling pathway. Metformin inhibited the expression of Twist1 by enhancing its gene promoter methylation slightly and a direct physical interaction without Twist1 methylation. Lower Twist1 expression increased the mRNA expression of *Runx2*[25]. In addition, Chava *et al*[26] extracted BMSCs from metformin-treated rats and demonstrated that metformin promoted osteogenic differentiation through AMPK directly mediating Runx2 phosphorylation at serine 118[26].

In addition to the two abovementioned signaling pathways, metformin promotes osteogenic differentiation through other mechanisms. Metformin promotes osteogenic differentiation through the serine/threonine kinase Akt (also known as protein kinase B or PKB) signaling pathway. Jia *et al*[27] treated periodontal ligament stem cells (PDLSCs) with metformin and found that metformin rescued osteogenic differentiation of PDLSCs, which was impaired by H2O2-induced oxidative stress by activating Akt and downstream nuclear factor E2-related factor 2 (Nrf2), an important transcription factor against oxidative stress[27]. Ma *et al*[19] treated hBMSCs with metformin and obtained the similar result. They stated that the Wnt/â-catenin signaling pathway probably participated in the osteogenic differentiation of BMSCs because metformin inhibited glycogen synthase kinase-3â, resulting in accumulation of cytosolic â-catenin and activation of the Wnt signaling pathway[19].

The effect of metformin on osteogenic differentiation may be influenced by the drug dose, cell origin, and glucose concentration. Several studies explored the effect of metformin on osteogenic differentiation of PDLSCs and found that 50 ìmol/L was the optimal concentration to exert effects[11,27,32]. Houshmand and Ma studied the same effect on BMSCs and hDPSCs, and found that 100 ìmol/L was the optimal concentration to promote cell osteogenic differentiation[19,33]. Moreover, Mu *et al*[12] found that metformin promoted osteogenic differentiation of murine preosteoblasts under high glucose conditions. In this study, metformin suppressed the phosphorylation of nuclear factor-kB by increasing silent information regulator (SIRT)-6 expression. High levels of SIRT6 will decrease mature osteoblast functions and delay maturation of bone matrix [12,34].

***Neuronal differentiation***

The nervous system has a complex structure and is the major controlling, regulatory, and communicating system in the body. However, unfortunately, when brain cells are damaged by trauma or disease, they are unable to automatically regenerate, which determines nervous dysfunctions and onset/progression of neurodegenerative diseases. Neurodegenerative diseases/neurodegenerative pathologies, including Huntington’s disease, multiple sclerosis, Alzheimer’s disease, Parkinson’s disease, and amyotrophic lateral sclerosis, represent a group of illnesses characterized by the following features: A decline in neuronal functions, brain atrophy, and often, abnormal deposition of proteins. Several studies have shown that metformin promotes the proliferation and neural differentiation of stem cells[17,35-37], indicating that metformin may be a promising drug for prevention and treatment of these diseases[38].

Ahn *et al*[36] treated hBMSCs with metformin and demonstrated that metformin may promote neuronal differentiation and neurite outgrowth by activating AMPK. Neuronal cells were characterized by an increase in the expression of neuron-specific genes *MAP-2, Tuj-1, NF-M, KCNH1,* and *KCNH5*[36]. Dadwal *et al*[35] extracted subependymal-derived neural precursor cells (NPCs) and plated them as single cells to form neurospheres in the presence of metformin. The results showed that metformin expanded the stem cell pools and facilitated neurogenesis in normal mice compared with CREB-binding protein (*CBP*) gene mutant mice, demonstrating that metformin directly promoted NSCs to differentiate *via* the atypical protein kinase C (aPKC)-CBP pathway[35]. Fatt *et al*[37] added metformin to NPCs extracted from the adult subventricular zone and found that metformin significantly enhanced neuronal differentiation by activating the AMPK-aPKC-CBP pathway[37].

***Myogenic differentiation***

Skeletal muscle is the largest organ of the body and plays an important role in essential life activities such as respiration, metabolism, mediating temperature, and movement. When empyrosis, trauma, and other factors cause damage to skeletal muscle, skeletal muscle can be regenerated. Skeletal muscle regeneration is dependent on a contribution from muscle-resident stem cells, named satellite cells marked by paired-box transcription factor 7 (Pax7)[39]. The effects of metformin on satellite cells are disputed. Several studies provide evidence that metformin maintains satellite cells in a low differentiation state and deplete skeletal muscle regeneration *via* calorie restriction, whereas others have stated that metformin alleviates muscle wasting post-injury[14,18,39].

A family of myogenic regulatory factors (MRFs), such as myogenic differentiation antigen (MyoD), myogenin, Mrf4, and myogenic factor (Myf5), plays an important role in myogenic differentiation[18,31]. Pavlidou *et al*[18] found that metformin (2-10 mmoL/L)-treated C2C12 cells had a reduced myogenic differentiation potential and significant decline in the expression of myogenic regulatory factors MyoD, myogenin, and myosin heavy chain[18]. In another study, they treated satellite cells with 2 mmoL/L metformin *in vitro*, resulting in retained expression of Pax7 for a longer time, whose delayed downregulation was accompanied by late expression of myogenic differentiation markers, indicating delayed differentiation. *In vivo*, metformin delayed regeneration of cardiotoxin-damaged skeletal muscle[14]. Conversely, Yousuf *et al*[39] injected metformin hydrochloride dissolved in phosphate buffered saline into mice after burn injury and found that metformin enhanced the proliferative activity of Pax7-positive satellite cells by activating AMPK. They attributed the contradictory conclusion between the results to different mouse mobility and the different nature of the injury[39].

The effect of metformin on myogenic differentiation of C2C12 cells is controversial and may be related to drug dose. Low doses of metformin (400 and 500 ìmol/L) promote myogenic differentiation and myotube formation, increasing the protein expression of Myf5 and MyoD, two important markers of early differentiation. Senesi *et al*[40] believed that metformin enhanced C2C12 differentiation by activating ERKs and decreasing p70S6 kinase[18,40]. Fu *et al*[41] inferred that metformin facilitated myogenic differentiation of C2C12s by activating AMPK. AMPKá1 phosphorylated histone deacetylase 5 (HDAC5) at Ser 259 and 498 in C2C12 cells, which acts as a conserved transcriptional repressor through an interaction with myocyte enhancer factor-2. Phosphorylated HDAC5 upregulates myogenin transcription and myogenesis[41]. Considering the paradoxical effect of metformin on myogenic differentiation, more studies in this field are needed.

***Adipogenic differentiation***

Considering the reciprocal relationship between osteogenic and adipogenic differentiation, various reasons, such as diabetic conditions and the use of thiazolidinedione, cause active adipogenesis in BMSCs/BMPCs, which consequently suppresses osteogenesis and damages bone health[42-44]. Metformin inhibits adipocyte differentiation of adipose-derived stem cells (ADSCs), BMSCs, BMPCs, and PDLSCs, manifesting as suppressed cell proliferation, lipid droplet generation, and expression of adipocyte genes such as peroxisome proliferator-activated receptor gamma (*PPARγ*), CCAAT/ enhancer binding protein alpha, and adipocyte lipid-binding protein[13,27,42,43,45,46].

Marycz *et al*[13] extracted ADSCs from metformin-treated rats to induce adipogenic differentiation. A reduction of lipid droplets in ADSCs and decreased proliferation potential demonstrated that metformin inhibited adipogenic differentiation[13]. Tolosa *et al*[42] extracted BMPCs from diabetic rats, treated them with/without metformin, and found that metformin partially abolished diabetic-related upregulation of PPARã expression[42]. Similarly, Molinuevo *et al*[46] treated BMPCs with both metformin and rosiglitazone and found that metformin abolished rosiglitazone-induced adipogenesis[46].

Metformin inhibits adipogenic differentiation through the AMPK signaling pathway. Wang *et al*[24] established LV-AMPKá 3T3-L1 cells stably overexpressing AMPKá through a lentiviral vector and treated them with puromycin to induce their adipogenic differentiation. The results showed that activated AMPK suppressed lipid droplet generation and adipocyte gene expression[24]. The other mechanism was inhibiting the mammalian target of rapamycin (mTOR)/ribosomal protein S6 kinase signaling pathway. Chen *et al*[44] found that metformin suppressed adipogenesis in C3H10T1/2 MSCs independently of the AMPK signaling pathway by measuring phosphorylation of a known AMPK substrate, Ser 79 of ACC. Lipid accumulation associated with adipogenesis in C3H10T1/2 cells was inhibited by incubation with the mTOR/p70S6 kinase inhibitor rapamycin[44]. In addition, because of the reciprocal relationship between osteoblast and adipocyte differentiation, metformin may inhibit adipocyte differentiation indirectly by promoting the expression of osteoblastic transcription factors[43].

***Chondrogenic differentiation***

Downregulation of chondrocytic differentiation has been described in various chronic skeletal diseases including osteoarthritis. Metformin appears to inhibit chondroblastic differentiation. Bandow *et al*[47] treated chondrogenic progenitor cells with metformin during chondrogenic differentiation. As a result, metformin inhibited chondroblastic differentiation by activating AMPK. In primary chondrocyte precursors, metformin decreased gene expression of sex determining region Y-box (*Sox*) 9 and *Sox6* along with other chondrogenic differentiation markers including collagen, type II, alpha 1 (col2a1), and aggrecan core protein (ACP). *Col2a1* and *ACP* promoter activities were directly repressed by AMPK-activated early growth response-1 (Egr-1), a transcriptional repressor in mouse chondrocytes independent of Sox9. Mutation of the putative Egr-1-binding site abrogated the inhibitory effects of an AMPK activator[47]. Sox9 plays an important role in various stages of chondrogenesis and is essential for chondrogenesis. Its gene deletion can lead to achondroplasia[48].

***Gastric parietal cell differentiation***

Metformin has been reported to reduce the risk of stomach cancer by up to 51% in diabetic patients following eradication of *Helicobacter pylori*[49]. A recent study showed that metformin promotes differentiation of gastric epithelial progenitor cells into acid-secreting PCs through AMPK activation. AMPK activation increased Kruppel-like factor 4 (KLF4) expression, facilitating progenitor cells to exit the cell cycle and differentiate toward the PC lineage. AMPK appeared to increase maturation of the PC lineage largely by peroxisome proliferator-activated receptor-ã coactivator-1á activation[50]. Considering that PC damage plays a crucial role in the occurrence and development of gastric cancer, metformin may have potential as an anti-gastric cancer drug by promoting gastric epithelial progenitor cells to differentiate into acid-secreting PCs.

***Role of metformin and stem cells in tissue injury healing***

Studies showed that metformin and stem cells also play an important role in tissue injury healing, which may relate to the effect of metformin on stem cell differentiation[51-53]. Stem cells have been demonstrated to promote tissue injury healing such as diabetic foot ulcer, burns, gastric ulcer, and ulcerative colitis (UC). They exert the repair effect through migrating to tissue injuries, differentiating into specific cells, reducing inflammation, and producing paracrine factors to promote angiogenesis[51,52]. Deng *et al*[53] found that bencher muscular dystrophy-endothelial progenitor cells (BMD-EPC) contributed to tissue repair in UC. Notably, stromal cell-derived factor-1 and its receptor CXCR4 also have been demonstrated to play an important role in the “homing” of BMD-EPC to injured sites and neovascularization in tissue repair[53]. While metformin exerts the repair effect through facilitating the process described above, metformin also promotes injury healing through insulin sensitization in diabetic foot ulcer[51,52]. The combination of stem cells and metformin appears to be a better synergistic option for the treatment of diabetic wounds (Figure 1).

**Metformin regulates stem cell aging and rejuvenating regeneration**

Aging can be considered as a developmental program that is beneficial early in life but not switched off upon its completion. From a stem cell-centered perspective, aging results in an impaired regenerative capacity to effectively maintain tissues and organs as well as depletion of stem cell pools in adult tissue, leading to tissue dysfunction and age-associated diseases. For example, the number and proliferation or differentiation ability of stem cells gradually decrease with age. Therefore, damaged tissues and organs cannot be repaired and regenerate in time, which directly leads to the occurrence of human aging and diseases[54]. Both extrinsic (local microenvironment and systemic circulation) and intrinsic factors (genomic instability, DNA damage, oxidative damage, and deteriorated mitochondrial functions) contribute to stem cell dysfunction during aging-related regenerative decline[55-57]. Anti-aging has been a research hotspot in recent years. In 2019, Fahy’s research on reversing “biological age” became the headline of *Nature*. He found that systemic treatment with a cocktail of growth hormone, dehydroepiandrosterone (DHEA), and metformin partially reverses DNA methylation age (DNAma) clocks. DNAma is a prominent biomarker of mammalian aging[58,59]. It was the first time that clinical research indicated the anti-aging effect of metformin. Thus, metformin is currently undergoing repurposing as an anti-aging agent[6,58,60,61].

Metformin rescues stem cells from aging by alleviating intrinsic undesirable changes. The anti-aging effect of metformin is closely related to its antioxidant effects in stem cells. Fang *et al*[62] demonstrated that chronic low-dose metformin treatment increases the lifespan of HMSCs through Nrf2-mediated transcriptional upregulation of endoplasmic reticulum-localized glutathione peroxidase 7 (GPx7) whose depletion results in premature cellular senescence[62]. Metformin also inhibits mTORC1 activity, initiating the process of autophagy[55,63]. Autophagy of stem cells maintains internal homeostasis in response to stress conditions and plays a crucial role in stem cell rejuvenation[64]. Na *et al*[65] found that metformin inhibited aging-related phenotypes in Drosophila midgut intestinal stem cells (ISCs) through the Atg6 (autophagy related gene 6; Beclin 1 in mammals)-related pathway, which was negatively regulated by the AKT/TOR pathway[65]. Metformin reduced age and oxidative stress-related accumulation of DNA damage marked by drosophila ãH2Ax foci and 8-oxo-dG by suppressing the AKT/mTOR pathway in Drosophila midgut ISCs[66,67]. Therefore, inhibiting the AKT/mTOR pathway may decrease DNA damage through activation of autophagy.

Furthermore, metformin is involved in metabolic-induced rejuvenation by mimicking the metabolic effects of calorie restriction. The mechanism may be related to stimulating AMPK, the principal energy sensor in cells, to reduce energy-consuming processes and increase insulin sensitivity[60,64,68,69]. It is commonly accepted that the quiescent state of stem cells, which has been linked to their metabolic state, is correlated with their regenerative capacity[14,57,70]. Pavlido’s research showed that metformin inhibited mTOR activity and reduced p70S6 phosphorylation that induced a low metabolic state associated with quiescence of satellite cells[14]. Similarly, Neumann *et al*[71] demonstrated that metformin rejuvenated aging oligodendrocyte progenitor cells by activating AMPK[71].

Activation of AMPK by metformin provides an efficient barrier to reprogramming somatic cells (mouse or human fibroblasts) to stem cells[72]. Cellular reprogramming is inducing somatic cells into a pluripotent cell state by expression of transcription factors octamer-binding transcription factor-4 (Oct4), Sox2, Klf4, and c-Myc[73]. Reprogramming reverses many aspects of aging by resetting metabolic signatures, mitochondrial networks, and other factors to a youthful state[69]. Metformin inhibits the reprogramming process by selectively impairing expression of Oct4, but not other reprogramming factors[72,74].

**Effect of metformin on cancer stem cells**

Cancer stem cells (CSCs) are cancer cells with the capacity to renew indefinitely, resulting in tumor formation. They possess stem cell properties including self-renewal, proliferation, and differentiation potential. CSCs are responsible for the clinical failure of the majority of currently available oncological therapies because they survive treatment with hormones, radiation, chemotherapy, and molecularly targeted drugs[75]. Therefore, how to eliminate CSCs has become a research focus in recent years. Many studies have explored the effect of metformin on CSCs and the results are inspiring. Metformin is expected to become an anti-cancer agent[7,9]. The studies showed that metformin inhibits CSCs *via* inhibition of self-renewal, metastatic, metabolic, and chemoresistance pathways.

Metformin inhibits pathways associated with self-renewal and metastasis in various CSCs. Saini *et al*[75] stated that the inhibition mechanism included hedgehog, Wnt, and transforming growth factor-â pathways. CSCs self-renew indefinitely, results in tumor formation, and has a potential role in tumor metastasis. Courtois *et al*[76] confirmed the anti-proliferative effect of metformin on gastric carcinoma cell lines by regulating the expression of genes implicated in cell-cycle regulation (*GADD45, p21, E2F1,* and *PCNA*)[76]. Zhao *et al*[77] demonstrated that metformin inhibits the proliferation of osteosarcoma stem cells (OSCs) by inducing G0/G1 phase cell cycle arrest[77].In a recent study, Barbieri *et al*[78] stated that metformin and other biguanides exert anti-proliferative effects in glioblastoma CSCs by interfering with the activity of the extracellular portion of the active transmembrane chloride ion channel. Chloride intracellular channel 1 (CLIC1) activity promotes cell cycle progression and cell division during G1/S phase transition, leading to accelerated growth in glioblastoma CSCs[78].

Metformin also inhibits pathways associated with CSC metabolism. Metabolic reprogramming refers to changes in the metabolic patterns of CSCs compared with normal cells, which makes the body provide sufficient resources for CSCs. A classic example of metabolic reprogramming is the Warburg effect. CSC-driven tumorigenesis through metabolic reprogramming is closely associated with the acquisition of stem cell-like properties in iPSCs. Several studies have demonstrated that metformin suppresses the expression of reprogramming factor Oct4, providing a barrier against malignant CSCs[69,72,79]. Saini *et al*[75] stated that the metabolic effect of metformin on CSCs mainly depends on inhibition of mitochondrial complex I[75]. It is generally believed that the therapeutic effect of metformin results from an inability of CSCs to turn on glycolysis for ATP production (Warburg effect) upon inhibition of oxidative phosphorylation[80]. A recent study highlighted the strong dependence on energy-producing pathways of colorectal cancer CSCs, suggesting that modulation of AMPK activity is an effective therapeutic approach[81]. Zhao *et al*[77] showed that metformin induced caspase-mediated apoptosis in OSCs by inducing mitochondrial dysfunction. In addition, metformin influenced the capacity of OSCs to self-renew *via* enhanced autophagy, which was suppressed by 3-methyladenine, an inhibitor of autophagy[77].

Metformin inhibits CSCs by impairing the chemoresistance of CSCs[82-84]. Tan *et al*[85] demonstrated that metformin regulates the miR708/cluster of differentiation 47 (CD47) axis to eradicate breast cancer stem cells and enhance chemosensitivity because of the critical role that CD47 plays in evasion of immunological eradication[85]. Bishnu *et al*[86] showed that continuous metformin treatment impeded acquirement of chemoresistance by reducing the CSC proportion through taurine generation and removing CSCs from quiescence. Maintaining a more proliferative cellular state also contributes to chemosensitivity[86].

**Cardiovascular protective effects of metformin related to ePCs**

The cardiovascular system is also called the circulatory system, consisting of the heart and blood vessels. It transports oxygen and metabolizes waste to maintain steady metabolism of the internal environment and normal life activities. Cardiovascular diseases are associated with impaired vascular remodeling and a lack of endothelial cell reconstructive functions[87]. EPCs, as a kind of precursor cell, have the functions of migration, proliferation, adhesion, and differentiation into endothelial cells involved in the generation of adult neovascularization[88]. Functionally impaired EPCs manifest as decreased EPC numbers in circulation, decreased angiogenic potential, and endothelial dysfunction[89-91]. To our knowledge, most studies have reported that metformin protects the cardiovascular system by improving EPC functions and angiogenesis[92,93].

Metformin improves EPC functions through the AMPK-endothelial nitric oxide synthase (eNOS)-nitric oxide (NO) signaling pathway. Li *et al*[93] treated EPCs from normal individuals with metformin and found that metformin promoted EPCs to differentiate into ECs. Yu *et al*[94] treated EPCs from diabetic rats and found that metformin increased EPC capacities for immigration and tube formation. In the two studies, metformin increased both phosphorylated AMPK and eNOS expression in EPCs and enhanced NO production[93,94]. The reduction in NO bioavailability due to reduced synthesis from eNOS can cause EPC dysfunction[95]. Li *et al*[93] also found that metformin improved EPC functions through AMPK-mTOR-autophagy-related and AMPK-mTOR-p70S6K pathways. The drug activated AMPK and inhibited mTOR[93].Metformin improved palmitic acid (PA)-induced EPC dysfunction by mediating microRNA (miR) 130a and phosphatase and tensin homolog (PTEN), which may be associated with activation of the phosphatidylinositol-3-kinase/AKT signaling pathway. Levels of miR-130a are lower and those of PTEN are higher in EPCs of diabetic patients[87].

Some studies have suggested negative effects of metformin on EPCs. Metformin attenuates EPC migration through the AMPK/mTOR/autophagy-related pathway. Metformin also activates AMPK phosphorylation and inhibits mTOR and Akt phosphorylation, decreasing matrix metalloproteinase (MMP)-2 and MMP-9 expression in EPCs, indicating that decreased activity of gelatinase and fibrinolysis may contribute to this phenomenon. However, how AMPK/mTOR-related autophagy regulates cell migration is controversial[93,96,97]. The inhibitory effect of metformin on EPC angiogenesis is mediated by down-regulating miR-221 expression, which is negatively correlated with the concentration of metformin and consequently increased p27 expression and activated autophagy[97]. Asadian *et al*[98] found that metformin inhibited EPC proliferation and angiogenesis following inhibited activation of the Tunica internal endothelial cell (Tie2)/AKT signaling pathway, which may be associated with the Tie2/Akt/eNOS signaling pathway[98,99]. Similarly, Montazersaheb *et al*[100] found that prolonged incubation with metformin decreased the angiogenic potential of hBMSCs by modulating the mTOR-related autophagy signaling pathway[100]. Considering the paradoxical effects of metformin on EPCs and angiogenesis, one explanation would be that metformin behaves differently according to diabetic conditions and drug concentration[92,100]. Hence, the role of metformin is debatable in EPC functions and needs to be validated by future studies.

**Antioxidant effects of metformin on stem cells**

Various diseases, such as diabetes, Alzheimer’s disease, and cardiovascular disease, are associated with oxidative stress, and metformin acts as an antioxidant at the cellular level *via* the mechanisms described below[38,54,101,102]. First, metformin decreases free radicals, including reactive oxygen species (ROS) and NO, and upregulates activities of antioxidant enzymes in stem cells, such as superoxide dismutase (SOD) and ER-located GPx7[13,27,40,62,103,104]. It significantly attenuates ROS production of BM-derived hematopoietic stem cells after total body ionizing radiation irradiation[105]. Low-dose metformin increases the nuclear accumulation of Nrf2 that binds to antioxidant response elements in the *GPX7* gene promoter to induce its expression[62]. Advanced glycation endproducts (AGEs) elevate during certain physiological and pathological states including inflammation, aging, diabetes, and neurodegenerative diseases. Human neural stem cells (hNSCs) treated with AGEs have decreased cell growth, but metformin rescues hNSCs from AGE-induced oxidative stress, normalizes ROS levels, and improves SOD activity by decreasing the levels of receptor for advanced glycation endproducts that is downstream of AMPK[104]. Furthermore, metformin protects mitochondria from oxidative damage. Cytosolic cytochrome c activates the caspase protein family, thereby initiating mitochondrion-mediated apoptosis.

Chiang *et al*[102]’s work focused on hNSCs exposed to amyloid-Aâ or AGEs, which had reduced expression of mitochondrion-associated gene, mitochondrial deficiency (lower displacement loop level, mitochondrial mass, maximal respiratory function, cyclooxygenase activity, and mitochondrial membrane potential), as well as increased activation of caspase 3/9 activity and cytosolic cytochrome c in common. Metformin abrogates these negative effects through the AMPK signaling pathway[102,106].

**Anti-inflammatory effect of metformin on stem cells**

The anti-inflammatory effects of metformin in neurodegenerative diseases, EPC dysfunction, and aging have also attracted attention in recent years[38,54,107]. Chung *et al*[106] demonstrated that metformin suppressed AGE-induced inflammation in hNSCs by activation of AMPK, which inhibited inhibitory nuclear factor kappa-B (NF-kB) kinase (IKK) activity and normalized expression of inflammatory cytokines including interleukin (IL)-1á, IL-1â, IL-2, IL-6, IL-12, and tumor necrosis factor á (TNF-á). Decreased NF-êB levels caused by inhibited IKK activity alleviated the inflammatory response *via* increased expression of inducible nitric oxide synthase (iNOS) and COX-2[15]. Han *et al*[107] showed that metformin decreases the expression levels of proinflammatory cytokines (IL-1b, IL-6, and TNF-á) by preventing high mobility group box 1 (HMGB1) release from the nucleus to cytoplasm in rabbit annulus fibrosus stem cells. HMGB1 has been proved to play a role in the development and maintenance of the inflammatory response[108]. Furthermore, the senescent phenotype induced by lipopolysaccharide is inhibited by metformin, indicating a correlation between its anti-inflammatory and anti-aging effects[107]. Considering the close relationship between oxidative stress and chronic inflammation, some researchers have suggested that the antioxidant, anti-inflammatory, and anti-aging effects of metformin are interactive[54,101].

**Metformin enhances immunomodulatory potential of stem cells**

Metformin has also shown potential to treat autoimmune diseases, such as rheumatoid arthritis and systemic lupus erythematosus, by regulating metabolism[109,110]. A recent study showed that metformin enhanced the immunomodulatory properties of ADSCs through the AMPK/mTOR/signal transducer and activator of transcription (STAT)-1 signaling pathway. Metformin increased AMPK and STAT1 phosphorylation in a dose-dependent manner, but decreased the phosphorylation of mTOR[111]. STAT1 plays an important role in cell survival and proliferation, whose overexpression strongly enhances cord blood-derived MSC-mediated T-cell suppression[112,113]. STAT1 inhibition of ADSCs by metformin significantly impairs induction of immunomodulatory markers, including indoleamine 2,3-dioxygenase, IL-10, and transforming growth factor-â, and inhibits T-cell proliferation *in vitro*[111].

**Adverse effects of metformin on stem cells**

Various studies show that metformin acts in a dose- and time-dependent manner[114-116]. The adverse effects of metformin on stem cells include decreasing proliferation activity, cytotoxic effects (morphology and ultrastructure changes), and apoptosis when the drug concentration is far more than the therapeutic doses administered to diabetic patients whose plasma concentrations of metformin are usually < 50 ìM. The levels of metformin that accumulate in tissues might be several times higher than that in blood[117]. Œmieszek *et al*[116] exposed ADSCs and BMSCs to 1, 5, and 10 mmoL/L metformin and then measured cell proliferation activity and characteristic features after 24, 48, and 72 h. They found that metformin inhibited cell proliferation in a dose- and time-dependent manner, and that 5 and 10 mmoL/L metformin had cytotoxic effects on ADSCs, causing abnormal morphological, ultrastructural, and apoptotic changes. The decrease in cell proliferation was associated with cytotoxic effects of metformin[115,116]. Consistent with this result, Jia *et al*[27] observed that a high concentration of metformin (2500 µM) slightly inhibited cell proliferation of PDLSCs[27]. They suggested that metformin induced hUC-MSC apoptosis in an AMPK-mTOR-S6k1-Bad (Bcl-2 family member)-dependent manner, which was reversed by compound C. Metformin greatly increased the rate of hUC-MSC apoptosis in a dose- and time-dependent manner without affecting autophagy or proliferation[118]. Compared with BMSCs and ADSCs, lower concentrations of metformin (100 ìM, 250 ìM, and 500 ìM) also caused hUC-MSC apoptosis. This phenomenon may due to different cell origins. The authors recently found that glucose modulates metformin-induced MSC apoptosis *via* the AMPK-mTOR-S6k1-Bad pathway in another study. High glucose (10, 15, and 30 mmoL/L) exerts a protective effect on metformin-induced apoptosis, which is inversely proportional to the glucose level[119,120]. However, Zafarvahedian *et al*[114] showed that glucose conditions do not affect metformin toxicity and hyperglycemia itself inhibits the proliferation of MSCs[114]. More evidence on whether the glucose level influences metformin-induced adverse effects in MSC is needed in the future.

**Other mechanisms not related to stem cells’ actions**

Metformin can affect many of these conditions by acting also through other mechanisms not related to stem cells’ actions. For example, metformin acts on liver cells to reduce liver gluconeogenesis and promote anaerobic glycolysis, on skeletal muscle cells to increase glucose uptake, and on intestinal epithelial cell to inhibit or delay gastrointestinal glucose absorption, which exerts hypoglycemic effects synergistically[1]. Metformin under physiological tolerance concentration can alleviate aging-associated inflammation *via* enhancing autophagy and normalizing mitochondrial function in T cells, which is a major source of inflammatory cytokines. Autophagy made it possible to prolong normal life *via* improving inflammation[121]. Besides, metformin can directly act on cancer cells. Liu *et al*[122] found that metformin can exert growth-suppressive effects on gallbladder cancer (GBC) cell lines *via* inhibition of p-Akt activity and the Bcl-2 family. Also, inhibition, knockdown, and upregulation of the membrane protein CLIC1 can affect GBC resistance in the presence of metformin[122]. Totally speaking, metformin exerts different effects by acting various mechanisms.

**Clinical perspectives**

Stem cells have become a research hotspot in recent years. Many studies have applied stem cells to the fields of regenerative medicine, such as tissue engineering, and stem cell medicine such as the treatment of refractory disease stroke, autoimmune disease, neurological disease, and cardiovascular disease[16,17,55,92,123,124]. After understanding the effects and mechanisms of metformin on stem cells, we can apply metformin with stem cells in these fields to improve the therapeutic effect.

Regenerative medicine, which uses stem cell therapies to create tissues and organs and repair them, has the potential to address donor organ shortages. The seed cells (usually stem and progenitor cells) and scaffolds (biological materials) play important roles in tissue engineering. A previous study has shown that metformin in scaffolds regulates seed cell differentiation and proliferation. For example, Zhao *et al*[32] seeded hPDLSCs on a calcium phosphate cement scaffold delivering metformin[32]. Shahrezaee *et al*[124] seeded BMSCs on a polylactic acid and polycaprolactone scaffold delivering metformin and implanted the construct into calvarial bone defects in a rat model[124]. Smieszek *et al*[22] cocultured ADSCs with sol gel-derived silica/zirconium dioxide delivering metformin. All of the above studies found that metformin promoted osteogenic differentiation of stem cells, suggesting that metformin has the potential to promote bone tissue engineering by affecting stem cell differentiation.

Metformin and stem cells also have broad application prospects for the treatment of refractory diseases. Stroke is a public health issue, resulting in neurological disabilities in many patients. NSCs and NPCs are expected to be a new treatment for neurological disabilities. Ould-Brahim *et al*[17] treated a rat endothelin-1 focal ischemic stroke model with metformin-preconditioned human iPSCs (hiPSC)-NSCs. The results showed that metformin preconditioning enhanced the differentiation of hiPSC-NSCs, accelerated gross motor recovery, and reduced the infarct volume under ischemic and hypoxic conditions[17]. Metformin also can improve endothelial dysfunction and neovascularization by targeting stem and progenitor cells[5,17]. For example, metformin directly improves the function of vascular endothelial cells (ECs) and increases blood flow[125]. Furthermore, the antioxidant and anti-inflammation effects of metformin on stem cells contribute to treatment effects on neurodegenerative and cardiovascular diseases[38,54,101,102,107].

Metformin has the potential to be an anti-aging agent associated with stem cells. Adult stem cells are affected by the same aging mechanisms that involve somatic cells, resulting in an impaired regenerative capacity to effectively maintain tissues and organs[55]. For example, skeletal muscle drives human movement, and aging and atrophy of muscle are major signs of human aging. In age-related muscle atrophy (sarcopenia) and several dystrophies, regeneration cannot compensate for the loss of muscle tissue due to depletion of the satellite cell pool or the functional loss of satellite cells. Pavlidou *et al*[14] demonstrated that metformin delayed satellite cell activation and differentiation by favoring a quiescent, low metabolic state, thereby alleviating depletion of the satellite cell pool and the functional loss of satellite cells[14]. Moreover, the antioxidant and anti-inflammatory effects of metformin on stem cells contribute to anti-aging.

Metformin also has the potential to be an anti-cancer agent by targeting CSCs. Metformin inhibited CSCs from three aspects, including the inhibition of self-renewal and metastatic pathways, inhibition of metabolic pathways, and inhibition of chemoresistance pathways.

However, there are still some limitations and issues for the application of metformin and stem cells. For example, the effect of metformin may be different depending on the cell origin. Drug delivery is also an issue. There is still a lack of clinical research on these issues. The mechanisms and effects of metformin on myogenic differentiation and angiopoiesis of stem cells are controversial. A high glucose environment may influence cell differentiation and apoptosis[20,117]. These issues remain to be resolved in future studies.

**CONCLUSION**

In summary, a large number of studies have demonstrated the pleiotropic effects of metformin on stem cells. These inspiring results provide new treatment possibilities in many fields including regenerative medicine, stem cell medicine, anti-aging, and anti-cancer.

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**Footnotes**

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**Figure Legends**



**Figure 1 Effect of metformin and stem cells on tissue injury healing.** TNF: Tumor necrosis factor; HMGB1: High mobility group box 1; LKB1: Liver kinase B1; AMPK: Adenosine 5′-monophosphate-activated protein kinase; GSK3β: Glycogen synthase kinase 3β; IKK: Inhibitory NF-κB kinase; Runx2: Runt-related transcription factor 2; HDAC: Histone deacetylase; mTOR: Mammalian target of rapamycin; MRFs: Myogenic regulatory factors; IL: Interleukin; NF-κB: Nuclear factor kappa-B; PEG: Polyethyleneglycol; SDF-1: Stromalcell-derivedfactor1; eNOS: Endothelial nitric oxide synthase; NO: Nitric oxide.

**Table 1 Summary of effect of metformin on stem cells and suggested mechanisms**

|  |  |  |  |
| --- | --- | --- | --- |
| **Role of metformin** | **Stem cell type(s)** | **Suggested mechanism(s)** | **Ref.** |
| Promoting osteogenic differentiation | BMPC; ADSC; UC-MSC; iPSC-MSC | LBK1/AMPK activation | [20-23] |
| BMSC | AMPK activation-Runx2 (serine 118) | [26] |
| MC3T3-E1 | AMPK/Gfi1/OPN axis; SIRT-6/NF-κB | [12,24] |
| hBMSC | Twist1 inhibition; GSK3β/β-catenin/Wnt signaling pathway | [19,25] |
| PDLSC | AKT/Nrf2 | [27] |
| ADSC; PDLSC; hDPSC | None | [8,11,32,33] |
| Promoting neuronal differentiation | NPC | aPKC/CBP | [35] |
| hBMSC | AMPK activation | [36] |
| NPC | AMPK/aPKC/CBP signaling pathway | [37] |
| hiPSC-NSC | None | [17] |
| Promoting myogenic differentiation | Satellite cell | RPS6-mTOR | [14] |
| C2C12 | ERK; AMPK (AMPKα1)/HDAC5 | [40,41] |
| Muscle progenitor cell | AMPK | [39] |
| Inhibiting adipogenic differentiation | MC3T3-E1 | AMPK/Gfi1/OPN axis | [24] |
| MSC | AMPK/mTOR/p70S6K | [44] |
| ADSC; PDLSC; BMSC; BMPC | None | [13,27,42,43,46] |
| Inhibiting chondrogenic differentiation | ATDC-5 | AMPK | [47] |
| Gastric PC differentiation | Gastric EPC | AMPK | [50] |
| Regulating stem cell aging and rejuvenating regeneration | HMSC | Nrf2/GPx7 | [59] |
| ISC | AKT/TOR/Atg6-related pathway; AKT/mTOR pathway | [62-64] |
| Satellite cell | mTOR /p70S6 | [14] |
| OPC | AMPK activation | [68] |
| Inhibiting CSCs | CSC | Hedgehog, Wnt, and TGF-β pathways | [72] |
| Glioblastoma CSC | C1CL1 | [74] |
| Colorectal cancer CSC | MIF/CD74 axis | [77] |
| Breast CSC | MiR708/CD47 axis | [81] |
| CSC | None | [73,76,82] |
| Improving EPC functions and angiogenesis | EPC | AMPK/eNOS/NO signaling pathway; AMPK/mTOR/autophagy pathway; AMPK/mTOR/p70S6K pathway | [89,90] |
| Antioxidant | ADSC, C2C12 | ROS&NO reduction/SOD activation | [13,40,99] |
| PDLSC | AKT-Nrf2 signaling pathway | [27] |
| HMSC | Nrf2/GPX7 | [59] |
| hNSC | AMPK activation | [98,100,102] |
| Anti-inflammatory | hNSC | AMPK/(IKK/NF-κB) | [15] |
| rabbit AFSC | HMGB1 | [104] |
| Immunomodulatory potential | ADSCs | AMPK/mTOR/STAT-1 signaling pathway | [107] |

BMPC: Bone marrow progenitor cells; ADSC: Adipose-derived stem cells; UC: Ulcerative colitis; iPSC: Induced pluripotent stem cell; MSC: Mesenchymal stem cell; LKB1: Liver kinase B1; AMPK: Adenosine 5′-monophosphate-activated protein kinase; BMSC: Bone marrow stromal cell; Runx2: Runt-related transcription factor 2; Gfi1: Growth factor independence-1; OPN: Osteopontin; NF-êB: Nuclear factor kappa-B; GSK3â: Glycogen synthase kinase 3â; PDLSC: Periodontal ligament stem cell; Nrf2: Nuclear factor E2-related factor 2; DPSC: Dental pulp stem cells; NPC: Neural precursor cell; aPKC: Atypical protein kinase C; CBP: CREB-binding protein; hiPSC: Human induced pluripotent stem cell; NSC: Neural stem cell; RPS6: Ribosomal protein S6 kinase; mTOR: Mammalian target of rapamycin; HDAC5: Histone deacetylase 5; MSC: Mesenchymal stem cell; EPC: Endothelial progenitor cell; HMSC: Human mesenchymal stem cell; GPx7: Glutathione peroxidase 7; ISC: Intestinal stem cell; OPC: Procyanidins oligomers; CSC: Cancer stem cell; TGF-β: Transforming growth factor-β; MIF: Macrophage migration inhibitory factor; eNOS: Endothelial nitric oxide synthase; IKK: Inhibitory NF-κB kinase; hNSC: Human neural stem cell; AFSC: Amniotic fluid or stem cells; HMGB1: High mobility group box 1; STAT-1: Statim-1; SIRT-6: Silent information regulator-6.