Autophagy and senescence, stress responses induced by the DNA-damaging mycotoxin alternariol

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Abstract

The mycotoxin alternariol (AOH), a frequent contaminant in fruit and grain, is known to induce cellular stress responses such as reactive oxygen production, DNA damage and cell cycle arrest. Cellular stress is often connected to autophagy, and we employed the RAW264.7 macrophage model to test the hypothesis that AOH induces autophagy. Indeed, AOH treatment led to a massive increase in acidic vacuoles often observed upon autophagy induction. Moreover, expression of the autophagy marker LC3 was markedly increased and there was a strong accumulation of LC3-positive puncta. Increased autophagic activity was verified biochemically by measuring the degradation rate of long-lived proteins. Furthermore, AOH induced expression of Sestrin2 and phosphorylation of AMPK as well as reduced phosphorylation of mTOR and S6 kinase, common mediators of signaling pathways involved in autophagy. Transmission electron microscopy analyzes of AOH treated cells not only clearly displayed structures associated with autophagy such as autophagosomes and autolysosomes, but also the appearance of lamellar bodies. Prolonged AOH treatment resulted in changed cell morphology from round into more star-shaped as well as increased β-galactosidase activity. This suggests that the cells eventually entered senescence. In conclusion, our data identify here AOH as an inducer of both autophagy and senescence. These effects are suggested to be to be linked to AOH-induced DSB (via a reported effect on topoisomerase activity), resulting in an activation of p53 and the Sestrin2-AMPK-mTOR-S6K signalling pathway.

Keywords: Alternariol; autophagy; senescence; Sestrin2; mTOR; lamellar bodies
**Abbreviations:** AOH (alternariol), AO (acridine orange), AMPK (AMP-activated protein kinase), ATM (ataxia telangiectasia mutated), Senescence-associate-β-galactosidase (SA-β-Gal), BSA (bovine serum albumin), ConA (Concanamycin A), Chloroquine (CQ), DMEM (Dulbecco’s Modified Eagle Medium), DSBs (DNA double strand breaks), EBSS (Earl’s balanced saline solution), ECACC (European Collection of Cell Cultures), FBS (fetal bovine serum), γH2AX (phosphorylated histone H2AX), HDAC (histone deacetylase), LC3 (microtubule-associated protein 1 light chain 3), mTOR (mammalian target of rapamycin), NAC (N-acetyl-L-cysteine), PBS (phosphate buffered saline), PFA (paraformaldehyde), ROS (reactive oxygen species), TEM (transmission electron microscopy).
1. Introduction

Mycotoxins are biologically active secondary fungal metabolites. They are often found as contaminants in food and pose a major risk for human and animal health. The immune system is a known target for several mycotoxins (Pestka 2010). The mycotoxin alternariol (AOH) is produced by the *Alternaria* genus, often found in fruit, vegetables and grain (Ostry 2008; Uhlig et al. 2013), as well as in processed fruit products such as juices and wines (Ackermann et al. 2011). AOH was present in 31% of samples of feed and agricultural commodities in Europe, with concentrations ranging from 0.02 – 7.1 µM (EFSA 2011). The highest levels of AOH are found in legume, nut and oilseed food (EFSA 2011). Currently there are no regulations on *Alternaria* toxins, including AOH, in food or feed (EFSA 2011). However, the estimated chronic dietary exposure of AOH at 1.9–39 ng/kg body weight per day exceeds the threshold of toxicological concern value of 2.5 ng/kg body weight per day used by the European Food Safety Authority in the risk assessment of AOH (EFSA 2011). This underlines the need for additional toxicity data.

AOH is found to be both genotoxic and mutagenic in several *in vitro* studies (Brugger et al. 2006; Fehr et al. 2009; Pfeiffer et al. 2007). AOH has been shown to induce both single as well as double stranded DNA breaks (Brugger et al. 2006; Solhaug et al. 2013). AOH also reduces cell proliferation by causing a long lasting cell cycle arrest. This arrest has been suggested to be caused by the DNA damage. In comparison, cell death, mainly necrosis, is only observed at substantially higher concentrations and at later time points (Bensassi et al. 2012; Solhaug et al. 2012). Furthermore, AOH is reported to induce reactive oxygen species (ROS) (Solhaug et al. 2012; Tiessen et al. 2013) resulting in oxidative DNA damage measured by the comet assay (Solhaug et al. 2012). However, addition of the antioxidant ascorbic acid did not reduce the effect.
of AOH on cell cycle, suggesting that DNA damage caused by the increased ROS levels was not directly linked to the cell cycle arrest (Solhaug et al. 2012). Interestingly, a study done by Fehr and coworkers (Fehr et al. 2009), identified AOH as a topoisomerase II poison, stabilizing the cleavable complex of DNA, thereby forming double stranded breaks (DSBs). The topoisomerase-effect of AOH may thus explain the increased levels of phosphorylated histone H2AX (γ-H2AX), a marker of DSBs (Solhaug et al. 2013), which might specifically contribute to the G2 cell cycle arrest (Solhaug et al. 2013). Furthermore, AOH is also suggested to interfere with the separation of chromosomes through an inhibition of the decatenatory activity of topoisomerase II (Solhaug et al. 2013), possibly leading to the abnormal nuclear morphology that is observer after AOH exposure in RAW264.7 cells (Solhaug et al. 2013).

Mycotoxins have generally not been associated with autophagy. Recently however, both the genotoxic mycotoxin patulin (Guo et al. 2013) and the estrogenic mycotoxin zearalenone (Wang et al. 2014) have been linked with autophagy. Autophagy can be stimulated by multiple forms of cellular stress, including ROS, DNA damage, protein aggregates and damaged organelles (Kroemer et al. 2010). Several forms of autophagy have been described: macroautophagy, chaperone-mediated autophagy and microautophagy. Recent reports suggest that macroautophagy in particular plays an important role in determining the cell fate following an initial insult (Kroemer et al. 2010; Rodriguez-Rocha et al. 2011). Macroautophagy, hereafter referred to as autophagy, is a re-cycling mechanism by which cells through lysosomal degradation re-use amino- and fatty acids. It involves sequestration of cytosolic proteins and organelles within double-membrane structures termed autophagosomes and their subsequent fusion with lysosomes, thereby generating the autolysosome. In the autolysosome, degradation occurs via various lysosomal hydrolases. Importantly, autophagy can be integrated with other cellular stress
responses through parallel stimulation, thus being central in the integrated stress response (Kroemer et al. 2010). Several signaling pathways have been implicated in the control of autophagy, including the mammalian target of rapamycin (mTOR) pathway. mTOR activity is controlled by AMP-responsive protein kinase (AMPK), which functions as a sensor for cellular nutrient and energy levels. Whereas mTOR activation stimulate protein synthesis, its down regulation is associated with autophagy (Jung et al. 2010). Recently, increased genotoxic stress has been suggested to increase autophagy, possibly through a p53-dependent induced expression of sestrins (Budanov and Karin. 2008). Sestrins have cytoprotective function based on their ability to act as an antioxidant to scavenge excessive ROS (Budanov. 2011). However, independent of its redox-potensial, Sestrin2 induced activation of AMPK is found to reduce mTOR activation and thereby induce autophagy (Hay 2008).

In general, autophagy is considered as a pro-survival mechanism. The relationship between autophagy and the various forms of cell death is still unclear (Marino et al. 2014; Ryter et al. 2014). Autophagy and apoptosis have been reported to be both co-incident and antagonistic depending on experimental context, and to cross-talk between signal transduction elements. Autophagy may also be a part of or modulate the outcome of other regulated forms of cell death such as necroptosis (Asare et al. 2009; Ryter et al. 2014). Senescence is another stress related response that has been suggested as an alternative outcome of autophagic activity (Gewirtz 2013). This is a biological state where cells have lost the ability to divide, but still remain metabolically active (Evan and d'Adda di Fagagna 2009).

AOH induces ROS production and interferes with topoisomerase activity, which both may result in DNA damage as well as cell cycle arrest. As such events may initiate autophagy and/or
senescence; we hypothesized that this mycotoxin could initiate similar events. Here we have used the RAW264.7 macrophage cell line, which is a well-known in vitro model to study various effects of mycotoxins on the immune system (Pan et al. 2014; Pestka 2010; Solhaug et al. 2012). We find that AOH induces autophagy possibly through the Sestrin2-AMPK-mTOR-S6K pathway. Interestingly, AOH also induces generation of lamellar bodies, and after prolonged exposure the cells enter senescence.

2. Materials and Methods

2.1. Reagents and chemicals

Bovine serum albumin (BSA), gels and buffers for Western blotting (NuPAGE system) were all purchased from Invitrogen (Life Technology, Carlsbad, CA, USA). Lysis buffer and senescence β-Galactosidase staining kit were from Cell Signaling (Beverly, MA, USA). Dulbecco’s Modified Eagle Medium (DMEM), Penicillin/Streptomycin, Fetal bovine serum (FBS), and Earl’s balanced saline solution (EBSS) were from Lonza (Verviers, Belgium). Bio-RAD DC protein assay was from Bio-Rad Laboratories Inc (Hercules, CA, USA). Super signal west dura chemo luminescence system was from Pierce, Thermo Scientific (IL, USA). AOH, Concanamycin A (ConA), Chloroquine (CQ), Hoechst 33342, valine, N-acetyl-L-cysteine (NAC), and Acridine Orange (AO) were from Sigma-Aldrich (St.Louis, MO, USA). [14C]-valine was from Perkin Elmer, Waltham, MA, USA.

2.2. Cell culture and treatments
Mouse macrophage cell line RAW264.7 was obtained from the European Collection of Cell Cultures (ECACC). The cells were cultured and treated as described previously (Solhaug et al. 2013). AOH was dissolved in DMSO and the final concentration of solvent in cell culture was 0.1%. Appropriate controls containing the same amount of solvent were included in each experiment. The concentrations of AOH used in this study are in the same range that causing DNA damage as previously published by us (Solhaug et al. 2012) and others (Brugger et al. 2006; Fehr et al. 2009; Pfeiffer et al. 2007). NAC was dissolved in complete medium complemented with HEPES (25 mM) and pH adjusted to 7.4. The cells were pre-treated with NAC for 1 h before exposure to AOH.

2.3. Morphological characterization

The number of normal round shaped cells and star-shaped cells in randomly selected fields was analysed by light microscopy (Leica DMIL, Solms, Germany) and expressed as percentage of all cells counted. A minimum of 300 cells were manually counted per incubation. Pictures were taken by Moticam 1000 (Motic, Hong Kong, China).

2.3. Lysosomal labeling with acridine orange

Lysosomes were analyzed by using the lysosomotropic weak base Acridine Orange (AO, Sigma-Aldrich), a metachromatic fluorophore that accumulates in lysosomes. The cells were plated on poly-L-lysine coated coverslips (BD Biosciences), stained with AO, 1 µM for 20 min at 37°C, and washed twice with PBS. Pictures were taken with a fluorescence microscope (Nikon Eclipse...
80i (Melville, NY, USA), equipped with a DS-Ri1 camera). For analysis with flow cytometry (Accuri C6, BD bioscience, San Jose, CA, USA), the cells were harvested, stained as above and analyzed by using 488\text{ex}:670\text{LP}_{\text{em}}.

2.4. Measurement of LC3II by flow cytometry

Following treatments, cells were harvested, washed once in PBS, fixed in 1% paraformaldehyde (PFA) in PBS for 15 min on ice, and post-fixed/permeabilized in 90% ice-cold methanol for at least 48 h at -20°\text{C}. For staining with antibody, 5 \times 10^5 cells were washed twice with 0.5% BSA in PBS and then incubated with LC3B antibody (Cell Signaling; 1:100) in 0.5% BSA/PBS overnight at 4°\text{C}. The cells were then rinsed twice in 0.5% BSA/PBS and incubated with secondary antibody conjugated to Alexa Fluor 647 (Molecular Probe; 1:2000) for 2 h at room temperature in the dark. The cells were then rinsed twice and analyzed by flow cytometry (Accuri C6) using 633\text{ex}:675/25\text{em}.

2.5. Western blot

RAW264.7 cells were washed twice in ice cold PBS and placed at -70°\text{C} until the next day. The cells were then scraped in lysis buffer (Cell Signaling) and centrifuged (6000 \times g, 10 min). Protein concentrations of the supernatants were quantified by using Bio-Rad DC protein assay kit. Western blotting were then performed as previously described (Solhaug et al. 2012), using the NuPage Novex system from Invitrogen and the following antibodies: LC3B, p-mTOR
(S2448), p-p70S6K (Thr389), p-AMPK (Thr172), β-Actin Rabbit mAb HRP conjugate and Anti-Rabbit IgG HRP-linked antibody (Cell Signaling), and Sestrin2 (Proteintech, Chicago, IL, USA).

2.6. Long-lived protein degradation assay

Cells were incubated for 30 h with 0.25 μCi/ml L-[14C] valine-supplemented medium, before AOH (30 μM) or vehicle (DMSO) was added and the incubation continued for 19 h. The cells were washed and chased for 5 h in fresh medium containing AOH or vehicle and 10 mM non-radioactive L-valine to allow degradation of short-lived proteins. Cells were thus incubated with AOH for a total of 24 h. Next, the cells were washed and chased for 2 h in complete medium without AOH in the absence or presence of the autolysosome inhibitor Concanamycin A (ConA). Incubation in Earl’s balanced salt solution (EBSS) for 2 h served as a positive control. Cellular protein was precipitated with TCA, and percent degradation was assessed as the acid-soluble radioactivity divided by the total radioactivity. Valine was used in the LLPD-studies as its presence does not affect autophagy in itself.

2.7. Immunofluorescence

For staining of endogenous LC3, cells were grown on glass coverslips, treated with AOH (30 μM) or vehicle for 24 h, fixed in 4% PFA, and incubated with an antibody against LC3 (#PM036, MBL), followed by Alexa488-labeled donkey anti-rabbit IgG (Jackson ImmunoResearch). The cells were stained with Hoechst 33342 (2 μg/ml in PBS) for visualization of nuclei. For high-throughput analysis pictures were taken using the automated Olympus ScanR system (Tokyo,
Japan) with an UPLSAPO 40×/0.95 objective. 64 images were automatically taken from each coverslip with identical settings and below pixel saturation. Quantification of LC3 spots was performed by the Olympus ScanR analysis software. In the automated analysis, the nuclear Hoechst-staining was used for identification of individual cells, and discrete LC3 spots were identified in the GFP channel upon manual setting of threshold values for gating of proper spot size, shape and intensity. The software then determined the number of spots and the total intensity of the spot pixels per cell using the same threshold values for all samples. LC3 spots were quantified from 400-2,000 cells per condition in each experiment.

2.8. Transmission electron microscopy (TEM)

Following drug exposure the cells were rinsed with 0.15 M Na cacodylate buffer and fixed by drop wise addition of glutaraldehyde (2.5%) for 1 h. After fixation, the specimens were rinsed several times with 0.15 M Na cacodylate buffer and post fixed with 1.5% osmium tetroxide for 1 h. After further rinsing with cacodylate buffer, the samples were dehydrated through a series of graded ethanol from 70 to 100%. The specimens were infiltrated in a mixture of acetone-Eponate (50/50) for 3 h, then in pure Eponate for 16 h. Finally, the specimens were embedded in DMP30-Eponate for 24 h at 60°C. Sections (0.5 μm) were cut on a LEICA UC7 microtome and stained with toluidine blue. Ultra-thin sections (90 nm) were obtained, collected onto copper grids and counterstained with 4% uranyl acetate then with lead citrate. Examination was performed with JEOL 1400 electron microscope (Tokyo, Japan) operated at 120 kV.

2.9. Viability
For measurement of necrotic/late apoptotic cells, the cells were plated on UpCell plates (Nunc, Rochester, NY, USA). These plates were unique in that the surface of these dishes is temperature-responsive and allows cells to detach without the use of cell scrapers at temperatures below 32°C. For harvesting, the dishes were put on ice for 10 min to allow detachment. Since necrotic and late apoptotic cells lose membrane integrity they cannot exclude PI and PI-staining is thereby detectable using flow cytometry. Briefly, cells were plated on UpCell dishes, after 24 h the cells were pretreated with the autophagy inhibitor for 1 h before AOH exposure for 24 h. After exposure the cells were harvested as described above. Cells were stained with PI (5 µg/ml) by adding the dye directly to the medium, incubated for 10 min in the dark at ambient temperature and analyzed immediately by flow cytometry (Accuri C6). Debris and doublets were excluded by gating and 10 000 cells were analyzed.

2.10. Senescence β-galactosidase staining

β-galactosidase activity is present only in senescent cells. RAW264.7 cells were fixed and stained according to the protocol provided by the manufacturer (Cell Signaling). The number of β-galactosidase positive cells in randomly selected fields was analysed by light microscopy (Leica DMIL, Solms, Germany) and expressed as percentage of all cells counted. A minimum of 300 cells were counted per incubation. Pictures were taken by Moticam 1000 (Motic, Hong Kong, China).

2.11. Statistical analysis
The data analyzes were performed using Sigma Plot version 12.0. Statistical significance (p < 0.05) was assessed using 1-way-ANOVA, followed by Dunnett’s post-test between control and treated samples or Tukey’s post-test between two treated groups.

3. Results

3.1. AOH induces morphological changes

To assess the effect of AOH on the morphology of RAW264.7 cells, the cells were treated with AOH (30 µM) for 18, 24 or 48 h before investigation under light microscopy. Fig. 1A shows that AOH induces changes in the cell morphology already after 18 h exposure. The cells changed from small and round-shaped cells into big star-shaped cells with an extensive vacuolization. After 24 h with AOH (30 µM) exposure, the changes were very distinct and about 40% of the cells had a star-shaped morphology (Fig. 1B). In accordance with our previous findings (Solhaug et al. 2012; Solhaug et al. 2013), little or no cell death was observed (Fig. 5).

3.2. AOH induces autophagy

To test if the increased vacuolization could be due to autophagy, we stained the AOH treated cells with Acridine Orange (AO). AO accumulates in acidic vacuoles like lysosomes and autolysosomes. Analysis done by fluorescence microscopy revealed that AOH treatment increased the numbers as well as the size of acidic vacuoles (Fig. 2A). The increased level of AO upon AOH treatment was further analyzed and quantified by flow cytometry. Significant accumulation of AO was found after 24 and 48 h, starting at 30 µM AOH (Fig. 2B) suggesting
increased autophagy. Next we wanted to test if AOH-induced ROS production was responsible for the accumulation of acidic vacuoles. Addition of the ROS scavenger N-acetyl-L-cysteine (NAC, 5-20 mM) did not change the AOH-induced enhancement of acidic vacuoles (Fig. 2C). These data suggests that AOH-induced accumulation of acidic vacuoles is not directly mediated by ROS, or indirectly through oxidative DNA damage.

LC3II is generated by conjugation of LC3 to phosphatidylethanolamine in the autophagosome membrane upon autophagy activation, and is considered as an indicator of autophagosome number (Klionsky et al. 2012; Mizushima et al. 2010). To investigate if the increased level of acidic vacuoles was due to enhanced initiation of autophagy, we determined the expression level of LC3II. Indeed, treatment with AOH enhanced the protein levels of LC3II, as determined by both flow cytometry (Fig. 3A) and western blotting (Fig. 3B). In further agreement with AOH-induced autophagy, AOH significantly increased the number and intensity of autophagosomes, as indicated by the typical punctuate pattern observed upon immunofluorescence staining of endogenous LC3 (Fig. 3C-E).

Enhanced levels of autophagosomes often reflect increased autophagic activity, but could also be due to reduced autolysosomal turnover. To distinguish between these two possibilities we next assessed the autophagic flux by measuring the degradation of long-lived proteins, which are considered to be bona fide autophagy substrates (Klionsky et al. 2012). In support of an increased autophagic activity, AOH significantly enhanced the degradation of long-lived proteins (Fig. 4). The rate of degradation was increased to ~50% of the level induced by amino acid starvation,
which is a potent inducer of autophagy often used as a positive control. AOH-induced protein degradation was almost completely reversed by an inhibitor of autolysosomal degradation, Concanamycin A (ConA) (Huss et al. 2002), further confirming enhanced protein degradation via AOH-induced autophagy (Fig. 4).

Next we wanted to examine if the induced autophagy protected against AOH-induced cellular stress, as cell death was barely detectable after AOH exposure (Solhaug et al. 2012). AOH was incubated together with the well-known autophagy inhibitor, chloroquine (CQ) (Klionsky et al. 2012; Poole and Ohkuma. 1981) and cytotoxicity was detected by the uptake of PI and quantified by flow cytometry. In accordance with previous findings 30 µM AOH induced little or no cytotoxicity (Control: 1.8% ± 0.54 vs AOH: 8.0% ± 5.6), whereas slight cytotoxic effects could be observed after exposure to CQ alone (Fig. 5). Most interestingly, CQ markedly potentiated the cytotoxic effects of AOH compared to the effects of these two compounds given separately. This finding suggests that AOH-induced autophagy acts as a pro-survival response.

To elucidate the possible mechanisms underlying AOH-induced autophagy, we examined the effect of AOH on the mTOR signaling pathway, a well-known negative regulator of autophagy (Jung et al. 2010; Rodriguez-Rocha et al. 2011). Indeed, the phosphorylation of mTOR as well as its substrate S6K was decreased in parallel with the AOH-induced increase in LC3II (Fig. 6). Furthermore, AOH increased the expression of Sestrin2 simultaneously with increased phosphorylation of AMPK, an established negative regulator of mTOR (Jung et al. 2010). Taken
together these data suggest that AOH induces autophagic activity that may be induced by the Sestrin2-AMPK-mTOR-S6K signaling pathway (Fig. 9).

3.3 AOH induces changes in the cellular ultrastructure associated with autophagy and changed lipid metabolism.

As revealed by phase contrast microscopy, AOH induced severe cellular retraction and appearance of vacuoles in the cytoplasm together with changed morphology (Fig. 1). A deeper insight into the subcellular alterations observed in AOH-treated cells was acquired by TEM. Control cells showed a normal ultrastructure with uncondensed chromatin and well preserved organelles (Fig. 7A). When treated with AOH (30 µM, 24 h), the most striking ultrastructure modification was an extensive vacuolization of the cytoplasm (Fig. 7 B). When treated cultures were observed in detail, autophagosomes and autolysosomes, common structures associated with autophagy were observed (Fig. 7 C-F). Interestingly, lamellar bodies, associated with changes in the lipid metabolism and phospholipidosis, were also observed in AOH-treated cells (Fig. 7 C-E).

3.4. AOH induces senescence

Since AOH-treated cells are arrested in G2 (Solhaug et al. 2013) with a flattened and enlarged morphology (Fig. 1), enhanced autophagy (Fig. 3), but apparently without increased cell death (Solhaug et al. 2012), we lastly wanted to explore if AOH induced senescence. Senescence associated β-galactosidase (SA-β-Gal) is commonly used as a senescence biomarker (Dimri et al. 1995; Rodier and Campisi 2011). As can be seen in Fig. 8, AOH increased the level of SA-β-Gal
in a time- and concentration-dependent manner, suggesting that AOH induced senescence. Interestingly, approximately 80% of the cells were positive for the senescence marker after prolonged AOH exposure (48-72 h, 30-60 µM).

4. Discussion

Our previous studies (Solhaug et al. 2013; Solhaug et al. 2012) revealed that exposure of RAW264.7 macrophages to the mycotoxin AOH increased the production of ROS and the level of DNA damage. Somewhat surprisingly, this DNA damage did not result in any increased cell death, neither necrosis nor apoptosis. However, the AOH-exposure resulted in an almost complete G₂ cell cycle arrest with high sustained levels of cyclin B₁, or possibly in a kind of G₁ arrest as most of the arrested cells had diploid or abnormal partly divided nuclei. The AOH-induced cell cycle arrest was found to be independent of ROS production. This indicates that the ability of AOH to interfere with topoisomerase activity thereby causing DSBs (Fehr et al. 2009) is of more importance for the cell cycle arrest. In the present study, we show that this AOH-induced cell cycle arrest is accompanied by increased autophagy and senescence. Such effects could have implications for immune cells thereby modulating the immune responses.

A number of chemicals have been reported to induce autophagy. However, several of the earlier studies suffer from inadequate methodology. Indeed, recent reviews on autophagy focus on the need for a more mechanistic approach when reporting effects on this process (Klionsky et al. 2012). Here we have employed a wide spectrum of well-known autophagy methods to show that AOH induces autophagic activity in RAW264.7 cells. Initially, AOH was found to increase the
number and expand the size of acidic vacuoles. Our more detailed examination of the induced process suggested that the increased vacuolization was due to increased initiation of autophagy as judged by increased protein levels of LC3II, as well as a clear induction of LC3-positive spots. Finally, in support of a real increase in the autophagic activity, AOH was shown to enhance the lysosome-dependent degradation of long-lived proteins.

The first cellular interaction explaining how autophagy is initiated is for most chemicals not known. The reason for this is that specific chemicals most often interact with many different cellular molecules and will then trigger various types of damage. Such interactions induces several stress responses including oxidative stress, DNA damage response (DSBs) and ER stress, which are known to induce of autophagy (Deegan et al. 2013; Niforou et al. 2014; Rodriguez-Rocha et al. 2011). However, similar cellular signaling pathway is also often involved in other cellular responses such as inflammation and apoptosis (Kroemer et al. 2010). The molecular mechanisms connecting these initial events and the final cellular responses are just beginning to be understood.

It is well known that p53 regulates several signaling pathways in response to certain type of cellular stress, and involves cell cycle arrest, apoptosis as well as metabolic changes (Vousden and Prives 2009). It has been suggested that both genotoxic and oxidative stress can stimulate autophagy through p53-dependent transcription of Sestrin 1/2 (Budanov and Karin 2008; Maiuri et al. 2009; Rodriguez-Rocha et al. 2011). Downstream of Sestrin2 the AMPK is activated and mTOR is suppressed (Budanov and Karin 2008), thereby releasing the negative regulation on
autophagy (Jung et al. 2010). Recently, we found that AOH induced ROS and DNA damage followed by p53 activation and a p53-dependent expression of Sestrin2 (Solhaug et al. 2012). Interestingly, the ROS scavenger NAC was found to only partially reduce the AOH-induced expression of Sestrin2 (Solhaug et al. 2012), suggesting that the DNA damage/p53-activation also was the result of other type of DNA damage, more specifically the interference with topoisomerase activity. Here, we find that NAC do not reduce the AOH-induced accumulation of acidic vacuoles. It seems therefore that the ability of AOH to interfere with topoisomerase activity, rather than its effect on ROS production, is a more likely explanation of the increased autophagy. Further linking this DNA damage and p53 response to autophagy, we find that AOH in addition to increase the expression of Sestrin2 also enhanced p-AMPK and decreased the activation of mTOR and its substrate S6K. As autophagy is found to be induced via the p53-Sestrin2-AMPK signaling pathway in several recent studies (Hay 2008; Wang et al. 2011), a similar link between these pathways and the observed AOH-induced autophagy seems reasonable. However, there are other DNA-damage initiated signaling pathways than p53-Sestrin2-AMPK-mTOR-S6K that also are linked to autophagy (Cam et al. 2014), which cannot be ruled out at this stage.

Autophagy can be induced as a response to various types of DNA damage (Rodriguez-Rocha et al. 2011). Most interestingly, also other DNA damaging agents that interfere with topoisomerase activity (e.g. camptothecin and etoposide) are found to initiate cell cycle arrest–autophagy (Abedin et al. 2007; Katayama et al. 2007) and senescence (Brel et al. 2011; Gewirtz et al. 2008; Zhao et al. 2010). Besides a role of p53, is has been suggested that the sequestration or degradation of mitotic proteins after DNA damage (in particular DSB) by autophagy might be
one way by which cells can maintain a robust checkpoint arrest in the M-phase (Dotiwala et al. 2012). However, this is likely not the case with regard to AOH as the AOH-induced cell cycle arrest is rather in G2/M transition or possibly secondary in G1 phase than in the M-phase (Solhaug et al., 2013). Interestingly, another novel link between DSB and autophagy has been suggested (Robert et al. 2011; Shubassi et al. 2012). Upon DNA damage, DNA repair factors become acetylated and are then together with the damaged DNA transported out of nuclei and degraded by autophagy. This probably allows the DNA repair machinery to be sequestered away from actively replicating DNA, which contains naturally nicks and breaks. Mistakenly “repairing” replication associated breaks and nicks could result in potentially harmful DNA damage. Both histone deacetylase (HDAC) inhibitors and rapamycin are found trigger autophagy, which then counteracts DNA repair (Robert et al. 2011). Interestingly, HDAC is also found to interact directly with DNA topoisomerase II (Tsai et al. 2000). This process might therefore be relevant for AOH induced DNA damage and autophagy as well.

ER stress is another cellular process often associated with autophagy (Deegan et al. 2013). Disturbances in the ER’s homeostatic environment disrupts the protein folding machinery and results in an accumulating of unfolded proteins in the ER lumen, thus activating the unfolding protein response (Deegan et al. 2013). Autophagy is therefore often induced and may help to digest misfolded proteins (Appenzeller-Herzog and Hall 2012). The mechanistic links between ER stress and autophagy are currently not well understood, but are suggested to involve inhibition of mTOR via Sestrin2 activation (Bruning et al. 2013). It has been suggested that the double evolvement of autophagosomes with membranes from the ER might explain the presence of lamellar bodies if multiple cycles of autophagy occur (Morissette et al. 2008). Thus, the AOH
increased autophagic activity may be linked to the enhanced level of lamellar bodies and induced cellular senescence. An accumulation of phospholipids in lysosomes and the appearance of lamellar bodies are also found to be commonly induced by drugs that alter the lipid metabolism (Anderson and Borlak 2006), as autophagy is also suggested as a mechanism to remove accumulated lipids (Kovsan et al. 2010; Morissette et al. 2009; Peropadre et al. 2011; Schmitz and Grandl 2009). However, judged by the TEM analyzes; AOH did not cause a marked increase in lipid vacuoles in RAW264.7 cells.

Senescence is known to be induced by several topoisomerase poisons (Gewirtz et al. 2008). Here we report that also prolonged exposure of AOH results in cellular senescence. Together with increased β-galactosidase activity, cellular senescence is generally accompanied by morphological changes, which may be quite striking (Kuilman et al. 2010). The cells are described as large, flat and more irregularly shaped. This is very similar to the appearance of the RAW264.7 cells observed after exposure to AOH. Furthermore, we found that the autophagy inhibitor CQ potentiated AOH induced cell death. This suggests that AOH induced autophagy as well as senescence can be considered as pro-survival mechanisms. In fact autophagy and senescence share a number of characteristics that collaterally protects the cells from toxicity of external stress. Interestingly, autophagy has also been suggested as an effector mechanism of senescence. This may be important for the rapid protein remodelling required for making an efficient transition from proliferative to a senescent state (Young et al. 2009). However, it is also found that an inhibition of autophagy is permissive for senescence (Gewirtz 2013). This type of relationship is logical if both autophagy and senescence act in a cytoprotective manner, as senescence then might serve as a backup response in the event that autophagy fails to provide
effective protection to the injured cells. Interestingly, Psedolaric Acid B (an anti-tubulin drug), induce mitotic catastrophe, G2/M cell cycle arrest followed by mitotic slippage into G1 followed by autophagy-dependent senescence (Yu et al. 2013). In our previous study (Solhaug et al. 2013) we also considered that AOH induced mitotic slippage. After prolonged AOH exposure, the cells were arrested in G2 (4N) with high levels of cyclin B1, but abnormal nuclear morphology (partly or completely divided). This suggests that the cells had gone through karyokinesis via mitotic slippage ending up as G1 like cells. These findings propose that the same triggering mechanisms causing sustained cell cycle arrest and autophagy following AOH treatment also might be linked to senescence. p21 is considered to be a critical component in stress-induced senescence (Darzynkiewicz et al. 2014; Gewirtz et al. 2008), and is found to be necessary for Adriamycin/topoisomerase II inhibitor-induced autophagy and senescence in MCF-7 cells (Goehe et al. 2012). Consistent with an activation of this pathway, we also found a robust accumulation of p21 after AOH exposure (Solhaug et al. 2012), in the same timeframe as the occurrence of abnormal nuclear morphology and senescence. Also, AOH-induced autophagy and cell cycle arrest seems to appear earlier than the induction of senescence, further support the notion that senescence is a consequence of DNA damage, cell cycle arrest and autophagy.

5. Conclusion

We found that the mycotoxin AOH induces autophagy in RAW264.7 macrophages, possibly through the Sestrin2-AMPK-mTOR-S6K pathway. The induction of autophagy is most probably associated to AOH-induced DNA damage, due to association with topoisomerase activity rather than the production of ROS. After prolonged AOH exposure the cells entered senescence. These
*in vitro* experiments suggest possible modulator effects of AOH on immune cells, and they should therefore be followed up by *in vivo* studies.

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References:


Figure legends:

**Fig. 1: AOH induces morphological changes in RAW264.7 macrophages.** Cells were treated with AOH (30 µM) for 18 – 48 h and analyzed by light microscopy (A). Cells were treated with AOH (15 – 60 µM), analyzed by light microscopy and quantified (B). More than 300 cells were counted per condition. The results are representative of 3 independent experiments. Results represent mean ± SD of 3 replicates. * indicates significantly different from control (1-way-ANOVA with Dunnett’s post-test).

**Fig 2: AOH increases the level of acidic vacuoles.** Cells were treated with AOH (30 µM) for 48 h, stained with AO and analyzed with fluorescence microscopy (A). Cells were treated with AOH at concentrations and time points as indicated, stained with AO and analyzed by flow cytometry. Results represent mean ± SD of 3 independent experiments. * indicates significantly different from control (1-way-ANOVA with Dunnett’s post-test) (B). Cells were treated with AOH (30 µM) with or without NAC (5-20 mM) for 24 h, stained with AO and analyzed by flow cytometry. Results represent mean ± SD of 5 independent experiments. * indicates significantly different from control (1-way-ANOVA with Dunnett’s post-test). AOH treated cells were not significant different from cells treated with AOH in combination with NAC (1-way ANOVA with Tukey’s post-test) (C).

**Fig. 3: AOH increases the expression of LC3II and generation of LC3 spots.** Cells were treated with AOH (30 µM) at time points as indicated and analyzed for LC3II expression by flow cytometry (A) or western blotting (B). The results are representative of 2 independent experiments. The numbers quoted above the blot were obtained following densitometric analysis.
of the bands. Cells were treated with AOH (30 µM) for 24 h, and then fixed and prepared for immunofluorescence staining of endogenous LC3 (green) and Hoechst staining of nuclei (blue). Representative images are shown. Scale bar 10 µm. (C). The mean number of LC3 spots per cell (D) and the mean intensity of LC3 spot pixels per cell (E) were quantified from 400-2000 cells per condition per experiment. The results represent mean ± SD of 3 independent experiments. * indicates significantly different from control (1-way-ANOVA with Dunnett’s post-test).

Fig. 4: AOH increases the degradation of long-lived proteins. Cells were treated with AOH (30 µM) for 24 h in the presence or absence of ConA (50 nM) for the final 2 h and analyzed for long-lived protein degradation. Incubation in EBSS for 2 h in the presence or absence of ConA (50 nM) served as a positive control. The data represent mean ± SD of 3 independent experiments. * indicates significantly different from control (1-way-ANOVA with Dunnett’s post-test). ** indicates significantly different from AOH or EBSS, respectively (1-way-ANOVA with Tukey’s post-test).

Fig. 5: Cytotoxicity induced by combined treatment with AOH and the autophagy inhibitor chloroquine. Cells were treated with AOH (30 µM) for 24 h in the presence or absence of CQ (12.5 µM). Light microscopy pictures of the treated cells (A) and cell viability measured by flow cytometric analysis of PI stained cells (B). The data represent mean ± SD of 6 independent incubations, and are representative for 3 independent experiments. * indicates significantly different from control (1-way-ANOVA with Dunnett’s post-test). ** indicates significantly different from AOH (1-way-ANOVA with Tukey’s post-test).
**Fig. 6: AOH induces protein expression associated with autophagy.** Cells were treated with AOH (15-60 µM) for 24 h and analyzed for Sestrin2, p-AMPK (Thr172), p-mTOR (S2448), p-S6K (Thr389), LC3II and β-actin by Western blot (A). The results for AOH 30 µM are quantified and the data represent mean ± SD of 3-4 independent experiments (B). * indicates significantly different from control (1-way-ANOVA with Dunnett’s post-test).

**Fig 7: AOH induces morphological changes associated with autophagy and lipid metabolism.** Cells were treated with AOH (30 µM) for 24 h and analyzed by TEM. Cytoplasm of an untreated control cell showing normal ultrastructure (A). Intense cytoplasmic vacuolization after a 24 h exposure to 30 µM AOH (B). Closer examination of autolysosomes (aly), lamellar body (la), autophagosome (a) and preserved mitochondria (m) in AOH treated cells (C-F).

**Fig 8: AOH induces senescence.** Cells were treated with AOH (15-60 µM) for 24-72 h and analyzed for β-galactosidase activity. Results represent mean ± SD of 3 independent incubations and are representative of 3 independent experiments. * indicates significantly different from control (1-way-ANOVA with Dunnett’s post-test).

**Fig. 9: A schematic model of the proposed mechanism of AOH-induced autophagy and senescence.**
A. Bar graph showing counts of LC3II over time.

B. Western blot analysis of LC3II and β-actin levels.

C. Immunofluorescence images comparing control (Ctrl) and AOH-treated cells.

D. Graph showing LC3 spots per cell (% of untreated ctrl) for Ctrl and AOH treatments.

E. Graph showing LC3 spot intensity per cell (% of untreated ctrl) for Ctrl and AOH treatments.
A

Control

AOH 15 μM

AOH 30 μM

AOH 60 μM

B

% cellular Senescence

24 h

48 h

72 h

Ctrl

AOH 15 μM

AOH 30 μM

AOH 60 μM
AOH

DNA damage (DSB)

↑ p53

↑ Sestrin2 → ↑ AMPK

↓ mTOR

↓ S6K

↑ Senescence ← ↑ Autophagy