We detail here a protocol using tandem-tagged mCherry-EGFP-LC3 (C-G-LC3) to quantify autophagic flux in single cells by ratiometric flow cytometry and to isolate subpopulations of cells based on their relative levels of autophagic flux. This robust and sensitive method measures autophagic flux rather than autophagosome number and is an important addition to the autophagy researcher’s array of tools for measuring autophagy. Two crucial steps in this protocol are i) generate cells constitutively expressing C-G-LC3 with low to medium fluorescence and low fluorescence variability, and ii) correctly set up gates and voltage/gain on a properly equipped flow cytometer. We have used this method to measure autophagic flux in a variety of cell types and experimental systems using many different autophagy stimuli. On a sorting flow cytometer, this technique can be used to isolate cells with different levels of basal autophagic flux, or cells with variable induction of flux in response to a given stimulus for further analysis or experimentation. We have also combined quantification of autophagic flux with methods to measure apoptosis and cell surface proteins, demonstrating the usefulness of this protocol in combination with other flow cytometry labels and markers.

**1. Introduction**

Measuring autophagy is difficult.\(^1,2\) The inadequacy of current methods for measuring autophagy is a stumbling block to advancement of the field and has been a source of debate and lamentation by many authors.\(^3-4\) All available methods for measuring autophagy have caveats, are rarely quantitative, and are sometimes subjective or susceptible to bias. Furthermore, the mechanistic complexity of the autophagic process presents multiple stages, structures, and proteins that can be used to interrogate the pathway.\(^5-7\) Perhaps the most important distinction is the difference between measurement of autophagosome formation (autophagy induction) and autophagic flux—the fusion autophagosomes with lysosomes and subsequent degradation of cargo. This distinction is not trivial as the number of autophagosomes in a cell can increase, either because their formation has been stimulated or because their degradation has been inhibited. Likewise, measurements of autophagosome formation do not provide information about whether autophagy has been induced or flux has been inhibited. It is therefore critical to assess the flux through the autophagic pathway before conclusions about autophagy’s role in a particular process can be made.

Autophagic flux is most frequently assessed using MAP1LC3B/LC3 and SQSTM1/p62 western blotting with and without lysosomal protease inhibitors to block autophagic degradation. In fact, the most popular method to measure autophagy, measurement of LC3 protein lipidation by western blot, is so frequently misused that a guide to interpretation of LC3 westerns was published in this journal several years ago.\(^8\) Other methods, such as visualization of LC3 puncta by microscopy and the “gold standard” of autophagosome visualization by electron
microscopy are also widely used to measure autophagy. However, none of these methods are amenable to straightforward measurement of autophagy in individual cells and none of them can directly measure flux of autophagic cargo through the autophagy pathway. An excellent method to measure flux is by using tandem-labeled mCherry-EGFP-LC3 (C-G-LC3) microscopy to count the number of autophagosomes and autolysosomes. The best techniques for high throughput, single-cell analysis of autophagy rely on microscopy and are therefore laborious and/or data intensive. The development of new, high-throughput methods for measuring autophagy is important if we are going to answer critical questions that cannot be addressed using available methods. We provide here a detailed protocol for our recently published method using mCherry-EGFP-LC3 to quantify autophagic flux by flow cytometry and to sort subpopulations of cells based on their relative levels of autophagic flux.

Flow cytometry is a powerful tool for quantitative, single-cell analysis and can be used for ultra-high-throughput analysis and sorting for multiple markers of interest. Several methods have been put forth for the measurement of autophagy by flow cytometry using lysosomotropic dyes, or fluorescent LC3 or SQSTM1. However, these methods have not gained traction primarily due to low dynamic range, low sensitivity and/or low specificity. Of these, the method with the highest dynamic range uses the detergent saponin to extract soluble cytoplasmic (nonlipidated) GFP-LC3, leaving behind the lipidated autophagosome-associated GFP-LC3 thereby quantifying the number of autophagosomes. However, the saponin extraction step is not amenable to other flow cytometric methods or staining protocols because it disrupts the integrity of the cell membrane. Furthermore, while this is a good method to measure the quantity of autophagosomes in a cell, it does not measure autophagic flux through the pathway and therefore suffers the same difficulties with interpretation that plague LC3 western blotting. We have therefore developed a method to measure autophagic flux by flow cytometry and have used it to successfully sort cells based on their relative levels of autophagic flux. We have adapted C-G-LC3, which is used as a reporter for autophagic flux by microscopy, for use as a ratiometric flow cytometry reporter (Fig. 1A). The basis for the utility of C-G-LC3 as a reporter for autophagic flux lies in the higher sensitivity of EGFP fluorescence to the acidic environment of the autolysosome relative to mCherry. Cells with higher flux are less green due to fusion of autophagosomes with lysosomes, which increases the mCherry/EGFP ratio in the cell. Using ratiometric flow cytometry to calculate the flux in each cell based on this ratio, we are able to not only quantify flux in individual cells but to sort cells based on their relative autophagic flux (Fig. 1B). This method has been extensively validated; it reliably and accurately quantifies autophagic flux induced by multiple stimuli and blocked by chemical and genetic inhibition of autophagy. This protocol provides in-depth, detailed steps to generate and validate reporter cells to measure autophagic flux, identify and properly set up an adequately equipped flow cytometer, and use it to quantify and sort cells based on their relative levels of autophagic flux (Fig. 1B).

2. Materials

2.1 Cells

Most mammalian cell lines or types should be suitable to measure flux; many of the cells we have used in our lab are listed in Table 1. Cells should be cultured in their normal growth medium. GP2-293 cells (Clontech, 631505) are used to generate C-G-LC3 retrovirus and should be maintained in DMEM (Mediatech, 10-013-CV), supplemented with 10% fetal bovine serum (Sigma, F6178).

2.2 Plasmids

The reporter we have used with the greatest success is mCherry-EGFP-LC3. The order of the 2 fluorophores is likely not critical but LC3 must be at the C terminus of the fusion protein so that it can be cleaved by ATG4, lipidated, and incorporated into the phagophore membrane. Any protein fused to the N terminus of LC3 will be cleaved off by ATG4 in order to activate LC3 for lipidation. It is also important that the green fluorescent protein be EGFP (another fluorescent protein should work as long as it has a pKₐ ≥ 6.0). The red fluorescent protein choice is not crucial; other RFPs should work as long as they have limited spectral overlap with GFP and a pKₐ ≤ 4.5. We tried using ECFP-EGFP-LC3 to measure flux but were not successful. This is perhaps due to the higher pKₐ of ECFP relative to mCherry, lower fluorescence when excited with a 405 nm laser and/or spectral overlap between ECFP and EGFP.

The protocol described here uses pBabe-mCherry-EGFP-LC3 (Addgene, 22418) to generate C-G-LC3 retrovirus to mediate expression of the reporter. To generate C-G-LC3 retrovirus, any plasmid expressing VSV-G envelope protein off of a CMV promoter may be used (e.g., pCMV-VSV-G, pMD2.G; Addgene, 12259).

2.3 Equipment

2.3.1 Flow cytometer

Measuring autophagy flux by flow requires a flow cytometer that can detect both EGFP and mCherry fluorescence. This requires both a blue (488 nm) laser and a yellow (560 nm) laser. The laser lines do not necessarily need to be on separate channels, they can be co-linear as long as proper filter sets are used to separate the green and red channels. We have successfully used several flow cytometers for measurement of autophagic flux including a MoFlo XDP-70 (Beckman Coulter) with a 552-nm laser for mCherry excitation on a separate laser line, a MoFlo XDP-100 (Beckman Coulter) with a 561-nm laser for mCherry excitation on a separate laser line, a Gallios (Beckman Coulter) with a co-linear 561-nm laser for mCherry excitation and an LSR-II (BD Biosciences) fitted with a 532-nm laser for mCherry excitation (all of these machines use a 488-nm laser to excite GFP). The 2 MoFlo sorters have been used extensively and successfully to sort cells with differential autophagic flux.

2.3.2 Flow cytometry software

When measuring flux by flow cytometry it is highly advisable to have software either during the acquisition of flow data or in the subsequent analysis that is capable of handling a ratiometric calculation as
A derived parameter. Namely, the quotient of the 2 fluorescence channels:

\[ A_f = \frac{F_{\text{mCherry}}}{F_{\text{GFP}}} \]

where \( A_f \) is autophagic flux, \( F_{\text{mCherry}} \) is mCherry fluorescence and \( F_{\text{GFP}} \) is GFP fluorescence. For cell sorting based on relative autophagic flux, the acquisition software must be able to calculate \( A_f \) and sort on that derived parameter. If the sorter the user would like to use cannot sort on ratiometric parameters, other methods can be used (such as creating diagonal gates in an mCherry and GFP dot plot) to work around this limitation, but they are not optimal and may not prove useful.

The 4 machines we have used to measure and sort cells on flux use software that can use a ratio of 2 parameters as a derived parameter. The 2 XDP machines use Summit software for acquisition, the Gallios uses Kaluza and the LSR-II uses FACSDiva. We have found that on Summit it works better to use a ratio of the linear fluorescent parameters while the other 2 software platforms use log ratios.

2.3.3 Other equipment

Tissue culture incubators, hoods, and other associated apparati are necessary to culture and maintain the reporter cells.
In addition, general lab equipment for the purification of plasmid DNA and other general lab reagents are also necessary. In addition, if the cells being analyzed or sorted are biohazardous, then appropriate protections should be used and any other users should be informed that dangerous materials are being used on the machine.

### 2.4 Chemicals

- Trans-IT LT1 (Mirus, MIR2300)
- Polybrene (Sigma, 107689)
- PBS (Mediatech, 21-040-CV)
- ANXA5/annexinV-APC (BD Biosciences, 550474)
- DAPI (Sigma, D8417)
- Earle’s Balanced Salt Solution (EBSS; Sigma, E2888)

### 3. Methods

#### 3.1 Generation of C-G-LC3 reporter cells (1–2 wk)

The first step of this protocol is the creation of a reporter cell line stably expressing C-G-LC3. The cells must be stably expressing C-G-LC3; transient transfection will yield highly variable expression and will not produce usable data. Furthermore, for the highest sensitivity and consistency, stable, individual clones should be chosen, or pooled clones flow-sorted for similar expression levels and tested by flow cytometry for consistent (low coefficient of variability) fluorescence from cell-to-cell (Fig. 1B). Some consideration must be given to the cells to be used and how to mediate expression of the reporter. The reporter we prefer is mCherry-EGFP-LC3 expressed either off of a CMV promoter via a plasmid or off of the viral LTR in the pBABE retrovirus. Both plasmid- and retrovirus-mediated expression have given us good results, though the latter is our method of choice because the low variability in expression from retrovirus reduces the time needed to create and characterize the reporter cells.

The method to introduce the C-G-LC3 construct into your cells of interest should be determined by the researcher using the best method to mediate efficient stable gene expression in the experimental system being used. We have found retrovirus-mediated expression to be the fastest method for creation of stable clones and it produces the most consistent and long-lasting expression. We have observed lost or diminished C-G-LC3 expression in many of our stable clones; the use of retrovirus to express C-G-LC3 minimizes this effect.

#### 3.1.2 Choosing a cell line

Selection of the cells to be used as reporters for autophagic flux is a critical step in this procedure. We have successfully used many cell lines and types including HeLa, HEK293, BJAB, Jurkat, GL261, and U87MG (Table 1). However, not all cell lines and types work equally well, primarily due to variability in the size of the cells and therefore the fluorescence. While using tandem tagged C-G-LC3 provides an internal control for the overall fluorescence of the cell, large differences in size increase the cell-to-cell variability in autophagic flux thereby decreasing the sensitivity and reproducibility of the assay. The best cells we have found for measuring and sorting cells for autophagic flux are lymphocytes, while the least sensitivity is found with MEFs and other primary cells. However, the suitability of individual cell lines and types to this method must be determined empirically. Generating several reporter cell lines that suit the needs of the particular experimental system could save time in the long run. We have also found that for experiments where the researcher wishes to see the effect of overexpression or knockdown of a gene of interest, it is crucial to make the reporter cells first, then manipulate the gene of interest rather than making several different C-G-LC3 reporter cell lines from pre-existing stable clones. The variability in expression of C-G-LC3 from clone to clone will make it difficult—or impossible—to quantitatively determine differences in flux using this method.

#### 3.1.3 Making C-G-LC3 retrovirus

GP2-293 cells are maintained in growth medium at 37 °C in 5% CO₂ at 30–80% confluence. For transfection, cells are plated in a 10-cm dish overnight such that they will be at approximately 50% to 70% confluent at the time of transfection. Cells are then transfected with Trans-IT 293 reagent according to the manufacturer’s protocol using a 1:1 ratio of pBabe-C-G-LC3 and pCMV-VSV-G. Sixteen hours following transfection, fresh growth medium is added to cells. Virus-containing medium is subsequently harvested at ~24 and 48 h thereafter. C-G-LC3 virus is aliquoted and stored at −80 °C. We do not filter or centrifuge the viral media as this can reduce viral titer considerably; freezing the viral media kills any remaining 293 cells, with nominal reduction in titer.

#### 3.1.4 Transduction of cell lines with C-G-LC3 retrovirus

Cells to transduce should be maintained in growth medium and plated at

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**Table 1.** Cell lines used to measure autophagic flux by flow cytometry

| Cells         | Type                      | Expression | Success |
|---------------|---------------------------|------------|---------|
| HeLa          | Human cervical cancer     | Transfection | ***     |
| HEK293        | Human embryonic kidney    | Transfection | **      |
| BJAB          | Human B-cell lymphoma     | Retrovirus  | ****    |
| Jurkat        | Human T-cell leukemia     | Retrovirus  | ****    |
| GL261         | Mouse glioma              | Retrovirus  | *       |
| U251          | Human glioma              | Retrovirus  | ***     |
| MCF10A        | Human mammary epithelium  | Retrovirus  | ***     |
| MEF           | Mouse embryonic fibroblast| Retrovirus  | *       |
| MCF7          | Human breast cancer       | Retrovirus  | ***     |
| MDA-MB-231    | Human breast cancer       | Retrovirus  | ***     |
| U87MG         | Human glioma              | Retrovirus  | ***     |
| BT16          | Human AT/RT brain tumor   | Retrovirus  | ***     |
| 794           | Human AT/RT brain tumor   | Retrovirus  | **      |
| DBTRG         | Human astrocytoma         | Retrovirus  | **      |
| NMC-G1        | Human astrocytoma         | Retrovirus  | *       |
| AM-38         | Human astrocytoma         | Retrovirus  | ***     |
-70–80% confluence in a 6-well plate. Viral media from 3.1.3 should be mixed with polybrene to obtain a final concentration on the cells of 8 ng/mL. After aspirating the growth medium from the cells, replace 1 mL of fresh growth medium and add 1 mL of C-G-LC3 viral medium. More or less viral medium may be used based on the viral titer and susceptibility of the cells to retroviral transduction and should be determined empirically. Cells can be treated from 2 h to overnight with C-G-LC3 retrovirus whereupon they should be fed and split accordingly.

3.2 Reporter cell characterization (1–2 wk)

3.2.1 Confirmation of C-G-LC3 expression

Following retroviral transduction, GFP and mCherry expression are tested by fluorescence microscopy at 48 h post-transduction. Many cell lines will only express low levels of C-G-LC3 and the expression may be nearly undetectable via the naked eye on an epifluorescence microscope—this does not mean the cells are not going to work for measuring flux by flow cytometry; the most effective cell line and clone we have used has nearly undetectable C-G-LC3 expression by the naked eye. If expression is hard to detect, a long exposure time on an epifluorescence camera setup or flow cytometry (using proper negative and positive controls—the same cells expressing GFP and mCherry separately and without any fluorescent protein expression) should be used to confirm expression. Cells should then be placed in selection media [growth medium + puromycin (~1 μg/mL, determined empirically) and selected for 48–72 h before sorting or dilutional cloning.

3.2.2 Sorting/clonal selection

Following selection, cells should either be cloned by limiting dilution for single-positive clones or flow sorted for mCherry and GFP double-positive cells. These clones or sorted +/- pools should then be grown up and tested for their ability to measure flux. Again, having the proper positive (EBSS-treated) and negative (untreated) controls is critical to validating each clone or pool.

Limiting dilution clones can be plated one cell per well into a 96-well plate to obtain single-cell clones. This method has proven successful in our lab, but we have found that many clones must be tested for their ability to measure flux in the dynamic range necessary to robustly distinguish between signal and noise. Each clone must first be tested for expression of mCherry and GFP (by either microscopy or flow) then tested for proper response to autophagic stimuli by flow cytometry. If there is any question about the expression of mCherry or GFP, positive control cells expressing each separately can be invaluable.

Flow sorting for pools of double-positive cells has yielded excellent results and avoids potential problems with using clonal populations that may have other alterations introduced or perpetuated by the cloning and selection process. Following puromycin selection, we flow sort mCherry and GFP double-positive cells using small gates. Generally, we will sort out 2 populations, a high-expression population and a medium-expression population. Then each population is expanded and tested for autophagy flux quantification.

3.2.3 Validation of clones or sorted pools

After confirming expression of mCherry and GFP in each clone they must then be tested for the ability to quantify flux induction. The stimulus used to induce flux depends on the experimental system, but in our experience, starvation is the most potent autophagy stimulus. After starving cells for a period of time (using several time points from 0.5–6 h is most informative) known to induce autophagy in the cells, they are transferred to a flow cytometry buffer and placed on ice for flow. We use regular growth media with serum (specific to the cells being used) as the flow buffer to avoid inducing flux, which can occur when using typical PBS-BSA flow buffers, but it should not matter if the cells are kept on ice. It should also be noted that removing adherent cells with trypsin and/or EDTA can cause changes in autophagic flux and should be done as quickly as possible; control and treated cells should be subjected to non-treatment buffers/media for the same period of time. Next, using a properly set up flow cytometer (3.3.1), treated cells are assessed for their ability to measure flux. A good clone or pool will look like Figure 2A; untreated cells will show a tight peak by the mCherry/GFP ratiometric parameter with the peak spreading further to the right depending on the potency of the stimulus. The mCherry-GFP dot plot should show a downward shift in the GFP signal with little to no movement in the mCherry signal. The best performing clones (high dynamic range) are sometimes difficult to keep on scale at the time point with the greatest change in signal.

3.3 Quantification and sorting cells by autophagic flux (4–6 h)

As stated above, sorting for autophagic flux requires a flow cytometer with software capable of ratiometric calculation of derived parameters and the ability to sort on such a parameter. Post-processing of data will allow the quantification of flux from a cytometer without ratiometric capability but there are difficulties with keeping the data on scale and if the voltages during acquisition are not set correctly (this is determined empirically) then data can be unusable unless the analysis software is capable of higher order mathematical parameters. Cells (preferably the same cell line) that express GFP and mCherry separately can greatly enhance your ability to troubleshoot. These are especially important if the cytometer you are using has the GFP and mCherry channels on the same laser line so that compensation can be set up correctly. An important limitation of this technique (this is true of any flow cytometry technique) is that it provides only a relative measure of flux, i.e., induced flux vs. control. These differences cannot be standardized nor can they be compared between different cell lines or from different experiments. Another limitation of the method is that it does not accurately measure reductions in flux below basal levels. It is very good for measuring induction of flux over basal and any inhibition of that induced flux can also be detected. Most cells have modest basal flux, and reductions from there can only be detected by observing the long-term increase in both GFP and mCherry fluorescence. This method relies on the reduction in GFP fluorescence when there is an increase in flux; however, when flux is inhibited GFP only increases modestly.
Figure 2. For figure legend, see page 1333.
3.3.1 Flow cytometer setup

Proper setup of the flow cytometer is a critical step of this protocol. Each cytometer and associated acquisition software have their own quirks, especially with regard to handling Thankof ratiometric parameters. It may prove valuable to get help from someone with experience using ratiometric parameters on the machine being used. Here, we will describe setup on the XDP-100 sorting flow cytometer (Fig. 2A).

Forward scatter/side scatter and doublet discrimination should be set according to the parameters most applicable to the flow cytometer being used, and detector voltages set for the particular cells being used. As depicted in Figure 2A, following FSC-Height and SSC-Height, SSC-Width is used to gate singlets on the XDP-100 sorting flow cytometer. These gates result in the mCherry-GFP dot plots and ratiometric histograms shown for control and EBSS treated cells, respectively.

We have found that on the XDP sorters, using linear input parameters rather than log is most useful for the derived ratiometric mCherry/GFP parameter. However, this may not work best for all machines and is not possible with some. It is sometimes difficult to keep both the negative and positive control conditions on scale for the ratiometric parameter, unless the software is capable of using derived, higher-level mathematical parameters. For most applications, the user must set the voltages/gain for both the mCherry and GFP parameters carefully to get the ratio parameter on scale and with good dynamic range. Generally, this means getting the GFP channel just below the middle of its range with the mCherry channel just above the middle of its range (Fig. 2A). This results in a low positive number that can be tweaked from experiment to experiment to optimize the scale and dynamic range. It should be noted that using voltages too low will decrease the dynamic range, but voltages that are too high increase the noise. Getting the balance just right sometimes takes some trial and error and varies greatly from cytometer to cytometer. We have found that this is easiest to do by first adjusting voltage using a positive control for flux (like EBSS starvation) to get those on scale, then using negative control (either basal or flux-inhibited) cells and adjusting to get those on scale. Once the voltage and gain are set on a particular machine they should be noted and used as a starting point for the next experiment, though they will frequently need to be adjusted.

3.3.2 Quantification of flux

Quantitation of autophagic flux by flow cytometry can be accomplished by other means, but we have found that using the ratio of mCherry to GFP fluorescence is the most consistent, robust, useful, and least arbitrary measurement of flux we have tested. After setting the cytometer up correctly to measure flux the user must then decide how to compare the different conditions of their particular experiment. The 2 most common ways are to look at differences in median fluorescence or by measuring the percent of cells that fall within a particular gate (Fig. 2B). Both measurements work well for assessing differences in flux; the choice between the two depends on the particular experiment being performed. For most experiments the entire curve will shift resulting in a wholesale change in both the median and % in a gate. However, certain treatments have a more graded effect across a population that change the shape of the curve resulting in a change in the % of gated cells without a substantial change in the median mCherry/GFP ratio.

3.3.3 Sorting cells for autophagic flux

C-G-LC3 ratiometric flow cytometry can be used to sort either cells with a differential level of autophagic flux in response to a given stimulus or cells with different amounts of basal autophagic flux. For instance, sorting for different levels of induced flux can be used to isolate cells that are either efficient inducers of flux in response to a given stimulus or poor inducers based on their levels of flux. When performing these types of experiments it is very important to have the proper controls (untreated vs. starved cells) to create sort gates so that the sorted cells are not a mixed population, i.e., the gates should be sufficiently far apart. It is also important that the stimulus be strong enough to move the mCherry/GFP histogram substantially in order to separate the high and low flux populations (Fig. 1A).

Sorting for high and low basal flux can require some minor changes to the voltages to facilitate the separation of the different populations. By increasing the voltage in the mCherry channel, or decreasing it in the GFP channel, the population at basal will move to the right of the mCherry/GFP ratio histogram. This will (usually) result in a spreading of the population to separate the low and high flux subpopulations. We have had success in separating completely divergent populations by sorting the top and bottom quartiles, but have the most success sorting the top and bottom 10–20%. Complete separation of the populations (such that if they are re-run through the flow cytometer they do not overlap) may not be possible in all cell lines or types. It is very important to run samples sorted for basal flux back through the flow cytometer to confirm that the samples have sufficiently differential flux such that separate subpopulations are actually being sorted, rather than an artificial difference in the detected (but not actual) fluorescence intensity. If there is significant overlap in the sorted populations, the gates can be widened and made more stringent to sort only the cells with highly divergent flux. It is also important to confirm that the FSC/SSC profiles of the 2 sorted populations are not different; this could indicate that the sorted cells actually differ in their size making it look like there are differences in flux that do not actually exist.

Figure 2 (See opposite page). Quantification of autophagic flux by flow cytometry. (A) Example of flow-cytometry data illustrating flow cytometer setup for measuring autophagic flux. Scatter and singlet gates should be used to eliminate debris, dead cells, and mitotic cells. Voltages and gain on GFP and mCherry detectors are set empirically to allow both negative and positive control (starvation) plots to fit the mCherry/GFP ratio histogram. (B) Comparison of this protocol to other previously published methods for flow cytometric quantification of autophagy. C-G-LC3 reporter cells were starved for 4 h in EBSS followed by flow cytometry with or without extraction of cytosolic (nonlipidated) LC3 by saponin. (C) Concurrent measurement of autophagic flux and apoptosis using the established flow cytometry markers ANXA5/annexinV and DAPI. C-G-LC3 reporter cells were treated with an apoptotic stimulus (Fas ligand) for 4 h and stained with ANXA5/annexinV-APC and DAPI followed by flow cytometry.
Sorted cells can then be used in whatever experiments the user desires. It should be noted that differences in flux can change quickly (on the order of minutes to hours) and the buffer used to sort the cells into should be chosen such that it does not alter the flux during the time the cells are being sorted or thereafter. Furthermore, there should be sufficient volume such that the sheath buffer from the cytometer does not dilute the sample substantially. We sort the cells into growth medium on ice, so that changes in flux are minimal while the cells are sorted, then quickly transfer them into the experimental system.

4. Results and Discussion

The availability of an accurate and robust flow cytometry marker for autophagic flux represents a powerful tool for both the measurement and sorting of cells based on autophagy. While the use of a genetically encoded reporter limits the use of the method to engineered cell lines, mCherry-GFP-LC3 is substantially better than other methods that we have tried to measure autophagy by flow cytometry. Our recent publication pioneered the quantification of autophagic flux by ratio-metric flow cytometry using mCherry-GFP-LC3 and exhaustively validated that the methodology can be used to not only measure autophagic flux but to sort cells based on differential basal autophagic flux and consequently to test if differences in autophagy affect responses upon exposure to other agents—in our case high and low autophagic flux determined the likelihood of cells dying in response to activation of death receptor induced apoptosis. In addition, we successfully combined quantification of autophagic flux with previously established methods for measuring apoptosis in live cells (using ANXA5/annexin-V-APC and DAPI; Fig. 2C) and cell surface markers (anti-PTPN13/CD95-APC). The capability to sort cells based on both autophagic flux and other flow cytometry markers makes this method particularly adaptable to the study of autophagy in stem cell research.

Furthermore, quantification and sorting cells based on autophagic flux can be adapted for high-throughput functional genomics screening, drug screening, and other flow-cytometry based screening platforms. So, until a flow cytometry reagent is devised that can quantify autophagic flux in cells without a genetic reporter, this may be the best method available.

Disclosure of Potential Conflicts of Interest

The authors report no potential conflicts of interest.

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