Keywords: autophagy, autophagy flux, autophagy response, Cyto-ID, RNA interference screen, small molecule screen, spectrophotometric assay

Abbreviations: 3-MA, 3-methyladenine; FBS, fetal bovine serum; GFP, green fluorescent protein; LAMP1, lysosomal-associated membrane protein 1; MAP1LC3B/LC3B, microtubule-associated protein 1 light chain 3 beta; MAP3K6, mitogen-activated protein kinase kinase kinase 6; MDC, monodansylcadaverine; mRFP, monomeric red fluorescent protein; MTOR, mechanistic target of rapamycin; NS, nonsilencing; RAB5A, member RAS oncogene family; RNAi, RNA interference; shRNA, short-hairpin RNA; SQSTM1, sequestosome 1; WNK2, WNK lysine deficient protein kinase 2

© Sujuan Guo, Yanping Liang, Susan F Murphy, Angela Huang, Haihong Shen, Deborah F Kelly, Pablo Sobrado, and Zhi Sheng.

Correspondence to: Zhi Sheng, E-mail: zhisheng@vtc.vt.edu

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The lack of a rapid and quantitative autophagy assay has substantially hindered the development and implementation of autophagy-targeting therapies for a variety of human diseases. To address this critical issue, we developed a novel autophagy assay using the newly developed Cyto-ID fluorescence dye. We first verified that the Cyto-ID dye specifically labels autophagic compartments with minimal staining of lysosomes and endosomes. We then developed a new Cyto-ID fluorescence spectrophotometric assay that makes it possible to estimate autophagy flux based on measurements of the Cyto-ID-stained autophagic compartments. By comparing to traditional autophagy approaches, we found that this assay yielded a more sensitive, yet less variable, quantification of the stained autophagic compartments and the estimate of autophagy flux. Furthermore, we tested the potential application of this autophagy assay in high throughput research by integrating it into an RNA interference (RNAi) screen and a small molecule screen. The RNAi screen revealed WNK2 and MAP3K6 as autophagy-modulating genes, both of which inhibited the MTOR pathway. Similarly, the small molecule screen identified sanguinarine and actinomycin D as potent autophagy inducers in leukemic cells. Moreover, we successfully detected autophagy responses to kinase inhibitors and chloroquine in normal or leukemic mice using this assay. Collectively, this new Cyto-ID fluorescence spectrophotometric assay provides a rapid, reliable quantification of autophagic compartments and estimation of autophagy flux with potential applications in developing autophagy-related therapies and as a test to monitor autophagy responses in patients being treated with autophagy-modulating drugs.

Introduction

Under stress conditions such as nutrient deficiency or drug treatments, cells often undergo macroautophagy (referred to as autophagy hereafter), a process involving the genesis of various subcellular compartments such as autophagosomes and autolysosomes. The cellular components that are engulfed in these vesicles are eventually degraded in autolysosomes. Such “self-digestion” usually provides cells with more nutrients for continuous survival; however, excessive digestion also results in cell death. Given its essential role in cell survival and death, deregulated autophagy is often central to the pathogenesis of many human diseases such as neurodegenerative diseases, cancer, infectious diseases, heart failure, and diabetes, to name a few.2,3 And as such, it has been widely used as a target for therapeutic intervention. Nevertheless, the lack of an accurate and quantitative approach to
measure autophagy has significantly discouraged such effort. For example, hydroxychloroquine\(^4\) and everolimus\(^5\) that target autophagy pathways were used in the clinic to treat cancer patients. The efficacy was encouraging but unexpectedly low. Many compounds have been found to control autophagy in cultured cells or even animals but their further application in clinic is limited.\(^2\) One of the major reasons is that the diagnostic tools monitoring autophagy and hence allowing for dose adjustment in clinical treatment are lacking.

To date, many autophagy-detecting approaches have been devised; only a few of them are suitable for quantitative and high throughput analyses, yet are still inaccurate and labor-intensive.\(^6,7\) For instance, LC3B (microtubule-associated protein 1 light chain 3 beta) serves as a specific marker for autophagosomes because the protein level of LC3B-II, the cleaved and lipidated form that is conjugated with phosphatidylethanolamine and subsequently incorporated onto the autophagosome membrane, positively correlates to autophagosome production.\(^8\) However, such correlation is generally—but not always—found in most experimental settings.\(^6\) In addition, quantification of LC3B-II protein requires a comparison to housekeeping genes such as actin, the expression level of which is sometimes subject to change during autophagy.\(^7\) LC3B conjugated with green fluorescent protein (GFP) is also frequently used in fluorescence microscopy. The pitfall for this approach is that ectopically expressed GFP-LC3B often forms aggregates that are difficult to distinguish from the GFP-LC3B puncta, the characteristic subcellular structures that refer to autophagosomes under a fluorescence microscope.\(^9\) Electron microscopy, the most conventional—yet still a standard—method in detecting autophagy, acquires images of early or late autophagic compartments in cells but often fails to provide quantitative data and is infeasible for clinical application.\(^7\) Other approaches such as staining using acidotropic fluorescent dyes or analyzing autophagy substrate degradation also have obvious caveats and low specificity, limiting their application in quantifying autophagy.\(^6,7\) Taken together, there is an unmet and increasing need for developing novel approaches that accurately and rapidly quantify autophagy.

Cyto-ID, a recently developed cationic amphiphilic tracer dye, labels autophagic compartments with minimal staining of lysosomes and endosomes,\(^10,11\) suggesting that it is a specific autophagy marker. Here we took advantage of this fluorescent dye and developed a Cyto-ID-based fluorescence spectrophotometric assay (abbreviated as the Cyto-ID assay) that permits high content performance and measures the size of autophagic compartments which makes it possible to monitor autophagy flux. Compared to other traditional autophagy-detecting approaches, this assay is considerably more sensitive, reliable, and time-saving thereby providing a platform for

**Figure 1.** The Cyto-ID dye specifically labeled autophagic compartments with minimal staining of lysosomes and endosomes. Colocalization of the Cyto-ID fluorescence dye, Hoechst 33342, and mCherry-LC3B (A), LAMP1-mRFP (B), or mRFP-RAB5A (C) in HeLa cells treated without or with 10 \(\mu\)M PP242 for 4 h.
drug screening and has a potential clinical application as a diagnostic tool to gauge autophagy responses in patients.

**Results**

**The Cyto-ID dye specifically labels autophagic compartments**

Previous reports showed that Cyto-ID stains autophagic cells through labeling autophagic compartments.\(^1\)\(^2\) During autophagy, autophagosomes also fuse with endosomes to establish another intermediate—amphisomes—before fusion with lysosomes.\(^1\) Thus, to validate that Cyto-ID marks most autophagic compartments including amphisomes and autolysosomes, we transfected HeLa (human cervical cancer) cells with plasmids encoding mCherry-LC3B (a phagophore, autophagosome or amphisome marker), LAMP1-RFP (a lysosome marker),\(^1\)\(^2\) or mRFP-RAB5A (an endosome marker)\(^1\)\(^3\) and then monitored the colocalization of Cyto-ID with these fluorescent proteins. In untreated HeLa cells (Fig. 1A to C, top panels), mCherry-LC3B was distributed evenly in cells, whereas LAMP1-mRFP or mRFP-RAB5A aggregated as small dots representing lysosomes or endosomes. By contrast, no or very weak Cyto-ID fluorescence was detected in these nonautophagic cells indicating that Cyto-ID has minimal staining of lysosomes or endosomes. However, Cyto-ID formed puncta (a characteristic of autophagic vesicles) and partially colocalized with mCherry-LC3B, LAMP1-RFP, or mRFP-RAB5A in HeLa cells treated with PP242, a selective inhibitor of MTOR (mechanistic target of rapamycin) that induces autophagy\(^1\)\(^4\) (Fig. 1A to C, bottom panel). These data indicate that Cyto-ID labels autophagic compartments such as amphisomes or autolysosomes.

**The Cyto-ID dye, but not EGFP-LC3, is suitable for spectrophotometry**

Similar to other fluorescent dyes, the Cyto-ID is sensitive to light.\(^1\)\(^0\) To compare the light sensitivity of Cyto-ID to EGFP-LC3 (a widely used autophagy marker), we imaged autophagic HeLa cells using the Cyto-ID or EGFP-LC3 at different bleaching times. We found that the sensitivity of Cyto-ID to the light was equivalent to that of EGFP-LC3 (Fig. 2A and B). To further determine the stability of Cyto-ID in spectrophotometry, we compared HeLa cells stained with Cyto-ID with those stably expressing EGFP-LC3 using a FilterMax F3 microplate reader. The fluorescence intensities of Cyto-ID remained unchanged in DMSO and PP242-treated cells at least 15 min after staining (Fig. 2C, right panel) and so did the intensities of EGFP-LC3 (Fig. 2C, left panel). However, the Cyto-ID, but not EGFP-LC3, was dramatically increased in PP242-treated cells compared to their respective controls indicating that the Cyto-ID dye is more suitable than EGFP-LC3 for autophagy assay using spectrophotometry.

**The Cyto-ID assay measures changes in the size of autophagic compartments induced by various autophagy activators**

We next developed an expeditious Cyto-ID-based spectrophotometric assay...
To minimize the background signal of Cyto-ID, we attempted to normalize the results from the Cyto-ID assay using relative cell densities measured by the MTS or Cell-Titer-Blue assay. We then measured Cyto-ID-positive autophagic compartments after induction by drugs or starvation in several cell lines using this assay. In K562 human leukemic cells treated with imatinib at the dose (2 μM) that has been shown to induce autophagy (detected by LC3B immunoblotting)15 Cyto-ID fluorescence was increased by more than 3 fold (Fig. 3B). In HEK293 (human embryonic kidney epithelial) or OVCAR8 (ovarian cancer epithelial) cells, PP242 treatment induced a 2- or 4-fold increase of Cyto-ID fluorescence, respectively, (Fig. 3C). In K562 or HEK293 cells treated with nutrient deprivation, Cyto-ID fluorescence was increased by 2- or 4-fold, respectively (Fig. 3D). To further validate that the elevated level of Cyto-ID fluorescence refers to autophagy induction, we treated cells with 3-methyladenine (3-MA), an inhibitor of autophagic sequestration.15 As expected, 3-MA blocked the induction of Cyto-ID fluorescence in K562 treated with imatinib (Fig. 3E) and HEK293 or OVCAR8 cells treated with PP242 (Fig. 3F). Together these results demonstrate that the Cyto-ID assay quantitatively monitors the changes in the size of autophagic compartments induced by various autophagy activators in both normal and malignant cells.

**The Cyto-ID assay provides an estimate of autophagy flux**

Autophagy is a dynamic, multistep process including autophagosome formation, autolysosome formation, and degradation of autophagic substrate (often denoted as autophagy flux). In fact, monitoring autophagy flux is the most relevant way for the estimation of autophagic activity.6,7 The commonly used assay for monitoring autophagy flux is the turnover of LC3B or SQSTM1/p62 using immunoblotting, which measures the content through autophagy flux; however, this approach is time-consuming and the results are often varied in different experimental settings and hard to interpret. Thus we tested whether the Cyto-ID assay provides a rapid analysis of autophagy flux, because the quantity of autophagic compartments usually correlates with the level of autophagy flux.

We first monitored the dynamic change of autophagy flux in K562 cells. The activated autophagy flux is characterized by the increased amount of autophagic compartments and the reinforced degradation of autophagic cargos and their receptors such as SQSTM1.7 As shown in **Figure 4A and B**, the levels of Cyto-ID...
and LC3B-II increased in K562 cells treated with imatinib over time, coinciding with the decreased level of SQSTM1. We then estimated the impaired autophagy flux by monitoring the accumulation of autophagic compartments induced by chloroquine, a lysosome inhibitor that impedes the fusion of autophagosomes and lysosomes and/or the activity of autolysosomes.\(^1^6\) As expected, chloroquine increased the level of Cyto-ID fluorescence by a factor of approximately 2.5-fold in all the cells tested (Fig. 4C). To further monitor the dynamic change of impaired autophagy flux, we treated K562 cells with chloroquine at different time points. The Cyto-ID fluorescence (Fig. 4D) or LC3B-II level (Fig. 4E) increased over time. Although we expected an increase of SQSTM1 due to the impaired autophagy flux, we found that the SQSTM1 protein level remained relatively unchanged (Fig. 4E). Taken together, our data demonstrates that the Cyto-ID assay provides a fast analysis that estimates autophagy flux.

The Cyto-ID assay determines the status of autophagy flux

Monitoring autophagy flux is difficult because the increase of autophagic compartments at a steady state during activated autophagy is often indistinguishable from the accumulation of autophagic compartments resulting from impaired autophagy flux.\(^6^,^7\) We then sought to determine whether the Cyto-ID assay could be used as a surrogate of the LC3B immunoblotting assay to distinguish activated autophagy from impaired autophagy flux. To test this hypothesis, we treated K562, HEK293, and OVCAR8 cells with imatinib (or PP242) and chloroquine. In principle, imatinib or PP242 (that activates the formation of autophagic vesicles), when combined with chloroquine (that leads to the accumulation of autophagosomes and/or autolysosomes), would substantially increase the levels of autophagic compartments. In order to minimize the cell damage caused by the conjunctive treatment, we intentionally used the low doses of imatinib, PP242, or chloroquine that failed to significantly induce Cyto-ID fluorescence (Fig. 5A and C). As expected, the combination of imatinib and chloroquine, but not each treatment alone, induced a 2.5-fold increase of the Cyto-ID fluorescence in K562 cells (Fig. 5A). Similar results were obtained in HEK293 and OVCAR8 cells (Fig. 5C). The synergistic effect was confirmed in K562 cells treated with a combination of imatinib and chloroquine when using the LC3B immunoblotting assay (Fig. 5B). However, LC3B-II level was not significantly enhanced in HEK293 and OVCAR8 cells treated with PP242 and chloroquine (Fig. 5D), suggesting that the LC3B immunoblotting assay might not be suitable for assessing autophagy flux in these cells. Thus, the Cyto-ID assay can be used to determine whether autophagy flux is activated or impaired.

The Cyto-ID assay yields a more sensitive, yet less variable, analysis for autophagy

We next sought to determine the sensitivity of the Cyto-ID assay by comparing...
Autophagy research are those using LC3B and allows a robust and accurate quantification for monitoring autophagy and estimating autophagy flux, we probed possible applications of this assay in basic science research e.g. screening drug targets or identifying small molecule inhibitors that target autophagy. We tested the Cyto-ID assay in 2 high-throughput screens. First, we integrated the Cyto-ID assay with a short hairpin RNA (shRNA)-based RNA interference (RNAi) screen; 88 shRNAs against 28 human kinase genes were introduced into K562 cells via viral transduction. The subsequent analysis using the Cyto-ID assay revealed that 4 shRNAs (directed against ANK1, NTRK3, WNK2, or MAP3K6) enhanced the Cyto-ID fluorescence by more than 2-fold (Fig. 8A). Among the 4 candidates, PP242 at various time points and measured autophagic compartments using either the Cyto-ID assay or LC3B immunoblotting. We found that both assays detected a significant induction of autophagy within 2 h as indicated by a more than 2-fold increase of Cyto-ID fluorescence (Fig. 7A) or LC3B-II (Fig. 7B) in HEK293 and OVCAR8 cells. Surprisingly, after HEK293 and OVCAR8 cells were treated with PP242 for 4 h, the LC3B-II protein level dropped remarkably (Fig. 7B). In contrast, an exponential increase of Cyto-ID fluorescence was detected throughout the entire time course in both cell lines (Fig. 7A). In K562 cells treated with imatinib, a time-dependent linear increase of the Cyto-ID fluorescence was detected (Fig. 4A). However, LC3B protein levels (detected by the LC3B immunoblotting assay) increased modestly in K562 cells with the same treatment (Fig. 4B). These data suggest that the Cyto-ID assay is equally sensitive to the LC3B immunoblotting assay, yet it is more stable.

Taken together, our results show that the Cyto-ID assay yields a more sensitive and reliable quantification of autophagic compartments and is therefore better than the conventional autophagy detecting approaches.

The Cyto-ID assay has a broad spectrum of applications in basic research and clinical diagnosis

Since the Cyto-ID assay is timesaving and allows a robust and accurate quantification for monitoring autophagy and estimating autophagy flux, we probed possible applications of this assay in basic science research e.g. screening drug targets or identifying small molecule inhibitors that target autophagy. We tested the Cyto-ID assay in 2 high-throughput screens. First, we integrated the Cyto-ID assay with a short hairpin RNA (shRNA)-based RNA interference (RNAi) screen; 88 shRNAs against 28 human kinase genes were introduced into K562 cells via viral transduction. The subsequent analysis using the Cyto-ID assay revealed that 4 shRNAs (directed against ANK1, NTRK3, WNK2, or MAP3K6) enhanced the Cyto-ID fluorescence by more than 2-fold (Fig. 8A). Among the 4 candidates,
MAP3K6 and WNK2 were further validated by a significant increase of LC3B-II using LC3B immunoblotting assay (Fig. 8B) and an efficient knockdown of the target genes (Fig. 8C). To further determine the stage of autophagy that these 2 kinases target, we treated K562 cells with a combination of chloroquine and shRNAs against WNK2 or MAP3K6. The combinations substantially increased the Cyto-ID fluorescence compared to each treatment alone (Fig. 8D) suggesting that these 2 kinases target the initiation rather than maturation stage. Moreover, our work further showed that these 2 kinases suppressed autophagy by activating MTOR as knocking them down remarkably repressed both activity and expression of MTOR (Fig. 8E). Of note, WNK2 has previously been identified as an autophagy modulator from another RNAi screen.19

We next applied the Cyto-ID assay in a small molecule screen to identify drugs that potently modulate autophagy. We first selected the concentrations of chemical compounds and treatment duration based on PP242-induced autophagy. By virtue of its strong capability to induce autophagy, we hypothesized that any drugs or chemicals that are stronger than PP242 should be regarded as potent autophagy modulators. In K562 cells, 10 μM PP242 increased the Cyto-ID fluorescence by 1.5-fold within 4 h (Fig. 9A); this setting was then used to screen 80 chemical compounds composed of FDA-approved drugs or bioactive natural products. Due to the auto green fluorescence, we excluded fluorescein from this library albeit it showed positive results. The Cyto-ID analysis showed that 2 compounds (sanguinarine and actinomycin D) led to a > 2-fold increase of Cyto-ID fluorescence (Fig. 9B). Our validation experiments showed that actinomycin D or sanguinarine enhanced the Cyto-ID fluorescence in a dose-dependent manner (Fig. 9C and D). Furthermore, combinations of these 2 chemical compounds with chloroquine showed significant elevation of the Cyto-ID fluorescence compared to each treatment alone (Fig. 9E and F). Thus, we validated that actinomycin D and sanguinarine are potent autophagy inducers. Notably, actinomycin D is known to be a potent autophagy modulator.20 Collectively, our results demonstrate that the Cyto-ID assay is suitable for high content screening to identify new genes or chemical compounds that modulate autophagy.

As described earlier, autophagy is an appealing drug target for many different human diseases but current autophagy-related therapies yield unsatisfactory outcomes in clinic. One of the major hurdles appears to be the lack of accurate and quantitative approaches to monitor autophagy. We therefore explored the potential of this autophagy assay to be used clinically to monitor the effectiveness of autophagy-related therapies by testing it in normal mice or a mouse leukemia model. To monitor autophagy in primary peripheral blood cells, we treated mice with either PP242 or chloroquine by intraperitoneal injection and measured autophagy at different time points. Two out of 3 PP242 or chloroquine-treated mice showed a significant increase of Cyto-ID fluorescence 4 h after drug administration. In contrast, no induction was observed in the untreated mice (Fig. 10A). However, we noticed different drug responses among mice and at various time points in the same mouse, further
Autophagy-related therapies in the clinic. To mimic clinical application, we generated leukemic mice and then treated them with either imatinib or chloroquine. In mice bearing BCR-ABL-driven leukemia, we found that both imatinib and chloroquine significantly increased the level of Cyto-ID fluorescence in mouse primary bone marrow cells (Fig. 10B). Taken together, our results demonstrate that the Cyto-ID assay can be extremely valuable as a diagnostic tool to monitor dose responses of autophagy-related therapies in the clinic.

Discussion

The caveats in current autophagy-detecting approaches (reviewed in refs. 6 and 7) make it difficult to quantitatively monitor autophagy or autophagy flux. Here we described a Cyto-ID-based fluorescence spectrophotometric assay that measures changes in the size of autophagic compartments rapidly and accurately. This assay also provides an estimate of autophagy flux and helps determine the status of autophagy flux at a steady state. The advantages of this assay are as follows:

1) the Cyto-ID fluorescence dye, unlike other dyes used in autophagy assays, specifically marks autophagic compartments with minimal staining of lysosomes or endosomes (Fig. 1) and thereby has much lower background staining in nonautophagic cells (Fig. 6A); 2) the Cyto-ID is a more appropriate fluorescent dye for spectrophotometry than GFP-LC3 (Fig. 2C); 3) the Cyto-ID assay is more sensitive and reliable than the MDC spectrophotometric assay (Fig. 6B) or LC3B immunoblotting assay (Fig. 7); 4) the Cyto-ID assay rapidly monitors autophagy flux (Fig. 4); 5) the Cyto-ID assay can be used as a surrogate for LC3B immunoblotting assay to distinguish activated and impaired autophagy flux at a steady state (Fig. 5).

LC3B is a well-established autophagy marker and widely used in monitoring autophagy. However, we found that Cyto-ID provides a more reliable detection of changes in autophagic activity than LC3B. One pitfall of the LC3B-based assays is that they measure autophagosome only, whereas the Cyto-ID dye labels most autophagic compartments (Fig. 1). Another problem with LC3B immunoblotting is that the LC3B-II protein level might not be as stable as we expected. Results shown in Figure 7B suggested that LC3B-II protein is not stable 4 h after PP242 treatment. Moreover, the Cyto-ID assay, unlike LC3B-based assays, provides precise numeric readouts that correlate with the formation of autophagic compartments, and therefore, it is perhaps the most accurate quantitative assay among all autophagy assays developed so far.

Autophagy flux is a dynamic process that is difficult to be quantitatively monitored. Current approaches such as monitoring the turnover of LC3 or SQSTM1 or other autophagy cargo receptors using immunoblotting are time-consuming, not quantitative, and hard to interpret. Although the band intensity of these proteins can be measured using image software, the pixel limit of these immunoblots often provides inaccurate quantification. While the degradation of autophagic cargos or cargo receptors presents direct evidence of autophagy flux, the quantity of autophagic compartments also correlates with the status of autophagy flux (activated or impaired). We found that the Cyto-ID assay measured the quantities of autophagic compartments at different time points during activated or impaired autophagy flux with no limitations (Fig. 4A, 4D, and 7A). Thus, this assay yields a fast and quantitative estimate of autophagy flux. Furthermore, similar to the LC3B immunoblotting assay that is frequently used in determining the activated or impaired autophagy flux, the Cyto-ID assay can be used as an improved surrogate for this purpose as it is faster and simpler.

There are several high-throughput studies that have been reported recently.19,21-27 In these studies, fluorescence microscopy using GFP-LC3 as a reporter was used as a standard approach to monitor autophagy. However, this approach demands a special fluorescence microscope that permits a high content analysis, and is time-consuming and expensive. In this report, we successfully integrated the Cyto-ID assay in an RNAi screen (Fig. 8) and a small molecule screen (Fig. 9) and identified autophagy-modulating genes or drugs, some of which have been revealed as autophagy regulators in previous reports.19,20 Given that the Cyto-ID assay is rather simple and
affordable, more researchers in different fields can easily access it for large-scale autophagy studies.

Monitoring autophagy-related therapies is also difficult in the clinic as many autophagy-detecting assays are time-consuming and/or expensive. Failure in measuring autophagy in clinical samples may result in the ineffectiveness of autophagy-related therapies. The Cyto-ID assay provides us an opportunity to circumvent such difficulty. We have shown that this assay measured the size of autophagic compartments and estimated autophagy flux in peripheral blood or bone marrow cells freshly isolated from normal or leukemic mice (Fig. 10). Since the Cyto-ID assay is rapid, quantitative, and inexpensive, we will further adapt this assay into a diagnostic tool and expect that this tool will be widely implemented into clinical treatments of diverse human diseases.

**Materials and Methods**

**Materials**

HeLa, HEK293, and K562 cells were purchased from the American Type Culture Collection (CCL2, CRL-1573, CCL-243). OVCAR8 cells were kindly provided by Dr. Michael Green at the University of Massachusetts Medical School. The Cyto-ID kit was purchased from Enzo Life Sciences (ENZ-51031-K200). The Cell-Titer 96® Aqueous One solution cell proliferation assay (MTS; G3580) and the CellTiter-Blue® Cell Viability Assay kits (G8080) were purchased from Promega. LC3B (2775), MTOR (2972), phospho-MTOR (2971), or ACTB/β-actin (A5441) antibody was purchased from Cell Signaling Technology or Sigma-Aldrich, respectively. SQSTM1/p62 (sc-28359) antibody was purchased from Santa Cruz Biotechnology, Inc. Plasmids mCherry-hLC3B-pcDNA3.1 (40827), LAMP1-RFP (1817), mRFP-RAB5A (14437), EGFP-LC3 (11546) were obtained from Addgene. PP242 (P0037), chloroquine (C6628) and MDC (30432) were purchased from Sigma-Aldrich. 3-methyladenine (3-MA; 189490) was purchased from EMD Millipore Corporation. Imatinib (I-5508) was purchased from LC Laboratories. Actinomycin D (11421) was purchased from Cayman Chemical Company. Sangunarine (310035) was purchased from Microsource Discovery Systems, Inc. The kinase shRNAs were purchased from Thermo Scientific (RHS4884). The chemical compounds for the small molecule screen was provided by the Virginia Tech Center for Drug Discovery (Blacksburg, VA).

**Cell culture**

HeLa and HEK293 cells were maintained in Dulbecco’s modified Eagle’s medium (Life Technologies Corporation, 12800-017) supplemented with 10% fetal bovine serum (FBS) (Atlas Biologicals, Inc., F-0500-A), streptomycin (100 µg/ml) and penicillin (100 IU/ml). OVCAR8 and K562 cells were maintained in RPMI 1640 media (Life Technologies Corporation, 1640-0020).
Corporation, 23400-021) supplemented with 10% FBS, streptomycin (100 μg/ml) and penicillin (100 IU/ml).

**Autophagy induction and autophagy flux inhibition**

To induce autophagy, HeLa, HEK293, and OVCAR8 cells were treated with PP242 (2 to 20 μM) for 0 to 24 h. K562 cells were incubated with imatinib (either 1 or 2 μM) or PP242 (10 μM) for 0 to 22 h. To induce autophagy by starvation, K562 or HEK293 cells were washed 3 times with PBS (137 mM NaCl, 2.7 mM KCl, 10 mM Na<sub>2</sub>HPO<sub>4</sub>, 1.8 mM KH<sub>2</sub>PO<sub>4</sub>) and incubated in Earle's Balanced Salt Solution (Life Technologies Corporation, 14155063) for various time points. To block autophagy flux, cells were incubated with chloroquine (2.5 to 20 μM) for 0.5 to 6 h.

**Fluorescence spectrophotometric assay**

To stain cells with Cyto-ID or MDC, we followed the manufacturer's protocol with modifications. In brief, cells were first washed with PBS (supplemented with 5% FBS) and then mixed with either Cyto-ID or MDC staining solution. The cells were then incubated at 37°C for 30 min (Cyto-ID) or 10 min (MDC) in the dark, then washed twice with PBS (without FBS) to remove the free dyes. The cells were then divided: one half for the measurement of Cyto-ID fluorescence; the other half for the MTS cell viability assay described below. In some experiments, cells were transfected with pEGFP-LC3 and the fluorescence of EGFP-LC3 was recorded using the method described below. To use the CellTiter-Blue assay to measure cell viability, the reagent was directly added into the cells after the Cyto-ID or MDC assay without further dividing the cells. To measure the Cyto-ID or EGFP-LC3 fluorescence, cells were first plated in a black round-bottom 96-well plate (Fisher Scientific, 06-443-2) and incubated at room temperature for 5 to 20 min in the dark. The Cyto-ID or EGFP-LC3 green fluorescence was read using the FilterMax F3 microplate reader (at excitation 480 nm, emission 530 nm; Molecular Devices, Sunnyvale, CA). MDC was read at excitation 350 nm and emission 520 nm. Based on the manufacturer's datasheet, this microplate reader reads a 96-well plate in 18 sec (less than 0.2 sec per well). Such a short exposure minimizes the light bleaching. The relative Cyto-ID or MDC fluorescence is defined as the ratio of Cyto-ID or MDC readings to MTS readings or CellTiter-Blue readings (see below). The fold change of the Cyto-ID or MDC fluorescence was defined as the ratio of the relative Cyto-ID or MDC fluorescence in the treated cells to that in the control cells.

**Cell viability assay**

To minimize the difference of cell density among each sample, we determined the relative number of live cells using 2 cell viability assays: MTS or CellTiter-Blue. Cells with various treatments in 100 μl culture media were plated (5000 to 10000 cells per well) in a 96-well plate. Either MTS (10 μl) or CellTiter-Blue (10 μl) was added to each well, then incubated at 37°C for 1 h. The absorbance values of MTS (490 nm) or the fluorescence intensity of CellTiter-Blue (excitation 560 nm, emission 590 nm) was measured using the FilterMax F3 microplate reader according the manufacturer's directions.
instructions. These readings were used to normalize the Cyto-ID fluorescence of each corresponding sample.

**Fluorescence microscopy and quantification of fluorescence**

To label autophagic compartments or normal vesicular structures such as endosomes or lysosomes, plasmids that encode mCherry-LC3B, LAMP1-mRFP, or mRFP-RAB5A were transiently transfected into HeLa or HEK293 cells using Effectene Transfection Reagent (QIAGEN, 301427). After 24 h, HeLa cells were treated with either vehicle or PP242 (10 \( \mu \)M) for 4 h. The cells were then stained with Cyto-ID and Hoechst 33342 and visualized under an inverted fluorescence microscope (Carl Zeiss Microscopy, LLC, Thornwood, NY). Photomicrographs were taken using a 40x objective. In some experiments, HeLa cells were transfected with pEGFP-LC3 and then treated with PP242. GFP fluorescence was recorded as described above. To quantify the intensity of the Cyto-ID or EGFP-LC3 fluorescence, we used ImageJ software (http://imagej.nih.gov/ij/) and randomly quantified 5 fluorescent dots (representing the autophagic compartments) at different bleaching times. The average intensities were shown in Fig. 2B.

**Immunoblotting and quantification of band intensity.**

Immunoblotting was performed as described in our previous reports.\(^{28,29}\) In brief, cells were lysed and the total proteins were quantified using the Bradford assay (Bio-Rad, 500-0006). Equal amounts of total protein (25 \( \mu \)g) in each sample were loaded onto a 15% SDS-PAGE gel. After transferring, the blot was incubated with a rabbit anti-LC3B (1:1000), a rabbit anti-MTOR (1:1000), a rabbit anti-phospho-MTOR (1:1000), a mouse anti-SQSTM1/p62 (1:200), or a mouse HRP-conjugated anti-actin (1:3000) antibody. Images were taken using a ChemiDoc™ MP System (Bio-Rad, Hercules, CA). The intensity of the band was quantified using Image Lab software (Bio-Rad) or ImageJ as described earlier. The fold change of LC3B-II is defined as the ratio of the intensity of the LC3B-II band normalized to that of actin.

**RNA interference screening**

K562 cells (3.75 \( \times \) 10\(^3\) cells per well) were plated in a 96-well plate. Viruses harboring 88 individual human kinase shRNAs or the control nonsilencing (NS) shRNA were added to each well. Cells were transduced by spin infection (900 x g for 90 min at 30°C). After 3 d incubation, the cells were subject to the Cyto-ID fluorescence spectrophotometric assay described above.

**Quantitative RT-PCR**

K562 cells (5 \( \times \) 10\(^5\)) were transduced with viruses containing NS or candidate shRNAs. Cells were selected by puromycin (1 \( \mu \)g/ml) for a week. Total RNA was extracted using Trizol (Ambion, Life Technologies™, 11596018) and cDNA was synthesized using reverse transcriptase (New England Bio Labs, M0253L). mRNA levels of ANK1, MAP3K6, NTRK3, or ANK2 were determined by quantitative PCR using a StepOnePlus™ Real-Time PCR system (Life Technologies, Grand Island, NY). Primers for each gene were ANK1-f (5′ACGCAAGGTCCACACTCATTTG), ANK1-r (5′CACCCAGACATCCATGTA3′), MAP3K6-f (5′CCGACATCATGAACTTG
Small molecule screening

K562 cells (2.5 x 10⁴ cells per well) were plated in a 96-well plate preloaded with 79 different small molecules (10 µM/compound/well; Microsource Discovery Systems, the Spectrum Collection) or vehicle control (DMSO). After 4 h incubation, the cells were analyzed using the Cyto-ID fluorescence spectrophotometric assay described above.

Mouse experiments

All animal studies were approved by the Institutional Animal Care and Use Committee (IACUC) of Virginia Tech. C57BL6 mice (Charles River Laboratories, 24101547) (n = 4 mice per treatment group) were administered with PP242 (60 mg/kg/d) by intraperitoneal injection or hydroxychloroquine (50 mg/kg/d by intraperitoneal injection). Peripheral blood (30 µL) was withdrawn from the tail vein before or 2, 4, and 8 h after the treatment. Red blood cells were lysed with ammonium chloride and the resulting white blood cells were subject to the Cyto-ID assay described above. To examine autophagy in a mouse model of BCR-ABL-driven leukemia, 32D/BCR-ABL cells (1.5 x 10⁵) were injected intravenously into 12 SCID-beige mice (n = 4 mice per treatment group) via the tail vein as described in our previous studies.⁹ After 6 d, mice were administered either imatinib (100 mg/kg/d) or hydroxychloroquine (50 mg/kg every other day) orally via gavage. Four or 5 d after treatment, mice from each treatment group were euthanized and their femurs and blood were collected. Bone marrow was extracted from the femurs and then white blood cells were isolated using Ficoll-Paque PREMIUM 1.084 solution (17-5446-02) purchased from GE Healthcare Life Sciences. Autophagy was monitored using the Cyto-ID-based fluorescence spectrophotometric assay described above.

Statistical analyses

The Student t test and the one-way ANOVA with the Tukey post-hoc test for multiple comparisons was used for statistical analyses. These tests were done using the SPSS software.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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