Integrative analysis of Paneth cell proteomic data from intestinal organoids reveals functional processes affected in Crohn’s disease due to autophagy impairment

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Summary statement

Using an integrative approach encompassing intestinal organoid culture, proteomics and protein-protein interaction networks, we link Paneth cell biological functions often found affected in Crohn’s disease to autophagy impairment.

Abstract

Crohn’s disease (CD) is associated with Paneth cell dysfunctions such as altered antimicrobial secretion that is dependent on autophagy which recycles cellular components. Patients carrying the CD risk allele in ATG16L1 – an important component of the autophagy machinery - have Paneth cell abnormalities, as reproduced in Atg16l1-deficient mouse models. However, the direct effect of Atg16l1-deficiency and autophagy-impairment in Paneth cells has not been analyzed. To investigate this, we generated a mouse model lacking Atg16l1 specifically in intestinal epithelial cells (Atg16l1ΔIEC) making these cells impaired in autophagy. Using a 3D intestinal organoid culture model that we enriched for Paneth cells, we compared the proteomic profiles of organoids derived from the wild-type (WT) and Atg16l1ΔIEC mice. We used an integrated computational approach combining protein-protein interaction networks, autophagy targeted proteins and functional information to identify the mechanistic link between autophagy-impairment and disrupted cellular processes. Of the 284 altered proteins, 198 (70%) were more abundant in autophagy-impaired organoids compared to WT organoids indicating a reduced protein degradation. Interestingly, the 284 differentially abundant proteins comprised 116 proteins (41%), which are potentially targeted by selective autophagy proteins such as p62, LC3 and ATG16l1. Our integrative analysis revealed autophagy-mediated mechanisms which degrade essential proteins belonging to key Paneth cell functions such as exocytosis, apoptosis and DNA damage repair. We performed validation experiments focusing on Paneth cell-derived lysozyme to confirm our inferred observation of down-regulated exocytosis. Our observations could explain how protein level alterations in CD as a result of autophagy impairment could affect Paneth cell functions.

Keywords:

Paneth cells; ATG16L1; intestinal organoids, quantitative proteomics; selective autophagy; Crohn’s disease
**Introduction**

Paneth cells, located at the bottom of the crypts of Lieberkühn in the small intestine, secrete various types of antimicrobial compounds (e.g. lysozyme, defensins) regulate the microbial composition of the intestine as well as growth factors that maintain the crypt-associated stem cell population (Bevins and Salzman, 2011). The secretory activity of Paneth cells strongly relies on pathways to release proteins like antimicrobials into the lumen as one of the gut barrier functions contributing to intestinal homeostasis (Bel et al., 2017); (Liu et al., 2013). Conventional protein secretion involves trafficking through ER and Golgi (Farquhar and Palade, 1981; Viotti, 2016). Paneth cell defects such as altered granule morphology and increased susceptibility to ER stress are seen in mouse models where autophagy is lost from intestinal epithelial cells (Liu et al., 2013; Wileman, 2013).

Autophagy is a pivotal recycling process that sequesters cytoplasmic misfolded proteins or damaged organelles as well as clears the cytosol from invading pathogens. These targets are captured in double-membrane vesicles called autophagosomes that are subsequently delivered for degradation to lysosomal compartments (Deretic et al., 2013; Glick et al., 2010; Todde et al., 2009; Wileman, 2013). Although initially considered as a non-selective process elicited upon starvation, stress or infection, recent studies have indicated that the cargoes of autophagy, be it organelles (such as mitochondria, peroxisomes, ribosomes, endoplasmic reticulum), pathogens or protein aggregates, are recognized in a very selective manner, termed selective autophagy (Fimia et al., 2013; Zaffagnini and Martens, 2016). Sequestration of selective autophagy-targets follows recognition by specific cargo receptors and involves the Atg12-Atg5-Atg16 complex, instrumental in the early stages of the autophagosome biogenesis by determining the site of LC3 lipidation (Fujita et al., 2008). Through LIR (LC3 interacting region) motifs, the lipidated LC3 adaptor not only targets various cargoes for sequestration but also recruits multiple autophagy receptor proteins such as p62, NDP52, NBR1, NIX and Optineurin. Cargo recognition by the autophagy receptors happens generally via ubiquitin-dependent or ubiquitin-independent mechanisms (Khaminets et al., 2016). The autophagy receptors bridge their cargoes (which are specifically targeted by the presence of receptor recognition motifs and degradation signals) with the autophagosomal membrane (Stolz et al., 2014). These events eventually result in the cargo engulfment by the autophagosome, which then fuses with the lysosome to form the autophagolysosome in which the contents are degraded by lysosomal enzymes (Glick et al., 2010; Todde et al., 2009). In addition to its recycling role, autophagy is involved in Paneth cell response to microbial exposure. Upon microbial challenge, lysozyme secretion by Paneth cells is conducted through the diversion of degradative autophagy towards a secretory process, named secretory autophagy pathway (Bel et al., 2017; Kimura et al., 2017), although various autophagy-independent secretory pathways have also been reported (Barlowe and Miller, 2013).
Alteration in secretory autophagy have been associated with many intestinal diseases. Severe gut pathologies including Crohn’s disease (CD), an inflammatory bowel disease (IBD) are associated with Paneth cell dysfunctions including disrupted antimicrobials production as observed in CD patients compared to healthy subjects (Liu et al., 2016; Perminow et al., 2010; Wehkamp et al., 2005). GWAS studies have identified mutations in autophagy-related genes associated with CD. In particular, mutation in the key autophagy gene ATG16L1 resulted in granule exocytosis abnormalities in Paneth cells with a negative effect on autophagy-mediated defense against bacterial pathogens (Cadwell et al., 2008; Lassen et al., 2014; Perminow et al., 2010; Wehkamp et al., 2005). Due to its critical function in the autophagy machinery, ATG16L1 is required for the proper functioning of autophagy in general (Kuballa et al., 2008; Mizushima et al., 2003) and in various intestinal cell types, including Paneth cells (Cadwell et al., 2008; Patel et al., 2013). In mice harbouring mutations in key autophagy genes such as Atg7 or Atg16l1, lysozyme levels were decreased, granule size was reduced, and exocytosis was abnormal in Paneth cells compared with wild-type mice (Cadwell et al., 2008; Conway et al., 2013; Lassen et al., 2014; Wittkopf et al., 2012). Specific mutations in Atg16l1 such as T300A affect the activity of Atg16l1 due to the gain of a Caspase 3 cleavage site without compromising the protein architecture (Salem et al., 2015). Even though the critical role of ATG16L1 in modulating autophagy in Paneth cells is known, the exact molecular mechanisms and cellular processes affected by autophagy-impairment as a consequence of Atg16l1 mutation remain to be elucidated.

In this study, we use the small intestinal organoid culture model which reproduces crypt-like and villus-like domains characteristic of intestinal morphology recapitulating many functions of the small bowel. Intestinal organoids contain cell types such as Paneth cells that cannot be examined in cell lines making them a unique model system to analyse Paneth cell proteins and functions (Sato et al., 2009). To increase the usefulness of the organoid model, we enrich both WT and autophagy-impaired organoids for Paneth cells by directing the lineage of organoid differentiation by following established protocols (see Methods section). Thereafter, we analyse the quantitative proteome of Paneth cell-enriched small intestinal organoids specifically lacking Atg16l1 in intestinal epithelial cells (Atg16l1IEC), and compare it to the proteomic profile of WT Paneth cell-enriched organoids. Given the known defects of autophagy in CD, the major autophagy impairment due to the loss of Atg16l1 could be considered as an extreme model of CD. In order to understand the possible mechanisms by which autophagy impairment could affect the altered proteins, we establish an in silico workflow (Figure 1) combining several computational approaches including protein-protein interaction networks, interaction evidence incorporating protein targeting by selective autophagy and information on functional processes. Using this integrative approach, we show that proteins with altered abundances in the autophagy-impaired Paneth cell-enriched organoids, could be substrates of selective autophagy and could be targeted by autophagy resulting in their degradation. The integrative approach also helped reveal novel mechanisms which mediate the modulation of several cellular processes by autophagy. Importantly, we confirm that autophagy dysfunction alters several cellular processes; for
example, cellular exocytosis which is known to be deleteriously altered in CD patients, was
downregulated in autophagy-impaired organoids. Taken together, our observations based on a
CD model of acute autophagy impairment in Paneth cells provides a mechanistic explanation of
Paneth cell dysfunction due to autophagy-impairment. The demonstrated involvement of novel
autophagy-dependent processes in Paneth cells enhances our understanding of disorders related
to autophagy dysfunction such as Crohn’s disease. Furthermore, it opens the door for the
development of new and/or supplementary therapeutic interventions for CD.

Methods

Animal handling
C57/Bl6 mice of both genders were used for organoid generation. All animals were maintained
in accordance with the Animals (Scientific Procedures) Act 1986 (ASPA).

Generation of Atg16l1<sup>flox/-</sup> VilCre (<sup>Atg16L1<sub>△IEC</sub></sup>) mice
Mice were generated using a Cre/LoxP system. Briefly, LoxP sites were inserted either side of
Atg16<sub>11</sub> exon 2, creating Atg16<sub>11</sub><sup>flox/+</sup> mice. Crossing these mice with PGK-Cre mice, expressing
Cre recombinase under the PGK promoter that is in all cell types, led to the excision of the
floxed exon 2 in one allele by Cre recombinase. This in turn introduced a stop codon, generating
Atg16<sub>11</sub><sup>+/−</sup> mice. Cell type-specific Atg16<sub>11</sub> deletion was induced using a Villin promoter to drive
expression of Cre recombinase only in villin-expressing cells. This was achieved by crossing
Atg16<sub>11</sub><sup>flox/flox</sup> mice with Atg16<sub>11</sub><sup>+/−</sup> VilCre mice to produce Atg16<sub>11</sub><sup>flox/-</sup> VilCre mice deficient in
Atg16<sub>11</sub> in intestinal epithelial cells. Heterozygous mice were used as wild-type controls (WT).
Transgenic mice were genotyped using end-point PCR and gel electrophoresis. Atg16<sub>11</sub> alleles
were firstly designated as WT (+), floxed (Fl) or knock-out (-) and the presence of the Cre
recombinase gene under the control of the villin promoter subsequently designated positive (Vil-
Cre) or negative. Combining the PCR results identified the Atg16<sub>11</sub><sup>Fl/+</sup> (WT) or Atg16<sub>11</sub><sup>Fl/-</sup> VilCre
(Atg16<sub>11</sub><sub>△IEC</sub> KO) mice. All primers used to validate the organoid models are listed in
Supplementary table 1.

Small intestinal organoid culture
Murine small intestinal organoids were generated as described previously (Sato et al., 2013).
Briefly, small intestine was opened longitudinally, washed in cold PBS then cut into ~5mm
pieces. The intestinal fragments were incubated in 30mM EDTA/PBS for 5 minutes, transferred
to PBS for shaking, then returned to EDTA for 5 minutes. This process was repeated until five
fractions were generated. The PBS supernatant fractions were inspected for released crypts. The
crypt suspensions were passed through a 70µm filter to remove any larger villus-containing
fragments, then centrifuged at 300xg for 5 minutes. For 3D organoid cultures pellets were
resuspended in 200µl phenol red-free Matrigel (Corning), seeded in small domes into 24 well-
plates and incubated at 37°C for 20 minutes to allow Matrigel to polymerise. Organoid media
(Advanced DMEM/F12 (Life Technologies)) containing growth factors including EGF (50 ng/ml, Life Technologies), Noggin (100 ng/ml, PeproTech) and R-spondin 1 (500 ng/ml, R&D) was then overlaid. For 2D organoid monolayers, pellets were resuspended in organoid media and overlaid onto coverslips coated with phenol red-free Matrigel (Corning). For the quantitative proteomics analysis, the Paneth cell population in 3D WT and Atg16l1ΔIEC organoids were enriched by addition of 3μM CHIR99021 and 10μM DAPT to the organoid culture media according to the previously published and well established protocol (Nakanishi et al., 2016; Yin et al., 2014) in order to shift differentiation mainly towards the Paneth cells phenotype.

**Western blot**

To confirm the absence of the Atg16L1 protein, impairment in autophagy and lysozyme quantification, specific protein expression was analysed by immunoblotting. On day 8 post crypt isolation, normally differentiated or Paneth enriched organoid pellets were lysed in m-PER lysis buffer (ThermoFisher Scientific) containing protease inhibitor cocktail (Roche). 3-30μg protein per well was separated using NuPAGE precast 4-12% polyacrylamide gel system (ThermoFisher Scientific). Immunoblotting was carried out using an X-Cell II blot module (ThermoFisher Scientific) onto polyvinylidene (PVDF) membranes (ThermoFisher Scientific). Membranes were probed for ATG16L1 (Abgent), LC3II (Sigma), β-actin (Sigma) and lysozyme (Dako) and protein bands visualized using Odyssey infrared imaging system (Li-Cor) at 700nm and 800 nm. Densitometry of lysozyme bands was performed using the FIJI/ImageJ package and expressed as arbitrary units (AU) from at least 3 biological replicates for each group.

**Lysozyme activity assay**

Lysozyme activity associated with Paneth cell antimicrobial defense was measured in 2D organoid culture medium using the EnzChekTM Lysozyme Assay Kit, according to manufacturer’s instructions (ThermoFisher Scientific). Briefly, 2D organoids were cultured from WT and Atg16l1ΔIEC mice as described in the relevant method section. Following a 20 hours post-seeding incubation, the organoid culture medium was collected. Remaining cellular debris were removed by centrifugation at 600xg for 5 minutes and filtration on 0.20μm PES filters. FITC-fluorescence, proportional to the lysozyme activity released by Paneth cells into the medium, was measured on a Fluostar Optima Fluorometer (BMG Labtech), and corrected for background fluorescence. Lysozyme activity expressed as U/ml was determined from standard curves on at least 3 biological replicates for each group.

**Reverse transcription and cell type specific RT-PCR**

To confirm WT and Atg16l1ΔIEC KO organoids retained the intestinal phenotype and expressed the intestinal epithelial cell-type markers, gene expression was analysed by RT-PCR. On day 8 post crypt isolation, normally differentiated organoid pellets were lysed in 500μl TRIzol (Life Technologies). RNA extraction was performed by chloroform extraction followed by precipitation in isopropanol and ethanol. Superscript II reverse transcription protocol was used...
with 200ng organoid RNA to generate single stranded cDNA. Gene specific primers for Lgr5 (stem cells), Villin (epithelial cells), Chromogranin A (enteroendocrine cells), Mucin 2 (goblet cells), CD24 (Paneth cells) and β-actin were used for amplification. Primers are listed in Supplementary table 1 (S1).

Sample preparation for proteomics
On day 8 post crypt-isolation, Paneth enriched 3D organoids were extracted from Matrigel using Cell Recovery Solution (BD Bioscience), washed in PBS and centrifuged at 300xg for 5 minutes. Organoid pellets were lysed by sonication in 1% (w/v) sodium deoxycholate (SDC) in 50mM ammonium bicarbonate. Samples were heated at 80°C for 15 min before centrifugation at 12,000xg to pellet debris. The supernatant was retained and proteins reduced with 3 mM DTT (Sigma) at 60°C for 10 min, cooled, then alkylated with 9 mM iodoacetamide (Sigma) at RT for 30 min in the dark; all steps were performed with intermittent vortex-mixing. Proteomic-grade trypsin (Sigma) was added at a protein:trypsin ratio of 50:1 and incubated at 37°C overnight. SDC was removed by adding TFA to a final concentration of 0.5% (v/v). Peptide samples were centrifuged at 12,000 x g for 30 min to remove precipitated SDC.

NanoLC MS ESI MS/MS analysis
Peptides were analysed by on-line nanoflow LC using the Ultimate 3000 nano system (Dionex/Thermo Fisher Scientific) system coupled to a Q-Exactive HF mass spectrometer (Thermo Fisher Scientific) essentially as described in Dong et al 2017 (ref[AS1]). Peptides were separated by an Easy-Spray PepMap® RSLC analytical column (50 cm × 75 μm inner diameter, C18, 2 μm, 100 Å) fused to a silica nano-electrospray emitter (Dionex). Column temperature was kept at a constant 35°C. Chromatography buffers consisted of 0.1 % formic acid (buffer A) and 80 % acetonitrile in 0.1 % formic acid (buffer B). The peptides were separated by a linear gradient of 3.8 – 50 % buffer B over 90 minutes at a flow rate of 300 nl/min. The Q-Exactive HF was operated in data-dependent mode with survey scans acquired at a resolution of 60,000. Up to the top 10 most abundant isotope patterns with charge states +2 to +5 from the survey scan were selected with an isolation window of 2.0Th and fragmented by higher energy collisional dissociation with normalized collision energies of 30. The maximum ion injection times for the survey scan and the MS/MS scans were 100 and 45ms, respectively, and the ion target value was set to 3E6 for survey scans and 1E5 for the MS/MS scans. MS/MS events were acquired at a resolution of 30,000. Repetitive sequencing of peptides was minimized through dynamic exclusion of the sequenced peptides for 20s.

Protein identification and quantification
Spectral data was imported into Progenesis QI (version 4.1, Nonlinear Dynamics). Runs were time aligned using default settings and using an auto selected run as reference. Peaks were picked by the software using default settings and filtered to include only peaks with a charge
state between +2 and +7. Spectral data were converted into .mgf files with Progenesis QI and exported for peptide identification using the Mascot (version 2.3.02, Matrix Science) search engine. Tandem MS data were searched against a database including translated ORFs from the *Mus musculus* genome (Uniprot reference proteome (reviewed), UP000000589, February 2017) and a contaminant database (cRAP, GPMDB, 2012) (combined 17,010 sequences; 9,549,678 residues. Precursor mass tolerance was set to 10 ppm and fragment mass tolerance was set as 0.05 Da. Two missed tryptic cleavages were permitted. Carbamidomethylation (cysteine) was set as a fixed modification and oxidation (methionine) set as a variable modification. Mascot search results were further validated using the machine learning algorithm Percolator embedded within Mascot. The Mascot decoy database function was utilised and the false discovery rate (FDR) was <1%, while individual percolator ion scores >13 indicated identity or extensive homology (p <0.05). Mascot search results were imported into Progenesis QI as XML files. Peptide intensities were normalised against the reference run by Progenesis QI and these intensities were used to highlight relative differences in protein expression between sample groups. Only proteins with 2 or more identified peptides were included in the dataset. Statistical analysis (one factor anova) of the data was performed using Progenesis QI to identify significantly (p < 0.05, absolute relative fold change ≥ 2, number of unique peptides ≥ 2) proteins with altered abundances. The proteomic dataset has been submitted to PRIDE (accession id : PXD010940).

**Interaction resources and computational methods to identify proteins targeted by autophagy**

In order to make the proteomic data comparable to human interaction networks, the human orthologs of the proteins with altered abundances from the Paneth cell organoids were identified using InParanoid (Sonnhammer and Östlund, 2015). To identify the autophagy targeted protein components, the interaction partners of the three autophagy receptor and adaptor proteins, namely p62, LC3 and ATG16L1, were retrieved from the manually curated section of the Autophagy Regulatory Network (Türei et al., 2015). To enhance the coverage and improve interpretations, the interactions retrieved from experimental data were complemented with the predicted targets of p62, LC3 and ATG16L1. The putative targets of p62 and ATG16L1 were inferred using in-house custom scripts written in the Python programming language. The predictions of p62 and ATG16L1 targets were based on the standard motif search and domain-domain interaction prediction methods (Korcsmaros et al., 2013), respectively. The p62 recognition motif was retrieved from (Jadhav et al., 2011). For the domain-domain interaction prediction method, the known set of interacting domain-pairs were obtained from the DOMINE database (Raghavachari et al., 2008). DOMINE captures information about interacting domain-pairs from experiments, structural studies and predictions. Domain annotations for all proteins were retrieved from UniProt (The UniProt Consortium, 2017). The targets of LC3 were downloaded from the iLIR database (Jacomin et al., 2016).
Functional analysis to identify affected processes
To assess the functional importance of proteins with altered abundances targeted by autophagy within a network context, Gene Ontology Biological Process terms (Ashburner et al., 2000) derived from UniProt (The UniProt Consortium, 2017) were used. Biological Process terms not related to the intestine (action potential, development of other organs/tissues etc) were discarded following a manual curation of the terms in order to maintain the intestinal context. (Supplementary table 2). We performed an additional, extensive manual curation to determine the role of all the proteins in these biological processes as well as to define their effect (activational or inhibitory) on these processes. Based on the directionality of the interaction between p62/LC3/ATG16L1 and the proteins with altered abundances, we inferred the direction of modulation of the assigned biological process terms. The aggregated trends of the processes were inferred as follows: the processes were considered to be either up-regulated or down-regulated if more than 70% of its interactions were classified as stimulatory or inhibitory, respectively. In cases where the singular proportion is less than 70%, the functional process was considered to have dual modulation (i.e., both up- and down-regulated). The networks were visualized in Cytoscape 3.5.1 (Franz et al., 2016).
Results

Atg16l1ΔIEC organoids are a valid model to study the role of autophagy in intestinal epithelial homeostasis

We have generated a mouse model that lacks Atg16l1 specifically in intestinal epithelial cells (Atg16l1ΔIEC) and have used self-organizing in vitro organoid cultures generated from small intestinal stem cells mice (Sato et al., 2009) to assess the impact of autophagy on Paneth cell functions. Organoids from both WT and Atg16l1ΔIEC mice formed viable budding crypts that expanded from a core organoid (Figure 2a) when cultured in media containing the in vivo niche components EGF, Noggin and RSPO1. The core organoids were enriched for Paneth cells using a well-established and published protocol, presented in details in the Methods (Nakanishi et al., 2016; Yin et al., 2014). Proteins from the differentiated organoids were over-represented (19/83, P-value = 0.0021) with Paneth cell markers determined and published in a previous study using single cell transcriptomics (Haber et al., 2017) thus supporting the enrichment of the differentiated organoids with Paneth cells.

Detection of mRNA transcripts by RT-PCR for lgr5, chrA, muc2 and cd24 revealed that Atg16l1ΔIEC organoids expressed markers for stem cells, enteroendocrine cells, goblet cells and Paneth cells at similar levels as WT organoids (Figure 2b), confirming expression of important cell types found in the in vivo small intestinal epithelium. We observed that the villin transcript shows a slight reduction in the KO organoids compared with the WT ones but remains indicative of the presence of enterocytes in both organoid models. In particular, we noted that the level of the Paneth cell marker CD24 was similar between WT and Atg16l1ΔIEC organoids (Figure 2b), suggesting that the number of Paneth cells was similar in both genetic backgrounds. Consistent with the intestinal epithelial cell-specific Atg16l1 deficiency, Western blot analysis confirmed Atg16l1ΔIEC KO organoids were deficient in the Atg16l1 protein. ATG16L1 was detected in WT organoid samples at 68kDa, but not in Atg16l1ΔIEC organoids even though a non-specific band is seen with the used antibody at 66kDa. In addition, we also observed LC3I to LC3II conversion in WT but not in the Atg16l1ΔIEC organoids thus indicating that ATG16L1 deletion leads to autophagy deficiency. As observed in previous studies, lack of Atg16l1 resulted in impairment of autophagy as confirmed by reduced levels of LC3II (Figure 2c) (Lassen et al., 2014; Cadwell et al., 2008; Patel et al., 2013). Altogether these observations validate Atg16l1ΔIEC organoids as a robust model for investigating the impairment of autophagy in epithelial homeostasis.

Alteration in the proteomic abundance profiles upon autophagy-impairment

In order to determine the functional significance of the Atg16l1 deficiency in Paneth cells, we established an integrated workflow (Figure 1) combining computational approaches to integrate and interpret the experimental readouts. We measured the protein levels in Paneth cell-enriched organoids derived from WT mice and mice harboring the Atg16l1 deficiency. Our proteomic experiments detected 283 mouse proteins corresponding to 284 human ortholog proteins with
altered abundances at the cut-offs used (p < 0.05, absolute relative fold change ≥ 2, number of unique peptides ≥ 2) (Supplementary table 3). Our initial functional analysis showed that proteins with altered abundance were related to at least 18 functional processes (Figure 3), and that the majority (70%) of all of these proteins were detected at levels twice greater than those found in WT organoids (Supplementary table 4), suggesting that the observed higher abundance could be a due to autophagy impairment.

Proteins potentially targeted by selective autophagy have altered abundances in Atg16l1ΔIEC Paneth cell organoids
Since the primary role of autophagy is to identify, target and recycle damaged proteins, altered protein levels in Atg16l1ΔIEC Paneth cell-enriched organoids reflect the possible effect of a disrupted autophagy process. To determine whether autophagy directly or indirectly affects the proteins that are differentially abundant in the autophagy-impaired background, we compared the altered proteins with the target proteins of known selective autophagy receptors and adaptors, such as p62, LC3 and Atg16L1 (Table 1). This network analysis and the subsequent functional investigations were performed using human data (by inferring the human orthologs of the differentially abundant mice proteins) due to increased data availability on human networks/ontologies and thereby an increased coverage. By incorporating information about the binding partners (using experimental evidence and structure-based predictions) of the human orthologs, we identified the autophagy targeting proteins which could potentially target the altered proteins in Paneth cell organoids. 116 proteins (41%; P-value 0.049) with altered abundance in autophagy-impaired organoids and more importantly, 85 proteins with increased abundance (43.14%; P-value 0.043) (Supplementary table 5) were found to be potentially targeted by at least one of the three autophagy-related proteins (p62, LC3 and ATG16L1). This indicates that the proteins with increased levels in autophagy-impaired Paneth cell organoids are targeted for degradation by selective autophagy in normal organoids where autophagy is functional and not compromised. Overlap analysis of the altered proteins individually targeted by p62 (upregulated in Atg16l1-deficient organoids as expected (Ichimura et al., 2008), LC3 or ATG16L1 indicates that only a small proportion (19%) of the altered proteins potentially targeted by autophagy are targeted by more than one of the three autophagy proteins (Figure 4, Supplementary table 5). This suggests that the autophagy machinery potentially mounts a coordinated effort to specifically target certain groups of proteins.

Identification of Paneth cell functional processes affected by autophagy-impairment
To determine which cellular functions could be affected due to the altered protein abundances upon autophagy-impairment, we analysed the protein functions using manual curation of experimental evidence and Gene Ontology Biological Process terms (Supplementary tables 6, 7). We identified altered functional processes, such as apoptosis, exocytosis, DNA repair etc that could be dependent on autophagy-mediated protein degradation (Supplementary tables 8).
Since post-translational regulators can elicit positive and negative effects on functional processes, we integrated an extensive literature curation evaluating the effect of each differentially abundant protein on associated functional processes (Supplementary table 9). For each functional process, we separately calculated an aggregated trend (see Methods for detail) to determine how the altered protein levels and the identified effect result in up- or down-regulation of process (Table 2). For example, HSPA4L, FGA and CSTB proteins, abundant after autophagy-impairment, are negative regulators of apoptosis; MAPK3 is also abundant after autophagy-impairment but it is a positive regulator of apoptosis. Therefore, this data could indicate that apoptosis might be down-regulated upon autophagy-impairment as 75% of the altered proteins that are negative regulators had increased abundances in the autophagy impaired organoids.

Figure 5 outlines the potential autophagy-dependent functional categories that were altered and their aggregated trends. Interestingly, based on the aggregated trends, we observed that 14 of the 16 altered functional processes were either uniquely up-regulated or bi-directionally modulated (both up- and down-regulated), while only two functional processes were uniquely down-regulated (Table 2). This suggests that the overall consequence of autophagy-impairment in Paneth cells is predominantly characterized by the (over)activation of various functions. These include processes such as DNA repair, endocytosis, immune response and mitochondrial organization. Some of the up-regulated functions such as endocytosis and immune functions have previously been directly associated with autophagy (Deretic et al., 2013; Levine et al., 2011; Tooze et al., 2014). The two uniquely down-regulated functional processes, apoptosis and exocytosis, have also been associated with autophagy (Brooks et al., 2012; El-Khattouti et al., 2013; Tooze et al., 2014). Thus our results extend existing interplay models (among processes) with an autophagy-mediated protein degradation control mechanism for even more precise regulation of key functions in Paneth cells, such as exocytosis.

Impact of autophagy-impairment on exocytosis in Paneth cells
As secretory cells, Paneth cells are reliant on high levels of protein biosynthesis and secretion, the latter being strongly reliant on functional autophagy. We assessed whether the autophagy impairment had an effect on the levels of proteins associated with exocytosis. Interestingly, our integrative analysis (using the workflow explained before) of the proteomic response revealed that exocytosis could be repressed in the absence of functional autophagy. Figure 6A shows the altered levels of exocytosis proteins in the Atg16l1ΔIEC Paneth cell organoids as well as the autophagy targeting proteins, which could be modulating them. This result is in agreement with the already established importance of autophagy in the exocytosis-mediated secretion of antimicrobial peptides (AMPs)(Brooks et al., 2012; Cadwell et al., 2008; Cadwell et al., 2009a; Cadwell et al., 2009b; Gassler, 2017; Tschurtschenthaler and Adolph, 2018). We also determined experimentally that lysozyme levels detected within organoids were significantly greater when autophagy was impaired (Figure 6B) than in WT organoids. However, levels of lysozyme
secreted into the culture medium were slightly reduced although not significantly different whether \textit{atg16l1} was present or absent (Figure 6C), suggesting a defective exocytosis pathway upon impaired autophagy. Detailed analysis of our proteomic data showed that proteins targeted by LC3, Atg16l1 and p62 (Figure 4) and involved in the inhibition of exocytosis were found to be more abundant when autophagy was impaired. The opposite effect was observed with autophagy targeted proteins involved in the activation of exocytosis being less abundant upon autophagy-impairment (Figure 6D). This agrees with the negative alteration of exocytosis of AMPs that we observed in our validation assays (Figure 6B,C).

The exocytotic pathways facilitating the secretion of proteins are mediated either by general mechanisms involving the endoplasmic reticulum and the Golgi apparatus (Barlowe and Miller, 2013) in an autophagy-independent manner, or, in the case of antimicrobial proteins through the recently discovered LC3- and autophagy-dependent secretory process (Bel et al., 2017; Ponpuak et al., 2015). So far it was neither clear whether these two exocytotic pathways are co- or independently-regulated, nor whether they share some of the proteins involved and target overlapping proteins. Here, we revealed that proteins with exocytosis functions having higher abundance levels upon impaired autophagy could be potential autophagy-targets; these include SRP14 (Signal recognition particle 14 kDa protein), GORASP1 (Golgi reassembly-stacking protein 1) and TMED3 (Transmembrane emp24 domain-containing protein 3; Figure 6D; Supplementary table 10). This unexpected result raises the question whether to revisit the autophagy relatedness of the ER/Golgi-specific pathway, which shunts lysozyme into secretory granules involved in exocytosis. Based on these observations, it is plausible that autophagy could have a direct effect on processes, which are thought to be autophagy-independent.

\textbf{Discussion}

Using a combinatorial approach integrating proteomic data generated from Paneth cell enriched organoids of a new Crohn’s disease mice model, interaction networks and validatory experiments, we revealed Paneth cell functional processes dependent on autophagy as a result of dysfunctional Atg16l1. Atg16l1 has been described as a pivotal autophagy protein in the last decade and it was shown that dysfunctional Atg16l1 leads to impaired formation of autophagosomes and poor degradation of long-lived proteins (Kaser and Blumberg, 2014; Saitoh et al., 2008). Our study focused on the role of autophagy in Paneth cell homeostasis, in particular on the consequences of impaired autophagy on Paneth cell functions. We used Paneth cell-enriched organoids derived from mice lacking the \textit{Atg16l1} gene specifically in intestinal epithelial cells. \textit{Atg16l1} is an important component of the autophagy machinery whose human ortholog was previously associated to Crohn’s disease. We determined quantitative proteomics profiles of Paneth cells-enriched organoids with functional or impaired autophagy. We developed and applied a computational systems biology approach based on the analysis of proteomics data generated from Paneth cells with functional and impaired autophagy and
integrated multiple types of already existing but so far unconnected disparate information (protein-protein interaction networks, information about proteins known to be targeted by autophagy and functional information about proteins displaying differential abundance when autophagy was impaired). Integration of these data with the interaction networks of selective autophagy receptors and adaptors, such as p62, LC3 and ATG16L1, helped relating the degradation of the altered proteins to their regulation by autophagy. Furthermore, by incorporating known functions and biological processes attributed to the affected proteins, we identified various cellular processes, which could be dependent on a functional autophagy process.

As recently reported for stem cell-enriched organoids, our study emphasises the robustness of systems-level approaches to fully capture the impact of major impaired cellular processes; in our case autophagy on homeostatic cellular functions (Lindeboom et al., 2018). The computational pipeline presented in this study enabled building regulatory networks of proteins displaying differential abundance upon autophagy impairment. To overcome the lack of mouse protein-protein interaction information involving the autophagy receptor and adaptor proteins, as well as to exploit the corresponding information already available in human datasets, we used the human orthologs of the mice proteins with altered abundances in mouse-derived Atg16l1ΔIEC organoids. Although the cross-species extrapolation could be a source of uncertainties and possible missing information, the identified processes and their direction of modulation concur to a certain extent with already existing knowledge about the effects of autophagy-impairment. Other notable limitations of our study include the inability of the proteomic measurements to distinguish between the two isoforms of LC3 - LC3I and LC3II, thereby hindering interpretations about the role of the isoforms.

Strikingly, our analysis revealed that when autophagy is impaired upon lack of Atg16l1, nearly three hundred proteins display increased or decreased abundance, encompassing at least 18 functional processes (Figure 3). This reinforces the ubiquitous role autophagy plays in many cellular functions. Among the altered proteins, several had previously been associated with pathologies affecting Paneth cells, such as ANXA1 and FGA, which were previously reported to be altered in inflamed mucosal tissue or epithelial cells from Crohn’s disease patients (Barceló-Batllori et al., 2002; Iskandar and Ciorba, 2012; Meuwis et al., 2007). We observed that when autophagy is impaired in Paneth cells, most of the differentially abundant proteins are present in greater abundance than in normal Paneth cells, thus suggesting that degradation through autophagy plays a key role in maintaining the intracellular concentrations of these proteins.

By integrating multiple levels of information such as protein-protein interactions, inhibitory/stimulatory relationships between the altered proteins and autophagy and functional processes associated with the altered proteins, our computational pipeline enhances our
understanding about the underlying mechanisms involved in autophagy-mediated degradation. Furthermore, by bringing together different levels of information, our approach helps explain the mechanistic underpinnings between the processes corresponding to the proteins with altered abundances and autophagy. Capturing such process level dependencies on cellular autophagy and their modulation would be difficult by using singular levels of information in isolation. For example, Zhang et al measured the proteome level changes in primary human fibroblasts which were impaired in autophagy as a means to explain the purported dependency of protein degradation on macroautophagy (Zhang et al., 2016). Patella et al identified proteomic alterations under conditions of autophagy blockage in endothelial cells to explain the potential role of autophagy in maintaining endothelial permeability (Patella et al., 2016). Similarly, various other studies have profiled the global proteomic changes in response to artificial impairment of autophagy by knocking out critical autophagy genes (Avin-Wittenberg et al., 2015; Mathew et al., 2014). Studies combining different -omic readouts have also been performed in various contexts to understand the role of autophagy in various processes and phenotypes (Chen et al., 2017; Kramer et al., 2017; Masclaux-Daubresse et al., 2014; Stingele et al., 2012). However, the studies aforementioned do not provide an explanation as to how the proteome level alterations are indeed dependent on autophagy from a mechanistic point of view. In this study, using networks and integration of heterogeneous datasets, we provide information on new mechanisms by which several cellular processes such as exocytosis, DNA repair and apoptosis are dependent on autophagy.

Upon microbial invasion or inflammation-mediated cellular damage, cells respond by activating apoptotic cell death. In general, autophagy and apoptosis are negatively correlated under most homeostatic conditions (Mariño et al., 2014), although altered cellular settings can drive autophagy to lead to programmed cell death. The interactions between the two processes are highly complex (Gump and Thorburn, 2011; Mariño et al., 2014). Interestingly, our observations showed a positive correlation between autophagy and apoptosis (apoptosis being inhibited in the autophagy-impaired organoids; Supplementary figure 1). This phenomenon is not unique and has been observed in the case of acute myeloid leukemia, where it was reported that autophagy can excessively degrade the cytoplasm, and lead to apoptosis (Wu et al., 2017). In this study, the observed down-regulation of apoptosis could prevent the perturbed Paneth cells from sacrificing themselves, which would then be compensated for by outcomes such as upregulation of DNA damage repair functions (Basu and Krishnamurthy, 2010; Nowsheen and Yang, 2012) (Supplementary figure 2). However, further experiments are needed to confirm the assumption about the role of DNA repair and apoptosis in Paneth cells and how the disruption of these processes could contribute to CD pathogenesis upon impaired autophagy.

ATG16L1 is not only known to be required for the normal functioning of autophagy but also has physiological relevance due to its mutated status in many CD patients. The intestinal epithelium in CD patients is characterized by a prolonged period of stress as a result of chronic inflammation and malfunction of antimicrobial innate defences. This is reflected in particular by
the alteration of exocytosis of antimicrobials as well as the manipulation of the genetic/epigenetic machinery and organelles by invading pathogens and various other causative agents (Fofanova et al., 2016; Sartor, 2006). In this study, the downregulation of exocytosis that we observed in autophagy-impaired organoids was illustrated by the lysozyme accumulation within organoid cells and the consequential alteration of lysozyme secretion into the extracellular milieu. These results support our computational analysis and concur with the previously reported autophagy dependency of exocytosis (Brooks et al., 2012; Cadwell et al., 2008; Cadwell et al., 2009a; Cadwell et al., 2009b; Gassler, 2017; Tschurtschenthaler and Adolph, 2018) (Figures 4, 6). Impaired autophagy can therefore have dramatic consequences on innate defense mechanisms against microbial invasion of the gut epithelium by de-regulating the protein degradation of key exocytotic proteins.

The Paneth cell-enriched organoids we derived from the Atg16l1ΔIEC mouse model could be perceived as a biased representation of the role autophagy impairment has in CD, as it does not consider other intrinsic factors such as mutations in non-autophagy related genes which have been shown to contribute to CD pathogenesis. Indeed, proteins like PARP-2, IFI35, S100A12, CRP and S100A8 have been shown to contribute to IBD pathogenesis (Cheluuvappa et al., 2014; de Jong et al., 2006; Jijon et al., 2000; Leach and Day, 2006; Vermeire et al., 2006). These biomarkers are indicative of an inflammatory state, some of them are mostly detected in the serum, but looking for indicators from faecal samples predict more accurately the state of CD. For example, inhibition of PARP dampens inflammation associated with colitis (Jijon et al., 2000), and elevated levels of CRP, S100A8 for example have been associated with Crohn’s disease. Yet in our study, these proteins were not found differentially abundant when autophagy was impaired, or were fluctuating in the opposite direction. These discrepancies could reflect the differences in the sample type as these studies did not focus on Paneth cells only. Complementary experiments and predictions would nonetheless help highlight some of the aspects of the molecular regulatory mechanisms that contribute to CD pathogenesis when autophagy is altered.

Our integrative analysis not only captured already known phenomena, namely the autophagy dependency of exocytotic functions associated with granule release, but also highlighted that the degradation regulation of many differentially abundant proteins, including those of exocytosis proteins occurring in an autophagy-dependent manner. The presented study therefore extended the list of proteins for which the degradation rate was already known to be regulated by autophagy. More interestingly, our analysis revealed additional cellular processes which could mediate the effects of autophagy impairment on Paneth cell functions. Taken together and using a mouse model where autophagy is impaired, and organoids to capture Paneth cells, we identified various cellular processes which are dependent on autophagy and the failure of which further explains and contributes to the pathogenesis of CD.
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Competing interests

None

Author contributions

EJ and ZM performed the experimental work with the organoids. LG and PS carried out the bioinformatic analysis. DD, JB, MJ, UM created and tested the mice model, SA carried out the proteomics study. IH performed the lysozyme assays. AW, PP, SC, TW, TK designed and supervised the experiments. EJ, LG, PS, IH and TK wrote the manuscript.

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Table 1

| Category                                      | Number of proteins |
|-----------------------------------------------|--------------------|
| Number of detected mice proteins              | 3832               |
| Number of differentially abundant mice proteins (DAPs) | 278                |
| Number of human orthologues of mice DAPs      | 284                |
| Number of autophagy targeted proteins         | 116                |
| LC3 targets (experimental, predicted, both)   | 49 (14, 19, 16)    |
| Atg16l1 targets (experimental, predicted, both) | 81 (0, 81, 0)     |
| p62 targets (experimental, predicted, both)   | 8 (6, 2, 0)        |

Table 2

| Altered functional category | Aggregated trend* |
|----------------------------|-------------------|
| Apoptosis                  | Down              |
| Cell cycle                 | Up + Down         |
| Chromatin organisation     | Up + Down         |
| Cytoskeleton and cell adhesion structures | Up + Down |
| DNA repair                 | Up                |
| Endocytosis                | Up                |
| Exocytosis                 | Down              |
| Gene expression            | Up + Down         |
| Immune response            | Up                |
| Metabolism                           | Up + Down |
|-------------------------------------|-----------|
| Mitochondrial organisation          | Up + Down |
| Post-translational modification     | Up        |
| Proteasome-ubiquitin system         | Up        |
| Signal transduction                 | Up + Down |
| Transport                           | Up + Down |

**Figure and table legends**

**Figure 1.** A schematic representation of the workflow to determine the functional effects of the autophagy impairment in $Atg16l1^{ΔIEC}$ in Paneth cell organoids.

**Figure 2.** $Atg16l1^{ΔIEC}$ organoids do contain the same intestinal epithelial cell types as WT organoids but lack $Atg16l1$ and LC3II both in the transcriptional and protein levels. A) Brightfield image of control (WT) and $Atg16l1^{ΔIEC}$ (KO) organoids after 7-days of growth. Magnification x10, scale bar 100µm. B) Cell specific gene expression in control (WT) and $Atg16l1^{ΔIEC}$ (KO) organoids was identical. Primers were specific for Lgr5, Villin, Chromogranin A (CHGA), Mucin 2 (Muc2), CD24 and β-actin. C) Western blots using anti-ATG16L1 and LC3 antibodies detected $Atg16l1$ and LC3II in control organoids but $Atg16l1^{ΔIEC}$ (KO) organoids were deficient in $Atg16l1$ and LC3II.

**Figure 3.** Percentage of higher and lower abundance proteins in different cellular functions. Proteins with higher abundance are marked with yellow and lower abundance proteins with blue background. “Other” category contains all of the proteins which have not fit to other functional groups.

**Figure 4.** Overlap between the proteins with altered abundances in $Atg16l1^{ΔIEC}$ Paneth cell organoids. Restricted to proteins potentially targeted by each of the three selective autophagy mediating proteins p62, LC3 and $Atg16l1$ under normal circumstances without any autophagy defects.
Figure 5. Potential autophagy dependency of the altered functional processes inferred from the proteomic profile of the Atg16l11ΔIEC Paneth cell organoids using our integrated approach. The autophagy dependency of the proteins with altered abundances (orange ellipsoids for proteins with increased abundance, blue ellipsoids for proteins with decreased abundance) are represented highlighting the effect of proteins in the processes (purple edge for activation and green edge for inhibition) as well. The aggregated trends of the altered functional processes as determined by integrative approach (see Methods section) are indicated (yellow rectangles for up-regulated functional processes; red rectangles for functional processes; white rectangles for functional processes, which are both up- and down-regulated). Proteins outside from the circle are grouped to the process which they are involved in to increase the clarity of the figure. The figure was created using Cytoscape.

Figure 6. Atg16l11ΔIEC deficiency in Paneth cell organoids and its impact on exocytotic proteins. (A) Proteins belonging to the functional category of exocytosis with lower abundance reflecting the impact of autophagy impairment on Paneth cell functions such as granule processing and release through exocytosis. (B) Western blot analysis for Paneth cell-derived lysozyme on cellular extracts from WT or Atg16l11ΔIEC organoids. (C) Lysozyme activity measured in culture medium of 2D WT and Atg16l11ΔIEC organoids as reporter of Paneth cell exocytosis. (D) Proteins with exocytotic functions that are found to be altered in their abundances and dependent on autophagy. Blue and orange bars correspond to proteins with decreased and increased abundances, respectively.

Table 1. Effect of the Atg16LIΔIEC mutation on the alteration of protein abundances in Paneth cell enriched organoids.

Table 2. The aggregated trends of the functional categories which correspond both to the proteins with altered abundances and are potentially targeted by selective autophagy in response to the Atg16LIΔIEC mutation in Paneth cell enriched organoids. * Aggregated trends were determined as described in the methods section.

Supplementary figure 1. Alterations describing the modulation of apoptosis in response to the Atg16LIΔIEC mutation.

Supplementary figure 2. Alterations describing the modulation of DNA repair in response to the Atg16LIΔIEC mutation.

Supplementary table 1. Primers used in the study.
Supplementary table 2. Discarded biological process terms not related to intestinal functions.

Supplementary table 3. Post-processed results from the measurements comparing the proteomic profiles of $Atg16L1^{ΔIEC}$ and WT Paneth cell enriched organoids.

Supplementary table 4. Human orthologs of the mouse proteins with altered abundances.

Supplementary table 5. Evidence suggesting the targeting of proteins with altered abundances by the autophagy proteins (p62, LC3, ATG16L1).

Supplementary table 6. Summary of abundance changes of proteins potentially targeted by autophagy and their assigned gene ontology terms.

Supplementary table 7. Functional evidence indicating the stimulatory/inhibitory effect of the proteins with altered abundances on their corresponding processes.

Supplementary table 8. The aggregated trends of the gene ontology terms corresponding to the altered proteins in Paneth cells upon the deletion of $Atg16l1$.

Supplementary table 9. Functional processes and their aggregated trends in Paneth cells upon the deletion of $Atg16l1$.

Supplementary table 10. Exocytotic proteins found to be altered in their abundances.

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**In vivo** tissue processing, Sequential mechanical disruption

*In vitro* culture

Organoid processing
Proteomic extraction and analysis

Differentially abundant proteins

Human orthologues of mice proteins

**Question:** How does the malfunction of autophagy influence the cell functions?

Filtering to potential autophagy targeted proteins

**Autophagy Regulatory Network (ARN)**

iLIR database

Target motif recognition

Domain-domain interaction prediction

Targets of ATG16L1, p62, and LC3

LC3 targets

p62 targets

ATG16L1 targets

Proteins targeted by autophagy

Gene Ontology of proteins

Functional processes affected by autophagy

Visualization and analysis

**Autophagy dependent functional landscape in Paneth cells**
Figure 5

[Diagram showing the regulation of various biological processes and proteins, with color coding for up-regulated, down-regulated, and activation or inhibition.]
Figure 6

A. Diagram showing interactions between autophagy proteins and exocytosis. The proteins include p62, ATG16L1, LC3, ANXA1, CDK1, GOKI, TMED3, GORASP1, SRP14, and GABARAPL2. The diagram uses symbols to indicate more abundant, less abundant, down-regulated, inhibition, and activation.

B. Bar graph comparing WT and Atg16l1\textsuperscript{ΔIEC} for densitometry. The WT and Atg16l1\textsuperscript{ΔIEC} have significantly different densitometry values.

C. Bar graph comparing WT and Atg16l1\textsuperscript{ΔIEC} for activity (U/ml). The graphs show no significant difference between WT and Atg16l1\textsuperscript{ΔIEC}.

D. Bar graph showing fold change (Log2) for inhibitors and activators, indicating the change in expression levels.