The ménage à trois of autophagy, lipid droplets and liver disease

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ABSTRACT

Autophagic pathways cross with lipid homeostasis and thus provide energy and essential building blocks that are indispensable for liver functions. Energy deficiencies are compensated by breaking down lipid droplets (LDs), intracellular organelles that store neutral lipids, in part by a selective type of autophagy, referred to as lipophagy. The process of lipophagy does not appear to be properly regulated in fatty liver diseases (FLDs), an important risk factor for the development of hepatocellular carcinomas (HCC). Here we provide an overview on our current knowledge of the biogenesis and functions of LDs, and the mechanisms underlying their lysosomal turnover by autophagy processes. This review also focuses on nonalcoholic steatohepatitis (NASH), a specific type of FLD characterized by steatosis, chronic inflammation and cell death. Particular attention is paid to the role of macroautophagy and macrophagolipophagy in relation to the parenchymal and non-parenchymal cells of the liver in NASH, as this disease has been associated with inappropriate lipophagy in various cell types of the liver.

Abbreviations: ACAT: acetyl-CoA acetyltransferase; ACAC/ACC: acetyl-CoA carboxylase; AKT: AKT serine/threonine kinase; ATG: autophagy related; AUP1: AUP1 lipid droplet regulating VLDL assembly factor; BECN1/Vps30/Tatg6: beclin 1; BCSL2/seipin: BCSL2 lipid droplet biogenesis associated, seipin; CMA: chaperone-mediated autophagy; CREB1/CREB: cAMP responsive element binding protein 1; CXXC3: C-X-C motif chemokine receptor 3; DAGs: diacylglycerol; DAPMs: danger/damage-associated molecular patterns; DEN: diethylnitrosamine; DGAT: diacylglycerol O-acyltransferase; DNL: de novo lipogenesis; E2BP1/NACSN (EH domain binding protein 1); EHD2/PAST2: EH domain containing 2; CoA: coenzyme A; CCL/chemokines: chemokine ligands; CCl4: carbon tetrachloride; ER: endoplasmic reticulum; ESCRT: endosomal sorting complexes required for transport; FAT: fatty acids; FFAs: free fatty acids; FFC: high saturated fats, fructose and cholesterol; FG21: fibroblast growth factor 21; FITM/FIT: fat storage inducing transmembrane protein; FLD: fatty liver diseases; FOXO: forkhead box O; GABARAP: GABA type A receptor-associated protein; GPAT: glycerol-3-phosphate acyltransferase; HFD: high fat diet; HSC: hepatic stellate cells; HSP/HS70: heat shock protein family A (Hsp70) member; ITCH/AIP4: itchy E ubiquitin protein ligase; Kcs: Kupffer cells; LAMP2A: lysosomal associated membrane protein 2A; LD: lipid droplets; LDL: low density lipoprotein; LEP: leptin; LEP/R/ORG: leptin receptor; LIPA/LAL: lipase A, lysosomal acid type; LPE/E/HSL: lipase E, hormone sensitive type; LR: LC3-interacting region; LPS: lipopolysaccharide; LSECs: liver sinusoidal endothelial cells; MAGS: monacylglycerol; MAPK: mitogen-activated protein kinase; MAP3K5/ASK1: mitogen-activated protein kinase kinase kinase 5; MAP1LC3/LC3: microtubule associated protein 1 light chain 3; MCD: methionine-choline deficient; MGLL/MGL: monoglyceride lipase; MLXIPL/ChREBP: MLX interacting protein like; MTOCRC1: mechanistic target of rapamycin kinase complex 1; NAFLD: nonalcoholic fatty liver disease; NAS: NAFLD activity score; NASH: nonalcoholic steatohepatitis; NPC: NPC intracellular cholesterol transporter; NR1H3/LXRx: nuclear receptor subfamily 1 group H member 3; NR1H4/FXR: nuclear receptor subfamily 1 group H member 4; PDGF: platelet derived growth factor; PIK3C3/VPS34: phosphati-didylinositol 3-kinase catalytic subunit type 3; PLIN: perilipin; PNPLA: patatin like phospholipase domain containing; PNPLA2/ATGL: patatin like phospholipase domain containing 2; PNPLA3/adiponutrin: patatin like phospholipase domain containing 3; PPAR: peroxisome proliferator activated receptor; PPARA/PPARA: peroxisome proliferator activated receptor alpha; PPARD/PPARD: peroxisome proliferator activated receptor delta; PPARG/PPARY: peroxisome proliferator activated receptor gamma; PPARC1A/PPCG1a: PPAR coactivator 1 alpha; PKAA/AMPK: protein kinase AMP-activated catalytic subunit; Ptlin3Kc: class III phosphati-didylinositol 3-kinase; Prkcs8; phosphoryldidylinositol 3-phosphate; PTEN: phosphatase and tensin homolog; ROS: reactive oxygen species; SE: sterol esters; SIRT1: sirtuin 1; SPART/SPG20: spartin; SOSTM1/p62.
An overview of general autophagic mechanisms

The term autophagy, which is derived from the Greek words αυτός (“auto”) and φαγεῖν (“phagein”), meaning self-eating, includes a number of lysosomal degradative and recycling pathways that control both quality and quantity of proteins, lipids and organelles in eukaryotic cells. Upon starvation, autophagic pathways are triggered to compensate for the lack of nutrients and to balance energy deficiencies. In addition, numerous other types of cellular stress can induce autophagic pathways, including the accumulation of cytotoxic protein aggregates that are selectively degraded through autophagy. Likewise, cellular organelles, including LDs, can be degraded in lysosomes through autophagic processes to generate metabolites. Autophagy is therefore important in maintaining cellular homeostasis in both cells and tissues and plays a crucial role during development, adaptation to stress, and immune and metabolic responses. The three different autophagic pathways macroautophagy/autophagy, microautophagy and chaperone-mediated autophagy (CMA) are discussed in the following subchapters.

Macroautophagy

Macroautophagy is a highly conserved lysosomal degradation mechanism [1]. Unlike microautophagy or CMA, macroautophagy requires the formation of double-membrane vesicles called autophagosomes, which sequester intracellular components such as proteins, macromolecular complexes and organelles, but also invading pathogens [2]. Macroautophagy takes place in almost all eukaryotic cells at basal levels. However, it is most often a response to cellular stress induced by a wide variety of physiological stimuli, including nutritional deficiencies, growth factor withdrawal, oxidative and mechanical stress, as well as pathophysiological situations, including drug treatment and radiation therapy.

Macroautophagy begins with the formation of a precursor structure called the phagophore [3], which then elongates and closes to generate an autophagosome. The latter then matures and finally fuses with lysosomes to expose the engulfed cytosolic material to acidic lysosomal hydrolases for degradation. To different extents, various cellular organelles, including the endoplasmic reticulum (ER), Golgi complex, mitochondria, recycling endosomes, but also the plasma membrane, have been shown to supply the lipids required for the assembly of autophagosomes [4], a process executed by the autophagy related (ATG) proteins [5]. ATG genes were first discovered and characterized by Yoshinori Ohsumi, Daniel Klionsky and Michael Thumm [5–7]. During the process of autophagy, induction of autophagosome formation requires the activation of the ULK1/Atg1 (unc-51 like autophagy activating kinase 1) kinase complex [8], which integrates upstream molecular signals principally via two key kinases, the mTOR (mechanistic target of rapamycin kinase) complex 1 (MTORC1) and the PRKA/AMPK (protein kinase AMP-activated catalytic subunit) [9–11]. ULK1 kinase complex activation is essential to initiate the de novo formation of the phagophore, which takes place in specialized subdomains of the ER called omegasomes [12]. The latter are considered as the “cradle” for the forming nascent autophagosome [13]. Specifically, the ULK1 complex phosphorylates and activates the class III phosphatidylinositol 3-kinase (PtdIns3K) complex, composed of BECN1/Vps30/Atg6 (beclin 1), PIK3R4/VPS15 (phosphoinositide-3-kinase regulatory subunit 4), ATG14, PIK3C3/VPS34 (phosphatidylinositol 3-kinase catalytic subunit type 3), AMBRA1 (autophagy and beclin 1 regulator 1) and NRBF2 (nuclear receptor binding factor 2) [14], which generates phosphatidylinositol-3-phosphate (PtdIns3P). In turn, the generation of PtdIns3P mediates the association of specific PtdIns3P-binding proteins, such as ZFYVE1/DFCP1 (zinc finger FYVE-type containing 1) [12] and WIPI (WD repeat domain, phosphoinositide interacting) proteins [15]. The four WIPI proteins, WIPI1, WIPI2, WDR45B/WIPI3 (WD repeat domain 45B) and WDR45/WIPI4 (WD repeat domain 45) are the mammalian members of the β-propellers that bind PROPPIN (β-propellers that bind phosphoinositides) and function as conserved PtdIns3P effectors at the nascent autophagosome [16,17]. Thereby, WIPI2, assisted by WIPI1 [16], recruits ATG16L1 [18], which assemble with the ATG12–ATG5 conjugate into a multimeric complex [18] and initiate phagophore formation. In contrast, both WDR45B/WIPI3 and WDR45/WIPI4 control phagophore elongation, whereby WDR45 specifically associates with ATG2 proteins [16,19]. The ATG12–ATG5–ATG16L1 complex acts as an E3-like ligation for the conjugation of the members of the MAP1LC/LC3 (microtubule associated protein 1 light chain 3) protein family to phosphatidylinositolamine [5]. The LC3 protein family comprises seven members, LC3A, LC3B, LC3B2 and LC3C, which are part of the MAP1LC3 subgroup, and GABARAP (GABA type A receptor-associated protein), GABARAPL1 (GABA type A receptor associated protein like 1) and GABARAPL2/GATE-16 (GABA type A receptor associated protein like 2), which belong to the GABARAP subgroup. LC3 proteins are important for autophagic cargo selection, phagophore elongation and closure. Complete autophagosomes finally contact lysosomes and fuse with them to form autolysosomes [20].

Macroautophagy can be either selective or nonselective. Nonselective macroautophagy involves random sequestration of cytoplasmic material by autophagosomes, whereas selective macroautophagy leads to the lysosomal degradation of specific cargoes and such pathways are named according to the
type of cargo that is targeted, including aggregated proteins (aggrephagy), mitochondria (mitophagy), peroxisomes (pexophagy) and LDs (lipophagy) [21,22]. Selective types of macroautophagy rely on the so-called autophagy receptors that physically link the cargo to the phagophore [21,22] by Ub (ubiquitin)-dependent or Ub-independent cargo recognition [22]. The vast majority of macroautophagy receptors physically interact with the phagophore by selective binding to LC3 proteins present on the inner membrane of the phagophore. Thereby, short linear peptide sequences known as LC3-interacting regions (LIRs) in mammalian cells or Atg8-interacting motifs in yeast, mediate the effective tethering of the cargo to the forming vesicle, guaranteeing selective sequestration [21,22]. Ub-dependent macroautophagy receptors like SQSTM1/p62 (sequestosome 1), possess a Ub-binding domain that allows them to recognize and be recruited onto ubiquitylated cargoes [21,22]. Ub-independent macroautophagy receptors, in contrast, are often membrane-associated and therefore present in the cargo that potentially can be targeted by selective autophagy [21,22]. While engagement of the Ub-dependent macroautophagy receptors is dictated by the ubiquitination of the cargo destined to degradation, those that are Ub-independent require additional regulation, which is often achieved through their phosphorylation [21–23].

**Microautophagy**

Microautophagy is a conserved process in which cytosolic components are directly engulfed and degraded by the vacuole in yeast or by endo-lysosomal compartments in mammals. Microautophagy cargoes include proteins and organelles, such as peroxisomes, mitochondria and portions of nucleus [24–26]. In mammalian cells two different mechanisms of lysosomal cargo sequestration have been reported, one involving protrusions of the lysosome limiting membrane and the other involving lysosomal invagination. However, given the fact that technical monitoring of microautophagy still presents a challenge, current investigations on microautophagy remain a difficult task [27]. Nevertheless, selective microautophagy-like uptake of protein cargo into late endosomes, referred to as endosomal microautophagy, has been described in mammalian cells [28,29], whereby cytosolic protein cargo are invaginated by late endosomes [27]. Such proteins harbor a KFERQ-like motif, which is recognized by the HSPA8/HSC70 [heat shock protein family A (Hsp70) member 8], to promote the translocation of the captured proteins into the lumen of a late endosome, or multivesicular body, in an endosomal sorting complex required for transport (ESCRT) III-dependent manner [30]. Subsequent fusion of the late endosomes/multivesicular body then delivers the translocated protein cargo into a lysosome for turnover [27].

**Chaperone-mediated autophagy**

CMA is a selective type of autophagy principally dedicated to the lysosomal degradation of cytosolic proteins [31]. The targeted proteins possess a KFERQ motif in their amino acid sequence [32], but unlike in endosomal microautophagy, are directly captured by lysosomes [27]. The KFERQ motif is recognized by the cytosolic chaperone HSPA8 and its co-chaperones, including STUB1/CHIP (STI1P homology and U-box containing protein 1), DNAJ/B1/HSP40 [DnaJ] heat shock protein family (Hsp40) member B1 and STI1P/HOP (stress induced phosphoprotein 1) [31], leading to the assembly of a cargo-chaperone complex [33]. This complex then bonds the cytosolic tail of the LAMP2A (lysosomal associated membrane protein 2A), a protein only expressed in mammals [34], which mediates the unfolding and translocation of the targeted substrate through the lysosomal membrane [33]. During this transport, substrates are translocated one-by-one into the lysosomal lumen [35], via a process involving the multimerization of LAMP2A into a translocation-competent complex and the lysosomal resident pool of HSPA8 [36]. The substrates are then rapidly degraded by lysosomal proteases and LAMP2A disassembles to become available for a new cycle of substrate binding and translocation [35].

**Lipid droplets**

LDs are conserved cellular organelles (Figure 1) that store neutral lipids [37] and fulfill critical functions in lipid and energy homeostasis [38]. In recent years, it has become apparent that LDs have a complex functional connection with autophagy [39], and that cellular energy levels can be elevated by employing free fatty acids (FFAs) gained from the selective breakdown of LDs through macroautophagy, referred to as macrolipophagy [40]. Macrolipophagy is a relatively new area in the field of autophagy, and will certainly provide us with exciting new information about its mechanism and regulation, but also its physiological relevance in the next few years. In fact, research on macrolipophagy has already begun to expand our understanding of the physiological relevance of autophagy. Macrolipophagy plays an important role in regulating lipid metabolism, and as part of lipolysis programs, can provide energy and the building blocks for membrane biosynthesis and lipid anabolism. In this latter context, macrolipophagy has an important role in the biosynthesis of the cholesterol-based lipidic hormone testosterone [41]. More recently, macrolipophagy-derived FFAs have been shown to undergo extracellular efflux via lysosomal exocytosis through a mechanism involving MCOLN1 (mucolipin TRP cation channel 1), a lysosomal Ca2+ channel that regulates lysosome-plasma membrane fusion [42]. These secreted lipids have been suggested to be important for cell-to-cell lipid exchange and signaling, but further studies are needed to better understand the significance of FFA efflux in tissue homeostasis and disease development [42]. Additionally, macrolipophagy is important for cell differentiation. For example, macrolipophagy-mediated lipolysis and subsequent generation of FFAs enables the shift from glycolysis to FFA β-oxidation, which generates the energy needed for neutrophil differentiation [43]. Finally, a study in the worm Caenorhabditis elegans showed that lipid signaling molecules, derived from lysosomal lipid hydrolysis, may modulate gene expression to extend lifespan [44].
In the following, we discuss the current knowledge about the structure, biogenesis, function and catabolism of LDs, before extending to macrolipophagy and other autophagic pathways used to degrade LDs or LD-associated proteins.

**The structure of lipid droplets**

LDs are spherical organelles with a hydrophobic core composed by neutral lipids such as triacylglycerols (TAGs), diacylglycerols (DAGs) and sterol esters (SE). This core is coated with a monolayer of phospholipids, free cholesterol and lysophospholipids, into which specific proteins are inserted (Figure 1). The limiting membrane of LDs has the major role in maintaining the morphology and stability of LDs and also facilitates the interactions with other organelles [45]. Multiple proteins constantly interact with the surface of LDs, including members of the PLIN (perilipin) protein family, which are among the best characterized LD-associated proteins [46]. This protein family has five members: PLIN1 to PLIN5 [46]. While PLIN1 and PLIN4 are mainly expressed in white adipose tissue, PLIN2/adipophilin, and PLIN3/TIP47, are ubiquitously expressed. The major expression tissues of PLIN5, in contrast, are the cardiac and skeletal muscles, the brown adipose tissue and the liver [46–48]. PLIN proteins have an important role in regulating LD structure, function and catabolism through lipolysis and CMA [49]. In addition to the PLIN protein family, other proteins such as lipases are associated to LDs [45]. Among these, is the PNPLA2/ATGL (patatin like phospholipase domain containing 2), the LIPE/HSL (lipase E, hormone sensitive type) and the MGLL/MGL (monocacylglycerol lipase), which together mediate lipolysis (see below). PNPLA2 is expressed in all tissues, with highest levels in adipose tissues. LIPE expression has also been found to be higher in adipose tissues, while MGLL is ubiquitously expressed [50]. Other lipases, including PNPLA3/adiponutrin (patatin like phospholipase domain containing 3) or PNPLA5 (patatin like phospholipase domain containing 5), which share significant sequence identity with PNPLA2, are also present on LDs and catalyze the hydrolysis of TAGs into DAGs [51]. ATG proteins have also been found on the surface of LDs. In particular, ATG2A, ATG14 and lipidated LC3 decorate LDs and are involved in the regulation of LD volume and/or distribution [52–54]. Interestingly, mutations in several LD-associated proteins that lead to a defect in expression and/or enzymatic activity, have been associated with human diseases [38].

**The biogenesis of lipid droplets**

LDs arise from the ER, where several neutral lipid biosynthetic enzymes are localized. Among these are the GPAT (glycerol-3-phosphate acyltransferase) enzymes, which catalyze the first step of the glycerol-3-phosphate pathway, the main cellular process that leads to the synthesis of TAGs in most cell types [55]. GPATs acylate glycerol-3-phosphate and acyl-coenzyme A (CoA) to generate lysophosphatidic acids, which are converted into phosphatidic acids by the 1-acyl glycerol-3-phosphate acyltransferase. Subsequently, LPIN (lipin) enzymes dephosphorylate the phosphatidic acids to generate DAGs, which are finally transformed into TAGs by diacylglycerol acyltransferases (DGATs) [56]. In the small intestine, liver and adipose tissue, TAGs can also be synthesized through the monoacylglycerol (MAG) pathway [57]. This pathway produces DAGs from MAGs and shares the final step with the glycerol-3-phosphate pathway. Other ER resident enzymes such as ACAT1 (acyetyl-CoA acyltransferase 1) and ACAT2, produce SE from sterols [38]. The process of de novo LD biosynthesis is divided in three major steps: (i) nucleation, (ii) growth and (iii) budding (Figure 1) [58]. Although the exact molecular mechanism of these different steps is still not yet fully understood, recent research has shed light on this important cellular process, and a consensus for the biogenesis of LDs is emerging. The nucleation step begins with the formation of an oil lens structure between the two leaflets of the ER membrane, which originates from the accumulation of the newly synthesized TAGs and SE [59,60]. The oil lens formation appears to be a lipid driven process, which does not rely on specific proteins as previously suggested [61]. It has been shown that the presence of non-bilayer lipids such as phosphatidic acids and DAGs in the ER bilayer increase the bilayer tension, impeding LD formation [61]. As a result, transformation of these lipids into TAGs has the opposite effect, i.e., inducing LD biogenesis. Nevertheless, two ER-resident protein families, the FITM/FIT (fat storage-inducing transmembrane protein) and the BSCl2/seipin (BSCl2 lipid droplet biogenesis associated, seipin) protein family, play an indirect role in the emergence of oil lens [62]. On the one hand, FITM proteins, in particular FITM2, are thought to be acyl-coenzyme A diphosphatases [63]. Their activity could promote the decrease of DAG level in the proximity of LDs’ budding site, at the cytosolic leaflet of the ER membrane, to help the unidirectional LD emergence [64]. On the other hand, BSCl2/seipins may regulate the local synthesis and distribution of phospholipids such as phosphatidic acid in the ER, and they interact and negatively control GPAT activity [65–68]. Normal levels and distribution of phosphatidic acid in the ER allow LDs to form and grow properly. Additional proteins that can modulate membrane rearrangements, including the ER membrane shaping proteins such as RTN (reticulon) and ATL/atlastin (atlastin GTPase) proteins or LD coat proteins such as PLIN3, play a role in biogenesis of LDs [38]. However, the exact mechanism by which those proteins promote the formation of LDs remains to be determined. The growth step is characterized by the phenomenon of ripening, a slow process that is promoted by the diffusion of molecules from the volume of smaller LDs toward larger LDs, and fusion [58]. This event is followed by the budding step, which is usually completed in mammalian cells, while in yeast, LDs remain associated with the ER (Figure 1) [59,69]. After budding LDs can still expand, either through LD-LD fusion, a process known as coalescence, transfer of TAGs from the ER to LDs via membrane contact sites, or local TAGs synthesis at the LD surface [70,71].
The functions of lipid droplets

An important function of LDs is to store non-esterified FFAs as inert TAGs, since an excess of FFAs in the cytoplasm would trigger the formation of harmful bioactive lipids, causing lipotoxicity [72]. Other main functions of LDs are to provide building blocks for the biosynthesis of cellular membranes and other lipid species, as well as for metabolic energy through FFA β-oxidation when nutrients are scarce [73]. LDs also play roles in the assembly of viruses such as hepatitis C virus, protein sequestration, and membrane trafficking and signaling [74].

One of the most interesting research areas on LDs is the characterization of membrane contact sites of LDs with other organelles, which are important for the fine-tuning of the regulation of LD functions. In addition to having membrane contact sites with ER, LDs physically communicate with peroxisomes to facilitate the transfer and the β-oxidation of very long chains and branched FFAs [75,76]. In addition, LDs often interact with mitochondria in a PLIN5-dependent manner in response to nutrient deficiencies in cultured cells, and to exercise in the skeletal muscles of animals [77,78]. These LD-mitochondria contact sites might help the transfer of FFAs for β-oxidation [73], although direct evidence for this event is lacking. Finally, LDs have also been functionally linked to macroautophagy. In this context, it has been shown that neutral lipid stores are mobilized from LDs to support the formation of the autophagosomes in both yeast and mammalian cells [51,79,80]. In summary, all these findings support our view on LDs, which are now considered to be highly dynamic organelles rather than inert fat bodies.

Catabolism of lipid droplets: neutral and acid lipolysis

The term lipolysis refers to the hydrolysis of the ester bonds in TAGs, to generate FFAs and glycerol. Cells are unable to uptake exogenous TAGs as they can only assimilate FFAs. Thus, lipolysis is essential to process dietary TAGs and lipoprotein-associated TAGs for the subsequent cellular uptake of FFAs from intestine and blood, respectively. In addition, lipolysis is required for the metabolism of TAGs endocytosed in association with lipoproteins from the blood, known as intracellular lipolysis. Intracellular lipolysis is also critical to catabolize TAGs stored in LDs in both adipose and non-adipose tissues. In contrast to adipose tissues, which secrete lipolysis products into the bloodstream, non-adipose tissues mobilize lipolysis products when cells require FFAs and glycerol to either generate energy or build new macromolecules [72,81].

Intracellular lipolysis can be divided into neutral and acid lipolysis, depending on the pH value and subcellular location where this process takes place. Neutral lipolysis occurs in the

Figure 1. The biogenesis of LDs. The process of de novo LD biogenesis can be divided in three main discrete steps: (A) Nucleation, (B) growth and (C) budding. The nucleation step is characterized by the formation of an oil lens structure in between the two lipid bilayers of the ER limiting membrane, which is catalyzed by ER resident proteins such as FITM1/FITM2 and BSCL2/seipin. The growth of the nascent LDs starts with an accumulation of TAGs and SE, and involves a ripening phenomenon that regulates their size. The partial (in yeast) or complete (in mammalian cells) detachment of the LDs from the ER membrane defines the LDs budding, LDs can also expand (D) by increasing their size through the coalescence of LDs and/or the local synthesis of TAGs. Several enzymes involved in the biogenesis and function of LDs, which are discussed in the review, are indicated.
cytoplasm, at neutral pH, and it is mediated by a series of cytosolic lipases that directly act on the TAGs and cholesteryl esters stored in LDs. Neutral lipolysis of TAGs requires the consecutive action of the three neutral lipases (i) PNPLA2, which preferentially catalyzes the hydrolysis of TAGs into DAGs, (ii) LIPE, which mainly mediates the processing of DAGs into MAGs, as well as the hydrolysis of the ester bonds of other lipids such as cholesteryl esters, and (iii) MGLL, which catalyzes the hydrolysis of MAGs into glycerol [72]. In contrast, acid lipolysis takes place at acidic pH inside lysosomes, and is mediated by lipases such as the LIPA/LAL (lipase A, lysosomal acid), which are believed to hydrolyze TAGs and cholesteryl esters, while acid phospholipases and proteases break down LD-associated phospholipids and proteins, respectively. Acid lipolysis hydrolyzes lipids delivered into lysosomes by both receptor-mediated endocytosis of lipoproteins and lipophagy [72,82]. In this context, the finding that LDs could be delivered into lysosomes for turnover by macroautophagy in 2009, was a ground-breaking finding in the field of both lipolysis and autophagy research [83], and added a new level of complexity to our view on the regulation of LD catabolism. In extension, the direct engulfment of LDs by endosomes or lysosomes/vacuoles, known as microlipophagy [84], as well as the degradation of LD-associated proteins by CMA, have been reported more recently [49].

Macrolipophagy

Similar to bulk nonselective autophagy, lipophagy can occur via both macro (Figure 2A) and microautophagic mechanisms (Figure 2B) depending on how LDs are transported into lysosomes/vacuoles [83,84]. Macrolipophagy involves the autophagosome-mediated sequestration of LDs and their subsequent delivery to lysosomes/vacuole for turnover [83], which were originally reported to occur in mouse hepatocytes that have been supplemented with oleate and then subjected to nutrient starvation [83]. Inhibition or loss of autophagy caused an increase in TAG and LD levels in vitro and in vivo, a decrease in TAG breakdown and a colocalization between ATG components and TAG or LD proteins [83]. Macrolipophagy has subsequently been described in many other mammalian cell types [41,85–93] and it therefore appears that this process contributes ubiquitously to the mobilization of lipids stored in LDs in higher eukaryotes [81].

As with macroautophagy, macrolipophagy requires ATG proteins, in particular those composing the highly conserved core machinery [83]. In mammals, sequestration of LDs appears to be through a piecemeal process, where phagophores sequester only a portion of the LDs [83]. Why phagophores may not engulf entire LDs is unclear, but it has been suggested that it is due to their enormous size [83]. In this context, it has been proposed that neutral lipolysis targets large LDs to (i) decrease their volume as well as to (ii) create small nascent LDs, generated by re-esterification of FFAs, which are then accessible for macrolipophagic sequestration [94]. Indeed, predominantly small LDs are targeted by the macrolipophagic machinery since lysosomal inhibition has been shown to lead to the accumulation of small LDs within autophagosomes [94]. Although the exact mechanism that allows lipophagic autophagosomes to preferentially sequester smaller LDs has yet to be deciphered, current evidence suggests that LDs could be selectively recognized in both an Ub-dependent and -independent manner.

Ub-dependent macrolipophagy

AUP1 (AUP1 lipid droplet regulating VLDL assembly factor) is a highly conserved and broadly expressed LD surface protein that is able to interact and thus recruit the UBE2G2/UBC7 (ubiquitin conjugating enzyme E2 G2) [95]. This could provide a possible molecular connection between LDs and ubiquitination as a putative lipophagy promoting signal, although a link with the subsequent Ub-dependent autophagic degradation of LDs remains uncertain (Figure 2A). Interestingly, Dengue virus induces deubiquitination of AUP1 and UBE2G2 [96], which is necessary to induce macroautophagy and subsequent β-oxidation of the resulting FFAs to generate energy necessary for Dengue virus replication and virus production [96].

Another study has shown that the localization of the adaptor protein SPART/SPG20 (spartin) on LDs is stimulated in several cell lines fed with oleic acid, and that this association with LDs occurs through the binding of SPART C-terminal region with PLIN3 [97] (Figure 2A). Importantly, depletion of SPART causes an increase in both number and size of LDs in HEK 293 T cells fed with oleic acid [97]. SPART recruits homologous to E6-AP C terminus (HECT) Ub ligases through an interaction between its PpXY motif and the WW domain in the HECT ligases [97]. One of those ligases is WWP1 (WW domain containing E3 ubiquitin protein ligase 1), which interacts and monoubiquitinates SPART, triggering its degradation and preventing its accumulation in LDs [97]. The role of WWP1 in LD turnover has not been clearly defined yet, but it may be that by controlling SPART levels via ubiquitination, it permits to regulate macrolipophagy. Another HECT E3 Ub ligase is ITCH/AIP4 (itchy E3 ubiquitin protein ligase), which is both activated and recruited to LDs by SPART (Figure 2A). In this case, however, ITCH does not modify SPART but rather polyubiquitinates PLIN2 (Figure 2A), and potentially other LD-associated proteins [98]. PLIN2 negatively affects the turnover of the LD pool of TAGs [99]. Although these observations indicate that SPART- and ITCH-mediated ubiquitination of PLIN2 could initiate Ub-dependent macrolipophagy, more direct evidence supporting this notion need to be collected [97,98].

Immunofluorescence and co-immunoprecipitation experiments performed in a separate study revealed that rapamycin-induced autophagy in myocytes cultured in the presence of an equimolar mixture of oleate and palmitate, triggers the binding of SQSTM1 to PLIN2 on the surface of LDs [90] (Figure 2A). Under these conditions, an enhancement of the macroautophagic flux and a concomitant decrease in lipid content was observed, suggesting that SQSTM1 via PLIN2 may direct LD sequestration into autophagosomes [90]. In line with this finding, both SQSTM1- and LC3-positive autophagosomes have been found to colocalize with LDs in mouse AML12 hepatocytes that were acutely exposed to ethanol to trigger lipophagy [100], which is induced as a cytoprotective response against liver injury.
Moreover, SQSTM1 and LC3 signals overlapped with another LD protein PLIN1, and both SQSTM1 and LC3 association with LDs were significantly reduced upon PLIN1 depletion [100]. These observations suggest that ethanol-triggered macroliophagy requires SQSTM1, and that PLIN1 could be another protein on the LD surface that is recognized by this selective macroautophagy receptor [100].

A recent study in *Drosophila* hepatocyte-like oenocytes and the human HepG2 hepatoma cell line further underlined the role of SQSTM1 in LD turnover by macroliophagy upon nutrient starvation [102], and revealed that HDAC6 (histone deacetylase 6) is also involved in this process [102]. HDAC6 is required for both the transport of protein aggregates via the microtubule network to aggresomes (pericentriolar cytoplasmic inclusion bodies) [103], and the ATG machinery recruitment to aggresomes for their selective autophagic degradation [104,105]. In particular, HDAC6 depletion prevented LD breakdown, as did the overexpression of a HDAC6 mutant lacking its Ub binding domain but also SQSTM1 variants that cannot oligomerize and bind to Ub [102]. These observations suggest that either macroliophagy employs the same machinery for the selective turnover of aggresomes by macroautophagy, or that SQSTM1-positive aggresome formation and recruitment of HDAC6 to these structures is essential for macroliophagy. Additionally, knockdown of huntingtin, a scaffold protein involved in selective types of autophagy that physically interacts with SQSTM1 to facilitate its association with LC3 [106], results in
significant LD accumulation in multiple cell types fed with oleic acid, further supporting the notion that huntingtin, SQSTM1 and ultimately Ub-dependent macrolipophagy are required for the lysosomal turnover of LDs [106]. Altogether, these data suggest that there could be a mechanism of Ub-dependent macrolipophagy that relies on SQSTM1, but more solid evidence is needed to support this concept.

**Ub-independent macrolipophagy**

Evidence suggests that LDs could be selectively targeted to macrolipophagy in a Ub-independent manner. Both PNPLA2 and LIPE, which are distributed on the phospholipid monolayer limiting LDs, contain several putative LIR motifs. Co-immunoprecipitation experiments have revealed that these proteins interact with LC3, and therefore could dock LDs onto the cytoplasmic surface of autophagosomes [89] (Figure 2A). Mutation of the LIR motif at amino acid positions 147–150 dissociates PNPLA2 from LDs and disrupts PNPLA2-mediated lipolysis [89], suggesting that the interaction with LC3 may be important for neutral lipolysis, but not necessarily showing that PNPLA2 mediates macrolipophagy [89]. Another possible scenario could be that autophagosomes recruit cytosolic lipases like PNPLA2 in a LIR motif-dependent manner, to consume large LDs until their size allows to be engulfed by an autophagosome. This assumption is supported by the recent study, which indicates that PNPLA2 targets large size LDs upstream of macrolipophagy [94]. Interestingly, it has been shown that PNPLA2 is necessary and sufficient for both autophagy and lipophagy induction in hepatocytes, thereby enabling control of hepatic LD catabolism and FFA β-oxidation [107]. In fact, while pharmacological inhibition and genetic knockdown of PNPLA2 reduces Atg gene expression, macroautophagic flux and lipophagy in hepatocytes, overexpressing PNPLA2 has the opposite effect and leads to an increase in TAG turnover [107]. The latter is mediated by an autophagy program because it is inhibited pharmacologically and genetically by chloroquine and ATG5 knockdown, respectively [107]. On the same line, blocking lysosomal lipid hydrolysis using the LIPA inhibitor LALisstat, prevents TAG turnover and reduction in LD content caused by PNPLA2 overexpression. Interestingly, ablation of the SIRT1 (sirtuin 1) deacetylase, a well-established activator of autophagy through its deacetylation and activation of key ATG proteins [108], abrogated the increase in expression of Atg genes, macroautophagic flux, lipophagy and TAG turnover upon PNPLA2 overexpression in primary hepatocytes, suggesting that SIRT1 is required for PNPLA2-mediated macrolipophagy [107]. However, the mechanism by which PNPLA2 regulates SIRT1 is still under investigation [107]. Altogether, these data suggest that PNPLA2 could be an Ub-independent macroautophagy receptor that targets LDs for degradation, and therefore could also be a mechanism of Ub-independent macrolipophagy.

**Small RAB GTPases in macrolipophagy induction**

Another group of proteins that have also been linked to macrolipophagy are members of the RAB protein family. RAB proteins are small GTPases that act as switches and in their GTP-bound form, recruit effector molecules for the regulation of intracellular vesicle trafficking [109]. RAB7 plays a fundamental role in the control of late endocytic membrane trafficking [110] and autophagosomal maturation events [111,112], and moreover, decorates the surface of LDs and regulates macrolipophagy in mammalian cells [113,114]. The recruitment of RAB7 to both LDs and autophagosomal membranes is enhanced by the stimulation of lipolysis mediated by β-adrenergic receptor activation, while the depletion or inactivation of RAB7 under the same conditions inhibits macrolipophagy [114]. Similarly, it was found that RAB7 is essential for macrolipophagy in hepatocytes exposed to nutrient deficiency, since the knockdown of RAB7 or the overexpression of a dominant-negative RAB7 mutant leads to an accumulation of LDs [113]. Therefore, macrolipophagy is likely to require RAB7 under several conditions. In this context, it has recently been shown that RAB10 localizes to LDs and autophagosomal membranes in a RAB7-dependent manner under starvation conditions [115], and similar to RAB7, depletion of RAB10 results in LD accumulation in starved hepatocytes [115]. Furthermore, it was found that RAB10 is necessary for the recruitment of the complex formed by EHBP1/NACSN (EH domain binding protein 1) and EHD2/PAST2 (EH domain containing 2) on LDs in order to initiate macrolipophagy. Thereby, EHBP1 serves as an adaptor protein for EHD2, a membrane-deforming dynamin-like ATPase, which has been suggested to promote the extension of the autophagosome membrane around LDs for engulfment [115].

**Macrolipophagy regulation**

The observation that both macroautophagy and lipolysis are regulated hormonally by insulin and glucagon, and that they are enhanced during starvation, led to the idea that macroautophagy may contribute to LD breakdown and to the first report on macrolipophagy [83]. Ever since, the regulation of macrolipophagy induction has been shown to be a complex cellular and physiological response triggered by starvation and FFA loading [83], as well as hormones such as thyroid hormone [116], adrenaline [114] and FGF21 (fibroblast growth factor 21) [117]. FGF21 is secreted by the liver and induces macrolipophagy in an autocrine/paracrine-dependent manner in hepatocytes [117,118]. Interestingly, exposure to cold induces autophagy in proopiomelanocortin neurons of the hypothalamus, which in turn activate macrolipophagy in the brown adipose tissue and liver through the sympathetic network [89].

The major cellular energy sensing pathways modulating autophagy, including the classic stress and nutrient-responsive pathways such as those downstream of MTORC1 and PRKA, also orchestrate macrolipophagy. MTORC1 inhibition drives macrolipophagy, reducing lipid accumulation in an autophagy-dependent manner in vivo [89], while PRKA can activate autophagy/macrolipophagy through various mechanisms, including stimulation of ULK1 [119], inhibition of MTORC1 [120] and activation of SIRT1 [121].

Macrolipophagy is also under major transcriptional control. The most notable transcription factors are the members of the microphthalmia family of basic helix-loop-helix –
leucine-zipper transcription factors family of transcription factors, which includes TFEB (transcription factor EB) and TFE3 (transcription factor binding to IGKM enhancer 3). Both TFEB and TFE3 are master regulators of lysosomal function and macroautophagy [122,123], but are also crucial in lipid catabolism. TFEB directly upregulates the expression level of PPARGC1A/PGC1a (PPARG coactivator 1 alpha), a central player in lipid metabolism during liver starvation responses. PPARGC1A, but also TFE3, regulate lipid metabolism in the liver through the downstream nuclear receptor PPARα (peroxisome proliferator activated receptor, alpha) [124]. The effect of TFEB on lipid metabolism requires macrolipophagy since TFEB overexpression in mice fed with a high-fat diet (HFD) and in which autophagy is specifically suppressed in the liver by Atg7 deletion, does not rescue the observed increase in liver lipid content whereas it does in the wild type mice [124]. In contrast, TFE3 alleviates FFA exposure-mediated hepatocellular steatosis through macrolipophagy and PPARGC1A-mediated FFA β-oxidation, since inhibiting these pathways using small interfering RNA against Atg5 and PPARGC1A, respectively, dramatically reduces TFE3-mediated alleviation of hepatocellular steatosis [125].

Another transcription factor family involved in macrolipophagy regulation is the FOXO (forkhead box O) family. FOXO1 modulates lipid metabolism not only by inducing expression of PNPLA2 in adipocytes [126], but also by upregulating LIPA and inducing macrolipophagy upon nutrient depletion [127]. In addition, simultaneous depletion of FOX1, FOX3 and FOX4 in the liver leads to steatosis and elevated levels of TAGs in mice [128], a phenotype that can be reversed upon overexpression of Atg14 [129]. Finally, CREB1/CREB (cAMP responsive element binding protein 1) has been shown to also promote macrolipophagy upon nutrient deprival, while fed-state sensing NR1H4/FXR (nuclear receptor subfamily 1 group H member 4) inhibits this response [130]. Mechanistically, CREB1 upregulates key Atg genes, including Atg7, ULK1 and TFE3, by recruiting the coactivator CRTC2 (CREB regulated transcription coactivator 2). In presence of nutrients, NR1H4 trans-repressed these genes by disrupting the functional CREB1-CRTC2 complex [130]. A separate study showed that PPARα and NR1H4 compete for the same DNA binding sites in the ATG gene promoters, with opposite transcriptional outputs and starvation-induced PPARα activation suppresses NR1H4-mediated inhibition of macrolipophagy [131]. Taken together, these studies suggest that transcription factors and co-activators induced by fasting, such as TFE3, TFE3, PPARA, PPARGC1A, FOXO1, FOXO3, FOXO4 and CREB, promote macrolipophagy, whereas nutrient-induced transcription factors like NR1H4, suppress macrolipophagy.

**Microlipophagy**

A series of recent studies revealed that turnover of LDs can also be mediated by microlipophagy (Figure 2B). Indeed, in yeast lipophagy appears to actually not involve a macroautophagic process, at least in the tested conditions, but rather proceed through the direct engulfment of LDs by vacuoles [84,132,133]. The molecular requirements for yeast microlipophagy, however, are contradictory.

There are studies reporting that microlipophagy induced by nitrogen starvation, gradual glucose depletion or acute glucose deprivation, requires the core ATG machinery, even though this process presents microautophagic morphological features and proceeds in the absence of autophagosomes [24,84,133,134]. These works, however, obtained contrasting results regarding the implication of specific Atg proteins known to be exclusively involved in selective types of autophagy [84,133]. There have also been studies reporting that microlipophagy does not require core Atg proteins [132,135]. In particular, Atg7-independent microlipophagy triggered by acute lipid imbalance resulting from phosphatidylcholine biosynthesis defects and ER stress, involves Atg39, a reticulophagy receptor [136], and Vps24 (vacuolar protein sorting 24), an ESCRT component [132] (Figure 2B). The existence of a core ATG protein-independent and ESCRT-dependent microlipophagy was independently confirmed in a study showing that in the absence of another ESCRT component Vps27, microlipophagy is compromised during gradual nutrient depletion [135] (Figure 2B).

Yeast microlipophagy also relies on local changes in the vacuolar membrane. Gradual nutrient depletion-induced microlipophagy was first shown to be mediated by liquid-ordered raft-like sterol-enriched vacuolar microdomains, whose formation and integrity are critical for LDs translocation into the vacuole [135]. Subsequently, it was revealed that the yeast NPC (NPC intracellular cholesterol transporter) protein orthologs, Ncr1 and Npc2, are essential for both the formation of these raft-like vacuolar microdomains and LDs engulfment by vacuoles through microlipophagy, during both gradual nutrient depletion and nitrogen starvation [134] (Figure 2B). Mammalian NPC proteins, NPC1 and NPC2, bind the cholesterol derived from endocytosed lipoproteins and they transport this lipid from the lumen to the limiting membrane of lysosomes [137]. Thus, it is likely that NPC proteins promote microlipophagy by increasing the sterol in the limiting membrane of the vacuole [134]. The formation of these raft-like vacuolar microdomains upon nitrogen starvation depends on at least two core Atg components, Atg7 and Atg8 [134] (Figure 2B). This finding could imply an indirect involvement of macroautophagy in microlipophagy, as delivery of organelles and thus lipids by autophagosomes will supply the amounts of sterols necessary for the formation of the raft-like vacuolar microdomains. Intriguingly, the core ATG machinery component Atg14, translocates from ER exit sites onto sterol enriched raft-like vacuolar microdomains in response to acute glucose starvation that leads to PRKA activation, and there, together with Vps30/Atg6, facilitates microlipophagy initiation, including vacuole docking and internalization of LDs [24] (Figure 2B). Cells that cannot activate PRKA or lack either Atg14 or Vps30/Atg6, do not deliver LDs into the vacuole and fail to thrive under acute glucose starvation [24]. Interestingly, atg14Δ and vps30/ata6Δ knockout strains form little or no raft-like vacuolar domains, suggesting that these two proteins and possibly the PtdIns3K complex that are part of, play a key role in the remodeling of the vacuolar membrane under nutrient deprivation conditions.
Ypt7, the yeast ortholog of RAB7, also associates with LDs and a recent study has revealed its involvement in the regulation of LD dynamics and membrane trafficking between the vacuole and LDs [139]. In the absence of Ypt7, FFA content and the number of LDs drastically increase when cells are deprived of nutrients [139]. It is well known that in the absence of Ypt7, fusion processes with the vacuole are impaired [140,141], including the one involving autophagosomes [142]. Thus, LD accumulation could be the consequence of an impairment of the vacuolar functions. However, it has also been observed by electron microscopy that LD morphology is significantly altered and the fusion between LDs appears to be blocked, revealing a key role for Ypt7 in LD dynamics as well [139].

According to the current evidence, different conditions lead to the induction of microlipophagy in yeast, and it appears that among few mechanisms of microlipophagy, some Atg machinery-dependent, exist. This is in line with the current view that there are at least three types of microautophagy, two taking place at the vacuoles/lysosomes and one at the endosomes [28]. Future studies are needed to unveil the precise mechanisms underlying microlipophagy and their regulation.

**Chaperone-mediated autophagy and the degradation of lipid droplet surface proteins**

Although CMA targets only proteins and not organelles such as LDs, several findings support a role for CMA in the catabolism of LDs by selectively breaking down proteins present on their surface [49,143]. In particular, PLIN2 and PLIN3, are the earliest LD-anchored proteins reported to be CMA substrates [49] (Figure 2C). Expression of CMA-resistant PLIN proteins, or CMA inhibition, e.g., in mouse liver, reduces the association of the neutral lipase PNPLA2 and core ATG proteins with LDs, which in turn causes an accumulation of LDs and a decrease in lipid β-oxidation [49]. Therefore, it is believed that CMA-mediated degradation of PLIN2 and PLIN3 stimulates macrolipophagy by promoting the recruitment of PNPLA2 and macrolipophagy machinery components on LDs. A follow-up study showed that PRKA phosphorylates PLIN2, an event that is required for CMA-mediated degradation of this protein [143]. More recently, it has been shown that PLIN5 is also a CMA substrate (Figure 2C) and that LD breakdown is impaired when CMA-mediated PLIN5 degradation is inhibited by LAMP2A depletion in mouse livers and HepG2 cells [144]. Overall, these studies present CMA-mediated degradation of PLIN proteins as an upstream regulatory event that is critical for both neutral lipolysis and induction of macrolipophagy, highlighting the important role of selective autophagy in the control of LD catabolism and the crosstalk between neutral lipolysis and lipophagy.

**Dysregulation of autophagy and lipid droplet function in nonalcoholic steatohepatitis**

The regulation of lipid metabolism by autophagy is critical for cellular and organismal homeostasis. Defects in autophagy and lipid metabolism are associated with a diverse set of human metabolic diseases, including liver diseases. In liver, the process of autophagy has been studied since the 1960's and a large body of evidence identified hepatic macroautophagy as an important intrinsic mechanism relevant to human liver diseases [145,146], such as HCC and nonalcoholic fatty liver disease (NAFLD). In this context, impaired lipophagy has been discussed as a risk factor for NASH, a type of FLD considered an increasing threat to human health worldwide [147,148]. Impaired lipophagy has also been found to be associated with disease progression in NAFLD patients [149]. Hence, lipophagy regulators such as FGF21 and the members of the FOXO and TFEB protein families (see section 3.1.4) are considered as possible targets for future rational therapies that may combat NASH through the modulation of macroautophagy [150–154]. In the following, we discuss insights into the functional relationship between autophagy, macrolipophagy and NASH.

**Nonalcoholic steatohepatitis**

NASH is a metabolic disease of the liver characterized by hepatic lipid accumulation (steatosis) and inflammation [155]. NASH is a more progressive and severe stage of NAFLD and is often accompanied by fibrosis (Figure 3) [155]. NASH is associated with obesity, insulin resistance, hyperlipidemia and the metabolic syndrome [156] and a diagnosis of NASH is indicative for the need of a liver transplant later in life, due to the likelihood of liver failure or HCC development [157]. In general, steatosis is a major human health concern with approximately 25% of the world population suffering from NAFLD, and in the United States of America NAFLD and NASH affecting the 30% and 5% of the population, respectively [158].

The pathophysiology of NASH is complex and involves genetic, lifestyle and environmental factors [157]. The progression from NAFLD to NASH (Figure 3), and potentially liver failure or HCC [159], is non-linear, partially reversible, and may progress slowly or quickly over several decades [160] or it may not progress at all [156]. Accumulation of LDs within hepatocytes occurs as a result of a dysregulated lipid metabolism, which is closely associated with a metabolic syndrome whose features include obesity, insulin resistance, dyslipidemia and hypertension [156]. Histological features of NASH include steatosis, hepatocyte ballooning, lobular inflammation and sometimes fibrosis [157].

Two hypotheses have been made to explain the pathogenesis of NASH. The **two-hit hypothesis** considers lipid accumulation in hepatocytes to be regarded as the **first hit**, which precedes and sensitizes the liver to a **second hit**, characterized by mitochondrial dysfunction, oxidative stress, and release of inflammatory cytokines and adipokines [161]. The less simplistic **multiple-hit hypothesis** contemplates multiple parallel hits leading to NAFLD and NASH. Hepatic fat accumulation, insulin resistance, mitochondrial dysfunction, ER stress, gut microbiota, endotoxins, genetic and epigenetic factors as well as an altered metabolic crosstalk between different organs are taken into account [162].
**NASH mouse models**

Several mouse models of NASH have been developed and they can be categorized into diet-, genetic- and chemically-induced models (Table 1); each presenting its own advantages and disadvantages [163]. At present, there is no ideal mouse model to study the complexity of NASH observed in patients [163], as each currently available mouse model present differences in regard to histology, altered molecular pathways and genetics [164]. Often, the mouse model is chosen depending on the disease stage and disease aspect that will be experimentally addressed.

Dietary mouse models partially mimic human diets that lead to the development of NAFLD, and animals are typically subject to a methionine-choline deficient (MCD) diet, a choline-deficient, a high-fat or a fructose diet (Table 1). Mice subjected to these diets display similarities to the histological and pathologic features of human NAFLD and NASH to different degrees [165]. Genetic NAFLD and NASH mouse models are often used in combination with one of the above-mentioned diets, as genetic models alone only result in the development of NAFLD, not NASH [166]. Chemical mouse models, in contrast, are more frequently used to study more severe NASH, fibrosis, cirrhosis and HCC. Mouse models are commonly utilized in the screening of potential NASH therapeutics.

Autophagy and lipophagy have been investigated in some of these mouse models, and results suggest that they play important roles in the development and progression of NAFLD and NASH [167–169]. The studies conducted so far mainly show a block in the macroautophagic flux in NASH triggered in mice through diet, genetics or chemicals [167–169]. Evidence indicates that nonselective macroautophagy, macrolipophagy and CMA are critically involved in the development of both NAFLD and NASH [49].

**Lipid metabolism in NASH**

The liver is a major site of lipid esterification and mitochondrial β-oxidation [170]. FFAs from adipose tissue, diet and de novo lipogenesis are stored within LDs in hepatocytes. Dysregulated de novo lipogenesis (DNL), neutral lipolysis and macrolipophagy play a role in the development of NASH.

**De novo lipogenesis in NASH**

De novo lipogenesis contributes to only 5% of lipids in healthy individuals compared with 26% in NASH [171]. FFAs derived from DNL are either stored within LDs as TAG or secreted from the liver as very-low density lipoprotein (VLDL). Various studies in both mice and humans have demonstrated that increased hepatic DNL is associated with NAFLD, even when fasting [171,172]. DNL is regulated by two transcription factors: SREBP1/SREBP1c (sterol regulatory element binding transcription factor 1) and

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*Figure 3.* Dysregulated lipid metabolism plays a role in the development of NASH. Altered lipophagy, lipolysis and de novo lipogenesis in the liver contribute to NASH development through fat accumulation, and subsequent inflammation and fibrosis. A “first hit” of increased FFAs leads to accumulation of LDs in hepatocytes, a condition known as NAFLD. Additional “hits”, including inflammation and insulin resistance contribute to steatohepatitis (NASH), characterized by hepatocyte ballooning, lobular inflammation and fibrosis. Liver-resident macrophages (KCs) may acquire a pro-inflammatory phenotype due to the excess hepatic fat burden and mediate the inflammatory response through the recruitment of immune cells from the periphery. In response to hepatic injury, normally quiescent HSCs become activated by various signals. Enhanced lipophagy of vitamin A-storing LDs provides energy for HSCs activation, leading to hepatic fibrosis through HSCs-mediated collagen deposition.
## Table 1. Mouse models of NASH.

| Model                      | Mechanism                                                                 | Pathophysiologial characteristics | Suitability for NASH modeling | Autophagy/lipophagy assessments |
|----------------------------|---------------------------------------------------------------------------|-----------------------------------|-------------------------------|---------------------------------|
| **Diet-induced models**    |                                                                           |                                   |                               |                                 |
| HFD                        | High fat intake (60% fat) leading to hepatic steatosis                    | • Obesity                         | • Severity of NASH less significant than the MCD animals, even after 28 weeks. |
|                            |                                                                           | • Insulin resistance              | • Only mild fibrosis          |                                 |
|                            |                                                                           | • Hyperinsulinemia                | • More suited for modeling NAFLD rather than NASH | Increased expression of Bec1, Pik3c3/Vps34, Atg12 and Atg4b mRNA was observed in 5 weeks HFD animals, while expression was significantly reduced in 10 weeks HFD animals. Decreased BECN1 and increased SQSTM1 levels indicated reduced autophagy in 10 weeks HFD mice. A transcription factor involved in lipid metabolism, USF1, suppressed autophagy via Atg gene downregulation [167]. |
|                            |                                                                           | • Hyperlipidemia                  |                               |                                 |
|                            |                                                                           | • Hepatic steatosis               |                               |                                 |
|                            |                                                                           | • Lobular inflammation           |                               |                                 |
|                            |                                                                           | • No hepatocyte ballooning        |                               |                                 |
|                            |                                                                           | • Mild fibrosis                   |                               | [258].                         |
| MCD                        | High sucrose (40%), high fat (10%) and no methionine and choline          | • No obesity (weight loss rather than gain) | • Time efficient (fibrosis seen within five weeks) | Reduced ATG protein levels in murine liver tissues after 8 weeks MCD [168]. |
|                            |                                                                           | • No insulin resistance          | • Histological characteristics of human NASH |                                 |
|                            |                                                                           | • Hepatic steatosis              | • Significant weight decrease rather than gain, and lack of insulin resistance |                                 |
|                            |                                                                           | • Lobular inflammation           | • It does not reflect the pathophysiology of human NASH with a metabolic syndrome |                                 |
|                            |                                                                           | • Hepatocyte ballooning          |                               |                                 |
|                            |                                                                           | • Fibrosis                       |                               |                                 |
| Choline-deficient diet     | Similar to MCD, but without choline. Proteins are replaced by equal amounts of L-amino acids | • No obesity (weight loss rather than gain) | • Longer time needed on this diet to develop NASH, but more severe than the MCD diet | Not analyzed                                                                                                                                 |
| Fructose diet              | High-fructose added to a diet high in fat and cholesterol                | • Obesity                        | • Histological characteristics of human NASH |                                 |
|                            |                                                                           | • Insulin resistance             | • Metabolic features of human NASH |                                 |
|                            |                                                                           | • Hepatic steatosis              |                               |                                 |
|                            |                                                                           | • Lobular inflammation           |                               |                                 |
|                            |                                                                           | • Hepatocyte ballooning          |                               |                                 |
|                            |                                                                           | • Fibrosis                       |                               |                                 |
| FFC diet [260]             | High saturated fat, high fructose and high cholesterol                   | • Obesity                        | • Histological characteristics of human NASH |                                 |
|                            |                                                                           | • Insulin resistance             | • Metabolic features of human NASH | Increase of LC3-II and SQSTM1 in liver lysates from mice fed a FFC diet indicated a block in autophagic flux and was accompanied by increased body weight, hepatic triglycerides, hepatic steatosis and increased mRNA expression of various proinflammatory cytokines [261]. |
|                            |                                                                           | • Hepatic steatosis              |                               |                                 |
|                            |                                                                           | • Lobular inflammation           |                               |                                 |
|                            |                                                                           | • Hepatocyte ballooning          |                               |                                 |
|                            |                                                                           | • Fibrosis                       |                               |                                 |

**Genetic models**

(Continued)
| Model                                      | Mechanism                                                                 | Pathophysiological characteristics | Suitability for NASH modeling                                                                 | Autophagy/lipophagy assessments                                                                 |
|--------------------------------------------|----------------------------------------------------------------------------|------------------------------------|------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|
| **The lep<sup>−/−</sup> knockout mouse**   | Overeating animals because of a mutation in the Leptin gene                | Obesity, insulin resistance, hyperinsulinemia, hyperglycemia, hepatic steatosis, no lobular inflammation, no hepatocyte ballooning, no fibrosis | Good model of hepatic steatosis <br> In combination with one of the diets, it induces NASH | The lep<sup>−/−</sup> mice displayed reduced autophagic flux, determined by detecting increased SQSTM1 levels and decreased LC3-I to LC3-II conversion. Metformin and caloric restriction increased autophagic flux through SIRT1, alleviating hepatic lipid accumulation in lep<sup>−/−</sup> mice [169].<br>Catalpol, a compound known to eliminate insulin resistance, was found to enhance autophagic activity through PRKAA2/AMPK activation and reduced steatosis in liver from lep<sup>−/−</sup> mice [262]. |
| **The lpr<sup>−/−</sup> knockout mouse**   | Animal overeating because of a mutation in the Lpr gene                    | Obesity, insulin resistance, hyperinsulinemia, hyperglycemia, hepatic steatosis, no lobular inflammation, no hepatocyte ballooning, no fibrosis | Good model of hepatic steatosis <br> More severe fibrosis than lep<sup>−/−</sup> mice when combined with a MCD diet <br> In combination with one of the diets, it induces NASH | Reduced LC3-II and increased SQSTM1 levels indicated an autophagy impairment in lpr<sup>−/−</sup> murine liver tissues. This inhibition was reserved when the lpr<sup>−/−</sup> animals were put on caloric restriction [244].<br>ATG7, ATG5, BECN1 and LC3-II levels were decreased in lpr<sup>−/−</sup> mice. PPARD/PPARβ, a member of the PPAR family that enhances fatty acid metabolism, was also reduced in lpr<sup>−/−</sup> mice and PPARD overexpression attenuated hepatic steatosis through an enhanced autophagic flux [263].<br>lpr<sup>−/−</sup> mice on MCD displayed a block in autophagic flux, detected by increased SQSTM1 and LC3-II levels. Treatment with polydatin, a precursor of the antioxidant resveratrol, restored autophagic flux by upregulating TFEB and restoring autolysosome acidification [264].<br>Enhanced autophagosome formation and autophagic flux were measured in pancreatic β-cells of diabetic lpr<sup>−/−</sup> mice upon FFAs stimulation [265].<br>lpr<sup>−/−</sup> mice treated with irbesartan, an AGTR (angiotensin II receptor) blocker that activates PPARG/PPARβ, displayed enhanced hepatic autophagic activity via PRKAA2/AMPK activation, and AKT and MTORC1 inhibition [249]. |
| SREBF1/SREBP1c knockout mouse              | SREBF1/SREBP1c overexpression leads to increased DNL                      | No obesity, insulin resistance, hepatic steatosis, lobular inflammation, hepatocyte ballooning, fibrosis | Histological characteristics of human NASH <br> Decreased adipose tissue mass is not representative of human NASH | Not analyzed                                                                                                                                 |
| **The pten<sup>−/−</sup> null mouse**      | Pten is a tumor suppressor gene                                            | No obesity, no insulin resistance (insulin hypersensitivity), hepatic steatosis, lobular inflammation, hepatocyte ballooning, fibrosis | Histological characteristics of human NASH <br> Insulin hypersensitivity and decreased body fat mass are not representative of human NASH | Not analyzed                                                                                                                                 |
| **Chemically-induced models**              | CCl<sub>4</sub> causes acute hepatic injury and tissue remodeling through ROS production | Hepatic steatosis, lobular inflammation, hepatocyte ballooning, fibrosis, HCC | CCl<sub>4</sub> treatment results in no insulin resistance or obesity, two metabolic features of human NASH <br> Must be used in combination with HFD or a choline-deficient diet to induce NASH | CCl<sub>4</sub> treatment upregulated the expression of BECN1, LC3-II and SQSTM1 in hepatic tissue. Additionally, CCl<sub>4</sub>-treated HepG2 cells showed a block in autophagic flux shown using the RFP-GFP-LC3 marker with an accumulation of autophagosomes [266]. |

(Continued)
MLXIPL/ChREBP (MLX interacting protein like). These two transcription factors are activated by insulin and carbohydrates, respectively [173], leading to the expression of genes involved in FFA and TAG synthesis. Evidence indicates that altered expression of SREBF1 or MLXIPL contributes to NAFLD through increased DNL [174]. For example, a twofold increase in SREBF1 expression has been found in NAFLD patient livers [172,175], while expression decreased with advanced NASH [176]. Expression of the lipogenic enzymes ACAC/ACC (acetyl-CoA carboxylase) 1 and FASN/FAS (fatty acid synthase), which are regulated by SREBF1, were also enhanced in NAFLD patients [172]. Severe NASH, i.e., fibrosis at stage 3–4, also termed burnt-out NASH, is associated with the loss of hepatic steatosis as well as other histological features of NASH [177]. Fat accumulation and SREBF1 expression have been negatively correlated with NASH progression and fibrosis stage [176], while the expression of DNL-associated genes regulated by SREBF1 such as FASN, ACAC1, ACAC2, DGAT1 and DGAT2 was reduced. The pathophysiological implications of SREBF1 reduction and reduced hepatic fat accumulation in severe NASH are currently unknown.

Furthermore, MLXIPL deficiency reduces DNL by decreasing the levels of key lipogenic enzymes, which results in a 65% reduction of the hepatic FFA biosynthetic rates [178]. Mice subjected to a high fructose diet were protected from hepatic steatosis when lacking MLXIPL, though they were more susceptible to liver damage due to increased cholesterol synthesis [179]. Conversely, MLXIPL overexpression in HFD fed mice caused increased DNL and hepatic TAG accumulation, while also displaying normal insulin levels and insulin signaling [180]. In liver biopsies from NASH patients, high MLXIPL expression correlated with steatosis and low MLXIPL expression with severe insulin resistance [180]. Altogether, these studies demonstrate that the expression of the lipogenic transcription factors SREBF1 and MLXIPL, which regulate DNL, is altered in NAFLD patients, but it is also stage dependent.

ACAs are regulated by SREBF1 and are the rate-limiting step in DNL [181]. Multiple studies have shown that ACAC inhibition reduces hepatic steatosis [159,182,183] and ACAC inhibitors are being investigated as a possible treatment for NASH [181]. Interestingly, ACAC inhibition reduced TGFβ1/TGFβ (transforming growth factor beta 1)-stimulated hepatic stellate cells (HSC) activation and reduced fibrosis in a HFD diethylnitrosamine (DEN)-induced model, in rat choline-deficient diet and murine lept−/− NASH models [184] (Table 1). This study implicated a role of ACAC not only in hepatocyte TAG accumulation, but also in the activation of HSCs.

**Table 1. (Continued).**

| Model | Mechanism | Pathophysiological characteristics | Suitability for NASH modeling | Autophagy/lipophagy assessments |
|-------|-----------|-----------------------------------|-------------------------------|---------------------------------|
| Streptozocin (together with HFD or a choline-deficient diet) | Streptozocin destroys pancreatic beta cells, leading to diabetes development | • Insulin resistance  
• Hepatic steatosis  
• Lobular inflammation  
• Hepatocyte ballooning  
• Fibrosis  
• HCC  
• Clear time course of the disease progression from NALFD to HCC  
• Reflects many aspects of human NASH, however it only accounts for NASH in the context of diabetes  
• Develop type I rather than type II diabetes  
• Must be used in combination with HFD or choline-deficient diet to induce NASH  
• A potential reduction in autophagic flux, detected by a decrease in LC3-II levels, was observed in HFD mice treated with streptozocin. Decreases in lipid droplet number as well as increases in autophagosome number, BECN1 and LC3-II levels were seen with exenatide treatment, a GLP1R (glucagon like peptide 1 receptor) agonist [251]. | |

DEN [267] (together with HFD or a choline-deficient diet) | Diethylnitrosamine (DEN) is a carcinogenic reagent that causes oxidative stress | • Obesity  
• Insulin resistance  
• Hepatic steatosis  
• Lobular inflammation  
• Hepatocyte ballooning  
• Fibrosis  
• HCC  
• Within 20 weeks HFD diet fed mice treated with DEN show insulin resistance, fibrosis and HCC  
• HCC develops rapidly  
• It has most of the key features of NASH  
• DEN-treated mice fed with a HFD developed NASH and showed decreased number of autophagosomes, a decreased LC3-I/LC3-II ratio and increased SQSTM1 levels, compared to non-obese DEN-treated mice on a low-fat diet [268]. | |

**Lipolysis in NASH**

Normally, circulating insulin prevents neutral lipolysis and promotes lipogenesis in adipose tissues after feeding [185]. Under starvation conditions, in contrast, catecholamine signaling promotes neutral lipolysis and FFAs are released for their redistribution to other organs to meet energy demands [185]. NASH is often accompanied by insulin resistance, which in turn increases neutral lipolysis in adipose tissues [186]. FFAs released in excess are transported to the liver where they contribute to lipid accumulation, leading to NAFLD [186]. Neutral lipolysis also occurs in hepatocytes. As mentioned, PNPLA2 is involved in the first step of neutral lipolysis, hence hydrolyzing TAGs into DAGs, and is highly expressed in adipose tissues and liver. A common single-nucleotide polymorphism in the PNPLA3 gene, a paralog of PNPLA2, increases the risk of developing NASH by three fold [187]. This isoleucine to methionine substitution at amino acid position 148 leads to a more pronounced sequestration of ABHD5/GCPI-58 (acylhydrolase domain containing 5) by PNPLA3, the shared co-factor of PNPLA2 and PNPLA3 [188], leading to a reduction of hepatic neutral lipolysis and an accumulation of fat in hepatocytes [189]. Studies in murine pnpla2−/−
models support the notion of an association between reduced hepatic neutral lipolysis and hepatic steatosis. Mice lacking PNPLA2 have increased levels of TAGs in both adipose and non-adipose tissues [190]. In particular, the *pnpla2*−/− mice showed a 2.3-fold increase in hepatic TAG accumulation [190]. Another study showed elevated hepatic steatosis in *pnpla2*−/− mice with a 3-fold increase in hepatic TAG content [191]. Furthermore, adenoviral-mediated knockdown of hepatic PNPLA2 in mice led to steatosis regardless of whether the animals were fed with a HFD or standard chow [192]. Conversely, PNPLA2 overexpression in primary hepatocytes prevents lipid accumulation and augments TAG turnover [192].

Altogether, even if dysregulated neutral lipolysis appears to play a role in the development of NASH, data obtained from *pnpla2*−/− cells or mice should be interpreted with caution knowing that PNPLA2 could act as a macrolipophagy receptor.

**Lipophagy in NASH**

**Hepatocyte macrolipophagy in NASH.** ATG proteins have been found to be dysregulated in the livers of NASH mouse models. In genetic and dietary-induced NASH models, macroautophagy was found to be downregulated in the liver, particularly by *Atg7* expression downregulation. Conversely, *Atg7* overexpression in *lep*−/− mice (Table 1) prevented accumulation of lipids in hepatocytes through the stimulation of macroautophagy [193]. In a HFD NASH mouse model, *Atg5*, *Atg12* and *Atg16L1* mRNA expression was found reduced [194]. Inhibition of autophagy by *Atg5* knockdown or 3-methyladenine in cultured rat hepatocytes, in combination with either FFAs supplementation or MCD medium, increased LD accumulation [83]. Altogether, these studies suggest that ATG proteins can be downregulated in NASH and that induction of macroautophagy in hepatocytes through ATG overexpression can reduce lipid accumulation.

In contrast, other studies have challenged this notion and implicate autophagy deficiency to be protective against HFD-induced obesity and insulin resistance. For example, hepatocyte-specific *Atg7* deficiency attenuated steatosis in mice on a HFD [195]. Furthermore, hepatocyte-specific *Rb1cc1/Fip200* (RB1 inducible coiled-coil 1) deficiency resulted in a decrease in liver TAGs and a decrease in LDs in fasted condition compared to hepatocyte autophagy efficient mice. However, Sirius Red staining and mRNA expression levels showed increased fibrosis and inflammation [196]. Additional studies also show autophagy deficiency through hepatocyte-specific *Atg7* or *Atg5* deletion to reduce hepaticLD accumulation and protect against steatosis, but in the context of starvation rather than models of NASH [53,197–199].

To date, many studies have shown autophagy deficiency to contribute to hepatic steatosis and induction of macroautophagy in hepatocytes to be beneficial in overcoming lipid accumulation. Intriguingly, few studies show autophagy deficiency to protect against HFD-induced steatosis. The discrepancies in these studies may be due to the different knockouts of *Atg* genes used. Further studies on how *Atg* genes participate in lipid metabolism might shed more light on the role of hepatocyte autophagy in the context of NAFLD/NASH.

A recent study highlighted the association of impaired autophagy with advanced disease in NAFLD patients. The number of autophagosomes and lipid-laden lysosomes (lipolysosomes) in liver increased with higher NAFLD activity score (NAS), while SQSTM1 correlated with NAS and fibrosis stage, indicating autophagy blockage [149]. Accumulation of SQSTM1 aggregates in hepatocytes of NAFLD patients has been confirmed in another study, where it also correlated with the number of autophagic vesicles, NAS and fibrosis [200].

The mechanism by which excess lipid accumulation may interfere with autophagy functions has been investigated in mouse studies. Electron microscopy examination of liver sections from mice on a HFD revealed a significant decrease in LD-containing macroautophagic intermediates, suggesting an impairment in lipophagy. Immunoblot analysis of isolated LDs showed that the association of LC3-II with LDs in HFD mice was more prominent in the fed rather than the fasted state [83]. This apparent macrolipophagy defect caused by an excess of lipids may be due to the altered lipid composition of autophagosomes and/or lysosomes, which in turn affects their fusion [201]. This scenario indicates a harmful cycle between lipid accumulation and impaired autophagy.

The defects in autophagy caused by an excess nutrient supply may be an indirect result of altered kinase pathways, acting upstream of autophagy. For example, western diet-induced NAFLD in mice increased the expression of the mixed lineage kinase domain-like pseudokinase, a mediator of necroptotic cell death, and resulted in liver injury and inflammatory responses through inhibition of autophagy. Another kinase, the MAP3K5/ASK1 (mitogen-activated protein kinase kinase kinase 5) is an upstream regulator of the MAPK8/JNK (mitogen-activated protein kinase 8) and MAPK14/p38 (mitogen-activated protein kinase 8) MAPK signaling cascades [202] and it was discovered that in human livers MAP3K5 expression negatively correlates with hepatic lipid accumulation and NASH scores, and positively correlates with ATG5, ATG7 and ATG12 levels, suggesting a macroautophagy-linked protective role of hepatic MAP3K5 against NASH and fibrosis onset [203]. However, other studies have implicated MAP3K5 in NASH development, whereby MAP3K5 is overactive in NAFLD, leading to inflammation and fibrosis through increased MAPK8 and MAPK14 MAPK signaling [204]. Furthermore, MAP3K5 inhibition reduces hepatic lipid accumulation, inflammation and fibrosis, supporting a negative role of MAP3K5 in NASH [205,206]. Although MAP3K5 is implicated in NASH, it remains unclear whether it has a negative or positive effect on NASH progression.

**Hepatic stellate cell macrolipophagy in NASH.** HSCs are the major player in hepatic fibrosis [207,208]. In their quiescent state, HSCs are a major store of vitamin A, which is stored within their LDs [209]. Upon liver injury and in response to hepatocyte- and Kupffer cell (KC)-derived cytokines and CCL/chemokines (chemokine ligands) or other stimuli [208], HSCs undergo a transition to become myofibroblast-like [210]. Once activated, vitamin A stores are depleted through macroautophagy [211]. Importantly, activated HSCs contribute to NASH progression through the deposition of type I and III collagen; leading to the formation of fibrotic tissues [209].

Evidence suggests a role for autophagy in the activation of quiescent HSCs in NASH. Increased macroautophagic flux has been demonstrated in activated HSCs, through the observation of
increased LC3-II and decreased SQSTM1 protein levels [211,212]. Pharmacological inhibition of macroautophagy with chloroquine or 3-methyladenine, as well as small interfering RNA-mediated Atg5 or Atg7 knockdown, reduced the mRNA expression of fibrosis-associated genes in murine HSCs [212]. Moreover, mice with autophagy-deficient HSCs displayed attenuated liver fibrosis, as shown by a reduction of α-smooth muscle actin expression. The β-oxidation of FFAs produced by macroautophagy is required for HSC activation [212] as exogenous oleic acid allows macroautophagy-deficient HSCs to become activated, reacquiring the fibrotic phenotype [212]. Additional studies have shown that HSC activation is either increased or decreased through the induction or inhibition of macroautophagy, respectively [213–217]. Recent studies unraveled a role of lipopolysaccharide (LPS)-associated activation of HSCs via macrolipophagy. Systemic LPS levels are elevated in human NASH because of the disrupted gut barrier, and is one of the factors that contributes to NASH progression through toll-like receptor 4-mediated activation of KCs and HSCs [218,219]. LPS was found to induce macrolipophagy in the murine LX-2 HSC cell line, while pharmacological inhibition of autophagy or Atg5 silencing reduced LDs decrease upon LPS-treatment [220]. Reactive oxygen species (ROS) levels are elevated in NASH patients [221] and have previously been linked to macroautophagy induction in response to chemotherapy [222]. ROS content of HSCs increased over a 7-day period and was abrogated upon treatment with ROS scavengers glutathione and N-acetylcysteine. Glutathione and N-acetyl-cysteine impaired autophagosome formation and macroautophagic flux, as well as binding between PtdIns3K complex and RAB25. Thus, the augmentation of ROS levels during HSC activation contributes to LD disappearance through increased PtdIns3K-RAB25 interaction and enhanced macrolipophagy, the effects of which can be avoided by ROS scavengers [223]. These studies implicate both ROS and LPS in the activation of HSCs through macrolipophagy.

All this compelling evidence shows that macrolipophagy leads to the activation of HSCs and subsequent fibrosis, rather than playing a protective role against NASH.

**Hepatic macrophage macrolipophagy in NASH.**
Macroautophagy has a role in macrophage polarization toward their anti-inflammatory M2 type [224]. In mice fed with a HFD for 12 weeks, macroautophagic flux was decreased in peritoneal and bone-marrow derived macrophages (BMDMs) [224]. Mouse bone-marrow derived macrophages, but also Atg5-deficient KCs, showed polarization anomalies with an increase in the pro-inflammatory M1 and a decrease in the anti-inflammatory M2 population [225]. While HFD and LPS were not sufficient to cause inflammation in wild type mice, the specific ablation of macroautophagy in macrophages promoted a pro-inflammatory macrophage phenotype in the liver of animals, without altering macrophage number and steatosis [225]. Myeloid cell-specific atg5−/− deficient mice that were subjected to repeated injections of carbon tetrachloride (CCL4) to induce liver injury, had increased hepatic levels of IL1A/IL1-α (interleukin 1 alpha) and IL1B/IL1-β (interleukin 1 beta), enhanced infiltration of leukocytes in the liver and hepatocyte apoptosis, as well as exacerbation of liver fibrosis [224]. This result highlights the protective, anti-inflammatory role of macrophage basal autophagy in liver injury and fibrosis [224]. In NAFLD, KCs acquire a pro-inflammatory phenotype following their activation by danger/damage-associated molecular patterns (DAMPs) and other factors released by increased hepatic fat burden. Alternatively, KCs may become activated due to excess lipid accumulation and defective lipid processing [226]. In biopsies from NASH, but not NAFLD patients, hepatic macrophages form crown-like aggregates around LDs derived from dead hepatocytes, and these structures are similar to those found in adipose tissue in obese patients [227]. In an animal model of early NASH induced by a HFD for 16 weeks, KCs had a distinct phenotype, characterized by an accumulation of LDs, a dysregulation of lipid metabolism at the gene level [228] and an increase of cholesterol, ceramides and DAGs [228]. Importantly, enhanced lipid accumulation in KCs from HFD animals was accompanied by a pro-inflammatory phenotype, as they secreted more IFNA1/IFN-α (interferon alpha 1), TNF/TNF- α (tumor necrosis factor), IL10/CSIF (interleukin 10), CCL2/MCP1 (C-C motif chemokine ligand 2), and CCL5/RANTES (C-C motif chemokine ligand 5). Inhibition of lipogenesis in these KCs by 5-tetradecylxoy-2-furoic acid could reverse this pro-inflammatory phenotype. Taken together, accumulation of LDs in KCs influences their polarization toward a pro-inflammatory phenotype, which is a key event in the transition from NAFLD to NASH and fibrosis. Nevertheless, evidence suggests that fat-laden M1 macrophages may aid in the resolution of inflammation in advanced NASH, by expression of anti-inflammatory mediators such as ANNXIN A1 [229]. Nonetheless, the relevance of using the M1/M2 macrophage classification is debatable; since M1 and M2 macrophages may represent the same subset of plastic macrophages during the course of disease, making it difficult to draw firm conclusions [226]. Thus, more studies are needed to elucidate the exact role of lipid dysregulation in the function of different hepatic macrophage subsets in the context of NASH assisting with the resolution of acute inflammation and the initiation of tissue healing through the stimulation of phagocytosis of apoptotic bodies [229]. Notably, apart from KCs being the main drivers of inflammation in NASH, they may also directly drive steatosis development, as KC depletion in rats has been shown to prevent HFD-induced steatosis [230]. In further support of this notion, in a HFD NAFLD model, LPS or FFA-driven activation of KCs mediated steatosis formation in hepatocytes via tumor necrosis factor signaling [231].

**Liver sinusoidal endothelial cell macrolipophagy in NASH.**
Although much less studied compared to the other major liver cells types, liver sinusoidal endothelial cells (LSECs) appear to play a role in NASH pathogenesis. Changes in LSECs have been observed in the HFD and choline-deficient mouse models, where KCs and HSCs were activated, leading to a progression from simple steatosis to early NASH [232]. A recent study showed that macroautophagy is altered in LSECs from NASH patients [233]. In particular, the number of autophagosomes in LSECs from NASH patient liver biopsies was half of that in the samples from patients with no liver abnormalities or simple steatosis [233]. Primary LSECs specifically lacking Atg5 isolated from animals on a HFD, had major changes in inflammatory pathways at the gene expression level and a modest upregulation of angiogenic pathways [233]. The same animals exhibited increased fibrosis, mainly in the perisinusoidal area, higher markers of liver injury and liver inflammation, accompanied by an infiltration of inflammatory
cells [233]. This data indicates that LSEC macroautophagy has a protective role in the context of NAFLD and NASH. In another study, LSECs from rats with CCl4-induced fibrosis displayed an augmented macroautophagic flux, which did not increase further in a more advanced fibrosis stage (6 weeks of CCl4 administration) [234]. This observation suggests a role for LSEC macroautophagy during early phases of liver injury [234]. Mildly fibrotic mice (1 week of CCl4 injections) with an LSEC-specific deletion of Atg7 exhibited endothelial dysfunction and an aggravation of fibrosis, due to activation of HSCs [234]. Overall, LSEC macroautophagy appears to elicit a protective effect in the context of NASH and liver injury, yet no studies have explored the role of LSEC macrolipophagy in this context.

**Macrolipophagy in non-liver tissues over the course of NASH**

Transition from simple steatosis to NASH can be triggered by intrahepatic and extrahepatic events (Figure 3). Extrahepatic events include adipose tissue dysfunction, intestinal dysbiosis and insulin resistance [155]. A recent study has highlighted the link between adipose tissue dysfunction and NASH pathogenesis [235]. Mice were subjected to a modified HFD and developed NAFLD or NASH of different severities, based on liver histology, alanine aminotransferase serum levels and hepatic TAG levels [235]. Gene expression profiling in liver by microarrays showed that among the mice who had the most severe histological features resembling NASH, there was a significant enhanced expression in genes involved in lipid metabolism, inflammation and fibrosis [235]. Additionally, these animals exhibited both high weight gain and high LEP/OB (leptin) levels in serum, and showed severe adipose tissue dysfunction and remodeling [235]. Adipose tissue dysfunction may thus promote progression of hepatic steatosis to NASH. Studies using adipose tissue from obese and non-obese patients showed that obesity is probably associated with an activation of autophagic flux in the visceral adipose tissue [236,237]. Direct evidence linking adipose tissue dysfunction, impaired lipid metabolism and NASH is missing, yet autophagy has a major role in adipocyte maturation via macrolipophagy and lipolysis-dependent mechanisms [238].

Increased lipolysis of the peripheral fat stored in adipose tissues reaches the liver as non-esterified FFAs, enhancing lipogenesis, reducing β-oxidation and downregulating the export of fat in the form of VLDL, thereby contributing to steatosis [239]. Increased plasma FFA levels result in an increase of intramyocellular lipids, which in turn leads to an impairment of insulin signaling, causing insulin resistance in the skeletal muscles of healthy subjects [240]. The size but not the number of individual subsarcolemmal LDs in human skeletal muscle fibers is associated with insulin resistance [241], but a direct association between LDs in skeletal muscle in the context of NASH is missing.

**Outlook**

LDs are dynamic cellular organelles that consist of a neutral lipid core which is surrounded by a phospholipid monolayer and LD-associated proteins, particularly of the PLIN protein family. LDs serve as platforms for energy storage and have multiple functions in lipid and energy homeostasis, membrane and hormone biosynthesis, cell signaling and differentiation. LDs are implicated in various metabolic diseases, such as obesity, atherosclerosis and FLD [242]. The latter is the world’s most common liver disease, characterized by excessive fat in the liver and either related to alcohol consumption or not. In this context, NASH is a subtype of FLD, which is also defined by the accumulation of liver fat and characterized by liver inflammation.

Intracellularly, fat is stored in LDs and there is a complex relationship between LDs and autophagy (Figure 1), as i) the autophagic machinery appears to be necessary for the formation of LDs, ii) LDs support the formation of autophagosomes, and especially iii) the autophagic machinery selectively breaks down LDs through (macro)lipophagy processes (Figure 2). In the context of FLD, an enhancement of lipophagic activity in the liver, possibly combined with drugs that act on the lipid biosynthetic pathways, could be viewed as a possible future therapeutic strategy to reduce the symptoms of FLD and stop its progression to NASH (Figure 3). However, as discussed in this review, autophagy modulation should be liver cell type-specific, since enhancement of autophagic activity may prove beneficial in hepatocytes, KCs and endothelial cells, while in HSCs it can lead to fibrosis [243].

In recent studies, strategies to increase autophagy, including exercise and dietary interventions [118,244,245], as well as the use of substances that modulate autophagy, including rapamycin [246], a novel TFEF inducer [154], herbal medicines [205,247] and marketed, FDA-approved drugs such as metformin [248], ezetimibe [151], irbesartan [249], pioglitazone [250] and exenatide [251], have shown that they counteract NASH in different animal models. Hence, stimulation of autophagy consistently resulted in improvements in NASH, demonstrated by decreased hepatic steatosis, inflammation, and fibrosis [118,151,154,205,244–251]. In this context, a new compound called MSL is of particular interest as it stimulates autophagy while acting on the TFEF pathway, and enhances the autophagic flux and the breakdown of LDs in the liver of Lep−/− mice as well as HFD-fed mice [154].

Even though the use of compounds that improve NASH by induction of autophagy is an extremely attractive hypothesis, it needs further investigation as potential therapeutic benefits could actually be traced back to modulation of additional pathways since none of these strategies act solely on autophagy [252]. In addition, the targeted modulation of autophagy in certain cells or tissues is a formidable challenge, as our molecular understanding of how autophagy is differentially affected in certain cell types of an organ, such as the liver, is limited. For this understanding, the elucidation of the molecular players that discriminate between autophagy and closely related mechanisms such as LC3-associated endocytosis or LC3-associated phagocytosis, is essential for the discovery of autophagy-selective therapeutic windows [253]. The development of small molecules that specifically modulate selective forms of autophagy, such as lipophagy in the liver, will be essential to understand the molecular mechanisms of lipophagy and pave the way for effective therapies for NASH and other human liver diseases.

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