mTOR controls neurotoxic lysosome exocytosis in inflammatory reactive astrocytes

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ABSTRACT

Inflammatory reactive astrocytes lose homeostatic functions and can be neurotoxic, potentially contributing to neurodegenerative diseases. However, the underlying cell biological mechanisms are not fully understood. Here, we demonstrate that lysosomes are remodeled and alkalinized in inflammatory reactive astrocytes, and that lysosome exocytosis drives astrocyte-mediated neurotoxicity. CRISPRi screens uncover mTOR as a regulator of neurotoxic lysosome exocytosis. These results pinpoint lysosome remodeling and exocytosis in inflammatory reactive astrocytes as a potential therapeutic target.
Reactive astrocytes are prevalent in many neurodegenerative diseases. Mounting evidence suggests that inflammatory reactive astrocytes contribute to disease progression by losing their normal homeostatic functions and secreting factors that are neurotoxic under certain conditions. However, the mechanisms that drive these functional changes in inflammatory reactive astrocytes remain unknown.

To identify the cellular pathways responsible for the functional changes that accompany inflammatory astrocyte reactivity, we re-analyzed our previously collected RNA-sequencing dataset of hiPSC-derived astrocytes (iAstrocytes) treated with the microglia-derived cytokines IL1α, TNF, and C1q (hereafter referred to as ITC) which induce inflammatory reactivity in vitro. As expected, many genes and pathways previously associated with inflammatory astrocyte reactivity were upregulated after ITC treatment, including SERPINA3 and C3, along with cellular pathways involved in NF-κB signaling, antigen presentation, and cellular response to cytokines (Supplementary Table 1). Interestingly, many genes encoding lysosome-localized proteins were downregulated after ITC treatment (Fig. 1b-c). These included many subunits of the vacuolar ATPase (V-ATPase) complex and lysosomal degradative enzymes, which coordinate the proper acidification of lysosomes and hydrolysis of autophagic substrates (Fig. 1c, Supplementary Table 1). We validated that many of these genes were also downregulated at the protein level in ITC-treated iAstrocytes via proteomics (Extended Data Fig. 1a,b, Supplementary Table 2). Further analysis of published RNA-sequencing datasets of hiPSC-derived reactive astrocytes confirmed that this lysosomal signature was broadly conserved across models of inflammatory reactivity, including those that use different differentiation protocols (Barbar et al.) and cytokine treatments (Perriot et al.). We also noted that a small subset of genes encoding lysosome-localized proteins involved in antigen presentation and inflammation (e.g. CTSS, HLA family) were strongly upregulated across models (Fig. 1c).

To determine if this lysosome signature is associated with changes in lysosome levels and function in inflammatory reactive astrocytes, we quantified levels of the lysosomal marker LAMP2 by immunoblotting, and found no significant difference after ITC treatment (Fig. 1d). Similarly, quantification of the lysosomal markers LAMP1 and LAMP2 by immunofluorescence did not show significant changes in response to ITC (Fig. 1e). However, there was a dramatic decrease in the fluorescent signal of LysoTracker Green, a cell-permeable, pH-dependent dye that accumulates in lysosomes (Fig. 1e). We hypothesized that ITC-treated iAstrocytes were deficient in lysosomal acidification, thereby preventing LysoTracker from accumulating in lysosomes. Using the genetically encoded lysosome pH sensor FIRE-pHLy, we found that the lysosomes of ITC-treated iAstrocytes were indeed less acidic than those of control iAstrocytes (Fig. 1f). Furthermore, ITC-treated iAstrocytes accumulated puncta of LC3B, an autophagosome marker degraded in the lysosome by acid-activated hydrolases (Fig. 1e). Using a GFP-LC3-RFP-LC3AG reporter, we verified that this accumulation in ITC-treated iAstrocytes reflected an impairment in degradative autophagic flux rather than an increase in the steady-state level of autophagic components (Fig. 1g).
To gain a more detailed understanding of lysosome remodeling in inflammatory reactive astrocytes, we investigated the lysosomal proteome by immunoprecipitating a LAMP1-3xHA lysosome tag (lyso-IP)\textsuperscript{11} introduced into iAstrocytes via lentiviral transduction (Fig. 1h, Extended Data Fig. 1c,d). In agreement with our RNA-sequencing and whole-cell proteomics data, we found that many proteins required for lysosomal degradation, including V-ATPase subunits and lysosomal hydrolases, were less abundant in the lyso-IP fraction of ITC-treated iAstrocytes compared to vehicle-treated iAstrocytes (Fig. 1i, Supplementary Table 2). We also found that CST3, a potent inhibitor of lysosomal proteinases, was highly enriched in the lyso-IP fraction of ITC-treated iAstrocytes. In addition to the above signature of decreased degradative capacity, we found that several mediators of inflammatory signaling – CCL2, TNFAIP3, and IL32 – were more enriched in the lyso-IP fraction compared to whole-cell proteomics in ITC-treated vs. vehicle-treated iAstrocytes (Fig. 1j, Supplementary Table 2). Given that IL32 has previously been reported to undergo likely lysosome-mediated secretion\textsuperscript{12} and that many proteins traditionally involved in exocytosis (eg. RAB11A, RAB27A, RAB3A) were enriched in the lyso-IP fraction upon ITC treatment (Fig. 1i,j, Supplementary Table 2), we hypothesized that inflammatory reactive astrocytes execute lysosome exocytosis, potentially as a mechanism to counteract the accumulation of autophagic substrates and/or as part of a non-canonical secretory pathway.

To test this hypothesis, we used TIRF microscopy to visualize lysosomes in iAstrocytes loaded with LysoTracker Green and expressing a lysosome membrane-targeted mCherry construct (Fig. 2a). Indeed, likely lysosome exocytosis events were observed as the trafficking of mCherry+/LysoTracker+ vesicles to the plasma membrane followed by an uncoupling of the mCherry and LysoTracker signals upon contact with the plasma membrane, after which the LysoTracker signal rapidly diffused, while the mCherry signal remained at the cell surface, suggesting the ejection of lysosomal contents (Fig. 2a-b, Supplemental Movies 1,2).

As an alternative and scalable approach to quantify lysosome exocytosis, we adopted a published flow cytometry-based protocol\textsuperscript{13,14} in which we stained non-permeabilized iAstrocytes with a fluorophore-conjugated antibody targeting a luminal epitope of LAMP1 that is exposed to the extracellular environment upon lysosome exocytosis (Extended Data Fig. 2a). With this approach, we found that ITC-treated iAstrocytes had almost twice the amount of cell-surface LAMP1 relative to vehicle-treated controls, despite equivalent levels of total LAMP1 protein between the two conditions (Fig. 2c,d). To test our hypothesis that the increased lysosome exocytosis in ITC-treated iAstrocytes was related to their deficit in lysosomal acidification, we treated iAstrocytes with bafilomycin A, a small molecule inhibitor of the V-ATPase complex. We observed a dose-dependent increase in lysosome exocytosis and concomitant decrease in lysosome acidification with increasing concentration of bafilomycin A (Fig. 2e). Together, these results suggest that lysosome remodeling involving decreased acidification and degradative capacity is associated with lysosome exocytosis in inflammatory reactive astrocytes.

Next, we turned our attention to understanding if lysosome exocytosis is involved in the known deleterious functions of inflammatory reactive astrocytes. Given that lysosome exocytosis contributes to the astrocyte secretome, we reasoned that lysosome exocytosis may play a role in the neurotoxicity of inflammatory reactive astrocytes. To test this hypothesis, we transferred astrocyte conditioned media (ACM) from vehicle-treated and ITC-treated iAstrocytes to hiPSC-
derived neurons\(^{15}\) (iNeurons), and subsequently assayed the levels of both lysosome exocytosis (measured by cell-surface LAMP1 in the astrocytes) and neurotoxicity (measured using caspase 3/7 activity reporter dye in the ACM-receiving iNeurons) (Fig. 2f). Consistent with previous reports\(^3,\ 4\), the ITC-induced increase in cell-surface LAMP1 was mirrored by a strong increase in apoptosis in iNeurons receiving ACM derived from ITC-treated astrocytes (Fig. 2g,h). When we treated astrocytes with vacuolin-1, a small molecule inhibitor of lysosome exocytosis, there was a dose-dependent decrease in both lysosome exocytosis in astrocytes and apoptosis in iNeurons (Fig. 2g.). Importantly, unconditioned media supplemented with ITC or vacuolin-1 did not substantially affect apoptosis in iNeurons, ruling out the possibility that residual amounts of these compounds were causing the apoptosis effects in ACM experiments (Extended Data Fig. 2b). In addition, vacuolin-1 did not induce excess cell death in iAstrocytes, ruling out the possibility that decreased astrocyte viability could explain the blunted neurotoxic activity (Extended Data Fig. 2c). Furthermore, to confirm that vacuolin-1 specifically inhibited lysosome exocytosis in astrocytes and did not perturb general vesicular trafficking to the plasma membrane, we co-stained astrocytes with an antibody against TFRC, a constitutively expressed plasma membrane receptor. We found no substantial effect of vacuolin-1 on cell-surface TFRC levels (Extended Data Fig. 2d). Lastly, we knocked down of genes reported to mediate lysosome exocytosis (\(\text{SYT11}\) and \(\text{VAMP7}\))\(^{16,\ 17}\). While \(\text{SYT11}\) and \(\text{VAMP7}\) knockdown resulted in only a small decrease in cell-surface LAMP1, this was sufficient to reduce ITC-induced neurotoxicity by \(~50\%) relative to control iAstrocytes (Figure 2i,j).

To test if increasing lysosome exocytosis would also increase astrocyte-mediated neurotoxicity, we treated iAstrocytes receiving vehicle or ITC with a low dose of bafilomycin A (10 nM) to induce lysosome alkalinization and lysosome exocytosis, and transferred the conditioned media to iNeurons as before. Bafilomycin A treatment resulted in a robust increase in cell-surface LAMP1 in both astrocytes treated with ITC or vehicle (Fig. 2k). Astrocytes treated with ITC and bafilomycin A were approximately twice as neurotoxic compared to astrocytes treated with ITC and DMSO (Fig. 2l). Interestingly, although bafilomycin A itself induced roughly the same level of lysosome exocytosis in vehicle-treated iAstrocytes as ITC-treated iAstrocytes, it did not cause a corresponding increase in neurotoxicity for iAstrocytes that were not treated with ITC, suggesting that ITC treatment alters the content of lysosomes in a way that increases neurotoxicity of lysosomal exocytosis. Neither ITC nor bafilomycin A (10 nM) induced apoptosis in iAstrocytes (Extended Data Fig. 2e) or iNeurons (Extended Data Fig. 2f). In summary, the above results suggest that lysosome exocytosis is a major mechanism underlying neurotoxicity of inflammatory reactive astrocytes.

To identify potential therapeutic targets to inhibit the lysosome alkalinization-exocytosis-neurotoxicity axis in inflammatory reactive astrocytes, we performed targeted CRISPRi screens in iAstrocytes (Fig. 3a). Briefly, we transduced iAstrocytes expressing CRISPRi machinery\(^5\) with a lentiviral sgRNA library targeting the “druggable genome”\(^{18}\). After treating with ITC or vehicle control, we stained iAstrocytes with either LysoTracker or a fluorophore-conjugated LAMP1 antibody to monitor lysosomal alkalinization and exocytosis, respectively, and used fluorescence-activated cell sorting (FACS) to collect sgRNA-transduced astrocytes with low or high levels of LysoTracker or LAMP1. We then conducted next-generation sequencing to identify sgRNAs that were over- or under-represented within their respective cell population relative to non-targeting control (NTC) sgRNAs (see Methods). Based on this analysis, we
identified candidate regulators (FDR < 0.05, see Methods) of lysosomal pH (and also potentially mass; LysoTracker screen) or lysosome exocytosis (LAMP1 screen). Complete screen results are provided in Supplementary Table 3.

In agreement with our hypothesis that lysosomal alkalinization drives lysosome exocytosis, we found a strong inverse correlation between the LysoTracker and LAMP1 phenotypes for many hit genes (Fig. 3b, Extended Data Fig. 3). Across all screens, genes involved in mTOR signaling were highly enriched among hit genes, including MTOR itself as one of our strongest hits (Fig. 3c,d). To validate our screening results, we transduced iAstrocytes with an sgRNA targeting MTOR or a non-targeting sgRNA. As expected, MTOR knockdown dramatically increased LysoTracker staining (Fig. 3e). Importantly, MTOR knockdown specifically decreased lysosome exocytosis in ITC-treated but not vehicle-treated iAstrocytes (Fig. 3f), suggesting that mTOR activity may control lysosome exocytosis specifically in inflammatory reactive astrocytes. Consistent with our finding that suppressing lysosome exocytosis prevents inflammatory reactive astrocyte neurotoxicity, MTOR knockdown in iAstrocytes also rescued the neurotoxicity of ITC-treated iAstrocytes (Fig. 3g).

mTOR is a member of both mTORC1 and mTORC2 complexes, which regulate their downstream targets via phosphorylation. Given that MTOR knockdown specifically blocked the ITC-induced increase of cell-surface LAMP1, we hypothesized that there would be increased mTOR activity in inflammatory reactive astrocytes. Immunoblots revealed that there were no significant changes in the levels of mTOR itself, but a clear trend towards increased phosphorylation of mTORC1 targets (p-S6 S235/236 and p-ULK1 S757) and to a lesser extent mTORC2 targets (p-Akt S473) in ITC-treated iAstrocytes (Fig. 3h-l).

To validate that mTOR activity is responsible for lysosomal changes in inflammatory reactive astrocytes using an orthogonal approach, we treated iAstrocytes with several mTOR inhibitors, including the pan-mTOR inhibitor PP242, in combination with vehicle and ITC. We found that PP242 suppressed mTOR activity (Fig. 3h), acidified lysosomes (Fig. 3m), decreased cell-surface LAMP1 (Fig. 3n), and blocked ITC-induced neurotoxicity (Fig. 3o) in a dose-dependent manner.

In summary, we uncovered lysosome remodeling – consisting of increased lysosome pH, altered lysosomal content, and increased lysosome exocytosis – as a novel functional phenotype of inflammatory reactive astrocytes and a key pathological process mediating their neurotoxic activity. Furthermore, we used pooled CRISPRi screens to identify mTOR as a central upstream regulator of the lysosome alkalinization-exocytosis-neurotoxicity axis in inflammatory reactive astrocytes (Fig. 3p).

Our model is consistent with recent findings by other groups: First, in vivo reports that Lamp1 is the most strongly upregulated astrocyte cell-surface protein in a mouse model of multiple sclerosis (MS). Second, the fact that mTOR inhibition blunts disease progression in mouse models of amyotrophic lateral sclerosis and MS that are driven by inflammatory reactive astrocytes. Last, in vitro data suggesting a role for astrocyte lysosome exocytosis in HIV-Tat-mediated neurotoxicity. Mechanistically, we have not yet resolved if the relationship between lysosome exocytosis and astrocyte-mediated neurotoxicity stems from direct toxicity of astrocyte
lyosome contents, or whether these contents contribute to autocrine-paracrine signaling in astrocytes to induce neurotoxicity through an indirect mechanism (Fig. 3p).

We hope that future work will bring clarity to how lysosome remodeling in astrocytes influences neurodegenerative diseases, particularly with respect to the disease-, region-, and context-specific functions of reactive astrocytes in vivo. Given our in vitro results, we envision that these studies could inform novel therapeutic approaches.

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AUTHOR CONTRIBUTIONS

B.R., K.L., and M.K. conceptualized and led the overall project, and wrote the manuscript with input from all co-authors. B.R. performed the majority of experiments and data analysis. K.L. developed the iAstrocyte platform, performed preliminary studies, and conducted the RNA-seq experiments. In consultation with J.E., F.M aided with sample preparation for whole-cell and lysosome-specific proteomics experiments, and performed mass spectrometry and preliminary computational analysis. I.V.L.R. performed sample prep for the cell-surface LAMP1 CRISPRi screens. K.A.H. guided TIRF experiments. M.L. and S.B aided with design and sample preparation for whole-cell and lysosome-specific proteomics experiments, respectively. M.Y.C. and A.W.K provided FIRE-pHLy reagents and guidance on lysosome pH experiments. S.F conducted LAMP2 immunofluorescence experiments and data analysis.

COMPETING INTERESTS STATEMENT

A.W.K is a member of the Scientific Advisory Board for Nine Square Therapeutics, Inc. J.E. is an advisor to Seer, Inc. M. K. has filed a patent application related to CRISPRi and CRISPRa screening (PCT/US15/40449) and serves on the Scientific Advisory Board of Engine Biosciences, Casma Therapeutics, and Cajal Neuroscience, and is an advisor to Modulo Bio and Recursion Therapeutics. None of the other authors declare competing interests.

DATA AVAILABILITY STATEMENT
Bulk RNA-seq data of hiPSC-derived astrocytes analyzed in this study were generated previously\(^5\) and are available on the Gene Expression Omnibus (GEO) under accession code GSE182307. Mass spectrometry results are in the process of submission to PRIDE.

**CODE AVAILABILITY STATEMENT**

The code used to analyze the data presented in this study will be shared upon request.

**METHODS**

**hiPSC culture**

Human iPSCs (male WTC11 background\(^24\)) were cultured in Essential 8 (E8) Medium (ThermoFisher Scientific cat. no. A1517001) on BioLite Cell Culture Treated Dishes (ThermoFisher Scientific) coated with Growth Factor Reduced, Phenol Red-Free, LDEV-Free Matrigel Basement Membrane Matrix (Corning cat. no. 356231) diluted 1:100 in DMEM/F12 (ThermoFisher Scientific cat. no. 11330032). Essential 8 Medium was replaced daily. When hiPSC colonies demonstrated mature morphology, the hiPSCs were either clump passaged with EDTA for routine maintenance or dissociated to a near single-cell suspension with Accutase Cell Dissociation Reagent (ThermoFisher Scientific cat. no. A11105-01) for applications requiring cell counting. For clump passaging with EDTA, hiPSCs were washed with Dulbecco’s phosphate buffered saline (DPBS; Milipore Sigma cat. no. D8537) and then incubated with Versene (ThermoFisher Scientific cat. no. 15040066) for 5-7 min at room temperature; the Versene solution was then aspirated and replaced with E8 + 10 nM Y-27632 dihydrochloride ROCK inhibitor (Tocris cat. no. 125410); hiPSC colonies were then gently detached mechanically using a cell scraper, resuspended gently, and passaged at 1:10-1:30 dilution in E8 + Y-27632, with Y-27632 removed the next day. For near single-cell dissociation, hiPSCs were washed with DPBS, incubated with Accutase for 5-10 min at 37 °C, and then gently triturated with a P1000 pipette tip; the cell suspension was then diluted with PBS, collected into conical tubes, and spun down at 300 g for 3 min; hiPSCs were then resuspended in E8 + Y-27632, counted, and plated onto Matrigel-coated plates at the desired density in E8 + Y-27632; Y-27632 would be maintained until the hiPSC colonies reached the appropriate size (> ~40 cells). Studies with hiPSCs at UCSF were approved by the The Human Gamete, Embryo and Stem Cell Research (GESCR) Committee. Informed consent was obtained from the human subject when the WTC11 line was originally derived\(^24\).

**Generation of iAstrocytes**

iAstrocytes were generated as detailed in Leng *et al.*\(^5\). Briefly, we generated hiPSCs (male WTC11 background) used for iAstrocyte differentiation by inserting a CRIPSRi machinery cassette (pC13N-dCas9-BFP-KRAB, Addgene plasmid no. 127968) into the CLYBL safe-harbor locus and a doxycycline-inducible *NFIA* and *SOX9* cDNA cassette into the AAVS1 safe-harbor
locus using TALEN-based editing. After recovering a monoclonal CRISPRi-NFIA-SOX9 line, we differentiated hiPSCs to the neuroectoderm lineage via an embryoid body (EB)-based neural induction protocol. After dissociating plating EB-derived cells, we selected and expanded those forming neural rosettes. We used fluorescence activated cell sorting (FACS) to enrich for pure CD133+/CD271- neural progenitor cell (NPC) populations for expansion.

For iAstrocyte differentiation, these NPCs were re-plated on 1:200 Matrigel-coated dishes and cultured with ScienCell Astrocyte Media (ScienCell Research Laboratories cat. no. 1801) + 2 μg/mL doxycycline (Millipore Sigma cat. no. D9891) to initiate iAstrocyte differentiation. A full media change was completed every other day (maintaining doxycycline at 2 μg/mL) until NPCs reached confluency, at which time they were dissociated with Accutase and split 1:10 onto new Matrigel coated dishes (or cryopreserved) for continued differentiation and expansion. Expansion was continued until day 20 (d20) of differentiation, yielding iAstrocytes. See Leng et al.5 for full details of iAstrocyte hiPSC generation and differentiation protocols.

**Induction of inflammatory reactivity in iAstrocytes**

For all iAstrocyte experiments in this study, d20 iAstrocytes were plated at 20,000 cells/cm² on 1:200 Matrigel-coated BioLite Cell Culture Treated Dishes (ThermoFisher Scientific) in ScienCell Astrocyte Media (without doxycycline) on d0. Full media changes were completed on d1, d3, and d5. On d5, media was supplemented with IL-1α (3 ng/mL; Peprotech cat. no. AF-200-01A), TNF (30 ng/mL; Peprotech cat. no. AF-300-01A), and C1q (400 ng/mL; Complement Technology cat. no. A099) to induce inflammatory reactivity according to Liddelow et al.3. All experiments were conducted 24 hours after cytokine treatment (d6).

**CRISPRi-mediated gene knockdown using lentiviral transduction of individual sgRNAs**

To knock down genes with individual sgRNAs, we cloned CRISPRi sgRNAs into pMK133425 (Addgene cat. no. 127965), as previously described26. We generated lentivirus of the resulting constructs by co-transfecting them with 3rd generation lentiviral packaging plasmids with TransIT-Lenti Transfection Reagent (Mirus cat. no. MIR6606) according to the manufacturer’s protocol. The next day, lentivirus was precipitated using Lentivirus Precipitation Solution (ALSTEM cat. no. VC150) according to the manufacturer’s protocol, resuspended in DPBS, and aliquoted and stored at -80 °C. iAstrocytes were transduced with lentivirus at the time of plating. When necessary, the functional titer of the lentivirus was determined by measuring BFP+ cells 48 hours after transducing iAstrocytes with a serial dilution of lentivirus. The protospacer sequences of the sgRNAs used in this study were designed based on our next-generation CRISPRi algorithm18 and are listed below:

NTC – GAGACGAGGACATGTGTAGC
MTOR – GGGACAGCGGGGAAGGCGGG
SYT11 – GGAAGGGGGAAGGTATCCAG
VAMP7 – GAGGGACGCGGGTCAGTGCA
Generation and lentiviral transduction of reporter constructs

To subclone the original FIRE-pHLy construct (pLBR10, Ubc:FIRE-pHLy)\(^9\) to make pLBR11 (GfaABC1D:FIRE-pHLy), we excised the FIRE-pHLy cassette from pLBR10 and linearized the backbone vector (pKL20; pHIV gfaABC1D:mRuby:WPRE) with BamH1-HF (New England BioLabs, cat. no. R3136) and ClaI (New England BioLabs cat. no. R0197). We then ran these reactions on a 1% agarose gel, gel extracted the relevant fragments, and ligated them together with T4 DNA Ligase (New England BioLabs cat. no. M0202) per manufacturer’s protocol to generate pLBR11.

Both pLBR11 (all FIRE-pHLy experiments), and GFP-LC3-RFP-LC3\(^{Δ}\) (reporter cassette from Addgene plasmid #84572)\(^10\) reporters were transduced into iAstrocytes with lentivirus at low multiplicity of infection (MOI) during plating to prevent high expression of exogenous lysosomal proteins.

Immunofluorescence imaging

For immunofluorescence imaging experiments, iAstrocytes plated in Greiner μClear 96-well plates (Greiner Bio-One cat. no. 655087) were fixed with 4% paraformaldehyde (diluted from a 16% solution; Electron Microscopy Sciences cat. no. 15710) for 15 min at RT. After washing three times with DPBS, we blocked and permeabilized cells with DPBS (Milipore Sigma cat. no. D8537) + 3% BSA (Milipore Sigma cat. no. A9647) + 0.1% Triton X-100 (Millipore Sigma cat. no. X100) for 30 min at RT. Primary antibodies against LC3 (1:50; rabbit polyclonal; Cell Signaling Technology cat. no. 2775), LAMP1 (1:50; mouse monoclonal; Abcam cat. no. ab25630), or LAMP2 (1:500, Abcam cat. no. ab25631) were added to blocking buffer and incubated with iAstrocytes overnight at 4 °C. Afterwards, the samples were washed with DPBS three times, incubated with pre-adsorbed secondary antibodies (1:500 goat anti-mouse IgG Alexa Fluor 488, 1:500 goat anti-rabbit IgG Alexa Fluor 555; Abcam cat. no. ab150119 and ab150086) for 1 hour at RT, washed three times with DPBS, incubated with 1 μg/mL Hoechst 33342 (ThermoFisher Scientific cat. no. H3570) and then washed two additional times before imaging on an IN Cell Analyzer 6000, using a 60X 0.7 NA objective, 2x2 binning, 100-400 ms exposure, an aperture width of ~1 Airy unit, and 9-16 fields per well.

LysoTracker staining and imaging

iAstrocytes were washed with DPBS and incubated with a 50 nM solution of LysoTracker Green DND-26 (ThermoFisher Scientific cat. no. L7526) in DPBS for 5 minutes at 37°C, washed two times with DPBS, and processed for downstream experiments. For imaging (Fig. 1e), iAstrocytes were imaged at 37°C and 5% CO₂ using an IN Cell Analyzer 6000 as above. For flow cytometry (CRISPRi screens), iAstrocytes were dissociated with Accutase for 10 mins at 37°C after LysoTracker staining, and resuspended in FACS buffer (DPBS + 1% BSA + 2 mM EDTA (Milipore Sigma cat. no. 324506)).

TIRF imaging
iAstrocytes were plated at 20,000 cells/cm² in 35 mm imaging dishes (Ibidi, cat. no. 81156) and simultaneously transduced with pLBR11 (GfaABC1D:FIRE-pHLy). Prior to imaging, the conditioned media of iAstrocytes was collected. iAstrocytes were then stained with LysoTracker as detailed above and re-incubated with the collected conditioned media for the duration of imaging. TIRF was performed on a Nikon Ti-E inverted microscope equipped with a Nikon TIRF illuminator, Nikon motorized stage with piezo Z inset, Andor iXon Ultra DU897 camera, and run by NIS-Elements (v5.20.00 build 1423). iAstrocytes were imaged using an Apo TIRF 100x/1.49 oil (DIC N2/ 100X I) objective, 488nm and 561nm lasers (Agilent MLC400 monolithic laser combiner) and Sutter Lambda 10-B with ET525/50m and ET600/60m emission filters (Chroma), respectively, for 120 seconds with 100 ms exposure (per channel).

Measurement of apoptosis in iNeurons treated with astrocyte-conditioned media

hiPSC-derived neurons (iNeurons) were generated from WTC11 hiPSCs with a stably integrated dox-inducible NGN2 cassette (i3N iPSCs) as previously described15. Conditioned media was collected from iAstrocytes and spun down at 300 x g for 5 min to remove dead cells, supplemented with CellEvent Caspase 3/7 Detection Reagent (ThermoFisher Scientific, cat. no. C10423) to a final concentration of 2 μM, and transferred to day 17 iNeurons after removing the original iNeuron media. As an unconditioned media control, we incubated ScienCell Astrocyte Media in identical conditions (37 °C, 5% CO₂, cytokine and/or small-molecule compounds) as their respective experimental groups but without iAstrocytes. After a ~24 hour incubation period, iNeuron viability was assessed by imaging on an IN Cell Analyzer 6000, using a 20X 0.45 NA objective, 2x2 binning, environmental control set to 37 °C and 5% CO₂, an aperture width of ~1 Airy unit, 200 ms exposure, and 4-9 fields per well. The level of apoptosis in iNeurons was quantified relative to an unstained negative control with CellProfiler (see Data Analysis section).

Antibody staining for flow cytometry

To measure cell-surface protein levels (i.e. LAMP1, TFRC), iAstrocytes were dissociated with Accutase for 10 mins at 37 °C, diluted with ice-cold DPBS + 1% BSA, and spun down at 300 x g for 5 minutes at 4 °C. After aspirating the supernatant, all cells were incubated in 1:20 Fc block (BD Bioscience cat. no. 564220) diluted in DPBS + 3% BSA for 10 minutes on ice, followed by the addition of 1:20 (1:40 final) primary antibodies against LAMP1 (AF488- or BV605-conjugated; BioLegend cat. no. 328610 and 328634), and/or TFRC (FITC-conjugated, Biolegend, cat. no. 334104) for an additional 30 minute incubation on ice in the dark. iAstrocytes were then washed with ice-cold DPBS + 1% BSA, spun down at 300 x g for 5 mins at 4 °C, and resuspended in DPBS + 1% BSA for flow cytometry.

Pooled CRISPRi screening

CRISPRi screens were conducted as described in Leng et al.5. Both LysoTracker and LAMP1 screens were conducted with the H1 (i.e. “druggable genome”) sgRNA sub-library from our
next-generation CRISPRi library\textsuperscript{18}. We generated lentivirus for the pooled H1 sgRNA library as previously described\textsuperscript{25}. For each screen (i.e. LysoTracker or cell-surface LAMP1, vehicle or ITC), iAstrocytes were plated at 20,000 cells/cm\textsuperscript{2} onto 4 Matrigel-coated 15-cm dishes, transduced with the H1 lentiviral sgRNA library with >70\% transduction efficiency, treated with vehicle control or IL-1α+TNF+C1q for 24 hours, stained for LysoTracker or cell-surface LAMP1, and then sorted into LysoTracker high vs. low or cell-surface LAMP1 high vs. low (top and bottom 35\% of cells on fluorescence histogram) populations. iAstrocytes were sorted with a BD FACSARia Fusion cell sorter at 5,000-10,000 events per second, and then pelleted via centrifugation at 300 x g for 10 minutes for genomic DNA extraction. sgRNA abundances were then measured using next-generation sequencing as previously described\textsuperscript{25}. LysoTracker screens were performed with two experimental replicates per condition, while LAMP1 screens were done with one replicate. Results from all CRISPRi screens in this study can be interactively explored at www.crisprbrain.org.

**Western blots**

iAstrocytes were dissociated with Accutase for 10 mins at 37°C, spun down at 300 x g for 5 mins, washed with ice-cold DPBS, lysed in RIPA buffer (Thermo Fisher Scientific, cat. no. 89900) with protease (Roche, cat. no. 04693159001) and phosphatase inhibitors (Sigma-Aldrich, cat. no. PHOSS-RO), and incubated on ice for 45 minutes with brief vortexing every 15 minutes. Samples were centrifuged at 18,000 x g for 20 mins at 4°C to pellet insoluble components of lysate. After the total protein was quantified using the Pierce BCA Protein Assay Kit (ThermoFisher Scientific, cat. no. 23225) in the supernatant, each sample was diluted to the same concentration, and denatured with NuPAGE LDS Sample Buffer (4X) (ThermoFisher Scientific, cat. no. NP0007) and reduced with DTT. Equivalent protein amounts (20-40 ug) were run on 10\% NuPAGE Bis-Tris gels (ThermoFisher Scientific, cat. no. NP0301) and transferred to nitrocellulose membranes (Bio-Rad, 1704271). We performed ponceau staining (Milipore Sigma, cat. no. P7170) to validate equivalent loading between lanes. Membranes were blocked with TBST (TBS + 0.1\% Tween-20) + 5\% BSA for 1 hour at RT on a shaker, after which they were incubated with primary antibodies against phospho-S6 (1:500; polyclonal rabbit; Cell Signaling Technology cat. no. 2211), S6 (1:1000; monoclonal rabbit; Cell Signaling Technology cat. no. 2217), phospho-ULK1 (1:500; polyclonal rabbit; Cell Signaling Technology cat. no. 14202), phospho-Akt (1:500; monoclonal rabbit; Cell Signaling Technology cat. no. 4060), pan-Akt (1:1000; monoclonal mouse; Cell Signaling Technology cat. no. 2920), LAMP1 (1:1000; mouse monoclonal; Abcam cat. no. ab25630), LAMP2 (1:1000; mouse monoclonal; Abcam cat. no. ab25631), CTSD (1 ug/mL, R&D Systems cat. no. AF1014), \(\beta\)-actin (1:1000, rabbit monoclonal; Cell Signaling Technology cat. no. 4970), or EGFP (1:200; mouse monoclonal; Santa Cruz Biotechnology cat. no. sc-9996) at 4°C overnight on a shaker. Blots were washed with TBS and incubated with secondary antibodies goat anti-mouse IRDye 800CW (1:10,000; LI-COR cat. no. 926-32210) and goat anti-rabbit IRDye 680RD (1:10,000; LI-COR cat. no. 926-68071) in blocking buffer (see above) for 1 hour at RT on a shaker. Afterwards, blots were washed with TBS, and imaged using an Odyssey Fc Imaging system (LI-COR, cat. no. 2800). To re-probe blots, we incubated membranes with 1X NewBlot Nitro Stripping Buffer (LI-COR, cat. no. 928-40030) for 5 minutes at RT on a shaker before re-
blocking and probing as above. Immunoblot bands were quantified with LI-COR ImageStudio software.
Source data for immunoblots is provided in Supplementary Figures 2-5.

Whole-cell proteomics sample preparation

iAstrocytes were washed and collected (detached gently using a cell scraper) in ice-cold DPBS, spun down at 300 x g for 5 mins at 4 °C, and stored at -80 °C. Cell pellets were lysed in 2.5% sodium dodecyl sulfate (SDS) 50mM Tris pH 8 and heated to 95 °C for 5 min, then subjected to probe sonication. Lysates were centrifuged at 14000g for 15 min before protein quantification using BCA assay (Pierce, Thermo Fisher Scientific) according to manufacturer’s instructions. Protein lysates were acetone precipitated by adding 5 volumes of ice-cold acetone, vortexed, incubated at -20 °C for 1 hr and pelleted by centrifugation at 21000g and 4 °C for 15 min. Protein pellets were resuspended in 1% sodium deoxycholate (SDC) 50 mM EPPS pH 8.5 and reduced by incubating at 37 °C for 20 min with rotation in the presence of 5mM DTT. Protein was then alkylated by incubating in the dark at 37 °C for 20 min in the presence of 25mM iodoacetamide before digestion overnight with LysC at an enzyme-to-protein ratio of 1:50 followed by 3 hr digestion with trypsin at an enzyme-to-protein ratio of 1:50. Digestion was stopped by acidification to 1% trifluoroacetic acid (TFA) and placed on ice for 10 min to promote precipitation of SDC. The SDC was then pelleted and removed by centrifugation at 21000g for 10 min. The supernatant containing acidified peptides were then de-salted on in-house made Stagetips[^27] packed with polystyrene divinyl benzene reverse phase sulphonate solid phase extraction material (CDS Empore, Fisher Scientific) as previously published[^27]. Stagetips were prepared by activating with 100% methanol, conditioning with 80% acetonitrile containing 0.1% TFA, equilibrated with 0.2% TFA, followed by sample loading, washing twice with 99% isopropanol containing 0.1% TFA, washing twice with 0.2% TFA and once with 0.1% FA. Peptides were eluted with 60% acetonitrile (ACN) containing 0.5% ammonium hydroxide. The desalted peptides were then flash frozen and dried by centrifugal evaporation.

Lysosome-specific proteomics sample preparation

pLBR09 (gfaABC1D: LAMP1-XTEN80-mEGFP-3XHA) used for lysosome immunoprecipitation was generated by subcloning pLBR08 (EF1α: LAMP1-XTEN80-mEGFP-3XHA). Briefly, pLBR08 was generated via Gibson assembly of PCR-amplified LAMP1 cDNA (from mTagRFP-T-Lysosomes-20 acquired from Nikon Imaging Center at UCSF, Addgene plasmid #58022) and PCR-amplified XTEN80-mEGFP-3xHA immunoprecipitation tag with a linearized backbone generated from ClaI and BspDI (New England BioLabs cat. no. R0557) digestion of pKL017 (pHIV EF1α:Clover:WPRE). pLBR09 was then generated by linearizing pKL020 (pHIV gfaABC1D:mRuby:WPRE) and excising the LAMP1-XTEN80-mEGFP-3XHA cassette of pLBR08 via restriction digest with Xbal (New England Biolabs cat. no. R0145) and ClaI. The resulting fragments were run on a 1% agarose gel and gel extracted (Zymoclean Gel DNA Recovery Kit, cat. no. D4007). These fragments were ligated together with T4 DNA Ligase per manufacturer protocol. iAstrocytes were plated at 20,000 cells/cm² per experimental
replicate and transduced with pLBR09 lentivirus during plating. We used three replicates each for vehicle and ITC conditions.

Lysosome immunoprecipitation was performed as previously described\textsuperscript{11}. Briefly, iAstrocytes were washed and collected (detached gently using a cell scraper) in ice-cold DPBS, spun down at 300 x g for 5 mins at 4 °C. We then resuspended the cell pellet in homogenization buffer (25 mM Tris HCl pH 7.5, 50 mM sucrose, 0.5 mM MgCl\textsubscript{2}, 0.2 mM EGTA with protease inhibitors). The iAstrocyte cell suspension was mechanically lysed with a 23G syringe, immediately transferred to isotonic buffer (2.5M surcrose, 0.2 mM EGTA, 0.5 mM MgCl\textsubscript{2}) and spun at 3000 x g for 10 mins at 4 °C. The supernatant was transferred to anti-HA magnetic beads (Thermo Scientific cat. no. PI88836) for the pulldown of intact lysosomes, with the final resuspension in 1X urea buffer (2M Urea, 1 mM DTT, 12.5 mM Tris). On-bead digest was performed by sequentially incubating the resulting protein samples with 5 mM (final concentration) IAA at 22°C and 225 rpm for 30 mins, 0.5 ug LysC at 22°C and 225 rpm overnight, 1 ug of trypsin at 22°C and 225 rpm for 4 hours, and 1% (final concentration) TFA. Acidified peptides were then de-salted according to the same protocol as whole-cell proteome samples. For western blotting (Extended Data Fig. 1c), NuPAGE LDS Sample Buffer (4X) was added to samples collected prior to the on-bead digestion. See Western blots section for further details.

**LC-MS Analysis**

Peptides were resuspended in 2% ACN with 0.1% TFA before loading onto a 25 cm x 75 \textmu m ID, 1.6 \textmu m C18 column (IonOpticks) maintained at 40°C. Peptides were separated with an EASY-nLC 1200 system (Thermo Fisher Scientific, San Jose, CA) at a flow rate of 300 nl min\textsuperscript{-1} using a binary buffer system of 0.1% FA (buffer A) and 80% acetonitrile with 0.1% FA (buffer B) in a two-step gradient. The following was performed for whole cell proteome samples, 3% to 27% B in 105 min and from 27% to 40% B in 15min. For lysosome-specific samples, 3% to 27% B in 52.5 min and from 27% to 40% B in 14.5 min. All samples were analyzed on a Fusion Lumos mass spectrometer (Thermo Fisher Scientific, San Jose, CA) equipped with a nanoFlex ESI source operated at 1550 volts, RF lens set to 30%, operated in data dependent acquisition mode with a duty cycle time of 1 sec. Full MS scans were acquired with a m/z scan range of 375-1500 m/z in the Orbitrap mass analyzer (FTMS) with a resolution of 240k for whole cell proteome and 120k for lysosome-specific samples. Selected precursor ions were subjected to fragmentation using higher-energy collisional dissociation (HCD) with a quadrupole isolation window of 0.7 m/z, and normalized collision energy of 31%. HCD fragments were analyzed in the Ion Trap mass analyzer (ITMS) set to Turbo scan rate. Fragmented ions were dynamically excluded from further selection for a period of 45 sec for whole cell proteome and 60 sec for lysosome-specific samples. The AGC target was set to 1,000,000 and 10,000 for full FTMS and ITMS scans, respectively. The maximum injection time was set to Auto for both full FTMS and ITMS scans.

**Drug treatments**

Vacuolin-1 (Milipore Sigma, cat. no. 673000), bafilomycin A1 (Milipore Sigma, cat. no. B1793), rapamycin (Milipore Sigma, cat. no. 553210), and PP242 (Milipore Sigma, cat. no. 475988) were resuspended per manufacturer’s instructions, aliquoted, and stored at -80°C. All compounds.
were added to iAstrocytes alongside vehicle or ITC treatments (i.e. 24 hour treatments) with the exception of bafilomycin, which was added to cultures for the 2-4 hours preceding experiments for purposes of completely alkalinizing lysosomes.

Data analysis

Analysis of published RNA-seq datasets

We used BioJupies\textsuperscript{28} to reanalyze bulk RNA-seq data and obtain differentially expressed genes from Perriot et al.\textsuperscript{8} (GSE120411; hiPSC-derived astrocytes treated with IL1β and TNF) and Barbar et al.\textsuperscript{4} (syn21861229; CD49f+ astrocytes sorted from cerebral organoids treated with vehicle control or IL-1α+TNF+C1q),

Proteomics

Raw files were searched against the human reference proteome (Swiss-Prot, downloaded on 20\textsuperscript{th} of April 2020, refer to PRIDE upload for full sequence) along with the default common contaminants list with enzyme specificity set to trypsin within MaxQuant version 2.0.1.0 using standard settings with match between runs and the MaxLFQ algorithm activated. Whole cell proteome (raw data file names begin with ‘Proteome’) and lysosome-specific (raw data file names begin with ‘LAMP1’) raw files were set as separate parameter groups to ensure that the MaxLFQ algorithm was performed only within each sample set. The resulting proteinGroups text file was then uploaded to Perseus version 1.6.7.0 where all reverse sequences, potential contaminants and proteins only identified by site were filtered out of the dataset. All protein abundance values were normalized by Log 2 transformation. The lysosome-specific raw data files were evaluated separate from whole cell proteomic files. Search results from lysosomal raw data files were filtered to include proteins that were quantified in at least two out of three biological replicates within at least the ITC (raw data file names include ITC) or vehicle (raw data file names include WT) triplicate set. Missing values were then imputed within each replicate by replacing from the normal distribution using default settings. A two sample students t-test was performed between the ITC treated and vehicle triplicates, with permutation-based FDR using default settings. The resulting T-test difference and significance were used to generate the lysosome-specific volcano plot.

Pathway enrichment analysis

We used Enrichr\textsuperscript{29-31} to perform enrichment analysis of gene lists.

Flow cytometry
Data from flow cytometry experiments were analyzed using FlowJo (version 10.7.1). Live cells were gated by plotting SSC-A vs. FSC-A and then single cells were gated by plotting FSC-H vs. FSC-A. For experiments involving CRISPRi knockdown, analysis was restricted to sgRNA-transduced cells (gating on the histogram of BFP fluorescence values). For antibody staining experiments where median fluorescence intensity (MFI) values were reported, the average MFI of unstained control samples were subtracted from the MFI of stained samples. An example gating strategy from the resulting flow cytometry data is show in Supplemental Figure 1.

**Fluorescence imaging analysis**

We used CellProfiler (v3.15)\textsuperscript{32} to quantify iAstrocyte immunostaining and neuron apoptosis (as measured by cleaved caspase 3/7 fluorescent dye) imaging data. For neuron viability, monochrome images of caspase 3/7 were thresholded (“Threshold” module, “Manual” setting) with respect to unstained (no caspase 3/7 dye) controls. The caspase 3/7 signal was then quantified as the total intensity for each image, using the thresholded image as a mask. Similarly, for iAstrocyte fluorescence microscopy images, the total image intensity of LAMP1, LAMP2, LC3, and LysoTracker signals were quantified after first thresholding images relative to either a no-primary antibody control (immunofluorescence—LAMP1, LAMP2, LC3) or no-dye control (LysoTracker). The total image intensity of LAMP1, LAMP2, and LC3 was then divided by the total intensity of Hoechst to correct for variation in cell number. All frames collected were averaged to return a single quantification per well, which we considered as a single data point for statistical tests.

**Statistics and reproducibility**

Sample sizes were determined by referencing existing studies in the field. Major findings were validated using independent samples and orthogonal approaches. Numbers of replicates are listed in each figure. Randomization was not relevant to our study because no animals or human subjects were involved. Statistics were computed in RStudio (version 4.0.5). Correction for multiple testing was performed where appropriate.

**References**


Fig. 1 | Lysosomes are remodelled in inflammatory reactive astrocytes. a, Schematic of the *in vitro* model of inflammatory astrocyte reactivity used in this study. b, Bar graph of Gene Ontology (GO) terms enriched among the top 1000 genes with decreased expression in iAstrocytes treated with IL1α+TNF+C1q (ITC) compared to vehicle control from RNA-Seq (n =...
3 per condition). **c,** Heatmap of the mean log2(fold change) of gene transcripts with GO annotations of lysosome, lysosomal membrane, or lysosomal lumen in response to inflammatory stimuli for astrocytes derived using protocols described in Leng et al.⁵ (iAstrocytes), Krencik et al.³³ and published datasets from Barbar et al.⁴ and Perriot et al.⁸ (see Methods). **d,** left, representative immunoblot of total LAMP2 protein in iAstrocytes treated with ITC or a vehicle control; right, quantification of blots from n = 3 independent wells. **e,** Representative images of vehicle- (top) and ITC-treated (middle) iAstrocytes stained for the indicated lysosome/autophagosome markers, with corresponding quantification (bottom, see Methods; n = 3 independent wells per condition; scale bar = 75 μm). **f,** Lysosome acidity of FIRE-pHLY-expressing iAstrocytes (measured by flow cytometry) treated with ITC relative to vehicle-treated controls (n = 3 (for LAMP2, n = 4) independent wells for each condition). **g,** Autophagic flux (measured with GFP-LC3-RFP-LC3ΔG reporter-expressing iAstrocytes) treated with vehicle, ITC, or 100 nM bafilomycin A (n = 3 independent wells for each condition). **h,** Workflow of lysosome proteomics (lyso-IP, see Methods). **i,** Volcano plot of the log2(fold change) and associated log-scaled p-values of protein abundances detected by lyso-IP from astrocytes treated with ITC relative to vehicle-treated controls (see Methods; n = 3 per condition). **j,** Scatter plot of the log2(fold change) of proteins detected by lyso-IP versus whole-cell proteomics from astrocytes treated with ITC relative to vehicle-treated controls (see Methods; n = 3 per condition). For bar graphs, individual data points represent individual wells, while error bars represent the standard error of the mean (s.e.m.). The two-sided Student’s t-test was used in panels **d-f.**
Fig. 2 | Lysosome exocytosis controls neurotoxicity of inflammatory reactive astrocytes. 

a, Representative TIRF microscopy images of a lysosome exocytosis event (arrows) in iAstrocytes.
b, Quantification of LAMP1-mCherry (lysosomal membrane) and LysoTracker (lysosomal lumen) fluorescence intensity over the course of the lysosome exocytosis event depicted in a. 
c, Cell-surface LAMP1 for iAstrocytes treated with ITC or a vehicle control measured by flow cytometry. Left, representative flow cytometry results, including isotype control staining; right, quantification of replicates, with the average median fluorescence intensity (MFI) value of IgG isotype controls (n = 2 for each condition) subtracted from the LAMP1 MFI values (n = 4 for each condition).
d, Total LAMP1 protein in ITC and vehicle-treated iAstrocytes. Left, Representative immunoblot; right, quantification of independent wells (n = 3 for each condition).
e, Bafilomycin A dose-response curves of LysoTracker and cell-surface LAMP1 measured by flow cytometry in iAstrocytes (n = 3 wells per condition). Values are quantified as the percent of the maximal value obtained in each individual experiment (LysoTracker or cell-surface LAMP1).
f, Experimental strategy for parallel measurement of lysosome exocytosis and neurotoxicity of astrocyte conditioned media (ACM) (see Methods).
g-l, Quantification of lysosome exocytosis in iAstrocytes (g, i, and k) and neurotoxicity of astrocyte (measured by
caspase 3/7 activity reporter dye intensity) in iNeurons (h, j, and l) after treating iAstrocytes with the lysosome exocytosis inhibitor vacuolin-1 (g,h), knocking down lysosome exocytosis machinery in iAstrocytes with sgRNAs targeting SYT11 and VAMP7 (i,j), or treating iAstrocytes the lysosome exocytosis-inducing compound bafilomycin A (k,l). For bar graphs, data points represent individual wells with error bars representing the s.e.m. The two-sided Student’s t-test was used in panels c-d and linear regression was used in panels i-l to compute p-values; for linear regression, p-values were corrected for multiple testing using Holm’s method.
Fig. 3 (Legend overleaf)
Fig. 3 | CRISPRi screens uncover mTOR as a regulator of lysosome exocytosis and inflammatory reactive astrocyte-mediated neurotoxicity.  

a, Workflow of FACS-based CRISPRi screens in iAstrocytes (see Methods).  
b, Pearson correlations of the gene scores (see Methods) of hit genes from the 4 CRISPRi screens completed (n = 2 per condition).  
c, Bar graph of Molecular Signature Database (MSigDB) terms whose constituent genes are most significantly enriched among the 20 strongest hit genes of any CRISPRi screen (46 genes total, see Methods).  
d, Heatmap of gene scores across all CRISPRi screens for genes from c, with mTOR-related hits starred.  
e-g, iAstrocytes were transduced with non-targeting control (NTC) or MTOR sgRNA and treated with IL1a + TNF + C1q (ITC) or vehicle as indicated (n = 3 independent wells per condition), and (e) LysoTracker staining (f) cell-surface LAMP1, and (g) neurotoxicity of astrocyte-conditioned media were quantified.  
h, Representative immunoblots of mTOR levels and activity in iAstrocytes treated with a panel of mTOR inhibitors in combination with ITC or vehicle control.  
i-l, Quantification of immunoblots (from n = 3 independent wells for each condition).  
m-o, Dose-response curves of (m) lysosome acidification (measured by flow cytometry of FIRE-pHLy-expressing iAstrocytes), (n) lysosome exocytosis (flow cytometry measurement of cell-surface LAMP1, and (o) astrocyte-mediated neurotoxicity (measured by caspase 3/7 activity reporter dye intensity) upon treatment of iAstrocytes with the pan-mTORC inhibitor PP242 in conjunction with vehicle vs. ITC (n = 3 independent wells for each condition).  
p, Model for a potential role of mTOR-dependent lysosomal remodelling and exocytosis in astrocyte-mediated neurotoxicity. For bar graphs, data points represent individual wells with error bars representing the s.e.m. Linear regression was used to compute p-values in all panels; p-values were corrected for multiple testing using Holm’s method.
Extended Data Fig. 1 | Further characterization of lysosome remodeling in inflammatory reactive astrocytes. a, Heatmap of the mean log2(fold change) (ITC-treated astrocytes relative to vehicle-treated astrocytes) of genes or proteins with GO annotations of lysosome, lysosomal membrane, or lysosomal lumen from RNA-seq or whole-cell proteomics. c, Representative immunoblot of vehicle and ITC-treated samples obtained for lysosome immunoprecipitation (lyso-IP) experiment (+pLBR09, IP) with untransduced and non-immunoprecipitated (+pLBR09, Input) negative controls. d, Quantification of the cathepsin D (lysosomal lumen to B-actin (cytosol) ratio from immunoblots shown in c.
Extended Data Fig. 2 | Further characterization of the lysosome alkalization-exocytosis-neurotoxicity axis in iAstrocytes. a, Cell-surface LAMP1 levels (median fluorescence intensity – MFI; measured by flow cytometry) in vehicle and ITC-treated iAstrocytes transduced with NTC or LAMP1 sgRNA. b, Apoptosis of iNeurons treated with unconditioned astrocyte media containing the maximum concentration of vacuolin used in this study (100 nM). c, Percentage of dead iAstrocytes (measured by flow cytometry as DAPI+ cells) in response to concentrations of vacuolin used in this study. d, Cell-surface TFRC levels (MFI; measured by flow cytometry) of the iAstrocytes shown in Fig. 2g,h. e, Percentage of dead iAstrocytes (measured by flow cytometry as DAPI+ cells) in response to concentrations of bafilomycin A used in this study. f, Apoptosis of iNeurons treated with unconditioned astrocyte media containing the maximum concentration of bafilomycin A used in conditioned media experiments 10 nM).
Extended Data Fig. 3 | LysoTracker and LAMP1 CRISPRi screening results. Scatterplots comparing gene scores (see Methods) between screens for (a) LysoTracker, (b) LAMP1, (c) vehicle-treated screens, and (d) ITC-treated screens.
SUPPLEMENTARY INFORMATION

Supplementary Fig. 1  Gating strategy for general flow cytometry experiments (Live and Singlets gates) and FIRE-pHLy experiments (FIRE-pHLy gate).

Supplementary Fig. 2  Source Data for (a) Fig. 1d and (b) Fig. 2d.
Supplementary Fig. 3 Source Data for Fig. 3h.
Supplementary Fig. 4 Source Data for Fig. 3h.
Supplementary Fig. 5 Source Data for Fig. 3h.
Supplementary Table Legends

Supplementary Table 1 | Re-analysis of previously published hiPSC-derived astrocyte RNA-seq data. Differentially expressed genes (DEGs) induced by ITC in iAstrocytes and Krencik et al. astrocytes were taken from Supplementary Table 1 from Leng et al.5. DEGs induced by ITC in human cerebral organoid-derived astrocytes from Barbar et al.4 and DEGs induced by IL-1β+TNF in hiPSC-derived astrocytes from Perriot et al.8 were called by analyzing the raw data from those studies with BioJupies28.

Supplementary Table 2 | Whole-cell and lyso-IP proteomics. “LFQ intensity”: log2-scaled sum of label-free quantification intensities from all peptides within the protein group from MaxQuant34. “Log2-fold change (ITC vs. Veh)”: Mean LFQ intensity of ITC samples minus mean LFQ intensity of Veh samples. “Log-scaled p-value”: -Log10(t-test p-value). “FDR (q-value)”: permutation-based false discovery rate calculated from t-test p-values35. “Majority protein IDs”: UniProtKB IDs of proteins that have at least half of the peptides that the leading protein has.

Supplementary Table 3 | Phenotype scores and p-values from CRISPRi screens. Gene_TSS: gene name followed by the transcriptional start site (TSS) targeted by sgRNAs for genes with more than one TSS. Phenotype score: log-scaled fold difference between sgRNA abundance in high bin vs. low bin. Gene score: –log10(P value)*(Phenotype Score).

Supplementary Movie Legends

Supplementary Movie 1 | Visualization of iAstrocyte lysosomes via TIRF microscopy. Time-lapse movie (30 fps) of iAstrocytes stained with LysoTracker Green (green, lysosome lumen) and transduced with FIRE-pHLy (red, lysosome membrane) (see Methods).

Supplementary Movie 2 | Visualization of individual lysosome exocytosis events in iAstrocytes. Time-lapse movie (5 fps) of highlighted area in Supplementary Movie 1. Arrow highlights a putative lysosome exocytosis event.