Abstract. Sirtuin 3 (Sirt3) is an important member of the sirtuin protein family. It is a deacetylase that was previously reported to modulate the level of reactive oxygen species (ROS) production and limit the extent of oxidative damage in cellular components. As an important member of the class III type of histone deacetylases, Sirt3 has also been documented to mediate nuclear gene expression, metabolic control, neuroprotection, cell cycle and proliferation. In ovarian cancer (OC), Sirt3 has been reported to regulate cellular metabolism, apoptosis and autophagy. Sirt3 can regulate autophagy through a variety of different molecular signaling pathways, including the p62, 5'AMP-activated protein kinase and mitochondrial ROS-superoxide dismutase pathways. However, autophagy downstream of Sirt3 and its association with OC remains poorly understood. In the present review, the known characteristics of Sirt3 and autophagy were outlined, and their potential functional roles were discussed. Following a comprehensive analysis of the current literature, Sirt3 and autophagy may either serve positive or negative roles in the regulation of OC. Therefore, it is important to identify the appropriate expression level of Sirt3 to control the activation of autophagy in OC cells. This strategy may prove to be a novel therapeutic method to reduce the mortality of patients with OC. Finally, potential research directions into the association between Sirt3 and other signaling pathways were provided.

Correspondence to: Dr Bing Wei or Dr Lei Zhan, Department of Gynaecology and Obstetrics, The Second Affiliated Hospital of Anhui Medical University, 678 Furong Road, Hefei, Anhui 230601, P.R. China
E-mail: 517275960@qq.com
E-mail: 499329901@qq.com

Contributed equally

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1. Introduction

Ovarian cancer (OC) is a life-threatening malignancy that represents 3.6% female malignancies worldwide (1). It currently ranks as the seventh most common type of gynecological cancer and 20th as the most common type of cancer worldwide. OC has the highest mortality of all gynecologic malignancies (1,2). According to a recent report, it was estimated that there were 295,414 new cases of OC diagnosed in 2018 and 184,799 cases of mortality resulting from this disease worldwide (3). Tumor debulking surgery followed by platinum and paclitaxel chemotherapy is currently the standard clinical treatment of OC (4). However, the survival rate of patients with advanced OC remains at ~30%, with the primary reasons being late discovery and chemoresistance. In particular, chemoresistance is mediated by both the tumor microenvironment and inherent resistance of OC cells to chemotherapy (5). Therefore, enhancement of responses to current treatment and the development of novel therapeutic strategies are urgently required to improve the survival rate.

Autophagy is a protective, catabolic process that operates to maintain intracellular homeostasis by recycling organelles and macromolecules (6). During this process, defective or aged organelles and other cytoplasmic components are enclosed by double-membrane vesicles to form autophagosomes. This then fuses with a lysosome where the vesicular contents are degraded into amino acids, lipids and carbohydrates by the lysosomal enzymes. The degradation products are in turn recycled to make new proteins and organelles (7). A basal level of autophagy is in operation under physiological conditions (8). However, downregulation or upregulation of autophagy induced by stress factors, including alterations in the levels of growth factors, hypoxia and cytotoxic damage, can result in cell death.
or cell adaptation in response (9). Autophagy also appears to serve contradictory roles in the development of cancer. Evidence exists reporting that inhibition of genes associated with autophagy can promote tumor development, whereas the expression of proteins associated with autophagy has also been demonstrated to result in inhibitory effects in several types of cancers (10-14). Therefore, autophagy can exert anti-tumor effects, but in contrast cancer cells may survive cellular stress in adverse microenvironments by utilizing autophagy, thereby promoting the development of tumors (15). In addition, autophagy is somewhat considered to be a double-edged sword in the clinical field of cancer. Promotion of autophagy can induce cell death, in a manner that is similar to apoptosis, whilst protective cellular autophagy has also been reported to be the major underlying cause of therapy resistance among cancer cells. Therefore, increasing the sensitivity of cancer cells to anticancer therapy by inhibiting autophagy remains a viable option (16).

Sirtuins are nicotinamide adenine dinucleotide (NAD⁺)-dependent histone deacetylases that are highly conserved among eukaryotic organisms, of which seven isoforms exist in mammals (17-19). They serve important roles in a number of biological and pathological processes. In particular, Sirt3 is localized to mitochondria, where it modulates the production of reactive oxygen species (ROS) to limit oxidative damage in cellular components (20). A wide range of important biological activities have been documented to be controlled by Sirt3, including the regulation of nuclear gene expression, metabolism, neuroprotection, cardiovascular disease, cancer cell cycle progression, and cell proliferation and apoptosis (21-24). Previously, the role of Sirt3 as a hallmark of cancer has been attracting significant research attention (25). Metabolism is an important caveat to cancer development (26), such that ATP is required for the maintenance of intracellular metabolic activity (27). A number of metabolic processes have been revealed to be regulated by Sirt3, including fatty-acid oxidation and oxidative phosphorylation (28,29). These observations suggested that Sirt3 could be a key regulator of cancer physiology.

Currently, there are a number of studies on autophagy and Sirt3 in cardiovascular diseases, neuronal diseases and hepatotoxicity (30-32). However, almost no article has conducted research on the relationship between autophagy, OC and Sirt3. Therefore, the present review primarily discussed the potential relationship between Sirt3 and autophagy in OC, with the aim to provide a possible novel direction for OC research and therapeutic strategies.

2. Sirt3 in OC

OC poses a significant threat to the health of women worldwide, and is a disease in which Sirt3 has been reported serve a regulatory role. Of note, this disease is gradually becoming the leading cause of mortality associated with gynecological cancer worldwide in both developing and developed countries (1). Several reports have suggested that OC is regulated by Sirt3 using a multitude of mechanisms, which is summarized in this section.

In a previous study, it was found that muscle tissues after exercise exhibit elevated expression levels of the Sirt3 protein, which gave rise to the hypothesis that the expression of Sirt3 is regulated by energy metabolism (33). Energy metabolism is also associated with the regulation of tumor growth and metastasis. Sirt3 is regarded as a tumor suppressor, due to a previous finding that its expression is reduced in tumors (34,35). The activation of cellular autophagy and apoptosis was demonstrated to be controlled by Sirt3 via the regulation of several signaling pathways during the development of OC. A previous study found that expression of the Sirt3 protein was significantly downregulated in OC tissues and in highly metastatic HO-8910PM cell lines (35,36). In addition, Xi et al (37) demonstrated that the activation of Sirt3 exerted a proapoptotic function in SKOV3 cells. These findings suggested that overexpression of Sirt3 can induce OC cell death. In terms of mitochondrial dynamics, a previous study revealed that stabilization of Sirt3 can increase mitochondrial biogenesis and cristae remodeling in OC tissues (38). Additionally, stabilization of optic atrophy protein 1, which increased resistance to apoptosis, was demonstrated to be regulated by increasing the expression of Sirt3 and prohibitin 2 (38). Activation of Sirt3 has also been found to enhance the sensitivity of OC cells to cisplatin (39), rendering Sirt3 to be a novel therapeutic target. In addition, Sirt3 was reported to be a favorable independent prognostic factor for overall survival for patients with serious OC in a previous study (40). In conclusion, Sirt3 serves an important role in the development of OC, with therapeutic and prognostic implications.

3. Autophagy in OC

Autophagy is a catabolic process that serves to maintain intracellular homeostasis by recycling damaged cellular organelles (41), which has been studied since the 1960s (42). Over the past decades, the molecular mechanisms underlying this process have been revealed gradually. It has been suggested that autophagy is a common phenomenon that occurs during both physiological and pathological conditions. According to the sizes of the substrates involved and degradation rate, autophagy can be divided into three sub-categories: i) Macro-autophagy; ii) microautophagy; and iii) chaperone-mediated autophagy (43). Although different types of autophagy utilize distinct mechanisms to degrade lysosomal proteins, common underlying characteristics remain (44). The autophagy pathway consists of the following six steps: i) Initiation of autophagy; ii) biogenesis of the phagophore; iii) expansion of the phagophore; iv) formation of the autophagosome; v) fusion with the lysosome; and vi) reformation of the lysosome (45). Autophagy is constitutively active at low levels in all cell types under physiological conditions, but can be potentiated by nutrient deprivation, hypoxia, endoplasmic reticulum stress, pathogenic toxicity and immune injury, to maintain intracellular homeostasis (46). Previous studies have demonstrated that autophagy serves an important role in the pathological processes of various diseases, including neurodegenerative and cardiovascular disease, cancer, infectious diseases and immune deficiency (47-50). Autophagy has been described as a double-edged sword, since it can exert both tumor suppression and growth promotion (51). In this section, the mechanism of autophagy in OC is discussed in detail.
Autophagy-related PI3K/AKT/mTOR signaling pathway. The autophagic process in OC is regulated by a number of factors. The PI3K/AKT/mTOR pathway has frequently been associated with the majority of human malignancies, studies have demonstrated that other signaling pathways related to oncogenesis are also caused by dysregulations in this signaling pathway (55-57). The reason for this dysregulation is manifold, including mutations in PI3K, AKT overexpression and the sustained activation of tyrosine kinase growth factor receptors (58). The PI3K/AKT/mTOR signaling pathway has been documented to regulate cell survival, proliferation, growth, transcription, angiogenesis and metabolism (55-57). In ~70% cases of OC, the PI3K/AKT/mTOR pathway has been revealed to be constitutively activated, which has been considered to be a therapeutic target (59). To verify if the PI3K/AKT pathway is involved in OC cell proliferation, Hu et al (60) treated OC cell lines with the specific PI3K inhibitor LY294002 and established a mouse model of OC. Proliferation of OC cells can be significantly inhibited by LY294002 treatment in vitro (61). In addition, other studies have found that AKT inhibitors can prevent the function of mTORC1/2 and AKT itself to inhibit the PI3K signaling cascade (61-63). In another study, Ichikawa et al (64) found that the cytotoxic effects of chemotherapeutic agents can be effectively enhanced by co-treatment with the selective non-competitive AKT inhibitor TAS-117 in vivo OC models. In OC, the PI3K/AKT/mTOR signaling pathway is frequently activated, which indicates that the inhibition of this signaling pathway could prove to be a potential avenue of treatment strategies, either as a monotherapy or in combination with other chemotherapeutic agents.

p53 signaling pathway. p53 is a key tumor suppressor that serves an important regulatory role in autophagy in mammalian cells (65). Expression of the p53 gene is activated by various intracellular events, including DNA damage, hypoxia and oncogene activation, to prevent cell damage and maintain cellular integrity. Numerous types of modifications, including acetylation, methylation, phosphorylation and ubiquitination, are involved in regulating the activation of p53 on a molecular level (66,67). p53 target genes negatively regulate mTOR activity, which in turn induces autophagy in the nucleus. p53 can promote autophagy by inhibiting mTOR via the 5’AMP-activated protein kinase (AMPK) pathway (68). In addition, p53 can also induce autophagy by activating damage-regulated autophagy modulator (69). Several studies have revealed that autophagy may be triggered by the inactivation of cytoplasmic p53, such that extranuclear p53 is an effective inhibitor of autophagy (70,71). A clinical study previously demonstrated the upregulated expression of p53 in OC, where it was revealed that at later tumor stages, the rate of p53-positive expression was higher (72). These results suggested that p53 serves an important role in the development of OC. Additionally, another study previously found that silencing the p53 signaling pathway may suppress proliferation whilst facilitating apoptosis and cisplatin chemosensitivity in OC cells (73). Therefore, these aforementioned findings suggested that the efficacy of chemotherapeutic treatments for OC can be improved by inhibiting the p53-induced autophagic process.

4. Relationship between Sirt3 and autophagy

Sirt3 belongs to the NAD+-dependent protein deacetylase family and is responsible for the majority of mitochondrial protein deacetylation (74). It has been reported to serve an important role in almost every aspect of mitochondrial biology, such as mitochondrial biogenesis and mitochondrial oxidative stress (30,75-77). Although Sirt3 is located in mitochondria, almost every major key component of autophagy is cytosolic. Therefore, a type of communication mechanism must exist between mitochondria and cytosol during the regulation of Sirt3 activity and autophagy. In previous years, accumulating evidence has suggested that there are a series of signaling pathways between Sirt3 and autophagy, including the p62, PI3K/AKT, AMPK-mTOR pathway and the mitochondrial ROS-superoxide dismutase 2 (SOD2) pathway. This section aimed to summarize the known information on its mechanism (Fig. 1).

Sirt3 and p62. p62 is the selective cargo receptor for autophagy that is indispensable for the degeneration of misfolded proteins. Reductions in p62 expression have been previously reported to activate autophagy (78). In addition, p62 is considered to be a marker of autophagic flux due to its differential expression profiles in association with other proteins linked to autophagy (79). As for Sirt3, a number of studies have indicated that there is a close relationship between Sirt3 and the p62-autophagy pathway. A previous study demonstrated that treatment with ANXA1sp, an annexin-A1 bioactive peptide, reduced the expression of p62 and concomitantly upregulated the expression of the mitochondrial protein deacetylase Sirt3 (80). This suggested that Sirt3 is involved in autophagy by downregulating p62. Tong et al (81) also reported a similar finding, where liraglutide treatment downregulated the expression of p62 to promote autophagy via the Sirt1/Sirt3-forkhead box O3-p62 pathway in mice that were fed a normal-fat diet or high-fat diet. Additionally, a previous investigation focusing on the effects of melatonin on diabetic cardiomyopathy revealed that melatonin can upregulate autophagy by increasing the expression of LC3-II whilst downregulating that of p62 (82), with the macrophage stimulating 1/Sirt3 signaling pathway being the likely associated mechanism. These aforementioned
pharmacological studies suggested that Sirt3 is a potent activator of autophagy. Supporting this, Xiang et al. (83) found that small interfering (si)RNA-mediated silencing of Sirt3 gene expression inhibited the process of autophagy and p62 degradation in human umbilical vein endothelial cells.

**Sirt3 and PI3K/AKT.** The PI3K/AKT pathway serves an important inhibitory role in autophagy, and is also involved in a variety of physiological and pathological processes (84,85). By regulating the PI3K/AKT pathway, Sirt3 can function as an autophagy suppressor. A previous report that investigated hepatocellular carcinoma found that the expression levels of Sirt3 was higher in adjacent non-cancerous tissues compared with those in hepatocellular carcinoma tissues (86). Furthermore, using Sirt3 knockdown, this previous study also demonstrated that Sirt3 may serve as a suppressor of autophagy in hepatocellular carcinoma by targeting the PI3K/AKT pathway. Another study revealed a consistent finding, where Sirt3 functioned as a growth suppressor in prostate cancer by inhibiting the activation of PI3K/AKT both in vitro and in vivo (87). This previous study also showed that the progression of prostate cancer may be downregulated via the Sirt3/AKT/c-Myc signaling axis (87). Wang et al. (88) demonstrated that glioblastoma multiforme cell growth was inhibited through the Sirt3/p53-mediated PI3K/AKT/ERK and mitochondrial signaling pathway. In addition, Sirt3 was reported to indirectly regulate AKT hyperactivation by regulating mitochondrial ROS production upstream of the ROS-mediated Ras-PI3K-AKT activation (89).

**Summary.** Sirt3 can regulate autophagy by either directly or indirectly regulating the PI3K/AKT pathway.

**Sirt3 and AMPK.** AMPK is a conserved intracellular energy sensor that mediates energy homeostasis by regulating lipid and glucose metabolism (90). Dysregulation of AMPK has previously been associated with accelerated aging (91). In previous years, a number of studies have attempted to unravel the mechanism underlying the regulation of autophagy. There is increasing consensus that AMPK and mTOR are regarded as the main regulators of autophagic degradation (92,93). A close mechanistic relationship has been reported to exist between Sirt3 and the AMPK-mTOR-autophagy pathway. Zhao et al. (94) previously found that induction of autophagy was directly controlled by Sirt3 via the AMPK-mTOR pathway during acute kidney injury. In addition, another study reported that Sirt3-induced autophagy protected against oxygen and glucose deprivation by regulating the AMPK-mTOR pathway (30).

Previous studies have found that AMPK can negatively regulate mTORC1 activity via two different mechanisms. AMPK can phosphorylate Thr1227 and Ser1345 residues to activate tuberous sclerosis complex (TSC) 2, thereby promoting the formation of the TSC1/TSC2 heterodimer to inhibit mTORC1 activity (95). By contrast, AMPK can also phosphorylate regulatory-associated protein of mTOR on its Ser722 and Ser792
residues to inhibit mTORC1 (96). Liver kinase B1 (LKB1) is a tumor suppressor and an upstream regulator of AMPK that has an essential role in the control of redox homeostasis (97). Incidentally, LKB1 can also be activated by Sirt3. A previous investigation into a rotenone-induced SH-SY5Y cell injury model found that the overexpression of Sirt3 promoted LKB1 phosphorylation, which activated AMPK and reduced the phosphorylation of mTOR. This observation suggested that the LKB1-AMPK-mTOR pathway can be regulated by Sirt3 (98). In addition, another previous study documented that activation of the LKB1-AMPK-mTOR-mediated autophagy signaling pathway can be induced by Sirt3 (99). Notably, AMPK can function upstream of Sirt3 during the regulation of insulin sensitivity in skeletal muscle via the AMPK-peroxisome proliferator-activated receptor γ coactivator 1c-Sirt3 autophagy signaling pathway in Sirt3-/-mice (100). Collectively, this indicated that Sirt3 and autophagy have a complex mutual regulatory relationship.

Sirt3 and ROS. ROS consists of a group of highly reactive chemical entities, including oxygen radicals, hydroxyl, peroxy, alkoxyl, non-radicals, singlet oxygen and hydrogen peroxide. These molecules are primarily produced from redox transactions as part of the oxidative phosphorylation system in mitochondria (100). A number of studies have demonstrated that ROS can act as a signal to trigger autophagy through autophagy-regulating protease 4, which serves as part of the autophagy process (101,102). Autophagy may be suppressed by Sirt3 via the regulation of mitochondrial ROS (mROS) production. Recently, a study investigating the potential antineoplastic properties of metformin combined with nelfinavir revealed that this drug combination can increase the expression level of Sirt3. This increment in Sirt3-mediated mROS production served a vital role in the autophagic mechanism within human cervical cancer cells (103). Upstream of ROS, SOD2 activity can moderately reduce cellular ROS levels. Sirt3-mediated deacetylation can significantly potentiate SOD2 activity to ultimately break down intracellular ROS (104). A previous study demonstrated that Sirt3 protein expression and activity was downregulated by cadmium (Cd), which can also concurrently promote the acetylation of SOD2 to suppress its activity, thereby increasing mROS production (31). In summary, mitochondrial-derived ROS-dependent autophagic cell death can be induced by Cd. Consistent with this notion, Cd-induced hepatotoxicity has been reported to be alleviated by the protective properties of melatonin (31). In this particular study, the activity, but not the expression of Sirt3, was revealed to be enhanced by melatonin treatment (31). Additionally, melatonin was also demonstrated to inhibit mitochondria-derived O₂ production, reduced the acetylation of SOD2 and suppressed autophagy (31). This finding suggested that melatonin exerts hepatoprotective effects by regulating mitochondria-derived O₂-stimulated autophagic cell death via the Sirt3-SOD2 pathway. According to the aforementioned findings, SOD2 activity and intracellular mROS homeostasis may underlie the Sirt3 downregulation of autophagy (31).

5. Potential relationship between Sirt3 and autophagy in OC

Autophagy is a fundamental catabolic process that has been reported to be involved in the progression of a variety of diseases. It can serve a protective role in OC cells from cell death since it may enhance resistance to cisplatin (105). A previous investigation revealed that cisplatin treatment activated autophagy, whereas Bcl-2-associated athanogene 3 attenuated cisplatin resistance by inhibiting autophagy (106).
However, recent studies have also suggested that autophagy can inhibit the growth of OC, since it has been found to promote OVCAR-3 cell death (107). By contrast, findings from another previous study suggested that inhibition of autophagy promoted the proliferation and invasion of OC cells via the PI3K/AKT/mTOR pathway (108). These findings demonstrated that autophagy is a double-edged sword in the regulation of OC physiology. Sirt3 serves an important role in the maintenance of intracellular homeostasis in OC. Previous studies have indicated that there is a close mutual regulatory relationship between Sirt3 and autophagy, which are linked by the aforementioned signaling pathways in the present review. Metformin-induced overexpression of Sirt3 promoted apoptosis and mitochondrial dysfunction whilst increasing the activation of AMPK in OC cell lines (109). In addition, metformin has been documented to promote autophagy in OC (110). S1, a novel Bcl-2 inhibitor, has also been reported to exert autophagic effects in OC by interrupting the interaction between Bcl-2 and beclin1 in OC to promote apoptosis (111); however, high doses and longer exposure of S1 can overpower the protective function of autophagy and induce apoptosis (112,113). Yang et al (114) previously found that S1 promoted JNK3 expression, thus increasing cell sensitivity to apoptosis. Sirt3 can also regulate autophagy via the glutathione S-transferase PI3K/AKT/mTOR pathway, such that Sirt3 knockdown has been demonstrated to alleviate S1-induced apoptosis (Fig. 2) (37).

6. Conclusion

Autophagy serves an important role in recycling damaged organelles and maintaining intracellular homeostasis. Sirt3 is a potential therapeutic target, since it has been previously reported to activate autophagy. In the present review, although the potential relationship between Sirt3 and autophagy in OC was explored, there remains an insufficient number of studies on this topic. The associated underlying mechanism in the Sirt3-induced autophagic process in OC remains unclear. With further study, novel insights into the molecular relationship between Sirt3 and autophagy may contribute to the development of novel therapeutic interventions for OC.

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YS and RH contributed to the conception of this manuscript and wrote the draft. YY contributed to the literature collection and preparation. YH contributed to the revision of this manuscript. LZ and BW provided funding and proofed the final version of the manuscript.

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Competing interests

The authors declare that they have no competing interests.

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