

Advanced segmentation and organelle analysis: A computer exercise using CellProfiler & CellProfiler Analyst software

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Background information:

The images in this experiment come from the [Broad Bioimage Benchmark Collection](#). They are 240 of 69,120 fields of U2OS cells treated with a panel of 1600 known bioactive compounds and imaged in five channels for a so-called Cell Painting assay- see Gustafsdottir et al, 2013 for more information. The compounds target a wide range of cell pathways, meaning that some cells and organelles will have very different morphologies both from each other and from the mock treated controls. This will give you an opportunity to try to find segmentation parameters that work across a wide range of conditions.

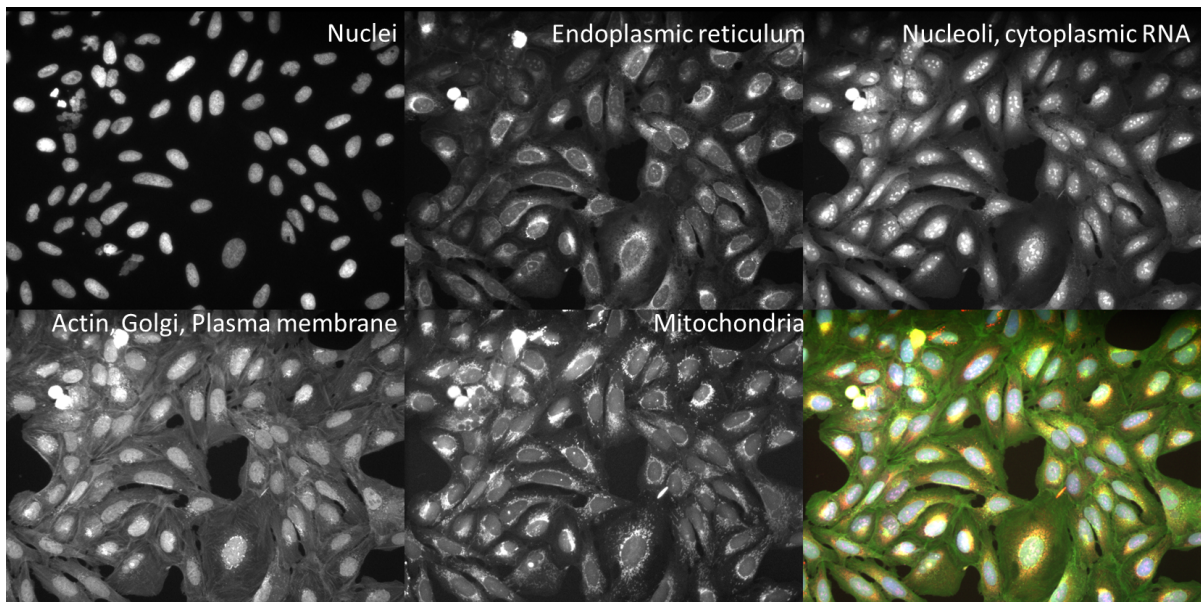


Figure 1: Images and channels from a CellPainting assay.

While in the traditional Cell Painting protocol we do not actually segment out any organelles (other than the nucleus), the large number of stained compartments make this an excellent set of images to find subcellular features. Finding the average or count of smaller objects inside a larger ‘parent’ object is a feature of many pipelines and an important skill to have in setting up a CellProfiler analysis.

Cell Painting generally consists of a few simple segmentation steps followed by adding as many measurement modules as can be reasonably included in a single pipeline; we have found that by doing this we can measure ~1500 features of each cell and from that create a ‘morphological profile’ that can be used to predict interesting biology including drug mechanisms of action, gene-pathway interactions, and more. See Bray et al 2016 and citations within for more information.

Goals of this exercise:

This exercise will give you practice at finding segmentation parameters that will be robust across whatever variability may exist in your sample. This is not always straightforward, so examining your segmentation across a wide range of images will be necessary.

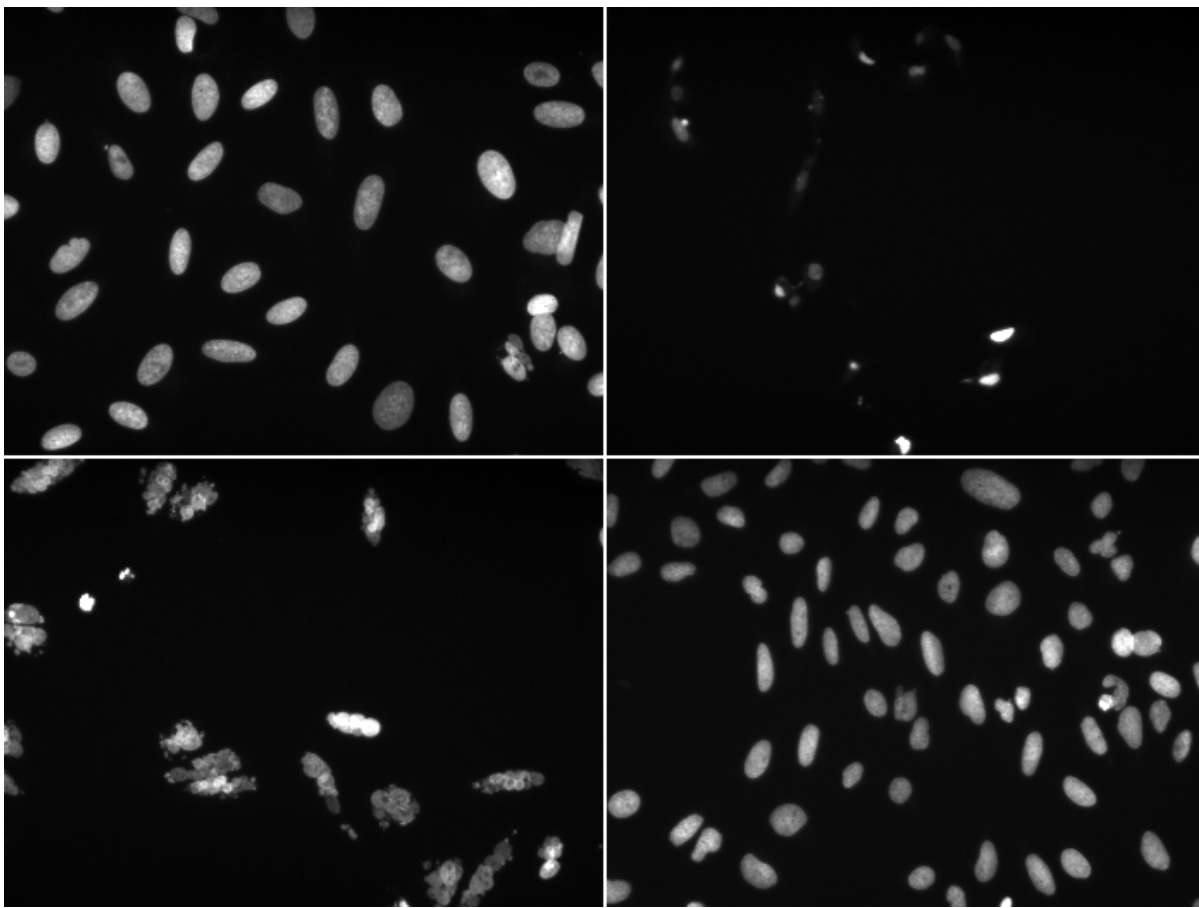


Figure 2: Examples of varied nuclei found in this data set.

This exercise will additionally show you some ways to pull out smaller features in your image by segmenting organelles within the cells and nuclei. You will also be shown how to use RelateObjects so that you can study the average counts, distances, and measurements of the smaller organelles inside their larger parent objects.

Materials necessary for this exercise:

These 1200 images (240 sites in 5 channels) represent 120 wells from a single 384 well plate, either mock treated with DMSO or treated with a variety of bioactive compounds. A CSV file containing associated drug treatment information has also been included.

It is additionally expected that you are generally familiar with CellProfiler, preferably after completing the Translocation tutorial or a similar introductory exercise.

1. Load images and metadata

- Start CellProfiler by double-clicking the desktop icon
- Drag and drop the ‘BBC022_Analysis_Start.cppipe’ file into the ‘Analysis modules’ box. 7 modules should pop up, and almost all of them will show errors. **This is the expected behavior.**
- Drag and drop the ‘BBC022_20585_AE’ folder into the ‘File list’ box. It should automatically populate. Notice that illumination correction images (with a file extension of ‘.npy’) are included in this data set.

2. Import metadata from the CSV

So that we can explore what cells treated with different drugs look like later in the exercise, we must add this information into CellProfiler from the CSV. Provided with this exercise is a CSV called ‘20585_AE.csv’ detailing drug treatment info for each image.

- In the ‘Metadata’ module, three metadata extraction methods should already be present and fully configured:
 - The first pulls Well, Site, and Channel metadata from all of the image files except for the illumination correction functions
 - The second pulls Plate metadata from the image folder
 - The third pulls Plate metadata from the illumination correction functions
- The fourth metadata extraction step requires you to tell CellProfiler the location of the CSV file. It is looking for it in CellProfiler’s Default Input Folder, which we must therefore configure.
 - Select the ‘View output settings’ button in the bottom left corner of the screen.
 - Set the ‘Default Input Folder’ to the location of ‘20585_AE.csv’ within the exercise folder
- Return to the ‘Metadata’ module and press ‘Update’. You should now see a number of columns in the Metadata window.
 - If you like, examine the CSV and how the ‘Match file and image’ settings are configured:
 - Image_Metadata_PlateID (from the spreadsheet) is matched to Plate (extracted from the folder name by the second extraction step)
 - Image_Metadata_CPD_WELL_POSITION (from the spreadsheet) is matched to Well (extracted from the file name by the first extraction step)

3. Examine the channel mappings in NamesAndTypes (optional)

The channel mapping here is a bit more complicated than anything we’ve worked with before- we have a single set of illumination correction images that map to each and every well and site. We can use the metadata we extracted in the last module to make that association possible.

- Two different ways of mapping images to channel names are demonstrated here. There are several others, and often you could create several correct mappings for a given set of images, but these may serve as a helpful example to refer to in your own work.
 - The ‘.tif’ image files are assigned a name by the Metadata extracted in the previous module (specifically ChannelNumber)
 - The ‘.npy’ illumination correction functions are assigned a name based on a unique string in the name (such as ‘IllumER’)

As there is only one set of illumination correction functions for each entire plate, the image sets cannot simply be constructed by using ‘Image set matching’ as ‘Order’.

- Scroll to the bottom of the ‘NamesAndTypes’ to see how the image sets are constructed
 - ‘Image set matching’ is set to ‘Metadata’
 - Each image channel is set to ‘Plate->Well->Site’.
 - Each illumination correction function is set to ‘Plate->(None)->(None)’
- Metadata based matching can be useful in any circumstance where a larger group of images needs to be mapped with a smaller one, such as every plate in an image set having its own illumination correction function or every movie in a series of timelapse movies being matched to its own unique cropping mask.

IllumER	IllumHoechst	IllumMito	IllumPh_golgi	IllumSyto	OrigER	OrigHoechst	OrigMito	OrigPh_golgi	OrigSyto	+/-	Green
Plate	Plate	Plate	Plate	Plate	Plate	Plate	Plate	Plate	Plate	+/-	Green
(None)	(None)	(None)	(None)	(None)	(None)	Well	Well	Well	Well	+/-	Green
(None)	(None)	(None)	(None)	(None)	Site	Site	Site	Site	Site	+/-	Green

Figure 3: A section of the ‘Image set matching’ dialog.

4. Examine the output of the CorrectIlluminationApply module (optional)

Since microscope objectives don’t typically have a completely uniform illumination pattern, applying an illumination correction function can help make segmentation better and measurements more even by compensating for this. Pay close attention to the top of the field of view to see the greatest effect.

- Enter test mode and hit ‘Step’ to run the CorrectIlluminationApply module.
- Briefly examine the output of the CorrectIlluminationApply module—you can see that the illumination correction functions show significant heterogeneity across the field of view.
 - These functions were created by averaging and smoothing all 3456 images from this plate, indicating the image captured is consistently dimmer in those regions for nearly all images.
- Also note that while the illumination correction functions for each channel are similar, they aren’t identical; each channel in your own experiments should therefore be illumination corrected independently.

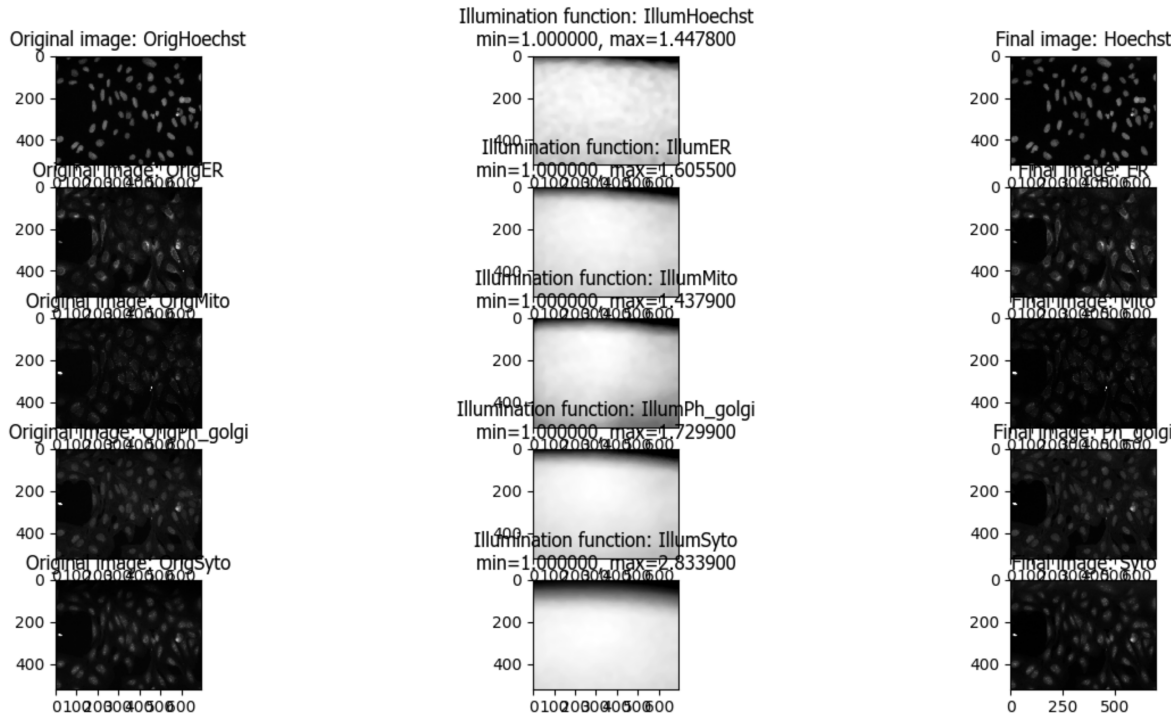


Figure 4: Application of the illumination correction functions.

5. IdentifyPrimaryObjects- Nuclei

Next we'll take a first pass at identifying nuclei and cells in our initial image.

- **After** the CorrectIlluminationApply module but **before** any others, add an IdentifyPrimaryObjects module (from the ‘Object Processing’ module category).
- Create objects called Nuclei by segmenting on the Hoechst channel. Hit ‘Step’ to run the module. How does your segmentation look?
- Use the magnifying glass at the top of the window to zoom in on an area that was segmented poorly, then update some of your parameters in IdentifyPrimaryObjects and hit ‘Step’ to rerun the segmentation.
- Adjust the segmentation parameters until you feel you’re ready to move on to identifying the cells around the nuclei; as you will test the parameters for robustness later, however, the identification should be good but doesn’t need to be perfect before you move on.

6. IdentifySecondaryObjects- Cells

- **After** the IdentifyPrimaryObjects module but **before** the EnhanceOrSuppressFeatures module, add an IdentifySecondaryObjects module.
- Create an object called Cells that is seeded on the Nuclei primary objects that you just created; use the Ph_golgi image.
- For the purposes of this exercise, you need not worry about excluding cell bodies that touch the edge of the image.
- Examine the segmentation and adjust the segmentation parameters until you feel you’re ready to test them on another image; they need not be perfect before you move on.

7. Test the robustness of your segmentation parameters across multiple compounds

It's (relatively!) easy to come up with a good set of segmentation parameters for a single image or a set of similar images; this data set however contains images from cells treated with many different classes of drugs, many of which have very different phenotypes. It's valuable to learn how to create a set of parameters that can segment cells that display a variety of morphologies since you may come across a similar problem in your own experiments!

- Go to Test->Choose Image Set to bring up a list of the images in your experiment.

Image_Metadata_SOURCE_COMPOUND_NAME	Image_Metadata_SOURCE	S.	Well	OrigER
NIFLUMIC ACID	Biomol International Inc.	1	A01	IXMtest_A01_s1_w22CA
NIFLUMIC ACID	Biomol International Inc.	2	A01	IXMtest_A01_s2_w246FF
Vitexin	Prestwick Chemical Inc.	1	A02	IXMtest_A02_s1_w2F2F5
Vitexin	Prestwick Chemical Inc.	2	A02	IXMtest_A02_s2_w28F2C
NICARDIPINE	Biomol International Inc.	1	A03	IXMtest_A03_s1_w2A057
NICARDIPINE	Biomol International Inc.	2	A03	IXMtest_A03_s2_w2B10C
6-Furfylaminopurine	Prestwick Chemical Inc.	1	A04	IXMtest_A04_s1_w2B174
6-Furfylaminopurine	Prestwick Chemical Inc.	2	A04	IXMtest_A04_s2_w2A259
LOPERAMIDE	Biomol International Inc.	1	A05	IXMtest_A05_s1_w271AC
LOPERAMIDE	Biomol International Inc.	2	A05	IXMtest_A05_s2_w21E2A

Figure 5: A section of the ‘Choose Image Set’ menu.

- Look at the column titled ‘Image_Metadata_SOURCE_COMPOUND_NAME’ to see what chemical was used in each well of the experiment. You may click on the column to sort the whole table by the values in it if you so desire.
- Choose a row where ‘Image_Metadata_SOURCE_COMPOUND_NAME’ is blank-this will be a mock treated well. Press the ‘OK’ button, then run that image in test mode for your first 3 modules (through your IdentifySecondaryObjects step). Examine the output – did your nuclear and cellular segmentation hold up compared to the first images you looked at? Once your segmentation is good, try it on one additional mock treated image.
- Test your segmentation on images from a few different compounds- you may choose ones you’ve worked with before, random ones, or some combination therein; if possible avoid using multiple compounds you KNOW have the same mechanism of action, though it’s alright if they occasionally do. Update your segmentation parameters until they work well on a few different compound wells, then go back to a mock treated well to make sure it still works well there.
 - You’re encouraged to explore the compound list on your own, but if you find yourself consistently ending up with images that look similar you can try adding images from the following list of wells-B18, C7, D6, D19, D22, E3
 - Some hints on what to play with:
 - In both IdentifyPrimaryObjects and IdentifySecondaryObjects adjusting the threshold limits can be a good thing for when wells are empty, confluent, or have bright debris in them
 - In both IdentifyPrimaryObjects and IdentifySecondaryObjects adjusting the threshold method may lead to better (or worse!) results.
 - In IdentifyPrimaryObjects, adjusting the declumping settings (make sure to turn ‘Use advanced settings?’ on) will probably be necessary for a robust segmentation
 - In IdentifySecondaryObjects, you will want to test the effects of using the various methods for identifying secondary objects (Propagation, Watershed-Image, Distance-N, etc) and, if using Propagation, the regularization factor.

8. IdentifyTertiaryObjects- Cytoplasm

- After the IdentifySecondaryObjects module but before the EnhanceOrSuppressFeatures module, add an IdentifyTertiaryObjects module.

- Create an object called Cytoplasm using the Cell and Nuclei objects you've created; 'Shrink smaller object prior to subtraction?' should both set to 'No'.

9. Examine the steps used to segment the Nucleoli

- The next 3 modules have to do with the creation of the Nucleoli objects. Look at the output from each to see how the image is transformed to aid in segmentation.
 - EnhanceOrSuppressFeatures is a module that helps enhance particular parts of an image- in this case, punctate objects or 'Speckles'. By specifying the feature size, you can enhance different parts of the object. As we are looking for nucleoli, we apply this to the RNA channel (Syto) image, and call the output 'FilteredRNA'. (See Fig 6 below)
 - MaskImage allows you to create a version of the 'FilteredRNA' image called 'SytoNuclei' where all of the pixels except the ones you specify are set to an intensity of 0- in this case, we set to 0 any pixel not inside a nucleus. By doing this, we can decrease the likelihood of detecting the cytoplasmic RNA dots.
 - IdentifyPrimaryObjects is used to find the Nucleoli- this is a Primary object segmentation because we are not using another object as a seed to grow around, but only segmenting based off the intensity in our 'SytoNuclei' image.
 - If you like, you can add an "OverlayOutlines" module at this point to overlay the identified nucleoli on the original Syto image to assure yourself that the segmentation not only matches the speckle-enhanced 'SytoNuclei' image, but also looks accurate on the unprocessed image as well. This is not necessary but can be a nice "sanity check".

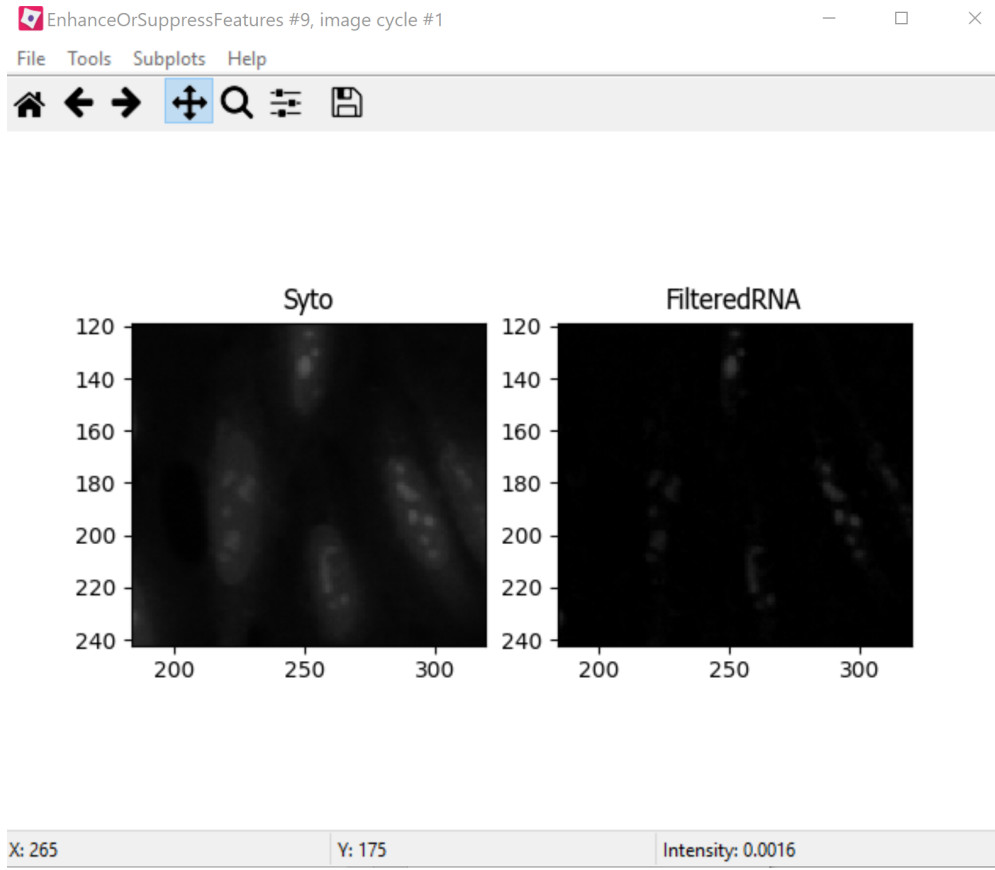


Figure 6: Enhancing the Syto image allows you to isolate nucleoli against the nucleoplasmic background signal.

10. Mask the Mito image by the Cytoplasm object

Now that you've seen an example of how to segment an organelle, you will do so for Mitochondria in the following steps.

- After the IdentifyPrimaryObjects module for Nucleoli but before the RelateObjects modules, add a MaskImage module (from the Image Processing module category).
- Call your output image ‘MaskedMito’.
- As you saw above with the Nucleoli example, mask the image via Objects, and use the Cytoplasm objects to create the mask.
 - You may even experiment with doing a similar EnhanceOrSuppressFeatures step before the masking as was used for the Nucleoli; you may get greater signal-to-noise, but possibly at the expense of “fragmenting” the Mitochondria objects in the later identification steps.

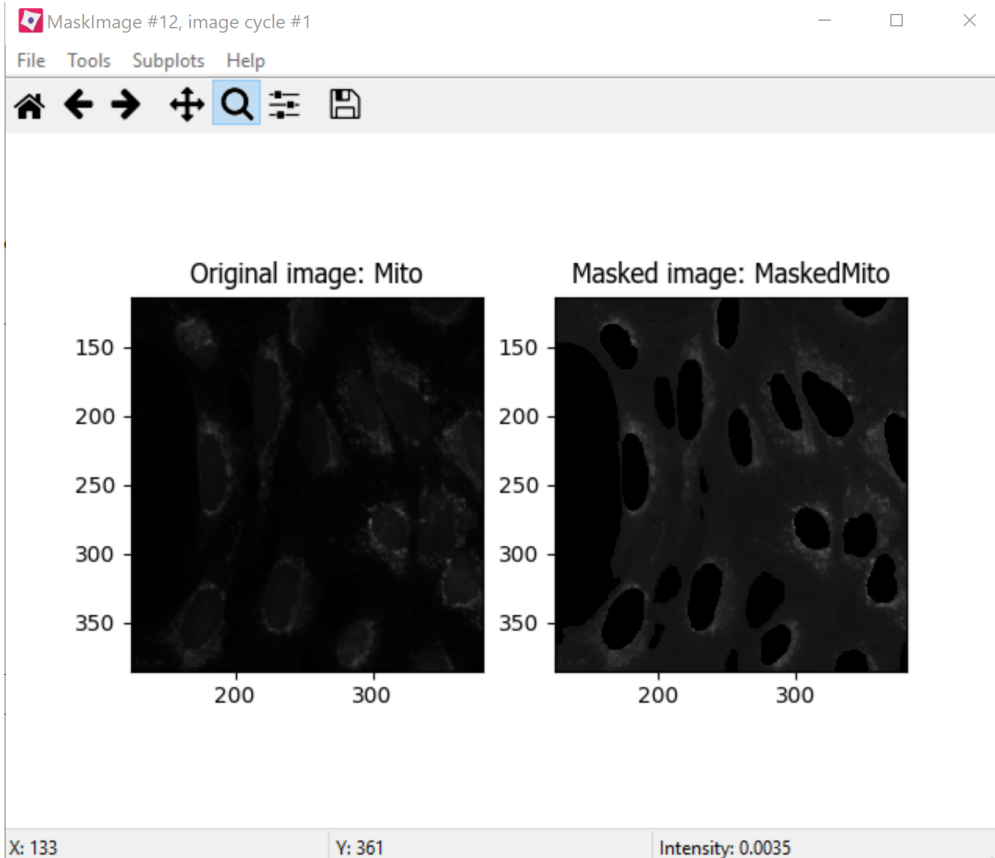


Figure 7: The MaskedMito image contains only the regions of interest.

12. IdentifyPrimaryObjects- Mitochondria

- After your MaskImage module but before the RelateObjects modules, add an IdentifyPrimary Objects module to identify Mitochondria from your MaskedMito image.
- You should consider using a wide range of pixel sizes here; 2-20 is a reasonable first place to start.
 - If you did use EnhanceOrSuppressFeatures in the previous step, using OverlayOutlines to compare the outlines with the original image is a good idea once again.

13. Add measurement modules to your pipeline

- After your segmentation of the mitochondria but before the RelateObjects modules, add as many object measurement modules as you would like.
- Some suggested modules to add- MeasureObjectSizeShape, MeasureObjectIntensity, MeasureGranularity, MeasureObjectNeighbors.
 - Which objects do you think would be valuable to measure with each of these modules? Which channels would you measure your objects in?
 - For a typical Cell Painting experiment you would add as many measurements as possible, but that isn’t necessary here; however, do make sure every object gets at least some measurements.
- While MeasureCorrelation, MeasureTexture, and MeasureObjectIntensityDistribution can produce valuable data for downstream profiling, they can be memory-intensive and/or slow so should not be added

for this example pipeline in the interest of pipeline run time. MeasureNeurons is not well suited for this pipeline.

14. Examine the settings of RelateObjects

- After your Measurement and before your Export modules you should find two RelateObjects modules. One relates Nucleoli to Nuclei, while the other relates Mitochondria to Cells.
- Relating the objects allows you to create per-parent means (ie, for this cell what is the average size of an individual mitochondrion) and calculate distances from the child objects to the edge and/or the center of the parent (ie how far is each nucleolus from the center of the nucleus).

15. Run the pipeline (optional)

- If you have time and/or if you'd like to play with the data in CellProfiler Analyst later, exit test mode, close the eyes next to each module, and run the pipeline
- The pipeline will create a database called BBBC022.db, containing the output of all of the measurements you have added to your pipeline
- Because you have different object counts for some of your different types of objects (the counts of Nuclei, Cells, and Cytoplasm will be the same, but the counts of Mitochondria and Nucleoli will not be), you will not be able to export the objects as a single data table but must instead use a different data table for each object. This will not affect the actual outcome of the experiment, but will mean that each object will get its own properties file and that you can only look at the measurement for one object at a time in CellProfiler Analyst.

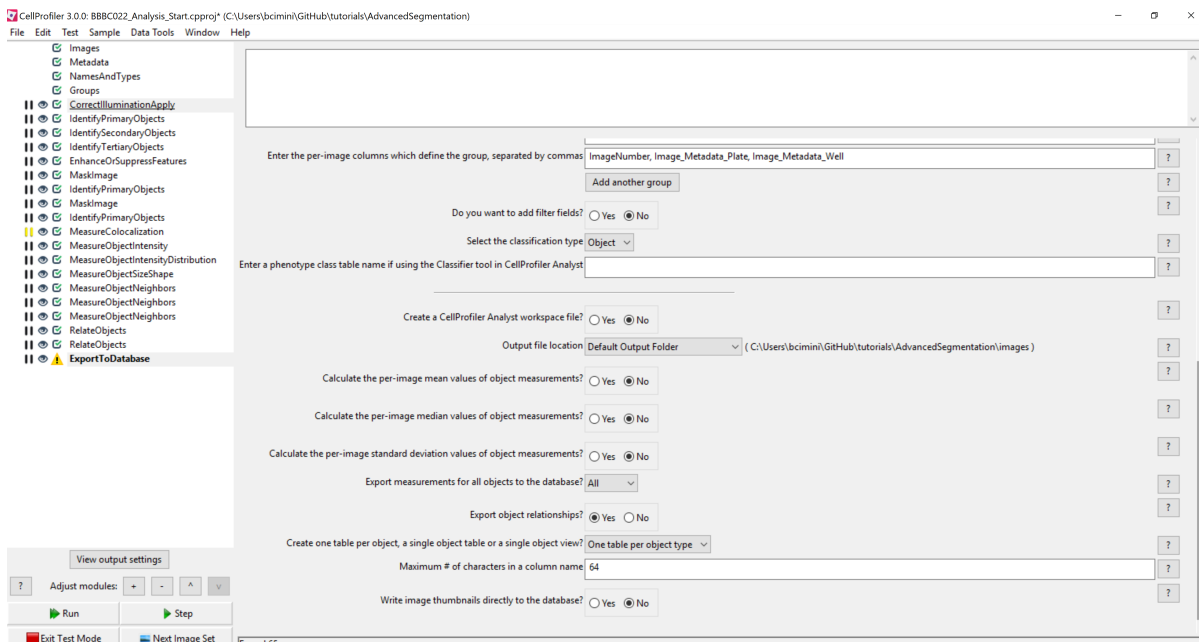


Figure 8: The ExportToDatabase module. The yellow warning symbol warns you that since you've chosen to make individual tables for each object, you will only be able to examine one object at a time in CellProfiler Analyst.