Review

Therapeutic Implications of Autophagy Inducers in Immunological Disorders, Infection, and Cancer

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Abstract: Autophagy is an essential catabolic program that forms part of the stress response and enables cells to break down their own intracellular components within lysosomes for recycling. Accumulating evidence suggests that autophagy plays vital roles in determining pathological outcomes of immune responses and tumorigenesis. Autophagy regulates innate and adaptive immunity affecting the pathologies of infectious, inflammatory, and autoimmune diseases. In cancer, autophagy appears to play distinct roles depending on the context of the malignancy by either promoting or suppressing key determinants of cancer cell survival. This review covers recent developments in the understanding of autophagy and discusses potential therapeutic interventions that may alter the outcomes of certain diseases.

Keywords: autophagy; immunity; cancer; activators

1. Introduction

The maintenance of cellular homeostasis in response to environmental stress is crucial for the survival of all cell-based life forms. Under metabolic stresses such as nutrient or growth factor restriction, there are physiological systems in place to eliminate damaged proteins, lipids, and organelles. Many of these intracytosolic components can be recycled to some extent in a ‘self-eating’ process called autophagy. The process can be induced by amino acid, glucose, or oxygen starvation, growth factor deficiency, and other forms of cellular stress accompanied by cytotoxic damage, and permits the reuse of damaged proteins and organelles [1,2]. Damaged or dysfunctional intracellular components including unfolded proteins are ubiquitinated for selective recognition by autophagy receptors [3]. Upon recognition, the damaged components are then exported to lysosomes for autophagic degradation and delivery to the cytoplasm where they are recycled. As a self-cleansing mechanism, autophagy is indispensible for ensuring the integrity of proteins and organelles, as well as energy homeostasis [2,4].

Autophagy is categorized into three major types. Microautophagy is a non-selective degradation process that involves the direct engulfment of small volumes by lysosomes. Chaperone-mediated autophagy (CMA) is a selective degradation process that recognizes and translocates specific proteins to the lysosome through lysosomal-associated membrane protein 2 (LAMP2), a receptor for the CMA pathway. Macroautophagy involves double-membrane bound autophagosomes and targets damaged components in the cytoplasm for degradation. This is the most well-characterized form of autophagy and the type primarily discussed in this review.
In normal tissues, low levels of basal autophagy catabolize damaged cellular components to maintain balanced cellular homeostasis. Multiple studies have linked autophagy and the pathology of various diseases including cancer, neurodegenerative diseases, infectious disease, autoimmunity, and aging [5,6]. Here, we briefly review the pivotal roles of autophagy in regulating immunity and cancer, and discuss the potential for autophagy inducers to be developed as therapeutic agents.

2. Role of Autophagy in Immunity

Autophagy as a fundamental process for maintaining cellular integrity and is intimately linked with immune responses. The deletion of autophagy signaling factors such as the autophagy-related gene (Atg) leads to defects in immune responses and related pathogenesis. T cell-specific deletion of beclin 1 (Atg6) in mice fails to induce T cell-dependent humoral immune responses and resistance to encephalomyelitis [7]. Mice lacking Atg16L1 in haematopoietic cells are prone to acute colitis induced by dextran sodium sulphate through endotoxin-induced inflammasone and IL-1β secretion, highlighting the crucial role of autophagy in endotoxin-induced inflammatory immune responses [8]. There have been multiple proposed mechanisms by which the immune system regulates autophagy, including the direct removal of microbes during infection, antigen presentation, lymphocyte homeostasis, and the control of inflammation [9,10].

Recognition of infected microbes by pathogen-recognition receptors such as TLRs, or NLRs (NOD-like receptor) induces autophagy to facilitate their elimination [6]. Activation of TLR4 or TLR7 by LPS or single-stranded RNA (ssRNA) leads to the formation of autophagosomes and eliminates mycobacteria, showing a strong association between autophagy and innate immunity via TLR signaling [11]. LPS-stimulated TLR4 triggers the activation of MyD88- and TIR domain-containing adapter-inducing interferon-β (TRIF)-dependent signaling which in turn mediates upregulation of nuclear factor kappa B (NF-κB) and activator protein 1 (AP1) signaling, leading to innate immune defenses such as pro-inflammatory cytokine production [12]. NLRP4, a member of the NLR family, detects bacterial infection and induces Beclin 1-mediated autophagy in response [13]. Recently, the role of autophagy in the expression of resistance to Mycobacterium tuberculosis was reported [14,15]. Mice with macrophage-specific deletion of Atg5 were more susceptible to infection against M. tuberculosis and developed severe inflammation in the lungs [16]. However, other autophagy signaling genes including Atg14, Atg12, Atg16L1, Atg7, and Atg3 had little effect on survival during M. tuberculosis infection, suggesting the possibility of a specific role for Atg5 against neutrophil-mediated inflammation in an autophagy-independent manner. These findings also imply that autophagy genes might not all play critical roles during mycobacterium infection [17].

In the adaptive immune response, autophagy also plays an essential role in antigen processing and presentation through major histocompatibility complexes (MHC). Traditionally, it is known that MHC-I presents antigens derived from intracellular proteasomal proteins to CD8+ T cells, whereas MHC-II displays extracellular antigens and presents to CD4+ T cells for lysosomal degradation [18]. Recently, the emerging role of autophagy for presentation of intracellular antigens by MHC-II has been reported [19,20]. Approximately 10–25% of MHC-II presents cytoplasmic and nuclear antigens via autophagy, and several mechanisms are proposed by previous studies [20,21]. Cytoplasmic antigens are degraded by cytosolic proteases and processed by factors in the CMA pathway including LAMP-2A and hsc70 to access lysosomes [22]. Via macroautophagy, intracellular antigens are sequestered and encapsulated by autophagosomes that bind to Atg8/LC3, which then interacts with MHC-II to form MHC-II compartments (MIIC) for lysosomal degradation [21]. MHC-II molecules are combined with invariant chains (Ii) in the endoplasmic reticulum (ER), and transported to MIIC by fusing with late endosomes and autophagosome-lysosomes. The antigen fragments captured by MHC-II are catalyzed by the MHC-encoded molecule HLA-DM in MIIC, which delivery of peptide fragment-MHC-II complexes to the cell surface for surveillance by CD4+ T cells [21,23].

Autophagy can also regulate the homeostasis of lymphocytes. More specifically, it is involved in the development, activation, survival, and differentiation of T lymphocytes [24]. The maintenance
of organelle homeostasis in T cells is one of the crucial roles of macroautophagy. The inhibition of macroautophagy by deletion of Atg7 in T cells induces defective survival, enhances mitochondrial contents, increases ROS production, and dysregulates the expression of apoptotic proteins [25,26]. Atg7-deficiency in T cells also induces the accumulation of content in the ER and calcium stores, showing the importance of autophagy in maintaining ER and calcium homeostasis in T lymphocytes [27].

Autophagy plays an important role for T cell survival. Loss of Atg5 in T cells leads to the accumulation of abnormal proteins and organelles in the cytoplasm and stimulates cell death via the generation of ROS and impaired mitophagy [26,28]. Disruption of Beclin 1 in T cells induces an increased accumulation of the pro-apoptotic proteins Bim, caspase-3 and -8, further support that under the absence of autophagy pathways, abnormal survival and apoptotic responses occur in T cells [29].

The full activation of T cells requires several signaling steps including the engagement of antigens with T cell receptors (TCR) via MHC, and subsequent engagement of co-stimulatory molecules including CD28 and IT2 signaling. Previous studies have demonstrated the importance of TCR engagement on CD4+ and CD8+ T cells for the induction of macroautophagy [30,31]. Full activation of T cells by TCR engagement upregulates the lipidated form of LC3 (LC3-II) in an mechanistic target of rapamycin (mTOR)-independent manner [24,30]. Selective degradation of specific proteins by p62-mediated macroautophagy is caused by modulation of Bcl10, which is downstream of the TCR. TCR engagement promotes ubiquitination of Bcl10 and binding with p62, which gradually induces degradation by autophagy. Silencing of p62 blocks Bcl10 degradation, and disruption of Atg3 in CD4+ T cells reduces TCR-mediated Bcl10 degradation, enhancing NF-κB activation [32]. In vivo studies using mice models with a deleted Atg7 gene in T cells revealed defective production of IL-2 and IFN-γ and decreased proliferation in response to T cell stimulation [31]. In contrast, there are several reports showing an opposite role for TCR activation in autophagy. T cell activation by TCR engagement can induce proliferation with very low levels of autophagy activity [33]. In addition, Bcl10-mediated autophagy regulates TCR activation of NF-κB [32,34], further demonstrating the complexity of TCR activation in autophagy modulation.

An emerging role for autophagy in the formation of memory CD8+ T cells has also been suggested in recent studies. The loss of Atg5 and Atg7 in mice triggers the formation of memory T cells with defects in survival during the contraction phase to memory phase transition, suggesting that autophagy is an essential process for antigen-specific effector CD8+ T cell memory formation [33]. Another study reported that autophagy in CD8+ T cells is dramatically reduced in aged mice, and deficiency in the Atg7 autophagy gene leads to a failure of memory formation in CD8+ T cells against influenza and MCMV infection [35]. In order to examine the mechanism responsible, metabolic and transcriptomic analyses were performed and it was shown that defects in autophagy (Atg7-deficient CD8+ T cells) caused dysregulation in lipid biosynthetic pathways in T cells during transition to the memory phase. Furthermore, this alteration in metabolic homeostasis may hinder growth factor signals required for memory T cell development [33]. In addition, Atg7−/− antigen-specific CD8+ T cells displayed problems in regulating metabolic markers and maintaining healthy mitochondria compared to their wild-type counterparts [35].

Autophagy also functions to control the balance of abnormal immune activation and inflammation [6]. Epidemiological studies in Caucasian and Chinese populations demonstrated a positive association between Atg5 expression and systemic lupus erythematosus (SLE) by meta-analysis [36]. Activation of the autophagy signaling proteins Beclin 1 and Atg7 has been reported in the pathogenesis of rheumatoid arthritis [37]. By genome-wide association studies, ATG16L1, NOD2, and IRGM were identified as major risk factors of inflammatory Crohn’s disease (CD) [38]. The variation (T300A) in ATG16L1 causes an autophagy-associated defect in the function of Paneth cells, which could be potentially linked with the pathogenesis of CD [39]. Loss of Atg16L1 in mice led to increased production of pro-inflammatory cytokines IL-1β and IL-18 by macrophages [8]. These studies indicate that autophagy might be crucial in the control of immunity and inflammation. Although
further mechanistic studies for autophagy-regulated immune-related molecules are needed, evidence suggests that beclin 1 might be disrupted by TLR adaptors, and NOD1/2 interacts with Atg16L1 and recruits LC3 [6,40].

As immunosurveillance plays a crucial role in carcinogenesis, the control of immune responses by autophagy can be a critical factor in determining the fate of cancer cells [9,41]. Tumors emerge by evading immune surveillance and altering their antigenic properties, as well as by suppressing anticancer immunity [42]. Recently, the activation of host immunity to treat cancer has been recognized as one of the most promising novel therapeutic approaches [43]. Because autophagy is deeply involved in the proper functioning of various immune cells, controlling the autophagy pathway may stimulate immune reactions against tumor cells [9]. Also, malignant cells with autophagy deficiencies have been reported to suppress the recruitment of immune cells after chemotherapy, reducing anticancer immunogenicity [44].

Taken together, these reports suggest that autophagy as a central modulator of immunity could be harnessed for the treatment of various immune-mediated inflammatory disease including bacterial and viral infections, transplantation rejection, cancer immunotherapy, and other inflammatory disorders.

3. Role of Autophagy in Cancer

Autophagy is clearly indispensable for homeostasis and survival in times of cellular stress in normal tissues. However, in some situations, it also enables cancer cells to circumvent programmed cell death [45]. Due to their unrestrained cell growth, cancer cells typically exhibit high energy demands which can be supported through alternative metabolic pathways, including autophagy. With defects in apoptosis, cancer cells can tolerate long-term metabolic stresses including glucose, cytokine, and oxygen deprivation, and can overcome such restrictions by relying on nutrient recycling mechanisms mediated by autophagy. Detecting environmental nutrient status is crucial for autophagy, and mTOR and AMP-activated protein kinase (AMPK) signaling act as key nutrient sensors in the autophagic machinery [46]. Nutrient and growth factor-induced mTOR complex 1 (mTORC1) activation inhibits autophagy by phosphorylating relevant target proteins such as unc-51-like kinase 1 (ULK1), ATG13, and ATG14L, thereby promoting cell growth [47,48]. Conversely, autophagy is induced by AMPK through the direct phosphorylation and activation of ULK1 on Ser 317 and Ser 777 under nutrient deprivation, whereas the hyperactivation of mTOR suppresses ULK1 by inducing phosphorylation at Ser 757 [49].

Autophagy has been reported to play dual roles in cancer, showing both anti-tumor and tumor-promoting effects depending on the carcinogenic stage and tissue involved [50]. Loss-of-function in the autophagy pathway via genetic mutation or knockdown of autophagy regulators such as beclin 1, Atg5, and Atg7, or constitutive activation of PI3K or mTOR signaling promotes cell survival and the proliferation of adenomas and carcinomas [1,51–53]. Liver-specific deficiency of Atg7 and mosaic deletion of Atg5 in mice causes the spontaneous development of liver adenomas marked by p62 accumulation, the induction of oxidative stress, and genomic instability [54]. Beclin 1 is also a key mediator of autophagy, and mutations in this gene can be found in 40–75% of human sporadic mammary, ovarian, liver, and prostate cancers [55–57]. The heterozygous loss of Beclin1 in mice causes the spontaneous development of lymphomas, colorectal carcinogenesis, and hepatitis B virus-induced hepatocellular carcinoma [56,58,59]. However, genetic deletion of single autophagy proteins such as Atg5 or Atg7 only induces the formation of benign liver tumors without additional tumors in other organs, and cells isolated from Beclin1+/− mice still exhibited autophagic activity [54,56]. Thus, it is difficult to determine the role of autophagy in tumor suppression based on observations from models of genetic deletion of a single autophagy protein.

Multiple reports have suggested that autophagy can be used to evade cell death in some cancer cells to survive during environments with limited energy. Various cancer cells have high levels of basal autophagy even under nutrient-rich conditions, highlighting a more complex role for autophagy in regulating cancer cell survival. Autophagosome formation is known to be critical for the survival
of some tumors, especially in hypoxic regions [60,61], playing a major role in long-term tumor survival and re-growth after chemotherapy by allowing residual tumor cells to recover or metastasize. Crosstalk exists between the autophagy and apoptosis pathways, and cancer cells with defects in the apoptosis pathway can feature upregulated autophagy dependence [62]. In apoptosis-defective cancer cells, blocking autophagy can trigger acute necrotic cell death arising from a failure to withstand metabolic stress, indicating that autophagy may represent a therapeutic target for selective cancer treatment [62–64]. Selective tumor treatment is possible by combining autophagy inhibitors with metabolic stress to sensitize tumor cells to cancer-specific necrotic cell death [63]. The inhibition of angiogenesis, growth factors, and related receptors common in many current cancer therapies often promotes intense metabolic stress on the glycolysis pathway [1]. Metastatic tumor cells have high energy demands to enable proliferation, migration, invasion, and angiogenesis, and residual cancer cells remaining after chemotherapy or the surgical removal of tumors also require favorable conditions for re-growth and survival during times of energy restrictions. In mouse models, the suppression of autophagy by chloroquine (CQ; a lysosome acidification inhibitor) promotes p53 activity by inducing apoptotic cell death in lymphoma cells [65]. Combination therapies involving anti-cancer agents such as tamoxifen, bortezomib, 5-FU, cisplatin, imatinib, rapamycin, and hormone therapy with autophagy inhibitors like 3-methyladenine (3-MA), BAF (bafilomycin A), CQ, or the knockdown of autophagy signaling mediators (ATG5, ATG6, and ATG7) by siRNA significantly inhibits cancer cell growth and metastasis in breast, colorectal, prostate, skin, and gastric cancers [66–69]. These results strongly suggest that inhibition of autophagy signaling may form part of a promising combination therapy for malignant, metastatic or residual cancer cells.

Despite the beneficial effects of autophagy inhibition toward sensitizing cancer cells, autophagy still acts as a barrier against the generation of abnormal proteins and organelles in normal cells. Precisely how autophagy upregulation suppresses tumorigenesis is only partly understood, but several potential mechanisms have been proposed. Loss-of-function in autophagy signaling mediators such as Beclin 1, Atg5, and Atg7 is strongly associated with tumor initiation and progression [57,70]. Conversely, over-stimulation of autophagy in tumors leads to excessive cellular damage and triggers autophagic cell death [71]. There are numerous potential benefits for autophagy activation in tumorigenesis. Autophagy limits genome instability and mutations by inducing apoptosis-independent cell death and eliminating damaged intracellular components. Under starvation conditions, autophagic cell death and the recycling of components is crucial for DNA damage repair and the maintenance of cellular homeostasis. Defects in endoplasmic reticulum (ER) chaperones and p62 accumulation in tumor cells induces reactive oxygen species (ROS) production and genome damage [71]. To eliminate damaged mitochondria, a specific form of autophagy in mitochondria, known as mitophagy, is activated by the PTEN/PINK1/PARK2 signaling pathway, and involves the ubiquitylation of damaged mitochondrial outer membrane proteins [72]. Moreover, as a crucial activator of nuclear factor erythroid-derived-2-like 2 (NRF2), p62 interacts with Keap1-NRF2 signaling. Liver-specific loss of autophagy function through Atg7 deletion induces the nuclear accumulation of NRF2, which stimulates NRF2-related gene expression and promotes tumorigenesis by p62 accumulation [54,73,74]. Defects in autophagy that promote oxidative stress and chronic tissue damage also lead to an upregulation of inflammatory cytokines and the promotion of tumor cell growth [75,76]. Increased inflammatory responses are associated with cancer progression under autophagy-restricted conditions, suggesting that combination therapies with anti-inflammatory agents and autophagy activators for cancer treatment might suppress cancer growth with a combination of calorie restrictive conditions and exercise [2]. Depending on the context, such as tumor type and stage, it appears that autophagy can have varying effects on cancer development. However, it is evident that autophagy plays a major role in cancer cell survival and therefore the development of autophagy-targeting agents is likely to be therapeutically useful if applied in the right context.
4. Activators of Autophagy for Immunity

4.1. BRD5631

Using a high-throughput screening system based on the assessment of GFP-LC3 punctae per cell, BRD5631 and other compounds with autophagy-activating activity have been identified [77]. The hit compounds identified promoted autophagy without inhibiting the mTOR signaling pathway. Autophagy is known to enhance the clearance of bacterial pathogens [9] and BRD5631 and two other selected compounds were observed to promote antibacterial autophagy. These compounds were also able to inhibit replication of *Salmonella* without showing direct bactericidal activity, and suppressed IL-1β secretion induced by lipopolysaccharide and muramyl dipeptide in macrophages. As these compounds work independently of the mTOR pathway, using these autophagy modulators may facilitate the regulation of immune responses in situations where inhibiting mTOR would cause undesirable side effects.

4.2. Spermidine

Spermidine is a natural polyamine found in various foods including soybean, broccoli, cauliflower, chicken, steak and potato [78]. Spermidine promotes cardioprotection and lifespan extension in yeast, flies, worms, and mice [79,80]. The primary mechanism responsible for these health-promoting effects is suggested to be the autophagy-enhancing ability of spermidine in an mTOR-independent manner [79–81]. In addition, spermidine has also been implicated in the control of immune responses. Mice lacking the autophagy gene *Atg7* in T cells exhibit defective memory CD8+ T cell formation in response to influenza and murine cytomegalovirus infection [35]. The study also showed that old mice had reduced memory CD8+ T formation and decreased autophagic flux. However, treatment with spermidine induced autophagy in T cells and increased influenza-specific CD8+ T cell responses in old vaccinated wild-type mice, but not in the old vaccinated T cell-*Atg7* knockout mice. While mTOR inhibition by rapamycin has been reported to stimulate the T cell memory response as well [82], spermidine was shown to activate autophagy and improve memory T cell generation without affecting the mTOR pathway. These results demonstrate that spermidine may be used to improve immunity through the activation of autophagy, especially in the aged population [35].

4.3. Trehalose

Trehalose is a disaccharide found in microorganisms, insects, invertebrates, and plants, and has been reported to induce autophagy via an mTOR-independent pathway [83]. Human cytomegalovirus (HCMV) can infect various types of cells and is a major cause of viral birth defects [84]. Additionally, HCMV has been linked with the development of atherosclerosis and brain tumors [85–87]. A previous study has shown that activating autophagy by trehalose may represent a potential therapeutic approach for treating HCMV disease [88]. Trehalose increases the number of autophagosomes and autolysosomes in HCMV-infected cells and suppresses HCMV replication and viral spread. It was also demonstrated that autophagy-inducing activity as well as inhibitory effects against HCMV were reproducible in three different cell types. Upregulating the host autophagy pathway as an antiviral approach using a natural compound is a promising approach, but autophagy inducers do not always provide benefit against viral infections. For example, trehalose-mediated autophagy was reported to impair the anti-viral response against human rhinovirus (HRV). HRV is a major cause of acute exacerbations of respiratory diseases, including asthma, chronic obstructive pulmonary diseases, and cystic fibrosis [89,90]. In the case of HRV infection, trehalose-induced autophagy led to the inhibition of anti-viral IFN-λ1 expression and promotion of HRV-16 replication in normal human tracheobronchial epithelial cells [91]. As observed in the case of trehalose, autophagy can function as both a pro-viral and an anti-viral depending on the virus and context [92]. Hence, activators of autophagy could be considered as therapeutic options against viral infections in a selective manner.
4.4. Resveratrol

The modulation of autophagy by resveratrol has been reported in various settings and studies have focused on its effects on immune function. A study involving a restrained mouse model with an induced splenic immunocompromised state examined the protective effect of resveratrol [93]. Restraint-stressed mice exhibited splenic damage, reduced CD4+ T-cell counts, and a decline in the number of splenocytes. Resveratrol administration reversed both the spleen index and overall splenocyte numbers induced by restraint and increased the number of CD4+ T cells. It was further confirmed that resveratrol induced autophagy in the mouse spleen. In another study, resveratrol was reported to suppress vascular endothelial inflammatory reactions [94]. Resveratrol prevented tumor necrosis factor α-induced inflammation by inducing autophagy in human umbilical vein endothelial cells. The study used siRNA knockdown to investigate the molecular mechanisms responsible and found that resveratrol activates the cAMP-PRKA-AMPK-SIRT1 signaling pathway to control autophagy.

4.5. Vitamin D3

The active form of vitamin D, 1α,25-dihydroxyvitamin D3 (1α,25-(OH)2D3), is a pleiotropic hormone that controls multiple functions in the body and has been found to be useful for the treatment of various diseases including rickets, osteoporosis, psoriasis, cancers, cardiovascular diseases, autoimmune diseases, and infectious diseases [95]. Tuberculosis is caused by the bacteria *Mycobacterium tuberculosis*, and is one of the top 10 causes of preventable death worldwide (the World Health Organization estimates that 1.8 million died from the disease in 2015). Progression of the disease is largely determined by the host immune response [96]. A previous study demonstrated that the induction of autophagy may contribute to 1α,25-(OH)2D3-mediated antimycobacterial immunity [97]. 1α,25-(OH)2D3 activates autophagy in monocytes and macrophages, and removal of *Mycobacterium tuberculosis* by 1α,25-(OH)2D3 in monocyte-derived macrophages and the THP-1 cell line was dependent on autophagy activation. Human cathelicidin was found to play a key role in 1α,25-(OH)2D3-induced autophagy by regulating the expression of autophagy-related genes such as Beclin-1 and Atg5. Human cathelicidin was also required for 1α,25-(OH)2D3-induced colocalization of mycobacterial phagosomes with autophagosomes in monocyte-derived macrophages. These findings suggest that vitamin D triggers autophagy to activate innate immunity against infection by intracellular *Mycobacterium tuberculosis*.

5. Activators of Autophagy for Cancer Therapy

5.1. Rapamycin and Rapalogs

Rapamycin is a naturally-occurring mTOR inhibitor, and its analogs (rapalogs) everolimus (RAD-001), temsirolimus (CCI-779), and deforolimus (AP-23573) inhibit mTORC1 to stimulate autophagy [46]. Everolimus was reported to suppress proliferation and enhance the anti-proliferative effect of paclitaxel when combined for the treatment of endometrial cancer cells. Co-treatment with the autophagy inhibitor chloroquine or transfection with shRNA targeting Atg5 reduced everolimus-mediated cell death, suggesting that everolimus induces autophagic cell death in endometrial cancer cells [98]. In another study, rapamycin was shown to sensitize A549 lung cancer cells to radiation therapy by inducing autophagy and delaying DNA damage repair [98]. Rapamycin activated autophagy, but not apoptosis in malignant glioma cells. Combination treatment with PI3K/Akt inhibitors synergistically enhanced rapamycin-sensitive and -resistant glioma cells to rapamycin by promoting autophagy [99]. Although it is well known that attenuating the mTOR pathway triggers autophagy, since mTOR controls various other cellular pathways including proliferation, protein synthesis, metabolism, and lipid synthesis [100], whether the induction of autophagy is solely responsible for any observed anti-cancer effects by rapalogs should be interpreted with caution. For example, everolimus-mediated autophagy was reported to contribute to a reduction
in the anti-proliferative effect caused by everolimus against aromatase inhibitor-resistant breast cancer cells. Additionally, simultaneous treatment of an autophagy inhibitor with everolimus improved the efficacy of everolimus treatment against breast cancer cell proliferation [101]. In the clinic, some partial responses have been seen with rapalog treatment in renal cell carcinoma and mantle-cell lymphoma [102,103]. However, in many cases, single treatment with rapalogs has shown only modest clinical outcomes in various cancer types [46,104–107]. Collectively, rapalogs represent potent inducers of autophagy via suppression of the mTOR pathway, however, their clinical application as autophagy activators requires further investigation.

5.2. Metformin

*Galega officinalis* (French lilac) contains high concentrations of guanidine, and has been used as a herbal treatment for the symptoms of diabetes in Europe for centuries [108]. Metformin was first synthesized as a derivative of guanidine. The biguanide metformin lowers glucose levels and improves insulin sensitivity, and is the most frequently prescribed antidiabetic drug in the world [109]. In addition to its usage as a medicine for diabetes mellitus, metformin has gained attention as a potential agent for cancer therapy [109]. Among the suggested mechanisms of action, metformin has been implicated to suppress several types of cancers at least in part by inducing autophagy. Metformin was reported to reduce the proliferation of melanoma cells and induce autophagy [110]. While short term treatment with metformin caused cell cycle arrest at G0/G1 phase, 96 h of metformin treatment induced apoptosis. Attenuation of autophagy through silencing *LC3* or *ATG5* led to a reduction in metformin-mediated anti-proliferation and apoptosis in melanoma cells. The administration of metformin inhibited melanoma tumor growth in mice and increased autophagy and apoptosis markers in tumor samples. Similar effects for metformin in an endometrial cancer cell line were observed [111]. Metformin induced cell cycle arrest and decreased cell viability in Ishikawa endometrial cancer cells, triggering autophagy. Knockdown of Beclin-1 or co-treatment with a pharmacological inhibitor of autophagy partially reversed the cytotoxic effects of metformin. In another study, metformin was reported to induce autophagic flux and enhanced TNF-related apoptosis-inducing ligand (TRAIL)-mediated apoptosis in lung adenocarcinoma cells [112]. Either treatment with an autophagy inhibitor or silencing of ATG5 partially reversed the sensitizing effect of metformin on TRAIL-induced tumor cell death. The primary molecular mechanism responsible for inducing autophagy is thought to be intimately associated with activation of the AMP-activated protein kinase (AMPK). Activated AMPK switches the cell from an anabolic to a catabolic state, and triggers autophagy by regulating mTORC1, ULK1, and p53 signaling pathways [113–115].

5.3. Obatoclax (GX15-070)

Obatoclax is a small-molecule indole bipyrrrole compound that targets Bcl-2 protein family members [116]. Obatoclax has been shown to exhibit anti-cancer activity against hematologic malignancies and overcomes glucocorticoid resistance in acute lymphoblastic leukemia [116–118]. The induction of autophagy was shown to be a primary mechanism of cell death elicited by obatoclax [118–121]. Obatoclax induces necroptosis by stimulating the assembly of necrosomes in autophagosomal membranes [119]. Preventing autophagosome formation by knockdown of Atg5 or Atg7 blocked obatoclax-mediated cell death. It was found that obatoclax triggers the interaction of Atg5 with components of the necroosome including Fas-associated death domain (FADD), receptor-interacting protein 1 (RIP1) and RIP3. Further examination showed that obatoclax-induced autophagic cell death through RIP1 and RIP3 [119]. Another study demonstrated that silencing of beclin-1 or treatment with an autophagy inhibitor blocked obatoclax-mediated cell death in B-cell lymphomas, further highlighting the importance of autophagy activation in the anti-cancer effects of obatoclax [122].
5.4. Liensinine, Isoliensinine, Dauricine and Cepharanthine

A number of natural alkaloid compounds have been discovered to induce autophagy in cancer cells. Liensinine, isoliensinine, dauricine, and cepharanthine promote autophagic activity by increasing autophagic flux and autophagosome formation [123]. The four alkaloids were shown to induce autophagy in various types of cancer and normal cells, showing a range of IC50 values against different cell types. Treatment with the alkaloids led to an increase in AMPK phosphorylation and subsequent downregulation of the mTOR pathway, suggesting that the compounds target the AMPK-mTOR pathway to promote autophagy. The study also used several cells with an apoptosis-resistant phenotype (e.g., caspase-3, -7, -8 deficient MEFs and Bax-Bak double knockout MEFs) to examine the cytotoxic effect of the alkaloids. Isoliensinine, dauricine, and cepharanthine induced cell death in apoptosis-defective or apoptosis-resistant MEFs.

5.5. Maprotiline and Fluoxetine

Antidepressants have been proposed to elicit various anti-cancer effects [124–127]. A previous study reported that the antidepressants maprotiline and fluoxetine inhibit the growth of Burkitt’s lymphoma cells by inducing autophagic-Type II programmed cell death [128]. The study found that the antidepressants promoted autophagic cell death in some Burkitt’s lymphoma cells, while causing apoptosis in another Burkitt’s lymphoma cell line, suggesting that the mechanism is cell-type specific. The autophagic cell death observed in the cell line tested required an increase in extracellular calcium influx. These results suggest that cancer cells that are apoptosis-resistant can be targeted by agents that induce autophagic cell death such as maprotiline and fluoxetine.

6. Conclusions

Autophagy is a homeostatic degradation process that captures, degrades, and recycles cellular proteins and organelles. Autophagy has dual functions in cancer cells, whereby its activation can promote survival by reducing the impact of various cellular and environmental stresses, and in other circumstances can cause cancer cell death. It can also protect against infections, control autoimmune diseases, and regulate inflammatory responses. Therefore, activators of autophagy may potentially contribute to the control of immunity and cancer (Table 1). However, with a plethora of complex pathways influenced by autophagy, the exact mechanisms of action and the outcomes involved must first be clearly elucidated. Future studies need to focus on dissecting the detailed circumstances of when activators and inhibitors of autophagy may elicit desirable therapeutic responses.

| Compound   | Mechanisms of Action                                                                 | Model (Related Disease)         | References |
|------------|-------------------------------------------------------------------------------------|---------------------------------|------------|
| Apigenin   | AMPK activation, mTOR inhibition, downregulation of TNF-α, IL-6 and IL-1β secretion | Human keratinocytes             | [129]      |
|            | Beclin-1 accumulation, conversion of LC3 protein, p62 degradation, enhanced ROS production | BCPAP cell (human papillary thyroid carcinoma) | [131]      |
|            | mTOR inhibition                                                                     | Human keratinocytes (UV-induced skin cancer) | [132]      |
| Arsenic trioxide | BNIP3 upregulation                                                                | U373-MG, U87-MG, and T98G cell (malignant glioma) | [133]      |
| BRD5631    | IL-1β suppression, mTOR-independent autophagy activation                            | HeLa cell expressing LC3, hiPSC-derived neurons | [77]       |
| Cepharanthine | AMPK activation                                                                   | MEFs and cancer cells            | [123]      |
| Compound         | Mechanisms of Action                        | Model (Related Disease)                                                                 | References |
|------------------|----------------------------------------------|----------------------------------------------------------------------------------------|------------|
| Curcumin         | Akt/mTOR inhibition                          | U87-MG and U373-MG (malignant gliomas)                                                 | [134]      |
|                  | Stimulation of LC3-II, down-regulation of p62| Primary human umbilical vein endothelial cells, rat model of carotid artery intimal injury | [135]      |
|                  | Upregulation of the JNK signaling pathway    | MG63 cell (human osteosarcoma)                                                         | [136]      |
|                  | Increased Atg8                                | Sf9 (Spodoptera frugiperda) insect cell (pest control)                                  | [137]      |
|                  | Suppression of AKT/mTOR/p70S6K signaling      | A375 and C8161 cells, A375-cell transplanted mice (human melanoma)                      | [138]      |
|                  | Iron deprivation-dependent autophagy induction| DU145 and PC3 cells (castration-resistant-human prostate cancer)                        | [139]      |
| Dauricine        | AMPK activation                               | MEFs and cancer cells                                                                   | [123]      |
| Delphinidin      | Atg5-dependent accumulation of LC3-II        | Atg5-deficient and normal mouse embryonic fibroblasts                                   | [140]      |
| Fisetin          | LC3-II accumulation (synergistic effect with paclitaxel) | A549 cells (lung cancer)                                                               | [141]      |
|                  | mTOR inhibition                               | PC3 cells (human prostate cancer), TU212 (laryngeal carcinoma)                           | [142,143]  |
| Galangin         | Inhibition of PI3K/Akt/mTOR signaling         | TU212 and M4e cells (human laryngeal cancer)                                            | [144]      |
| Genistein        | Activation of LKB1/AMPK pathway, inhibition of mTOR signaling | Human vascular smooth muscle cells (atherosclerosis)                                   | [145]      |
| Gossypol         | Accumulation of Beclin 1                      | Human pharynx, tongue, and salivary gland cancer cell lines (Head and neck carcinoma), SALTO cells and Balb/c mice transplanted SALTO cells (mouse salivary gland cancer) | [146]      |
| Grandifloracn    | Akt/mTOR inhibition                           | PANC-1 cell (human pancreatic cancer)                                                  | [147]      |
| Guttiereon K     | Akt/mTOR inhibition                           | HeLa cell (human cervical cancer)                                                      | [148]      |
| Isoliensinine    | AMPK activation                               | MEFs and cancer cells                                                                   | [123]      |
| Kaempferol       | Accumulation of LC3-II, ROS-mediated JNK activation | SH-SY5Y and primary neurons cells (Parkinson’s disease), HeLa (human cervical cancer) | [149,150]  |
|                  | AMPK activation, mTOR inhibition              | SK-HF1 cell (human hepatic cancer), palmitic acid-stressed RIN-3F cells and murine pancreatic islets (type 2 diabetes) | [151,152]  |
| Liensiniane      | AMPK activation                               | MEFs and cancer cells                                                                   | [123]      |
| Magnolol         | PI3K/PTEN/Akt inhibition                      | H460 cells (human lung cancer)                                                         | [153]      |
| Maprotiline and Fuoxetine | Calcium influx-mediated autophagy       | DG-75 cells (human Burkitt’s lymphoma)                                                 | [124,128]  |
| Metformin        | Downregulation of Src/CEBPD pathway, AMPK activation | Huh-7 cells (hepatocellular carcinoma)                                                 | [154]      |
|                  | AMPK activation, inhibition of pro-inflammatory pathway (NLRP3, TNF-α, IL-6) | SH-SY5Y cells, probenecid-induced mice model (Parkinson’s disease)                     | [155]      |
|                  | Downregulation of c-FLIP and decrease in p62  | A549 cell (human lung adenocarcinoma)                                                   | [112]      |
Table 1. Cont.

| **Compound** | **Mechanisms of Action** | **Model (Related Disease)** | **References** |
|--------------|--------------------------|----------------------------|---------------|
| Obatoclax    | Activation of Bax and Bak| Phase I clinical study (lymphocytic leukemia), rituximab/chemotherapy-sensitive, -resistant cell lines, and primary tumor-cells derived from patients (human lymphoma) | [116,122] |
| Oridonin     | GLUT1-mediated and AMPK-independent autophagy induction | HCT-15, COLO205, HCT116, RKO, SW480, and SW620 cell line (Colorectal cancer) | [156] |
| Pentagalloylglucose | mTOR inhibition | Atg7−/− MEF and knock out mice (HSV-1 infection), PC3 and TRAMP-C2 (prostate cancer) | [157,158] |
| Pterostilbene | JNK1/2 activation, inhibition of Akt, ERK1/2, and p38 | accumulation of LC3-positive vacuolar structures | [159] |
| Quercetin    | H-RAS degradation | A549 cell (lung cancer) | [161] |
|             | Reduced p62 protein expression | Rothenone-induced rat model (Parkinson’s disease) | [162] |
|             | Ameliorated ER stress and oxidative stress | | [163] |
| Quercetin    | Inhibition of PI3K/Akt/mTOR and STAT3 signaling | | [164] |
| Rapamycin    | Increased LC3-II/LC3-I ratio | MPC5 and MSC1097 mouses podocytes MPC5 and mesangial cells MSC1097 (Nephropathy), acute spinal cord injured rat model (neurodegenerative disease) | [166,167] |
|             | mTOR inhibition, increased LC3-II, Atg5, and Atg7, and p62 reduction | Cecal ligation and puncture-induced mice model (septic encephalopathy) | [168] |
| Resveratrol  | cAMP/PRKA/AMPK/SIRT1 activation | Human umbilical vein endothelial cell (endothelial inflammation, atherosclerosis) | [94] |
| Resveratrol  | ATG4 restoration | SH-SY5Y cells hyper-expressing the mutant polyQ Huntingtin protein (Huntington’s Disease) | [171] |
|             | Suppression of microRNA-383-5p | Human podocytes, db/db mice (diabetic nephropathy) | [172] |
| Resveratrol  | Inhibition of mTOR/ULK1 signaling | MCF7 cell (human breast cancer), PC3 and DU145 cells (human prostate cancer) | [173,174] |
| Resveratrol  | STAT3 inhibition | CAOV-3 and OVCAR-3 cells (human ovarian cancer) | [175] |
| Resveratrol  | AMPK activation, mTOR inhibition | Cisplatin-resistant human oral cancer cells, destabilization of the medial meniscus in mice | [176,177] |
Table 1. Cont.

| Compound        | Mechanisms of Action                                      | Model (Related Disease)                              | References |
|-----------------|-----------------------------------------------------------|------------------------------------------------------|------------|
| Rottlerin       | Inhibition of PI3K/Akt/mTOR signaling                     | Cancer stem cells                                    | [178–180] |
| Saikosaponin-d  | Inhibition of SERCA, leading to increase in Ca2+ and activation of CaMKKβ | HeLa and MCF-7 (human cervical and breast cancer)    | [181]      |
| Salvianolic acid B | Akt/mTOR inhibition                                    | HCT116 and HT29 cell (human colorectal cancer)       | [182]      |
| Salvigenin      | Increased LC3-II/LC3-I, Atg7, and Atg12 in the presence of H2O2 | SH-SY5Y cell (human neuroblastoma, neurodegenerative disorders) | [183]      |
| Spermidine      | Increased titin phosphorylation, suppression of inflammation | Cardiomyocytes, mice, and Dahl salt-sensitive rats fed a High-salt diet (cardiovascular disease) | [80]       |
|                 | Modulation of SIRT1-independent deacetylation pathway     | HCT 116 cells, C. elegans, and cytoplasts (longevity) | [81]       |
|                 | Inhibition of the caspase 3/Beclin 1 cleavage             | PC12 cells and cortical neurons (neurodegenerative disease) | [184]      |
|                 | Increased cholesterol efflux                             | High-fat fed ApoE−/− mice (atherosclerosis)          | [185]      |
| Trehalose        | mTOR-independent autophagy induction                     | Human foreskin fibroblasts, human aortic endothelial cells (human cytomegalovirus, atherosclerosis) | [88]       |
|                 | Impaired IFN-λ1                                          | Human primary airway epithelial cells (human rhinovirus, asthma) | [91]       |
| Vitamin D3      | mTOR-independent autophagy induction, reduction of mutant huntingtin aggregates | COS-7, SK-N-SH, HeLa, and HeLa cells expressing EGFP-LC3, Atg5-deficient and wild type mouse embryonic fibroblasts, PC12 (huntington’s disease and Parkinson’s disease) | [83]       |
|                 | Increased LC3, and Beclin 1                              | ApoE−/− and wild-type mice (atherosclerosis)         | [186]      |
|                 | Human cathelicidin-dependent activation of Beclin-1 and Atg5 | Human monocytes and macrophages (tuberculosis)       | [97]       |
|                 | Increased LC3, and Beclin 1                              | MIN6 mouse insulinoma β-cell and streptozotocin-induced mice model (type 1 diabetes) | [187]      |
|                 | Up-regulation of LC3-II/LC3-I and PR-39 mRNA expression   | Porcine small intestinal epithelial cell, pig model (rotavirus infection) | [188]      |
| Vorinostat      | HDAC inhibition                                         | U937 and SUDHL6 cell (hematological cancer)          | [189]      |

BNIP3, Bcl-2/adenovirus E1B 19-kDa-interacting protein 3; JNK, c-Jun N-terminal kinase; CEBPD, Transcription factor CCAAT/enhancer-binding protein delta; HSV-1, Herpes simplex virus type 1; LC, microtubule-associated protein 1A/1B-light chain; SERCA, sarcoplasmic/endoplasmic reticulum Ca2+ ATPase; CaMKKβ, Ca2+/calmodulin-dependent kinase kinase-β; HCMV, Human cytomegalovirus; TFEB, transcription factor EB; HDAC, Histone deacetylase.

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