RESEARCH ARTICLE

Oligopeptide transporter SIc15A modulates macropinocytosis in *Dictyostelium* by maintaining intracellular nutrient status

Yiwei Zhang^{1,2,*}, Hui Tu^{1,2,*}, Yazhou Hao^{1,2}, Dong Li^{1,3}, Yihong Yang¹, Ye Yuan^{1,2}, Zhonglong Guo⁴, Lei Li⁴, Haibin Wang^{1,‡} and Huaqing Cai^{1,2,‡}

ABSTRACT

Macropinocytosis mediates non-selective bulk uptake of extracellular fluid. It is the major route by which axenic Dictyostelium cells obtain nutrients and has emerged as a nutrient-scavenging pathway in mammalian cells. How environmental and cellular nutrient status modulates macropinocytic activity is not well understood. By developing a high-content imaging-based genetic screen in Dictyostelium discoideum we identified Slc15A, an oligopeptide transporter located at the plasma membrane and early macropinosome, as a novel macropinocytosis regulator. We show that deletion of slc15A but not two other related slc15 genes, leads to reduced macropinocytosis, reduced cell growth and aberrantly increased autophagy in cells grown in nutrient-rich medium. Expression of SIc15A protein or supplying cells with free amino acids rescues these defects. In contrast, expression of transportdefective SIc15A or supplying cells with amino acids in their di-peptide forms fails to rescue these defects. Therefore, SIc15A modulates the level of macropinocytosis by maintaining the intracellular availability of key amino acids through extraction of oligopeptides from the early macropinocytic pathway. We propose that SIc15A constitutes part of a positive feedback mechanism coupling cellular nutrient status and macropinocytosis.

This article has an associated First Person interview with the first authors of the paper.

KEY WORDS: *Dictyostelium*, Macropinocytosis, Nutrient status, Oligopeptide transporter

INTRODUCTION

Macropinocytosis is a cellular process that mediates non-selective bulk uptake of extracellular fluid. Studies in the model system *Dictyostelium discoideum* and immune cells have greatly facilitated our understanding of the cellular mechanisms that govern macropinocytosis (King and Kay, 2019). During this process, sheet-like extensions of the plasma membrane give rise to

[D] Y.Z., 0000-0001-8994-1808; H.T., 0000-0002-1898-2685; Y.Y., 0000-0001-9564-9563; Z.G., 0000-0002-3879-8738; H.C., 0000-0001-8434-6639

Handling Editor: Mahak Sharma Received 9 October 2021; Accepted 2 March 2022 micrometer-sized vesicles called macropinosomes. In the initial step of macropinosome formation, membrane extensions evolve into macropinocytic cups, which are organized around patches of active Ras, Rac and phosphatidylinositol (3,4,5)-trisphosphate (PIP₃), with actin-polymerizing activities that are driven, respectively, by the SCAR/WAVE complex and formin proteins at the periphery and base of the patches (Buckley et al., 2020; Fujii et al., 2013; Junemann et al., 2016; Mylvaganam et al., 2021; Veltman et al., 2016; Yoshida et al., 2009). A 'tent-pole mechanism', in which filopodia-like F-actin extensions twist to constrict the forming macropinosomes, has also been proposed (Condon et al., 2018). Once formed, macropinosomes can either recycle to the cell surface or mature by trafficking through the endolysosomal system where their contents are digested and extracted (Buckley and King, 2017; Donaldson, 2019; Freeman et al., 2020).

Macropinocytosis was originally recognized as a means for cells of the innate immune system, such as macrophages and dendritic cells, to survey environmental antigens for presentation to B cells and T cells (Lin et al., 2020; Norbury et al., 1995; Sallusto et al., 1995). More recently, studies in mammalian and *Dictyostelium* cells have uncovered an important metabolic function of macropinocytosis (Charpentier et al., 2020; King and Kay, 2019; Palm, 2019). Various types of cancer cells exploit macropinocytosis to survive in nutrientpoor environments by ingesting extracellular macromolecules and breaking them down in the lysosome to fuel cell growth (Commisso et al., 2013; Kim et al., 2018; Palm et al., 2015, 2017; Ramirez et al., 2019; Wyant et al., 2017; Yao et al., 2019). Axenic laboratory strains of Dictvostelium utilize macropinocytosis to obtain nutrients. including glucose and amino acids, from liquid medium (Bloomfield et al., 2015; Williams and Kay, 2018). These strains were isolated by selecting cells that performed macropinocytosis at a high rate in liquid culture. The mutations that allow axenic growth have been mapped to three separate loci, one of which is a gene encoding the RasGAP neurofibromin (axeB; Nfl in mammals) (Bloomfield et al., 2015; Clarke and Kayman, 1987).

As a means of acquiring nutrients, the extent of macropinocytosis is intrinsically connected to the nutritional status of the cell. In pancreatic ductal adenocarcinoma cells (PDACs), glutamine deprivation or restriction of extracellular proteins as the source of essential amino acids induces macropinocytosis (King et al., 2020; Lee et al., 2019). Amino acid shortage has also been shown to promote macropinocytosis in human placental trophoblasts (Shao et al., 2021). In axenic *Dictyostelium* cells, the presence of bacteria as a nutrient source promotes phagocytosis and suppresses macropinocytosis, whereas switching the feeding strategy to liquid medium prominently increases the rate of macropinocytosis (Kayman and Clarke, 1983; Williams and Kay, 2018). Although these observations demonstrate that cells can calibrate their macropinocytic activity according to changes in nutrient conditions, how this is achieved in cells and what the full complement of regulators are remain to be elucidated.



¹National Laboratory of Biomacromolecules, Institute of Biophysics, Chinese Academy of Sciences, Beijing 100101, China. ²College of Life Sciences, University of Chinese Academy of Sciences, Beijing 100049, China. ³Division of Life Sciences and Medicine, University of Science and Technology of China, Hefei 230027, China. ⁴State Key Laboratory of Protein and Plant Gene Research, School of Life Sciences and School of Advanced Agricultural Sciences, Peking University, Beijing 100871, China.

^{*}These authors contributed equally to this work

[‡]Authors for correspondence (huaqingcai@ibp.ac.cn; wanghaibin@ibp.ac.cn)

We developed a forward genetic screen in *Dictyostelium* to seek novel macropinocytosis regulators. In this study, we further characterized one of the genes isolated from the screening, which encodes an oligopeptide transporter we named Slc15A. We show that, opposite to the strategy of mammalian cells, Slc15A constitutes part of a positive feedback mechanism coupling cellular nutrient status and macropinocytosis. Furthermore, Slc15A modulates the level of macropinocytosis by acting in the early macropinocytic pathway instead of the lysosome where nutrient extraction is thought to take place in canonical models. Our results demonstrate that macropinocytosis and its regulation are far more sophisticated than we currently understand.

RESULTS

Screening for macropinocytosis regulators identified Slc15A

To screen for macropinocytosis regulators, we generated a pool of restriction enzyme-mediated insertion (REMI) mutants in the axenic *Dictyostelium* strain Ax2. Individual mutant clones were incubated with 70 kDa tetramethylrhodamine isothiocyanate dextran (TRITC dextran, hereafter referred to as TD) in liquid growth medium and then subjected to image-based high-content analysis (Fig. 1A). Cell boundaries and intracellular vesicles containing TD were identified in the digital phase-contrast channel and fluorescent channel, respectively (Fig. 1A). The ratio of the total intensity of TD spots to the total cell area was used to quantify the activity of macropinocytosis. In addition to Ax2 cells, cells deleted of class-1 phosphoinositide 3-kinases (PI3Ks) were used as controls. Consistent with previous reports (Hoeller et al., 2013), cells deleted for genes encoding PI3Ks exhibited significantly reduced macropinocytosis under these experimental conditions (Fig. 1B).

From the screen, we obtained a series of macropinocytosis mutants (Fig. 1C). This report focuses on the gene we named *slc15A* (dictyBase gene ID: DDB_G0272550), which encodes a protein homologous to the mammalian solute carrier family 15 proteins (Fig. S1A). SLC15 proteins are proton-coupled oligopeptide transporters (POTs) mediating the cellular uptake of a wide range of di- and tri-peptides and peptide-like drugs (Daniel et al., 2006; Smith et al., 2013). Phylogenetic analysis revealed that the SLC15 family in eukaryotes constitutes two clades, and *Dictyostelium* Slc15A is

closely related to the mammalian SLC15A1 and SLC15A2 (Fig. S1A). Sequence alignment showed that Slc15A shares 45–46% similarity to human SLC15A1 and SLC15A2 (Fig. S1B).

The REMI insertion site was identified at position 795 of the *slc15A* genomic locus (numbered from the ATG translation start site, Fig. S1B). To confirm whether this insertion is, indeed, responsible for the observed phenotype, we generated *slc15A* knockout (*slc15A*⁻) cell line in Ax2 (Fig. S2A). The knockout cells exhibited similar phenotypes as the REMI mutant. When incubated with TD, substantially fewer and less bright TD-containing vesicles were observed in *slc15A*⁻ cells compared to Ax2 cells (Fig. 2A,B). Similarly, the macropinocytic uptake of 70 kDa fluorescein isothiocyanate-dextran (FITC dextran, FD) was significantly impaired in *slc15A*⁻ cells (Fig. S2B). Quantification of macropinocytosis over time corroborated these findings. The rate of macropinocytosis and the steady-state plateau were reduced by ~50% in *slc15A*⁻ cells compared to that in control (Fig. 2C). In contrast, fluid-phase exocytosis was not affected (Fig. 2D). Furthermore, the macropinocytosis defects associated with *slc15A* deletion could be fully rescued by expression of GFP- or RFPtagged Slc15A (Fig. 2A-C and Fig. S2B). We also generated a slc15A⁻ cell line in the non-axenic Dictyostelium strain DdB (Fig. S2C). Macropinocytic activity was considerably lower in DdB compared to Ax2 (Fig. S2D), and deletion of slc15A further decreased this activity (Fig. 2E,F). These experiments confirm that Slc15A is a novel regulator of macropinocytosis.

SIc15A is specifically required for macropinocytosis regulation

To begin elucidating the function of Slc15A, we examined the distribution and dynamics of several macropinocytic marker proteins in Ax2 and *slc15A*⁻ cells. Rab5A, Rab7A, the V-ATPase subunit VatB, the PIP₃/PI(3,4)P₂ sensor PHcrac, the PI(3,4)P₂ sensor TAPP1 or the PI(3)P sensor $2 \times FYVE$ were fused with GFP or RFP and used to mark macropinosomes at different stages (Buckley and King, 2017; Dormann et al., 2004; Egami et al., 2014; Maekawa et al., 2014; Rupper et al., 2001; Swanson, 2014). We observed that the overall organization of the macropinocytic pathway appeared unaffected by *slc15A* deletion. Marker proteins

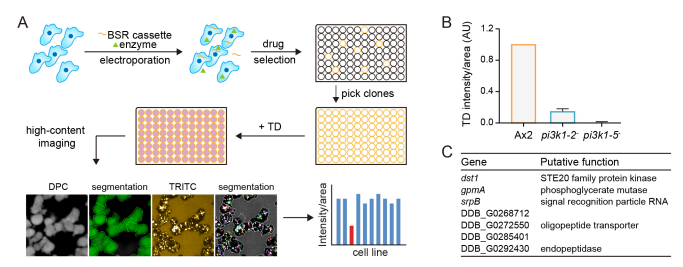


Fig. 1. Screening for macropinocytosis regulators. (A) Schematic of the genetic screen. Mutant clones were generated by REMI, seeded in 96-well plates in growth medium and incubated with TD. Images were captured in the digital phase contrast (DPC) and TRITC channels using a high-content screening system. The ratio of the total intensity of TD spots to the sum of cell area was used to quantitatively measure macropinocytosis. (B) Cells of the Ax2 strain deleted for PI3K1 and 2 (*pi3k1-2* null) or PI3K1-5 (*pi3k1-5* null) were analyzed for TD uptake by using the screening system. Data are from three independent experiments; mean ±s.d. (C) List of mutants obtained from the screen.

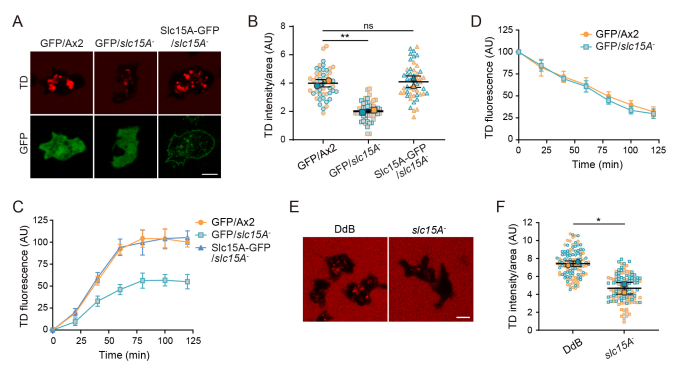


Fig. 2. Slc15A is involved in the regulation of macropinocytosis. (A,B) Confocal images and quantification of TD uptake in Ax2 strain wild-type expressing GFP (GFP/Ax2) and *slc15A*⁻ cells expressing GFP or Slc15A-GFP (*GFP/slc15A*⁻ or Slc15A-GFP/*slc15A*⁻, respectively). Data are from two independent experiments with at least 20 cells quantified per experiment (each experiment is shown in a different color); mean±s.d. (C,D) Quantification of TD uptake in DdB and exocytosis (D) by fluorimetric analysis. Data are from four independent experiments; mean±s.d. (E,F) Confocal images and quantification of TD uptake in DdB and *slc15A*⁻ cells. Data are from two independent experiments with at least 50 cells quantified per experiment is shown in a different color); mean±s.d. (E,F) Confocal images and quantification of TD uptake in DdB and *slc15A*⁻ cells. Data are from two independent experiments with at least 50 cells quantified per experiment (each experiment is shown in a different color); mean±s.d. (E,F) Confocal images and quantification of TD uptake in DdB and *slc15A*⁻ cells. Data are from two independent experiments with at least 50 cells quantified per experiment (each experiment is shown in a different color); mean ±s.d. Statistical significance was determined by one-way ANOVA with Dunnett post-test (B) and two-tailed unpaired *t*-test (F). Scale bars: 5 µm.

exhibited similar localization in Ax2 and $slc15A^-$ cells (Fig. 3A), and the sequential accumulation of PIP₃, PI(3,4)P₂, and PI(3)P proceeded with similar kinetics (Fig. 3B and Fig. S3). Measurement of macropinosome acidification also did not reveal any apparent defect in $slc15A^-$ cells (Fig. S4). However, time-lapse imaging and quantification by using PHcrac-GFP to mark macropinocytic cups and newly internalized macropinosomes revealed a significant reduction in the rate of macropinosome formation in $slc15A^-$ cells. Despite the production of similar-sized vesicles, the number of macropinosomes successfully generated per cell per minute was reduced by ~40% in $slc15A^-$ compared to Ax2 cells (Fig. 4A-C, Movie 1). In many instances, macropinocytic cups formed in $slc15A^-$ cells collapsed and receded to the plasma membrane without vesicle formation (Fig. 4A).

As macropinocytosis is the main pathway by which axenic cells obtain nutrients from liquid medium (Hacker et al., 1997), its defect often manifests as slow cell growth. Indeed, the generation time in shaken suspension increased from ~8 h for Ax2 cells to ~15 h for *slc15A*⁻ cells (Fig. 4D). Expression of Slc15A-GFP rescued this defect (Fig. 4D). Lack of nutrient uptake has also been shown to upregulate autophagy. Imaging the autophagy marker Atg8a tagged to GFP (GFP-Atg8a) (King et al., 2011) revealed that, although nonnutrient development buffer (DB) induced autophagy to a similar extent in Ax2 and $slc15A^{-}$ cells, the latter showed significantly increased autophagy in the commonly used HL5 growth medium (Fig. 4E,F; Movies 2 and 3). The increase of Atg8a structures in *slc15A*⁻ cells was not caused by a block of autophagy flux (Fig. S5). Therefore, *slc15A* deletion likely impairs macropinocytosismediated nutrient uptake, resulting in reduced cell growth and aberrantly increased autophagy in cells cultured in nutrient-rich medium.

In addition to liquid medium, *Dictyostelium* cells could utilize bacteria as the nutrient source through phagocytosis. We investigated whether Slc15A is involved in the regulation of phagocytosis. When plated on bacterial (*Klebsiella aerogenes*) lawns, the plaque growth of *slc15A*⁻ cells was indistinguishable from that of Ax2 cells, indicating that bacterial uptake and digestion were not affected (Fig. S2A). Quantifying bacteria phagocytosis by using flow cytometry analysis and assaying the proteolytic activity of bacteria-containing phagosomes confirmed this conclusion (Fig. 4G,H). Together, these experiments indicate that Slc15A is specifically involved in the regulation of macropinocytic, but not phagocytic growth.

SIc15A promotes macropinocytosis by maintaining intracellular availability of amino acids

To determine whether Slc15A modulates macropinocytosis through its putative function as an oligopeptide transporter, we generated a Slc15A variant bearing mutations (Slc15A^{mut}-GFP) in the two conserved residues R68 and E438 (Fig. S1B). The corresponding mutations R57H and E595R in human SLC15A2 and SLC15A1, respectively, have previously shown to abolish transport function (Terada et al., 2004; Xu et al., 2009). Although Slc15A^{mut}-GFP was expressed at a level similar to its WT counterpart, it failed to rescue the TD uptake and cell growth defects (Fig. 5A-C). This suggests that the oligopeptide transport activity is required for Slc15A to regulate macropinocytosis.

The standard HL5 medium contains amino acids, peptides and proteins derived from peptone and yeast extract. We therefore speculated that deletion of *slc15A* causes reduced peptide import from HL5 and, thus, a lack of amino acid-nutrients in cells, which in

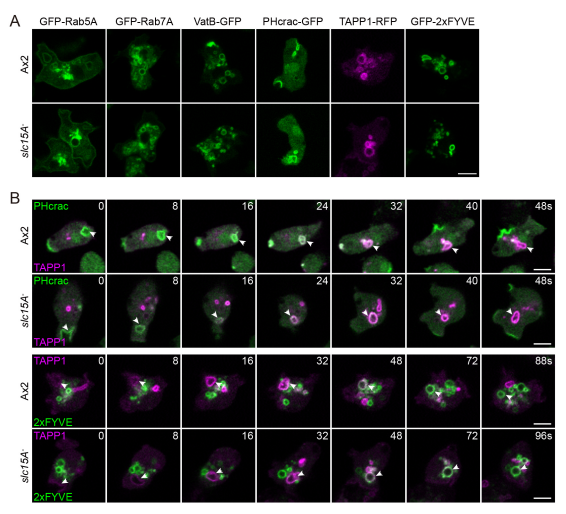


Fig. 3. Localization of macropinocytic marker proteins in Ax2 strain wild-type and *slc15A* deletion cells. (A) Confocal images of the indicated marker proteins in Ax2 and *slc15A⁻* cells. (B) Top: Time-lapse images showing the sequential accumulation of PHcrac-GFP and TAPP1-RFP in Ax2 and *slc15A⁻* cells. Bottom: Time-lapse images showing the sequential accumulation of TAPP1-RFP and GFP-2×FYVE in Ax2 and *slc15A⁻* cells. Scale bars: 5 µm.

turn might downregulate macropinocytosis. Were this true, providing $slc15A^-$ cells with free amino acids, which bypass the need for peptide transport, should rescue the macropinocytosis defect. To test this, we replaced HL5 with SIH medium, a synthetic medium containing a defined composition of free amino acids sufficient to sustain cell growth. Growing $slc15A^-$ cells in SIH medium fully rescued the macropinocytosis defect (Fig. 5D,E and Fig. S6), and cell growth and autophagy regulation were also restored (Fig. 5F,G; Movies 2 and 3). These experiments support our hypothesis that reduced amino acid availability underlies the macropinocytosis defect of $slc15A^-$ cells.

To determine whether specific amino acids within the SIH medium are crucial for the rescuing effect, we subjected $slc15A^-$ cells to SIH medium lacking arginine, glutamate or lysine, as these three amino acids had been shown to be needed for the upregulation of macropinocytosis when the feeding strategy was switched from growth on bacteria to liquid medium (Williams and Kay, 2018). Intriguingly, removing arginine (SIH-R) or lysine (SIH-K), but not glutamate (SIH-E), abolished the ability of SIH medium to rescue the macropinocytosis defect caused by slc15A deletion (Fig. 6A,B). Ax2 cells cultured in SIH-R or SIH-K also exhibited reduced macropinocytosis (Fig. 6A,B). These results indicate that slc15A deletion likely causes reduced availability of key amino acids, including arginine and lysine.

To further test whether Slc15A-mediated oligopeptide transport contributes to the intracellular pool of arginine and lysine, we subjected cells to medium containing synthesized dipeptide of arginine (R-R) or lysine (K-K). Supplementing R-R or K-K restored the macropinocytosis ability of Ax2 cells grown in SIH-R or SIH-K. However, it failed to do so for *slc15A*⁻ cells, confirming that the knockout cells are defective in dipeptide utilization (Fig. 6C,D). These experiments collectively indicate that Slc15A promotes macropinocytosis by functioning as an oligopeptide transporter to maintain the intracellular availability of key amino acids.

SIc15A functions in the early macropinocytic pathway

Extracellular proteins taken up by macropinocytosis are thought to be delivered to the lysosomes, degraded into amino acids, and then exported to the cytosol to fuel metabolism (Verdon et al., 2017; Wyant et al., 2017). However, we noticed that Slc15A did not appear to localize to lysosomal vesicles when expressed from extrachromosomal plasmids. To determine the localization more accurately, we generated Slc15A-GFP and Slc15A-RFP knock-in (KI) cells (Fig. S2E). We found that Slc15A localized to the plasma membrane in these cells and, when the membrane deformed to produce macropinosomes, remained associated with newly generated macropinosomes (Fig. 7A,B). Imaging Slc15A together with TAPP1 and 2×FYVE revealed that Slc15A and TAPP1

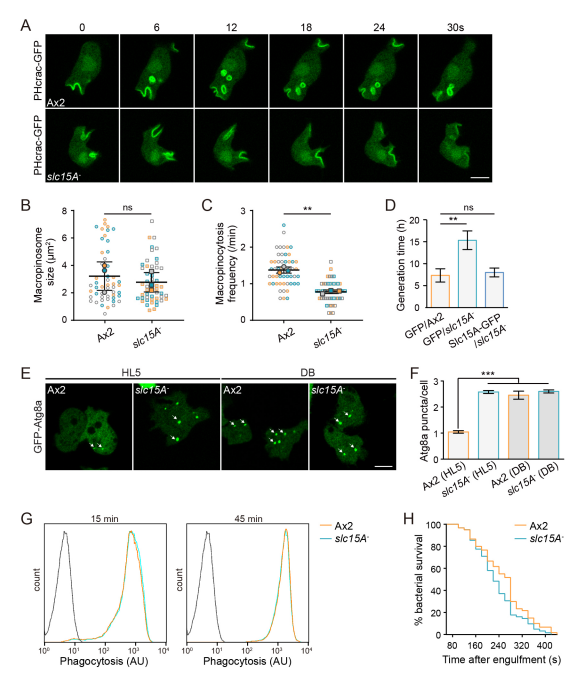


Fig. 4. Macropinocytosis is specifically impaired in *slc15A* **deletion cells.** (A) Time-lapse images of PHcrac-GFP in Ax2 strain wild-type and $slc15A^{-}$ cells. (B,C) Plotted is the size of newly enclosed macropinosomes labeled by PHcrac-GFP and rate of macropinosome formation. Data are from three independent experiments with at least 15 cells quantified per experiment (each experiment is shown in a different color); mean±s.d. (D) Cell growth in HL5 medium measured by generation time. Data are from three independent experiments; mean±s.d. (E) Confocal images of GFP-Atg8a in cells cultured in HL5 or non-nutrient DB for 1 h. (F) Plotted is the average number of Atg8a puncta per cell. Data are from three independent experiments with at least 30 cells quantified per experiment; mean ±s.d. (G) Quantification of bacterial phagocytosis. Cells were incubated with mCherry-expressing *E. coli* for the indicated time points and phagocytosis was measured by flow cytometry. The black lines show the background fluorescence. (H) Quantification of bacterial survival. Cells were incubated with GFP-expressing *E. coli*. The Kaplan-Meyer graph was based on the persistence of bacterial GFP-fluorescence within cells after phagocytosis. Per cell line, 60 bacteria were followed. Statistical significance was determined by two-tailed unpaired *t*-test (B,C) and one-way ANOVA with Dunnett post-test (D,F). Scale bars: 5 μ m.

colocalized on macropinosomes during cup closure and initial trafficking (Fig. 7A, Movie 4); the signal of Slc15A gradually decreased as the macropinosomes moved further into the cells, which was accompanied by the gradual accumulation of 2×FYVE (Fig. 7B). These experiments indicate that Slc15A recycles from the macropinosomes in the early stages of the pathway. In line with this finding, we found little overlap between Slc15A and the lysosomal/ post-lysosomal marker LmpA (Sattler et al., 2018) (Fig. S7). Thus,

Slc15A likely mediates oligopeptide transport across the plasma membrane or the early macropinosomal membrane but not efflux from the lysosome.

Given that the SLC15 family proteins rely on an electrochemical H^+ gradient for transport activity (Smith et al., 2013; Steel et al., 1997), we examined whether the presence of Slc15A on nascent macropinosomes overlapped with macropinosome acidification. By incubating Slc15A-RFP KI cells with FD, we found that newly

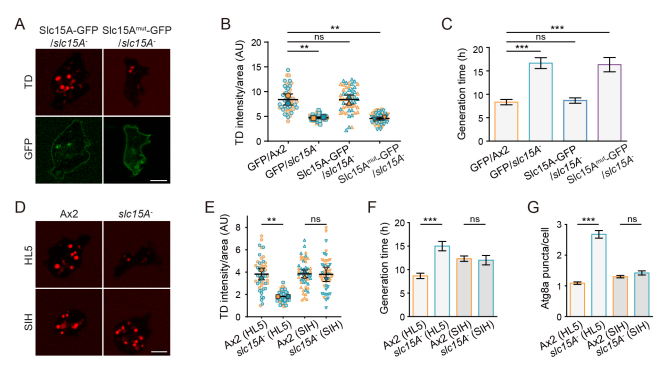


Fig. 5. The oligopeptide transport activity is required for SIc15A to modulate macropinocytosis. (A) Confocal images of TD uptake in *slc15A*⁻ cells expressing Slc15A-GFP or Slc15A^{mut}-GFP bearing R68H and E438R mutations. (B) Quantification of TD uptake in cells transformed with the indicated plasmids. Data are from two independent experiments with at least 20 cells quantified per experiment (each experiment is shown in a different color); mean±s.d. (C) Cell growth in HL5 measured by generation time. Data are from three independent experiments; mean±s.d. (D,E) Confocal images and quantification of TD uptake in Ax2 and *slc15A*⁻ cells grown in HL5 or SIH medium. Data are from two independent experiments with at least 20 cells quantified per experiment is shown in a different color); mean±s.d. (F) Cell growth in HL5 or SIH measured by generation time. Data are from two independent experiments with at least 20 cells quantified per experiment (each experiment is shown in a different color); mean±s.d. (F) Cell growth in HL5 or SIH measured by generation time. Data are from three independent experiments with at least 20 cells quantified per experiment (each experiment is shown in a different color); mean±s.d. (F) Cell growth in HL5 or SIH measured by generation time. Data are from three independent experiments; mean±s.d. (G) Plotted is the average number of GFP-Atg8a puncta per cell in cells grown in HL5 or SIH. Data are from three independent experiments with at least 30 cells quantified per experiment. Statistical significance was determined by one-way ANOVA with Dunnett post-test (B,C) and two-tailed unpaired *t*-test (E–G). Scale bars: 5 μm.

generated macropinosomes quickly acquired an acidic environment, indicated by the decrease of FD fluorescence before Slc15A-RFP was evidently recycled (Fig. 7C). Quantification revealed that Slc15A associated with newly internalized macropinosome for 115 ± 31 s (from 15 macropinosomes), whereas macropinosome acidification was initiated almost immediately after cup closure, with a significant drop of pH occurring within 60 s (Fig. S4). Imaging Slc15A together with VatB also revealed that the recruitment of V-ATPase to macropinosomes started before recycling of Slc15A (Fig. 7D). We speculate that the rapid acidification of nascent macropinosomes facilitates the function of Slc15A by providing a steeper proton gradient. However, at this stage we cannot rule out the possibility that Slc15A also functions at the plasma membrane.

DISCUSSION

In summary, we identified Slc15A as a macropinocytosis-specific regulator. We demonstrated that Slc15A is not directly involved in macropinosome formation or maturation; instead, it sustains the macropinocytic activity of cells by mediating oligopeptide transport from growth medium and maintaining the intracellular availability of key amino acids (Fig. 8). The *Dictyostelium* genome encodes two other putative SLC15 family proteins (Fig. S1A). However, these two proteins did not localize to the macropinocytic pathway and their disruption did not affect macropinocytosis (Fig. S8), confirming a unique function of Slc15A.

Characterization of Slc15A allowed us to gain fresh insights into the regulation of macropinocytosis. First, the localization study suggested that Slc15A functions earlier in the macropinocytic pathway. Thus, different from the canonical model, the lysosome is not the only cellular compartment where internalized nutrients are extracted. Although it is not yet feasible to pinpoint precisely where Slc15A-mediated transport occurs, we speculate that nascent macropinosomes are the more probable compartments because their acidic environment could promote the function of Slc15A. The characteristic substrate multispecificity of SLC15 proteins might further facilitate nutrient extraction in the form of oligopeptides before reaching lysosome (Ito et al., 2013). Intriguingly, the defects in $slc15A^{-}$ cells were alleviated by feeding cells with free amino acids, which might be delivered to the cell via amino acid transporters. This ability to retrieve nutrients in different forms and, possibly, at different stages along the macropinocytic pathway likely promotes metabolic flexibility. Second, our study indicates that Slc15A contributes to a positive feedback mechanism coupling cellular nutrient status and macropinocytosis (Fig. 8). Disrupting Slc15A, or depleting arginine or lysine from the growth medium all led to reduced macropinocytosis. Taking into account previous studies that analyzed macropinocytosis during a switch in feeding strategy (Williams and Kay, 2018), our experiments convincingly demonstrate that the intracellular availability of key amino acids can, in turn, promote the macropinocytic activity of Dictyostelium cells, constituting feedback regulation. Interestingly, seemingly opposite strategies are taken by Dictyostelium and mammalian cells to cope with amino acid stress conditions: Dictyostelium cells downregulate macropinocytosis, whereas mammalian cells upregulate the process (Lee et al., 2019; Shao et al., 2021). Such a difference might reflect different survival strategies. Starvation in Dictyostelium ultimately leads to multicellular development, a process that benefits from reduced macropinocytosis (Veltman et al., 2014, 2016), whereas

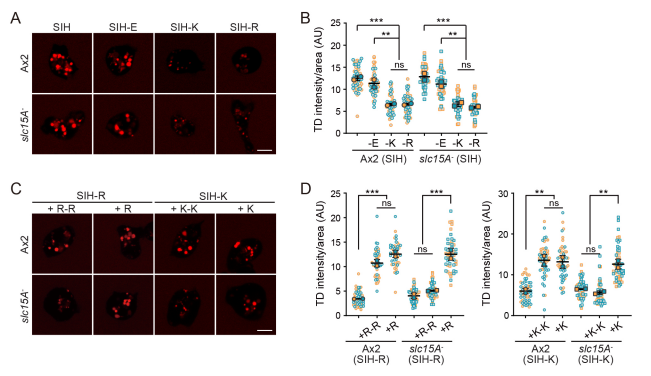


Fig. 6. SIc15A functions to maintain nutrient availability and promote macropinocytosis. (A,B) Confocal images and quantification of TD uptake in Ax2 strain wild-type and $slc15A^-$ cells cultured in normal SIH or SIH lacking amino acids glutamic acid (E), lysine (K) or arginine (R) – SIH-E, SIH-K or SIH-R, respectively. (C,D) Confocal images and quantification of TD uptake in Ax2 and $slc15A^-$ cells cultured in SIH-R or SIH-K or SIH-R, respective single or dipeptide amino acid as indicated. For plots shown in B and D, data are from two independent experiments with at least 20 cells quantified per experiment (each experiment is shown in a different color); mean±s.d. Statistical significance was determined by one-way ANOVA with Tukey post-test. Scale bars: 5 μ m.

mammalian cells lacking this alternative route might resort to macropinocytosis upregulation and protein-scavenging instead.

Future studies are needed to elucidate how intracellular nutrient levels are sensed and transduced to regulate macropinocytosis in *Dictyostelium*. In other systems, amino acid-sensing is mediated by activation of the mechanistic target of rapamycin complex 1 (mTORC1) pathway (Wolfson and Sabatini, 2017). It is possible that mTORC1 senses the nutrient stress condition in *slc15A⁻* cells to downregulate macropinocytosis. However, pharmacological inhibition of mTORC1 or depletion of the mTORC1-specific component Raptor does not affect macropinocytosis in *Dictyostelium* (Rosel et al., 2012; Williams and Kay, 2018), suggesting that the feedback regulation likely involves so-farunidentified pathways. Additionally, as SLC15 orthologues exist widely in eukaryotes, it will also be of great interest to investigate whether the SLC15 family of proteins is involved in the regulation of macropinocytosis in other species.

MATERIALS AND METHODS

Cell growth

We used cells of the axenic *Dictyostelium* strain Ax2, routinely cultured in HL5 medium (HLF3, Formedium,). Cells carrying expression constructs were maintained in HL5 containing the antibiotics G418 (10–20 µg/ml) or hygromycin (50–100 µg/ml). Development buffer (DB) contained 5 mM Na₂HPO₄, 5 mM KH₂PO₄, 2 mM MgSO₄ and 0.2 mM CaCl₂. The growth rate was measured by seeding cells at 0.5×10^5 /ml and measuring cell density for four consecutive days. DdB cells were used as the non-axenic wild-type strain. Cells of the non-axenic *Dictyostelium* strain DdB were cultured in SorMC buffer (15 mM KH₂PO₄, 2 mM Na₂HPO₄, 50 µM MgCl₂ and 50 µM CaCl₂, pH 6.0) containing *K. aerogenes* (OD₆₀₀=2) or grown on *K. aerogenes* lawns as described before (Paschke et al., 2018).

REMI screen

REMI mutants were generated as described previously (Kuspa, 2006; Shaulsky et al., 1996). In brief, a DNA fragment containing a blasticidin resistant cassette (BSR) was digested with BamHI and co-transformed into cells with the DpnII restriction enzyme. Cells were dispensed into 96-well dishes in HL5 medium and blasticidin (10 µg/ml) was added the next day to select mutant clones that had DNA-containing BSR inserted into their genome. Individual mutant clones were seeded in 96-well plates in growth medium, incubated with 70 kDa tetramethylrhodamine isothiocyanate dextran (TRITC dextran, hereafter referred to as TD; Sigma, T1162) for 1 h, and then analyzed using a PerkinElmer Phenix Opera high-content screening system. Images were captured in the digital phase contrast channel and TRITC channel with nine fields per well using a 63× water lens. Image analyses were performed using the built-in Harmony software. For gene identification of REMI mutants, genomic DNA was digested with PsiI and ligated to circularize fragments; inverse PCR was performed using a pair of outward-facing primers located within the BSR sequence as described previously (Keim et al., 2004). The obtained PCR bands were then sequenced to identify the insertion sites.

Molecular biology

To make knockout constructs for *slc15A*, *slc15B* or *slc15C* in Ax2, a BSR cassette (Faix et al., 2004) was inserted into pBlue-Script II SK+ to generate pBlueScript-BSR. Thereafter, 5' and 3' arms were PCR-amplified from genomic DNA by using primers listed in Table S1, and cloned upstream and downstream of the BSR cassette, respectively. The resulting disruption cassette was PCR amplified and electroporated into Ax2. Gene disruption was confirmed by resistance to blasticidin ($10 \mu g/ml$) and PCR amplification. To make knockout construct for *slc15A* deletion in DdB, 5' and 3' arms were PCR amplified with primers listed in Table S1 and cloned upstream and downstream of the NeoR cassette in pDM1082, respectively. Transfection and selection were performed as described before (Paschke et al., 2018). Gene disruption was confirmed by resistance to G418

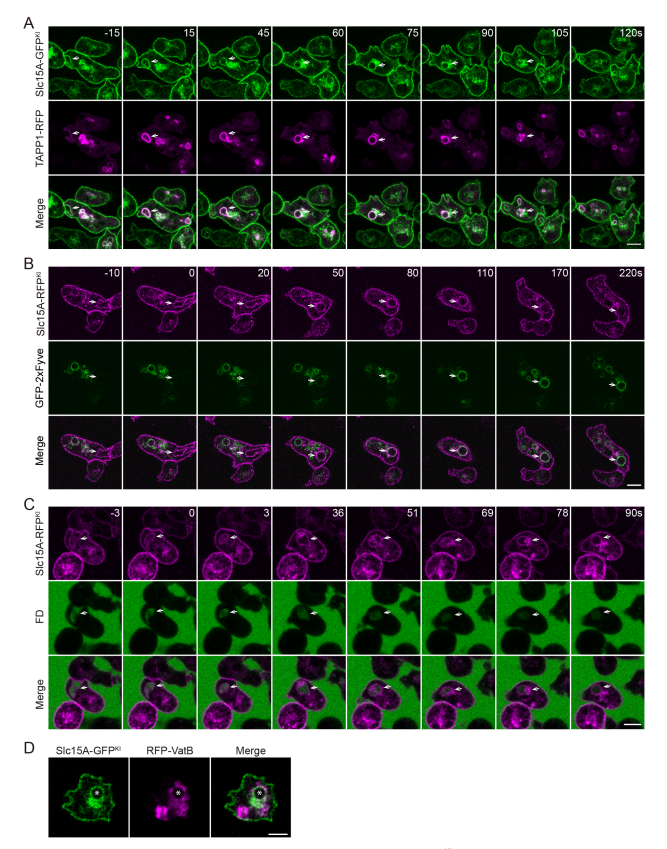


Fig. 7. SIc15A functions in the early macropinocytic pathway. (A) Time-lapse images of Slc15A-GFP^{KI} and TAPP1-RFP in Ax2 strain wild-type cells during macropinocytosis. Arrows indicate a newly generated macropinosome with which Slc15A and TAPP1 associated for ~100 s. (B) Time-lapse images of Slc15A-RFP^{KI} and GFP-2×FYVE in Ax2 cells during macropinocytosis. Arrows indicate a newly generated macropinosome that had been converted from Slc15A positive to 2×FYVE positive. (C) Time-lapse images of Slc15A-RFP^{KI} in cells incubated with FD. Arrows indicate a newly generated macropinosome that quickly acquired an acidic environment – indicated by the decrease of the fluorescence of FD. (D) Confocal images of Slc15A-GFP^{KI} and RFP-VatB. Asterisks indicate a nascent macropinosome marked by both Slc15A-GFP and RFP-VatB. Scale bars: 5 μm.

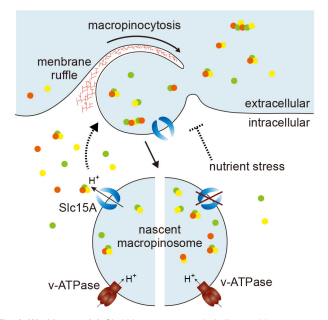


Fig. 8. Working model. Slc15A, a proton-coupled oligopeptide transporter located at the plasma membrane and early macropinosome, mediates oligopeptide transport from growth medium to the cytosol, thereby maintaining the intracellular availability of key amino acids and promoting macropinocytosis via a feedback regulation. Disruption of *slc15A* reduces oligopeptide transport and triggers a nutrient stress condition, which in turn impairs macropinocytosis and leads to defects in cell growth and autophagy regulation. Colored dots indicate amino acids and peptides.

 $(10 \ \mu g/ml)$ and PCR. Slc15A-GFP knock-in cell was generated by replacing the region of *slc15A* between +1920 and +1922 with a GFP-hygromycin cassette from pDM1355 (Paschke et al., 2018). GFP was replaced with RFP to generate the Slc15A-RFP knock-in cassette. Knock-in clones were selected by hygromycin resistance and confirmed by PCR and western blotting.

To generate constructs expressing GFP- or RFP-fusion proteins, DNA fragments encoding *Dictyostelium* Slc15A, Slc15B, Slc15C, Atg8a, Rab5A, Rab7A, VatB, LmpA or human TAPP1 were PCR-amplified using primers listed in Table S1 and cloned into *Dictyostelium* pDM expression vectors (Veltman et al., 2009) containing a multiple cloning site. To generate construct expressing GFP-2×FYVE, a fragment containing 2×FYVE domain of *C. elegans* EEA1 was released from a *C. elegans* expression vector (Li et al., 2009) and cloned into pDM317. To generate Slc15A^{R68H/E438R}-GFP, arginine (R) at position 68 and glutamic acid (E) at position 438 were mutated to histidine (H) and R, respectively. To generate constructs expressing RFP-GFP tagged in tandem to Atg8a (RFP-GFP-Atg8a), *atg8a* was first cloned into pDM317 using primers listed in Table S1 and inserted in between RFP and Atg8a.

Macropinocytosis and exocytosis assays

Macropinocytosis assay by microscopy imaging was routinely performed by incubating cells with 500 ng/µl TD (Sigma T1162) in HL5 for 1 h. For experiments presented in Fig. 2E, bacterially grown non-axenic cells were adapted in HL5 medium supplemented with 10% fetal bovine serum (Gibco 10091-148) for 24 h before incubating with TD in fresh medium. For experiments presented in Fig. 6A, cells cultured in SIH medium (Formedium) were shifted to SIH-K/R/E for 48 h before measuring macropinocytosis. For experiments presented in Fig. 6C, cells cultured in SIH were shifted to SIH-R or SIH-K for 48 h, followed by the addition of R (87.5 mg/l), R-R dipeptide (87.5 mg/l), K (156 mg/l), or K-K dipeptide (156 mg/l) for 24 h before measuring macropinocytosis. To quantify TD uptake, the medial optical section of cell was imaged using a Zeiss 880 inverted microscope equipped with a 63×1.4 oil immersion objective. Fluorescence intensity was quantified by ImageJ and normalized to the cell area.

Macropinocytosis dynamics was recorded in Ax2 and $slc15A^-$ cells expressing PHcrac-GFP by time-lapse imaging. The rate of macropinocytosis was quantified by counting the number of enclosed macropinosomes formed in 5 min. Macropinosome size was determined by measuring the area of newly formed macropinosomes immediately after ruffle closure with the freehand line tool in ImageJ.

Fluorimetric analysis of macropinocytosis and exocytosis was performed as described before (Rivero and Maniak, 2006). For macropinocytosis measurement, cells were shaken at a density of 5×10^6 /ml in growth medium, and TD was added to a final concentration of 1 mg/ml. Aliquots of 300 µl of cells were taken at each time point and mixed briefly with Trypan Blue solution (2 mg/ml in 20 mM citrate and 150 mM NaCl, pH 4.5) on ice. Cells were washed once with 1 ml ice-cold Sörensen buffer (14.6 mM KH₂PO₄, 2.0 mM Na₂HPO₄, pH 6.1) and resuspended in 0.5 ml Sörensen buffer. For exocytosis measurement, cells were first incubated with 1 mg/ml TD for 180 min, and were then washed and resuspended in fresh HL5 medium. Aliquots of 300 µl of cells were taken at each time point, washed and resuspended in 0.5 ml ice-cold Sörensen buffer. In both experiments, fluorescence was measured immediately using a Tecan Spark fluorescence spectrophotometer (544-nm excitation and 574-nm emission). Obtained values were normalized to protein content.

Phagocytosis assays

Expression of mCherry or GST-GFP was induced in *E. coli* strain BL21 (DE3) with 0.3 mM isopropyl β -D-1-thiogalactoside at 16°C overnight. Harvested bacteria were washed with Sörensen buffer. For flow cytometry analysis, cells were shaken at a density of 4×10^6 /ml in Sörensen buffer, and mCherry-expressing bacteria were added to a final density of 2×10^9 /ml. Aliquots of 0.5 ml were taken at each time point. Cells were washed with ice-cold Sörensen buffer. The total fluorescence intensity per cell was determined by a BD Biosciences Influx flow cytometer and data were analyzed by FlowJo. For bacteria killing assay, GFP-expressing bacteria were resuspended in HL5 medium to a concentration 2×10^8 /ml. 10 μ l of bacteria culture in 400 μ l HL5 was added into 8-well coverslip chambers before addition of 10^5 cells. Images were acquired every 14 s for a total of 100 frames.

Imaging

All microscopy experiments were performed at 22°C. To image the localization of fluorescent proteins in cells, 10^5 cells were plated in an 8-well coverslip chamber (Lab-Tek; NalgenNunc) and allowed to adhere. Images were taken on a Zeiss 880 inverted microscope equipped with a $40 \times /0.95$ or $63 \times /1.4$ oil-immersion objective. To image GFP-Atg8a signal, cells were incubated in HL5 or submerged in DB for 1 h before imaging. For experiments presented in Fig. S5A, the HL5 medium was supplemented with 2.5× protease inhibitor cocktail (Roche 11836170001) under the plus PIC condition.

For experiments presented in Fig. 7C, cells were plated in an 8-well coverslip chamber and allowed to adhere. Before imaging, the medium was replaced by fresh HL5 medium containing 1 mg/ml FD. For experiments presented in Fig. S4, cells were incubated in HL5 medium containing 1 mg/ml FD and 500 μ g/ml TD. FD and TD signals contained within the newly generated macropinosomes were quantified by ImageJ. The ratio of FD to TD was used to measure macropinosome acidification with the maximum ratio normalized to 1.

Quantification and statistical analysis

SuperPlots were generated as described previously (Lord et al., 2020). Statistical analysis was performed using GraphPad Prism. ***P<0.001, **P<0.01 and *P<0.05 (all figures).

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Competing interests

The authors declare no competing or financial interests.

Author contributions

Conceptualization: H.C.; Formal analysis: H.C., H.W., Y.Z., H.T., Z.G., L.L.; Investigation: Y.Z., H.T., H.W., Y.H., D.L., Y. Yang, Y. Yuan; Writing - original draft: H.C., H.W.; Writing - review & editing: H.C., H.W., Y.Z., H.T., L.L.; Supervision: H.C.; Project administration: H.C.; Funding acquisition: H.C., H.W., Y. Yang.

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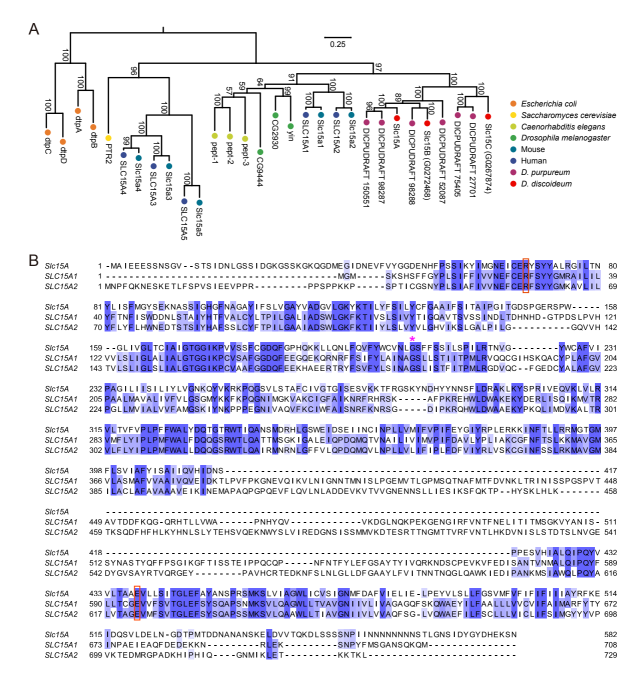


Fig. S1. Conservation of SLC15 family proteins. (A) Phylogenetic tree of the SLC15 family. Shown is a maximum likelihood tree built with LG+F+G4 model. Bootstrap values are from 1,000 iterations. In addition to Slc15A, *Dictyostelium* genome encodes two other proteins that are homologous to human SLC15 proteins. We named them Slc15B (DDB_G0272468) and Slc15C (DDB_G0267874). (B) Sequence alignment of Slc15A with human SLC15A1 and SLC15A2. Red boxes mark the two conserved residues, R68 and E438, in Slc15A. Mutation of the corresponding amino acid, R57H in SLC15A2 or E595R in SLC15A1, was shown to abolish the oligopeptide transport function of the protein. The magenta asterisk marks the position of REMI insertion.

J. Cell Sci.: doi:10.1242/jcs.259450: Supplementary information

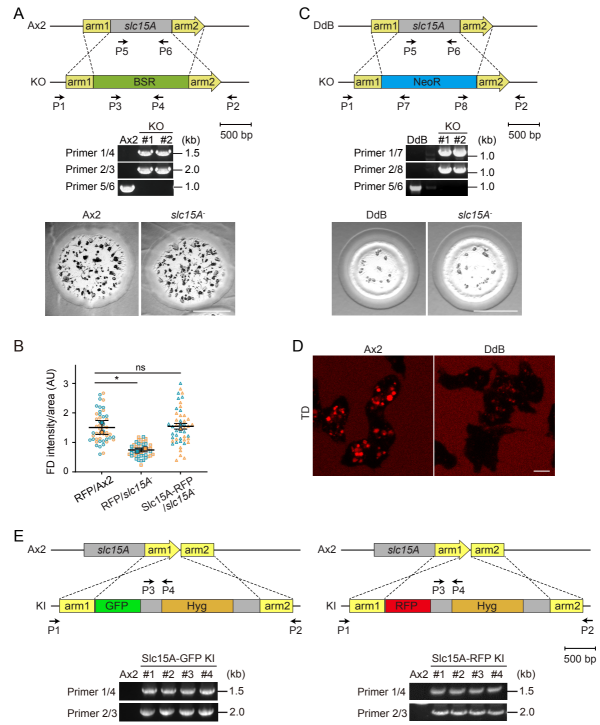


Fig. S2. Generation of *slc15A* **deletion and knock-in cells.** (A) Generation of *slc15A*⁻ cell line in the Ax2 background. Top: A blasticidin resistant cassette (BSR) was used to replace part of the open reading frame of *slc15A*, and targeted clones were verified by PCR. Bottom: Ax2 and *slc15A*⁻ cells were plated clonally with *K. aerogenes* on standard medium agar. Representative images were captured after 5 d. Scale bar, 5 mm. (B) Quantification of FD uptake in Ax2 and *slc15A*⁻ cells expressing RFP or Slc15A-RFP. Data are from two independent experiments with at least 20 cells quantified per experiment (each experiment is shown in a different color). Mean ± SD. Statistical significance was determined by one-way ANOVA with Dunnett post-test. (C) Generation of *slc15A*⁻ cell line in the DdB background. Top: A neomycin resistant cassette (NeoR) was used to replace part of the open reading frame of *slc15A*, and targeted clones were verified by PCR. Bottom: DdB and *slc15A*⁻ cells were plated clonally with *K. aerogenes* on standard medium agar. Representative images were captured after 4 d. Scale bar, 5 mm. (D) Confocal images of TD uptake in Ax2 and DdB cells. Scale bar, 5 µm. (E) Design of knock-in constructs. Slc15A-GFP^{KI} and Slc14A-RFP^{KI} cells were generated by targeted in-frame integration of GFP or RPF at the C-terminus of Slc15A. Targeted clones were verified by PCR.

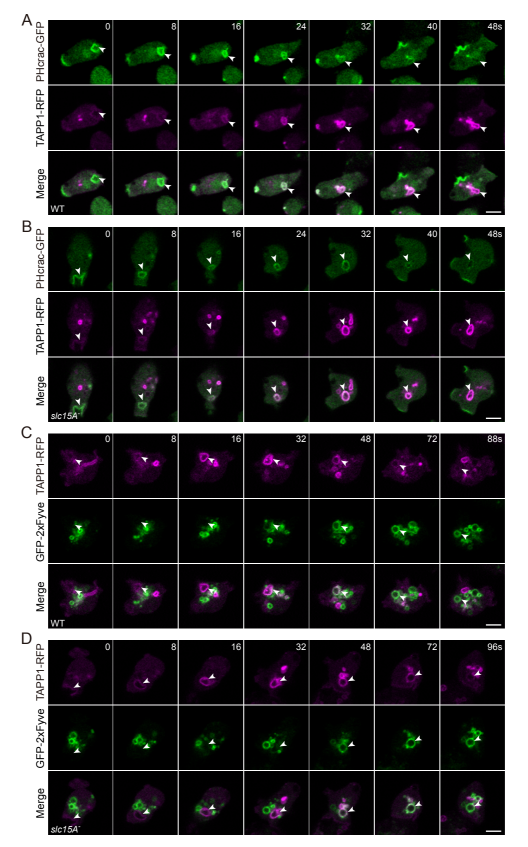
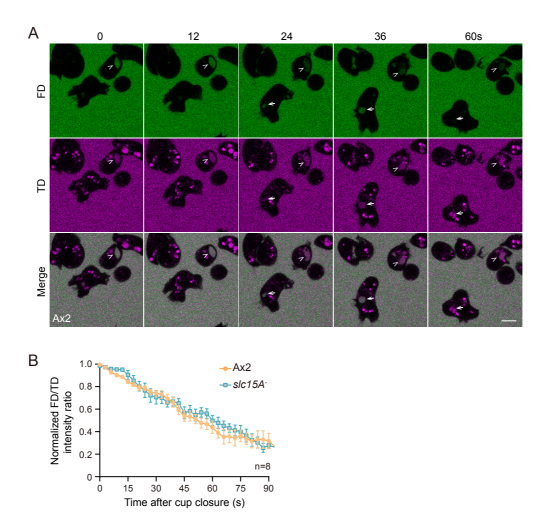
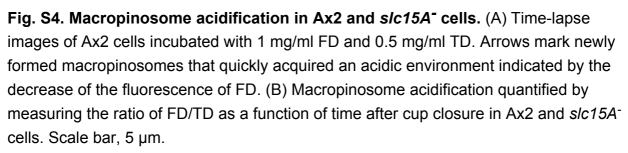


Fig. S3. PIP conversion during macropinocytosis in Ax2 and *slc15A*⁻ **cells.** (A-B) Timelapse images showing the sequential accumulation of PHcrac-GFP and TAPP1-RFP in Ax2 and *slc15A*⁻ cells. (C-D) Time-lapse images showing the sequential accumulation of TAPP1-RFP and GFP-2xFYVE in Ax2 and *slc15A*⁻ cells. Scale bars, 5 μm.





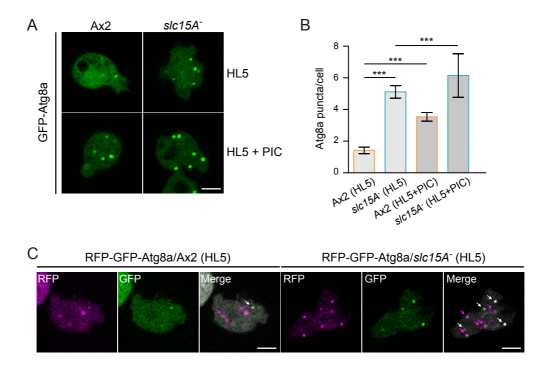


Fig. S5. Autophagy flux in Ax2 and *slc15A⁻* **cells.** (A) Confocal images of GFP-Atg8a. Cells were cultured in HL5 in the absence or presence of a protease inhibitor cocktail (PIC). (B) Quantification of average Atg8a puncta per cell. At least 25 cells were quantified per condition. Mean ± SD. The addition of PIC further increased the number of GFP-Atg8a puncta in *slc15A⁻* cells, indicating that autophagy flux was not blocked. (C) Confocal images of Atg8a fused to a tandem RFP-GFP tag. The presence of magenta-only puncta marked by the magenta arrows indicates that autophagy flux was not blocked in *slc15A⁻* cells. White arrows point to Atg8a structures with both GFP and RFP signals. Scale bars, 5 µm.

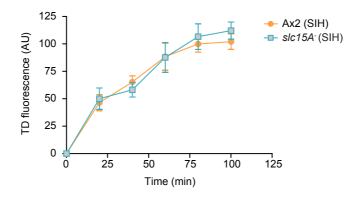


Fig. S6. Macropinocytosis in cells grown in SIH medium. Quantification of TD uptake in Ax2 and $s/c15A^{-}$ cells by fluorimetric analysis. Data are from four independent experiments. Mean \pm SD.

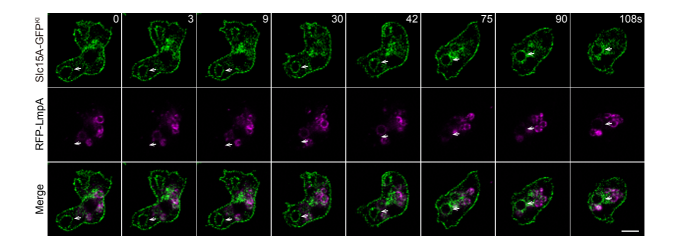


Fig. S7. SIc15A does not colocalize with the lysosomal/postlysosomal marker

LmpA. Time-lapse images of Slc15A-GFP^{KI} and RFP-LmpA. The arrows point to a newly generated macropinosome marked by Slc15A. Scale bar, 5 μ m.

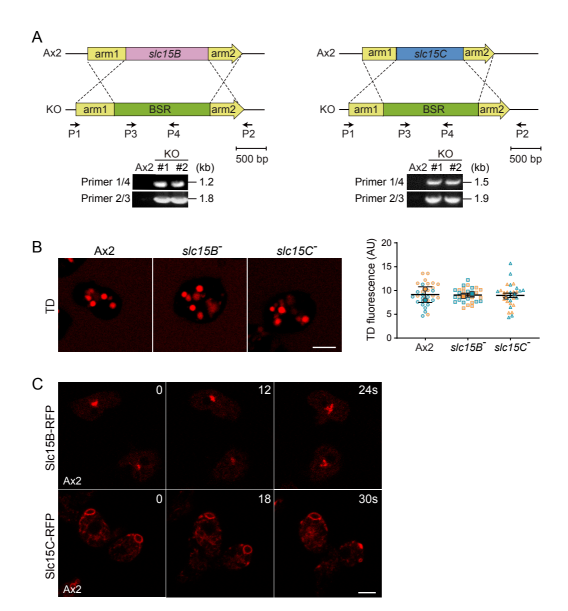
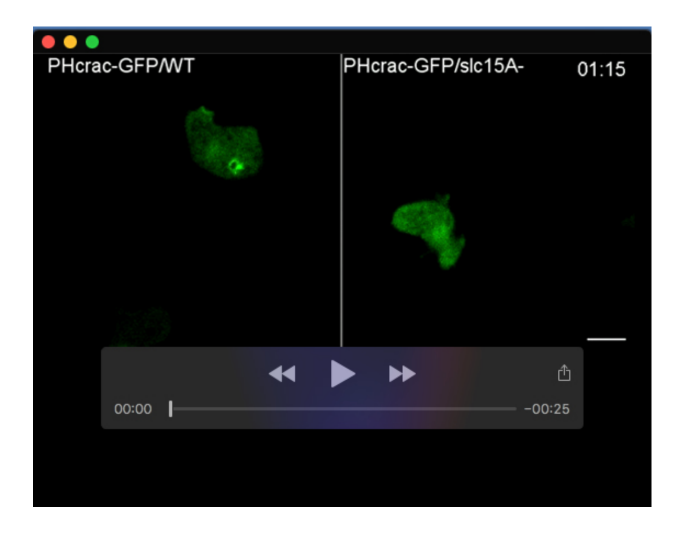


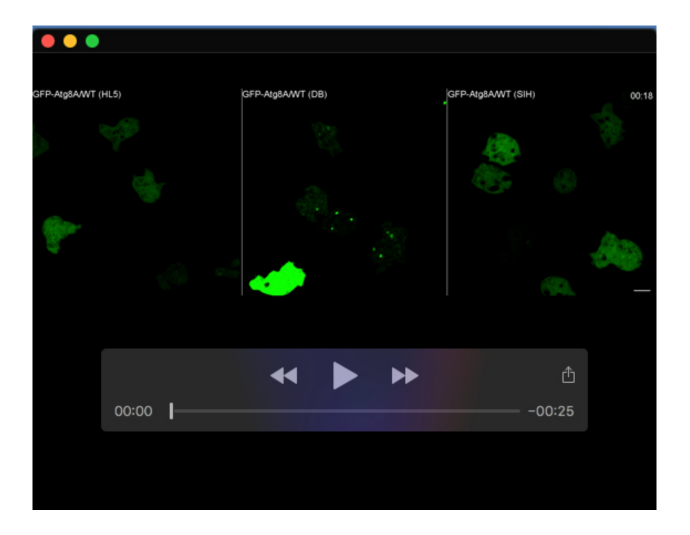
Fig. S8. Characterization of SIc15B and SIc15C. (A) Design of knockout constructs. The BSR was inserted via homologous recombination to replace part of the open reading frame of *slc15B* or *slc15C*. Targeted clones were verified by PCR. (B) Confocal images and quantification of TD uptake in Ax2, *slc15B*⁻, and *slc15C*⁻ cells. Data are from two independent experiments with at least 15 cells quantified per experiment (each experiment is shown in a different color). Mean \pm SD. (C) Time-lapse images of Slc15B-RFP and Slc15C-RFP expressed in Ax2 cells. Slc15B localizes to the Golgi and Slc15C localizes to the contractile vacuole structure. Scale bars, 5 µm.

Table S1. Plasmids and primers used in this study. Each primer is designated as forward (F) or reverse (R).

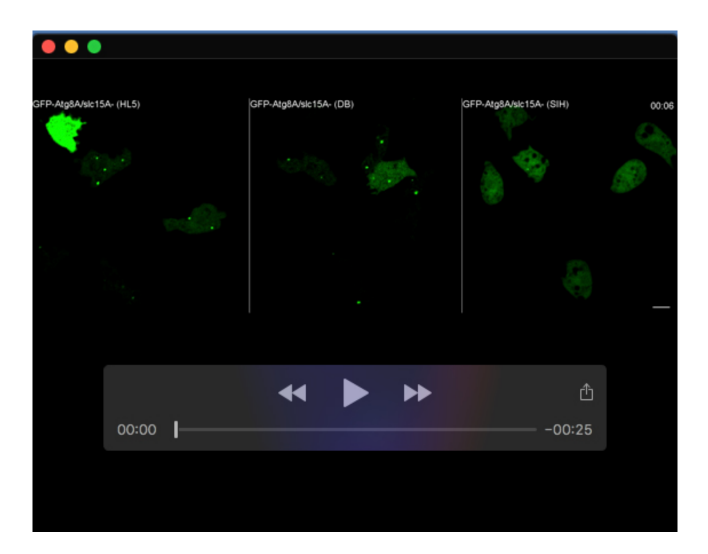
Usage	Plasmid backbone	Sequence, 5' - 3'
Expression in Dictyostelium	cells	
SIc15A-GFP	pDM323	F: GCTCTAGAATGGCGATAGAAGAAGAATCATCAAATTCT
	pbillozo	R: ATAAGAATGCGGCCGCATTTGATTTTCATGATCATAACCATAATCA
SIc15B-RFP	pDM324	F: GCTCTAGAATGGGTGGAATCGACGAAGATGG
		R: ATAAGAATGCGGCCGCTTTTGCATTTGATTTATTTGAATTTATTG
Slc15C-RFP	pDM324	F: GCTCTAGAATGAAAACAGACGATAAATATGCAAG
		R: ATAAGAATGCGGCCGCAATTAAATTATTTTCCAACTCTATAGATC
SIc15A ^{R68H/E438R} -GFP	pDM323	R68H F: AATTTGTGAACATTATTCATACTATGCTTT
		R68H R: TGAATATAGTTCACAAATTTCATTACCCAT
		E438R F: GTCTTGACTGCCGCTCGTGTTTTATTATCA
		E438R R: ACACGAGCGGCAGTCAAGACAACATATTGTG
GFP-Rab5A	pDM317	F: CGGGAGCTCATGAATAATAATAATAAGATATTTC
	F=	R: CGGACTAGTGTTACAACATTTGTTTTCTTTCC
GFP-Rab7A	pDM317	F: CGGGAGCTCATGGCCACAAAGAAAAAGG
		R: CGGACTAGTACAACAACCTGATTTAGCTGG
TAPP1-RFP	pDM324	F: CGGGAGCTCATGCCTTATGTGGATCGTCAGAATC
		R: CTAGCTAGCCACGTCACTGACCGGAAGGCTCGC
GFP-Atg8a	pDM317	F: CTAAGATCTAGTGCTGGTGGTATGGTTCATGTATCAAGCTTTAAA
		R: CTAACTAGTTAAATCACTACCAAAAGATTTTCACC
RFP-GFP-Atg8a	pDM449	F: CGGGAGCTCATGGTTCATGTATCAAGCTTTAAAAAC
		R: CGGACTAGTTTATAAATCACTACCAAAAGTATTTTC
		GFP F: TGCTCTAGAATGGGTAAAGGAGAAGAACTTTTCAC
		GFP R: GGAAGATCTGGATCTGAGTCCGGACTTGTATAG
VatB-GFP	pDM323	F: CGGGAGCTCATGGTTGGATTTGAAGATCATATCG
		R: CGGACTAGTGTTAGTTGAATCTACAGTACCCTTTG
RFP-LmpA	pDM318	F: CGGGAGCTCATGGTAAAAAGAGGGTGTTGCCATAGAAAAATG
		R:CGGACTAGTTGGTTCTGTTAAACGACGACGTAATTCTTCTTCTAAACGACTTG
		TTCATTATTAATAATTGCTTGATAAC
Generation of knockout cell		
<i>slc15A</i> knockout in Ax2	pBluescript-BSR	Insert 1 F: ACGGTCGACATGGCGATAGAAGAAGAATCATC
		Insert 1 R: ACGGAATTCAACACCATCAGCAACATAAGCAC
		Insert 2 F: ACGGGATCCGTGATTGCATTTTACATTTCAGC
		Insert 2 R: ACGACTAGTCAATACTATTACCCAAAGTTGAG
		Insert 1 F: GGGGCCGGCATGGCGATAGAAGAAGAATC
<i>slc15A</i> knockout in DdB	pDM1082	
		Insert 2 F: GGAAGATCTTGATTGCATTTACATTTC
		Insert 2 R: CGGACTAGTTCAATACTATTACCCAAAGTTGAG
		Insert 1 F: ACGCGTCGACATGGGTGGAATCGACGACGACGATGGT
<i>slc15B</i> knockout		Insert 1 R: CGGAATTCCTGCAATATATGCTCCAAATAATGTA
	pBluescript-BSR	
		Insert 2 R: ATAAGAATGCGGCCGCGCATTTGATTTATTTGAATTTATTGTGTAT
	pBluescript-BSR	
<i>slc15C</i> knockout		
		Insert 2 R: ATAAGAATGCGGCCGCAATTAAATTATTTTCCAACTCTATAGATC
Generation of knock-in cell		
s/c15A GFP and RFP knock-in	pDM1355	Insert 1 F: GGACTAGTCGTACCAATCTTTGAATATGGT
		Insert 1 R: GGACTAGTATTTGATTTTCATGATCATAACC
		Insert 2 F: GGAGCCGGCTCAATTTAAAAATAAAAACACTTTGTA
		Insert 2 R: GGAGCCGGCATGCAAATTCTCCATTACCTCA
Expression in <i>E.coli</i>		
mCherry	pET-15b	F: CGCCATATGATGGTGAGCAAGGGCGAGGAG
,	r=1 100	R: CCGCTCGAGGGGCCCCTTGTACAGCTCGTCC
GST-GFP	pGEX 4T-1	F: CCGGAATTCATGGTGAGCAAGGGCGAGGAG R: CCGCTCGAGCTTGTACAGCTCGTCCATGC



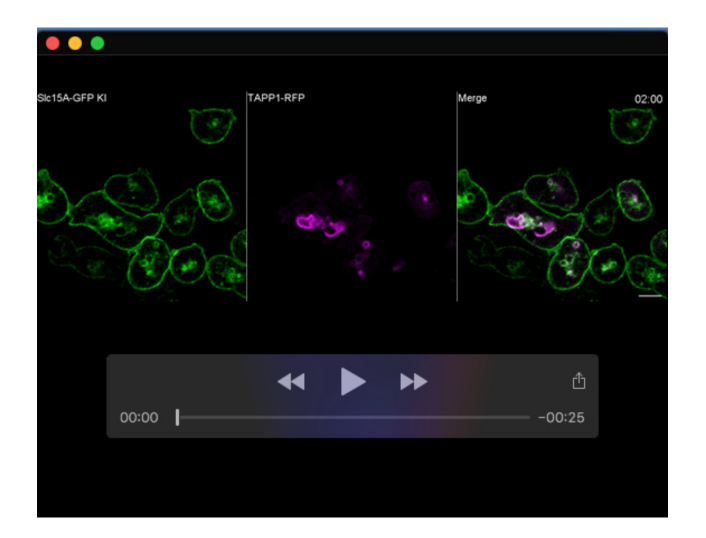
Movie 1. Time-lapse imaging of PHcrac-GFP dynamics in Ax2 and *slc15A*⁻ cells. Corresponds to figure 4A. Images were captured at 3 sec per frame and played back at 15 frames per second. Scale bar, 5 μ m.



Movie 2. Time-lapse imaging of GFP-Atg8a in Ax2 cells cultured in HL5 medium, SIH medium, or non-nutrient development buffer (DB). Corresponds to figures 4E-F and 5G. Images were captured at 6 sec per frame and played back at 1 frame per second. Scale bar, $5 \mu m$.



Movie 3. Time-lapse imaging of GFP-Atg8a in *slc15A*⁻ cells cultured in HL5 medium, SIH medium, or non-nutrient development buffer (DB). Corresponds to figures 4E-F and 5G. Images were captured at 6 sec per frame and played back at 1 frame per second. Scale bar, 5 μ m.



Movie 4. Colocalization of SIc15A-GFP^{KI} with the PI(3,4)P₂ sensor TAPP1-RFP. Corresponds to figure 7A. Images were captured at 15 sec per frame and played back at 6 frames per second. Scale bar, 5 μ m.