PCP Auto Count: A Novel Fiji/ImageJ plug-in for automated quantification of planar cell polarity

User guide

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Contents

Overview	2
Software requirements and plugin installation	3
Required image parameters:	4
Image Pre-processing	5
Brief overview of PCPA workflow and algorithmic processes	8
User options guide	12
1. Exclude chunks by size.	12
2. Exclude chunks near the image border.	12
3. Doublets/oblong chunks are split or excluded.	12
4. Plastic wrap	13
5. Exclude caves by size	13
6. Cave selection criteria applied	13
7. Orientation of angle measurements:	13
8. Customizable data output	13
Other PCPA functionalities	15
Counting of cells when researchers do not need PCP information	15
PCP Auto Count plugin batch processing (analyzing multiple images together)	15
Rose Diagram generation of non-PCPA data	15
References	16
Figures	17

Overview

Planar cell polarity is the systematic organization of cells in biological tissue, usually of epithelial origin, where cells exhibit coordinated directionality that arises due to a number of developmental factors. Planar cell polarity is an integral identity aspect for some cell types (e.g. cochlear hair cells, vestibular hair cells, fly ommatidia, etc.) and is required for proper functioning on an inter- and intracellular level. For investigators interested in planar cell polarity, the quantification of cellular direction is a tedious task where investigators must often manually draw directional arrows on every cell in a sample field. We have developed a Fiji (Fiji is Just ImageJ)[1] based plugin that automates the process of cell polarity measurements, called PCP Auto Count (PCPA).

Because planar cell polarity directional indicators differ by tissue type, we have elected to use generalized descriptive terms to explain the development and functioning of PCPA. Large planar surfaces (often the apical surface of a cell) are hereafter called "chunks" while prominent inclusions indicative of direction, often representative of cilium or groups of cilia, are hereafter called "caves." These chunk and cave configurations may also arise from combinations of various features, including the morphology of the cell or feature of interest (e.g. v-shaped features such as filiform papillae of the tongue or stereocilia bundles of mammalian outer hair cells). The following explanation guide provides examples using inner ear cochlear hair cells and vestibular hair cells. To visualize the cells' apical surfaces and find the directional indicator for each hair cell (called a fonticulus), we have elected to use BII-spectrin staining, which will bind to the spectrin-rich cuticular plate of a hair cell while leaving the actin-rich kinocilium unlabeled. The resulting immunohistochemistry produces the spheroid shape of the cell with staining absent from the kinociliar area. Validation of PCPA on other tissues, cell types, and staining methods is reported in Stansak et al. (2024; in prep). Our results suggest that PCPA should have broad applicability for any users who wish to measure the directionality of any cells or structures that adhere to a "chunk and cave" morphology. Readers are directed to consult Stansak et al. (2024; in prep) for additional information.

The goal of PCPA is to first identify cells in binary images, usually photomicrographs, then apply algorithmic processes to determine directionality for cells meeting user defined selection criteria, thus allowing for automated calculation of planar cell polarity angles. To accomplish this, PCPA identifies all white pixels in an image and creates grouped aggregates of adjoining pixels. User defined exclusion criteria is used to filter out artifact pixels and the remaining pixel aggregates are designated as regions of interest. While aggregates in the images herein generally represent "cells", other features of interest may be used in planar cell polarity studies, and other users may have non-biological images from which they seek to determine the orientations of certain features. Thus, we have elected to refer to aggregated pixel groups as "chunks." After the identification of chunks, the algorithm will then identify black pixels within the bounds of the chunk which we have termed "caves". PCPA will then assign the cave that best meets its user defined selection criteria as the directional cave of interest. An angle measurement (in degrees) is then calculated for each chunk by projecting a ray originating from the center of the chunk to the center of the cave.

Software requirements and plugin installation

Required software:

Fiji (Fiji is Just ImageJ), available from: https://imagej.net/software/fiji/

PCPA operates as a plug-in within the existing open-source image processing software, Fiji (Fiji is Just ImageJ)[1]. FIJI is compatible with Windows, MacOS, and Linux operating systems. More information about Fiji can be found at the link above.

Plugin installation:

- 1. Open Fiji
- 2. Select "Help > Update..."
- 3. Select "Manage Updated Sites"
- 4. Select "Add Unlisted Site"
- 5. Enter the following information into the new line-item's corresponding columns (without quotations)
 - a. Name: "PCP Auto Count"
 - b. URL: "https://sites.imagej.net/PCP-Auto-Count/"
- 6. Select "Apply and Close"
- 7. Select "Apply Changes"
- 8. To finish installation please restart Fiji
 - a. The PCP Auto Count menu will be found under "Analyze > PCP Auto Count"
 - b. The stand-alone rose diagram plugin will be found under "Plugins > Utilities > Rose Diagram"

Validation of the PCPA plugin suite has been published in Stansak et al. (2024; in prep)

PCP Auto Count is an open-access plugin. Source code can be found at

https://github.com/WaltersLabUMC/PCP-Auto-Count.git

A video demonstration of PCPA can be found at https://youtu.be/PTFIGv5Laa0

Required image parameters:

- \bullet A minimum image resolution of 1024 pixels along the longest edge of the image is strongly recommended for good performance when using images that contain approximately 10-100 cells
 - For images that are at lower resolution, users can try resizing images in Fiji through the following menu pathway: "Image > Adjust > Size..."
 - Note that while such size adjustments were generally beneficial in our development and validation testing of PCPA, images that have been resized may have artifacts or pixel mean gray values that may alter PCPA performance.
- A two-dimensional, binary (i.e. black and white pixels) image
 - Pre-processing steps which we have found to benefit PCPA performance on micrographs
 of inner ear tissues (e.g. thresholding) may not work on RGB, CMYK, or other color
 composite images.
 - Instructions on converting color images to grayscale and/or binary are explained in the "Image Pre-processing" section below.
- File type
 - While PCPA will run on any images that can be opened with Fiji, images should be saved
 at the time of acquisition as a file type that prevents data compression. Most inbuilt
 microscopy software defaults to save images in lossless format (e.g. CZI, ICS, OME-TIFF,
 LIF, ND2, etc.). Other lossless formats such as TIFF, PNG, and BMP are also acceptable.
 - JPEG and other formats that alter resolution or compress are not recommended

Recommended imaging parameters:

- We have found that PCPA works most effectively on micrographs acquired using a 40X or higher primary objective lens. Samples imaged at lower objective magnification may or may not have sufficient detail to convert well to binary images and users should use their best judgment based on their specific images and experimental goals.
- Image acquisition tips:
 - High signal-to-noise ratio antibody labeling is desirable
 - Samples used to demonstrate PCPA utility in this manual were immunolabeled with βII-Spectrin and were imaged to make the βII-Spectrin positive cellular features as bright as possible, such that nearly all of the pixels that comprise each chunk are at the level of complete saturation.

Image Pre-processing

The PCP Auto Count plugin requires users to convert their images to binary (i.e. all pixels are assigned 1 of 2 values: black or white). Several preprocessing steps are recommended to create a binary image with sufficiently removed background. Depending on staining practices, imaging parameters, and type of sample used, each user will need to decide which preprocessing steps result in the best binary images to use with the PCP Auto Count plugin.

Below are recommended preprocessing steps which can be carried out using existing functions within FIJI (Fiji is Just ImageJ). Each step includes a brief explanation of purpose and function, plus a few listed alternatives. This guide is not meant to be an exhaustive list of preprocessing measures users may take, rather, a starting point which showcases Fiji features our laboratory has used to generate binary images from several types of biological samples as reported in Stansak et al. (2024; in prep). Users are encouraged to develop a preprocessing protocol that works best for their project goals. All example images in the following guide were obtained using a Zeiss LSM 880 confocal microscope with a 40X or 63X oil immersion objective at 2048 x 2048 pixel resolution.

Step 1: Flatten z-stack and crop image.

PCPA is designed to work on two dimensional images. If a z-stack or time-series set of images are contained within the same window, PCPA will only analyze the window that is visible at the time the plug-in is run. Users are therefore advised to generate a 2D projection image from their z-stack and ensure images are grayscale or monochromatic. The easiest way to check if an image is monochromatic is to select "Image > Type" and verify your image is set as 8-bit. Crop out extraneous areas of the tissue which will not be quantified (Fig 1A-1B).

Image preprocessing, particularly the thresholding step necessary for creating binary images, works best for images where the mean gray value (i.e. intensity or "brightness") is relatively uniform across the image. In the case of cochlear samples used for illustrative purposes in this manual, we found that cochlear inner hair cells (IHC) showed significantly brighter fluorescence for our protein of interest (β II-Spectrin) than cochlear outer hair cells (OHC). The signal difference between the cell types caused issues during the thresholding step; to address this we cropped our image to analyze IHC and OHC

Optional steps:

- For images with low signal-to-noise ratios, Fiji's enhance contrast ("Process > Enhance Contrast...") may help. Otherwise, manual adjustments to brightness and contrast may also be beneficial.
- Selecting Equalize histogram will spread pixel intensities over a broader range and also decrease spread between brightest and dimmest pixel intensities. This differs from using linear contrast enhancements (Image > Adjust > Brightness/Contrast), which will alter spread of pixel values but does not affect relative pixel value in relation to brightest and dimmest pixels.
- Spherical aberration caused by microscope lenses can lead to decreased fluorescence intensities
 for features near the edges of the field of view. The Pseudo Flat Field Correction tool available in
 the BioVoxxel Toolbox (www.biovoxxel.de) plug-in for Fiji/ImageJ
 (https://imagej.net/plugins/biovoxxel-toolbox) can help normalize signal-to-noise ratio across
 the image

Step 2: Remove background signal.

Removal of background signal can be accomplished in several different ways using inbuilt Fiji functionality or Fiji plug-ins.

- Recommended: Fiji's Subtract Background ("Process > Subtract Background..."; Fig 2)
 - Subtracts background using a rolling ball radius to remove background staining while preserving low intensity pixels.
- <u>Alternative</u>: If your image has high variability of fluorescent intensity between cells that is not sufficiently addressed using Fiji's subtract background function, users may wish to try more nuanced background subtraction applications, such as the Biovoxxel toolbox plugin for Fiji (www.biovoxxel.de; Discussed in Step 1) which features a convoluted background subtraction feature that increases contrast by subtracting a modified filter from the entire image.
 - The convoluted background subtraction tool is heavily recommended over Fiji's Subtract Background if users applied Biovoxxel's Pseudo Flat Field Correction to adjust for spherical aberration (discussed in Step 1 optional steps).

Step 3: Threshold image and make binary.

Fiji's threshold feature allows users to select what pixel intensity defines the cutoff boundary for creating a binary image. The selected pixels will be assigned a value of 1 ("white") and all pixels not selected will be assigned a value of 0 ("black"). Choose one of the following methods, or a comparable method of the user's choosing.

- Recommended: Fiji's Manual Threshold Selection ("Image > Adjust > Threshold...")
 - This method gives users greater control than Fiji's Auto Threshold function. User generated threshold selection is recommended when signal-to-noise ratios and image quality vary across sample images in a data set.
 - Users should adjust the threshold bar to incorporate as many cells as possible, while
 preventing cells from dilating to the point of touching surrounding cells (Fig 3A, 3A', &
 3C). PCPA has a limited ability to "separate" two cells that touch but this function
 cannot account for more than two cells touching (See: "Doublet Splitting" in the "PCPA
 workflow and algorithm" section of this document).
 - Users should also ensure the directional point of interest is not obliterated during the
 thresholding process (Fig 3C). A directional point of interest that is not completely
 enclosed by the cell body can be addressed using user features outlined below (See:
 "Plastic Wrap" and "Cave Size Exclusion" sections in the "PCPA workflow and algorithm"
 section of this document)
- Alternative: Fiji's Auto Threshold function: "Image > Adjust > Auto Threshold"
 - Auto threshold works best on images with high signal-to-background ratios that are
 consistent across the entire image. Users may choose any type of auto threshold
 method which best compliments their images, but for the purpose of this document the
 method is selected as Default (Fig 3B & 3B').

Step 4: Remove outlier pixels.

Fiji's Remove Outliers function examines each pixel and converts it to the median color of the surrounding pixels in a user defined radius. This function is useful for removing small pixels or groups of pixels that are remaining artifacts left after the thresholding function is performed (Fig 4).

• Recommended: Fiji's Remove Outliers "Process > Noise > Remove Outliers..."

- Optional: Fiji's erode function ("Process > Binary > Erode").
 - The erode function removes a small number of pixels from the edge of objects in a binary image. This can be useful for images where cells touch their neighbors after thresholding the image (Fig 5).
- Optional: Fiji's dilate function ("Process > Binary > dilate")
 - The dilate function adds a small number of pixels to the edge of an object in a binary image. This function can be useful for samples where small holes occur after thresholding the image due to uneven staining intensity within a cell.

Brief overview of PCPA workflow and algorithmic processes

The general algorithmic processes used by PCPA are described below, in order of operation used by PCPA during analysis (Fig 6). All steps marked as optional are dependent on whether or not a user selected the setting prior to running PCPA. If no user settings are indicated for a step marked optional, PCPA will skip that step and continue on to the next process in its workflow. Optional steps are explained in terms of theory in this section. To see user options explained in practical terms, please see the User Options Guide chapter.

Chunks are identified

PCPA processes pixels on a Cartesian (x, y) coordinate system where the topmost, leftmost pixel in the image is assigned the coordinates (0, 0) with x values increasing left to right and y values increasing top to bottom. PCPA will begin parsing pixels starting from the top of the first column (coordinates: 0, 0) and subsequently descend the column until it encounters a white pixel or the image border is reached and the column terminates (Fig 7A). If PCPA encounters any white pixels in Column 0, PCPA records the XY coordinates and moves down the column to the next pixel. If the next pixel is also white PCPA will categorize the XY coordinates of the two touching pixels as belonging to the same group which we have termed a "chunk". Abutting white pixels will continue to be added to the same chunk until PCPA encounters a black pixel, in which case the chunk is considered complete for that row and the next white pixel encountered in the column will become the start of a new chunk (Fig 7B). At the termination of Column 0, all subsequent columns will be evaluated in the following manner: PCPA will descend from the top of the column until it encounters a white pixel or the image border is reached and the column terminates. In the case of encountering a white pixel, PCPA will check if the pixel abuts a white pixel in the column directly to its left (Fig 7C & 7D). If so, the pixel of the current column will be assigned to the chunk from the previous column (where: end1 >= start2 and end2 >= start1). If the pixel in the current column does not abut a white pixel to its left, then PCPA will create a new chunk designation This operation will recurse until PCPA has evaluated all columns in the image.

Optional step: Exclude chunks by size.

If users have set minimum or maximum size exclusion settings for chunks, PCPA will use these parameters to exclude all chunks that meet exclusion criteria from subsequent processes (Fig 8A & 8A').

Optional step: Doublets/oblong chunks are split or excluded.

Natural biological variability, insufficient magnification during imaging, and a variety of other factors can cause instances of cells crowded into close proximity. In mouse models, for instance, outer hair cells stained with β II-spectrin do not tend to overlap or touch, whereas inner hair cells typically have less space between cells, causing oversaturated pixels to touch. PCPA has a limited ability to separate a maximum of two cells that overlap. These user options are not able to recurse beyond splitting a "doublet" of cells. Users can set a threshold ratio to detect doublet cells by specifying a ratio of chunks that are taller than they are wide, and/or chunks that are wider than they are tall. If a chunk meets doublet criteria PCPA will encase the chunk in a bounding box, then split the bounding box down the middle, where direction of splitting will be based on whether the chunk violates "taller-than-wide" (horizontal split) or "wider-than-tall" (vertical split) criteria (Fig 8B-8B"). Once a doublet is split into two boxes, PCPA will then treat all pixels contained within each box as an independent set and apply user-defined chunk and cave criteria to pixels within each of the boxes. While PCPA is unable to split more than two merged chunks, PCPA user settings allow for size exclusion criteria to be applied (either

separately or in conjunction with Doublet Splitting) so that users are able to exclude large conglomerates of merged chunks from further analysis.

Optional step: Exclude chunks near the image border.

Removing chunks a set number of pixels from the image border allows the user to filter out cells that were not completely captured within the microscope's field of view during image acquisition. As the center of mass of a chunk is used to determine the angle of orientation, it is generally wise to exclude chunks at the image border which may have a distorted center position due to cropping. This setting creates an exclusionary border around the outer edge of the image and any chunk, regardless of size, touching this exclusion zone will be filtered out of subsequent steps (Fig 8A & 8A').

Optional step: Plastic wrap

PCPA defines "caves" as all aggregates of black pixels that are fully contained within the body of a chunk (i.e. completely surrounded by white pixels constituting the chunk). Depending on cell morphology and imaging quality, it is possible that true directional markers may not be fully enclosed within the cell body, and thus could be incorrectly ignored by PCPA's definition of "cave." If PCPA cannot find a cave that meets selection criteria, the chunk will be excluded from subsequent PCP analysis.

In order to overcome this limitation, we have developed a feature called Plastic Wrap, which is included in PCPA's user options menu. To begin Plastic Wrap, PCPA draws a square or rectangular box that encompasses all pixels of the chunk, termed a bounding box. Starting at the lowest value coordinates (0, 0) of the bounding box, PCPA scans Row 0 from left to right until it encounters a white pixel. PCPA then checks directly to the right of the beginning pixel for an adjacent white pixel. If encountered, the abutting pixel is added to the plastic wrap coordinates. PCPA will continue chaining white pixels that abut to the right of the previous pixel until PCPA detects an orphan pixel (i.e. a white pixel with no abutting white pixel in the current scanning direction). In this case, the algorithm will attempt to find the nearest white pixel by plotting an exploratory line from the orphan pixel to the last column of Row 0, using slope intercept formula. If no white pixels that are part of the chunk being assessed fall within the exploratory line, PCPA will next plot an exploratory line from the orphan pixel to the last column of Row 1. This pattern will recurse down the rows of the last column until PCPA encounters a white pixel of the chunk that falls within the exploratory plotted line. PCPA will then include the found pixel as the end of the exploratory line, and any part of the line that bridges the gap between the starting pixel and the terminating pixel will be added to the pixel mass. The terminating pixel now becomes the end of the chain, and PCPA will continue using a similar strategy as outlined above to either add abutting pixels to the chain or designate a new orphan pixel at the end of the chain. In the case of an orphan pixel, PCPA again sends out an exploratory line, which will now start at the last column of the orphan pixel's row, rather than start at the last column, row 0 (Fig 9B-E). Connecting orphan pixels to terminal pixels in an exploratory line occurs in the same manner as described above (Fig 9F).

Plastic Wrap will continue in a similar fashion until it encounters an orphan pixel that directly abuts the bounding box. Next, PCPA will change its chaining strategy from moving across row 0 to moving down the last column, following a similar manner as described above, and again terminate when the orphan pixel abuts the bounding box. Then, PCPA will move across the bottom row from left to right, following a similar manner as described above, and again terminate when the orphan pixel abuts the bounding box. Finally, PCPA will move up Column 0 from bottom to top, following a similar manner as described above. Once Plastic Wrap has connected the last orphan pixel to the pixel that began the Plastic Wrap chain,

the operation is finished. All pixels added as part of joining an orphan pixel and a terminal pixel are considered as part of the chunk mass for all subsequent calculations (Fig 9G). In this way, indentations in chunks can become fully enclosed caves and directionality can be determined from the most prominent indentation in a chunk (or the indentation that most closely aligns to the user specified direction).

Chunk centroids are calculated

The centroid of each chunk is calculated using the same programming logic as Fiji Image J's center of mass function, which calculates the average of the X and Y coordinates of all pixels in the chunk. The chunk centroid value is used as the vertex for planar cell polarity calculations of each cell.

Cave candidates are identified

PCPA will perform a similar operation as described in the chunk identification section to locate all caves within each chunk and record their (x, y) coordinates. To be recognized as a "cave," inclusions of black pixels must be enclosed within the body of the chunk (this includes black pixels enclosed by using the Plastic Wrapping user option).

Optional step: Exclude caves by size

If users have set minimum or maximum size exclusion settings for caves, PCPA will use these parameters to exclude all caves that meet exclusion criteria from subsequent processes

Cave selection criteria is applied

In the absence of user input for cave exclusion size and cave directionality, PCPA will assign the largest cave as the cave of interest for subsequent planar cell polarity calculations. While the majority of users will be best served by setting the cave of interest selection criteria as largest cave, we have afforded users the option to apply a directionality filter (northmost, southmost, westmost, easternmost, highest position, lowest position, leftmost position, or rightmost position) to the cave selection process. Whichever cave best meets the largest cave criteria, or user-defined direction criteria, will be designated the cave of interest for the chunk (Fig 10).

Cave centroids are calculated

The centroid of each cave is calculated using the same programming logic as Fiji ImageJ's center of mass function, which calculates the average of the X and Y coordinates of all pixels in the cave of interest. The cave centroid value is used as the point the ray is drawn through for planar cell polarity calculations of each cell.

Planar cell polarity is calculated and data is output

The values obtained for the centroid of the chunk and the centroid of the cave are used to calculate the inverse tangent of the slope of the line that connects these two points, and this value is used to draw directional arrows on the output image. The inverse tangent of differences in x and y is used to determine the angle measurement in degrees for each cell. Users can customize the reference for angle measurements by setting the axis scale (-180 $^{\circ}$ /180 $^{\circ}$ or 360 $^{\circ}$), axis direction (clockwise or counterclockwise) and direction of 90 $^{\circ}$ axis (north, south, east, or west). The following summary statistics (based on calculations from Fisher [2]) are calculated for the angle measurements and presented as a results table (Fig 14):

- · Processed Count: Count of analyzed chunks
- Bad Count: count of chunks that were excluded from PCP angle calculations due to violating set size parameters or because they did not have a recognizable cave.
- Total Count = Processed Count + Bad Count
- Processed % = $\frac{Processed\ Count}{Total\ Count} * 100$
- Resultant Mean Length (RML) = $\sqrt{\bar{c}^2 + \bar{s}^2}$

$$\bar{C} = \frac{\sum_{1}^{n} \cos \theta_{n}}{n}$$

$$\overline{\mathcal{S}} = \frac{\sum_{1}^{n} \sin \theta_{n}}{n}$$

- Variance (V) = 1 RML
- Standard Deviation $(\sigma) = \sqrt{-2\ln(RML)}$

User options guide

To set processing parameters and PCPA output options, users should select the following Fiji menu pathway: "Analyze > PCP Auto Count > Measure With Options...". Detailed explanations of the PCPA options menu are found below. PCPA will remember last user settings applied and for subsequent uses, if the user does not wish to change any options, users may select "Analyze > PCP Auto Count > Measure Now." It is recommended that users familiarize themselves with the tools provided below in order to maximize accuracy of PCPA measurements.

Exclude chunks by size (Fig 11A)

Under the "Chunk Detection" tab, users can set size exclusions to disregard chunks falling above and/or below a specified number of pixels by entering values in "Remove noise of this many pixels or fewer" and/or "Remove 'conglomerates' of this many pixels or more" fields. Chunks removed from further analysis via minimum or maximum size exclusions can be optionally indicated on the output image with pseudocoloring (see output description below and Fig 11A').

To set a minimum size requirement, users will select the checkbox then enter a numerical value into the "Remove noise of this many pixels or fewer" field. Fairly low minimum size requirements will remove noise (i.e. lone white pixels or small aggregates of pixels) similar to preprocessing steps outlined in the preprocessing section of this manual. Larger minimum size requirements can be set to exclude objects that are not cells or features of interest. Validation with mouse hair cell micrographs suggests a minimum size requirement to approximately 33% of the size of your average cell size will filter out irrelevant aggregates from most images, and values ~ 50-75% of the size of your average cell can allow investigators to filter out immature or fragmented cells (Fig 11A). Setting a maximum size requirement is achieved by selecting the checkbox then entering a numerical value into the "Remove 'conglomerates' of this many pixels or more" field.

Exclude chunks near the image border (Fig 11A)

Under the "Chunk Detection" tab, users may optionally set an exclusion zone near the image border to filter out cells that were incompletely captured by the microscopy field of view (Fig 11A). Users may find a small (e.g. 1-10 pixel) exclusion zone to be sufficient for excluding incompletely imaged cells, though cropping, resizing, or reorienting of images may require larger values. Border excluded cells can be optionally indicated on the output image with pseudocoloring (Fig 11A').

Doublets/oblong chunks are split or excluded (Fig 11B)

Users should endeavor to avoid allowing cells to touch during the thresholding preprocess. However, PCPA does contain limited ability to address merged chunks (termed "doublets"), though this operation is not able to address instances where more than two cells touch. Under the "Chunk Detection" tab users have the option to split chunks by setting a ratio of wider-than-tall and/or taller-than-wide. Depending on settings used, PCPA will split the doublet vertically or horizontally, respectively, into two equally sized cells that are each treated as independent chunks for further processing steps (Fig 11B'). Users may alternatively choose to exclude doublet chunks from further analysis. It is important to note that the success of the doublet splitting is optimal when the orientation of the doublet is vertical or horizontal; accuracy of doublet splitting decreases as the orientation of the doublet approaches a 45°, 135°, 225°, or 315° axis. If triplets or larger conglomerates of cells are present, Doublet Splitting is no longer possible and users should apply maximum size exclusion criteria (explained above) to remove conglomerates

Plastic wrap (Fig 12)

While PCPA was initially developed to detect fully enclosed caves, such as what is commonly seen in β II-spectrin labeled hair cells, biological and experimental variability often led to images with cells that did not have a "cave" that was completely enclosed by white pixels (Fig 12A'). Furthermore, a wide array of sample types in planar cell polarity studies have morphology more consistently described as indentations rather than holes.

In default mode, a chunk will be excluded from subsequent analysis if PCPA cannot find a cave that is not fully enclosed. PCPA has an optional, but recommended, feature called Plastic Wrap which will enclose concave features by following the perimeter of the cell and adding white pixels to bridge any gaps in the arc of the cell's perimeter. Under the "Cave Detection" tab, users should select the box labeled "Apply Plastic Wrap to chunks before cave detection" to use the Plastic Wrap feature (Fig 12A).

Exclude caves by size (Fig 12)

For any chunks that are not smoothly rounded (or perfectly rectilinear) there will often be a variety of small, concave inclusions of black pixels. The noise removal step described above in image preprocessing often removes many of these. However, persistent inclusions can be filtered out of the cave selection process by setting a lower size limit for cave selection (Fig 12B). Conversely, cell types that lack a common geometrical shape (such as is seen in basal cochlear cells which resemble a boomerang or horseshoe shape) can cause large artificial caves to be created during the Plastic Wrapping process. To control for this, users can also set an upper size limit for cave selection under the "Cave Detection" tab (Fig 12B).

Cave selection criteria applied (Fig 12)

PCPA's default mode is to assign the largest cave that does not exceed optionally set size limits as the cave of interest for subsequent planar cell polarity calculations. While the majority of users will be best served by setting the cave of interest selection criteria as largest cave, we have afforded users the option to apply a directionality filter to the cave selection process, found under the "Cave Detection" tab. The directional options are: northmost, southmost, westmost, easternmost, highest position, lowest position, leftmost position, or rightmost position (Fig 12C). The compass directions are determined using the angle calculation of the chunk and potential cave centroids whereas the body relative directions (leftmost, etc.) are determined using x or y values of the potential caves only. These directionality features should be used with minimum cave size requirements so that incidental inclusions that may best match directionality settings are discarded (Fig 12C').

Orientation of angle measurements (Fig 13)

Under the "Angle Measurements" tab, users can choose the orientation of the axis used to create PCP measurements. This allows users to standardize orientation across images by measuring all images on the same axis, or to customize the axis for each image measured. The latter approach allows users to standardize data collection across a variety of images that are not in the same anatomical orientation, without manually rotating the images prior. Users can customize how angle measurements are calculated by setting the axis scale (-180°/180° or 360°), axis direction (clockwise or counterclockwise) and direction of 90° axis (north, south, east, or west; Fig 13).

Customizable data output (Fig 14)

Under the "Output" tab, users can select which output boxes they wish to view after PCPA is ran. The output options include:

• Results Table

- Label: number assigned to a specific cell
- Chunk Centroid X: chunk centroid coordinates on the X-axis, calculated as:

$$\bar{x} = \frac{\sum_{1}^{n} x}{n}$$

• Chunk Centroid Y: chunk centroid coordinates on the Y-axis, calculated as:

$$\bar{y} = \frac{\sum_{i=1}^{n} y}{n}$$

• Cave Centroid X: cave centroid coordinates on the X-axis, calculated as:

$$\bar{x} = \frac{\sum_{1}^{n} x}{n}$$

• Cave Centroid Y: cave centroid coordinates on the Y-axis, calculated as:

$$\bar{y} = \frac{\sum_{i=1}^{n} y}{v}$$

- Vector length: square root of distance between chunk centroid and cave centroid
- Angle^o: PCP angle measurement (in degrees), calculated as the inverse tangent of the slope between the chunk centroid and cave centroid

• Chunk Summary

- Processed Count: count of successfully analyzed chunk-cave pairs
- Bad Count: count of chunks that were excluded from PCP angle calculations due to violating set size parameters or because they did not have a recognizable cave.
- Total Count = Processed Count + Bad Count
- Processed %: chunks processed as a percentage of total chunks
- RML: Root mean resultant length
- Variance
- Mean Angle
- Standard Deviation

Rose diagram

- Overlay: contains cell label, angle measurement, and directional line. Each aspect of the overlay can be optionally removed (Fig 14B). The overlay can be superimposed over the original image ("Image > Overlay > Add image..." select "zero transparent") to allow investigators to view the planar cell polarity line, label, and/or angle on the original image.
 - Arrows: directional arrows
 - Labels: number assigned to a specific cell; corresponds with chunk summary table
 - Angles: calculated angle of chunk-cave pair (in degrees)
- Image: the image fed into PCPA with overlays for various characteristics, including chunk label, angle measurement, directional arrow, Plastic Wrap, doublet splitting, and color coding of excluded chunks (Fig 14D). Color coding can be customized for accessibility under the "Colors" tab (Fig 14E).

Other PCPA functionalities

Counting of cells when researchers do not need PCP information

If users wish to count cells, but do not need or wish to view PCP angle measurements, Users will proceed in a similar manner as if they wished to obtain PCP angle measurements. On the PCPA user options menu, users should still set chunk size exclusions, split doublets, and exclude cells on the image border as needed for their images. It is not necessary to use Plastic Wrap. On the "Output" tab, we recommend selecting "Chunks Summary," "Overlay," and "Image" outputs (Fig 15A-B). For "Overlay Options" select "Labels." PCPA will then analyze the image as if it were collecting PCP information, including excluding chunks it cannot find a cave for. The output for the Chunk Summary table will then show how many cells had PCP angle measurements calculated ("Processed Count") and how many cells did not have angle measurements calculated (i.e. chunks lacking a cave; "Bad Count") (Fig 15C). To obtain total cell count (sum of "Processed Count" and "Bad Count"), users should refer to the "Total Count" value. All data fields beyond "Total Count" are related to PCP information, and can be ignored if only the cell count is desired. Please note that chunks excluded for violating chunk size requirements or border proximity are not included in the "Bad Count" or "Total Count" value.

PCP Auto Count plugin batch processing (analyzing multiple images together)

PCPA can analyze multiple images together (batch processing). This option considers each image of the batch to be a continuous set of data and all output metrics will be calculated based on data aggregated from all batched images. To batch process multiple images, open all desired images in Fiji then select "Analyze > PCP Auto Count > Measure All Open Images With Options..." if the user wishes to change any processing or output options for PCPA (Fig 16). If users wish to batch process images using last used settings, select "Analyze > PCP Auto Count > Measure All Open Images Now..."

Rose Diagram generation of non-PCPA data

Included in the PCPA package is an option for users to generate a rose diagram and summary statistics using data provided by the user. This action is independent from running PCPA data, so users can create rose diagrams and summary statistics tables from any circular data they wish (Fig 17). To use the standalone rose diagram function, select "Plugins > Utilities > Rose Diagram. Users can then enter or paste angle measurements into the input box, as one angle per line. Users are given options to input the axis orientation used for the data set, and to customize the rose diagram bar size and angle markers. Users may also obtain summary statistics (root mean length, variance, mean angle, and standard deviation) for the data set by checking the box for Angle Summary

References

- 1. Schindelin J, Arganda-Carreras I, Frise E, Kaynig V, Longair M, Pietzsch T, et al. Fiji: an open-source platform for biological-image analysis. Nat Methods. 2012;9: 676–682. doi:10.1038/nmeth.2019
- 2. Fisher NI. (1993). "Descriptive Statistics," in Statistical analysis of circular data. (New York, NY, USA: Cambridge University Press), 15-37.

Figures

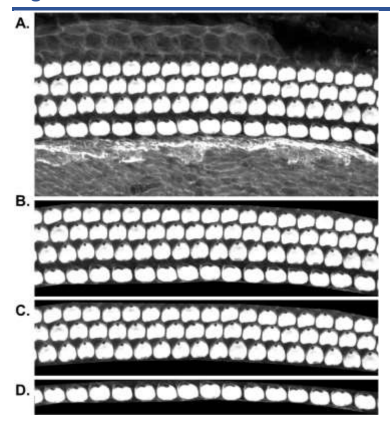


Figure 1. Image acquisition and field of view cropping. (A) Maximum intensity projection of z-stack images from P4 cochlea (63X oil immersion objective; 2048 dpi; 8-bit). **(B)** The original image was cropped to include only IHC and OHC. **(C-D)** The outer hair cells and inner hair cells were saved as separate images and preprocessed separately.

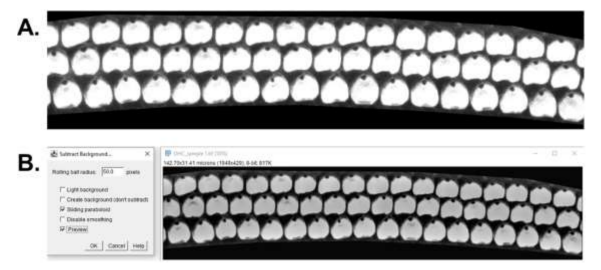


Figure 2. ImageJ's Subtract Background function. **(A)** Original cropped image of P4 cochlea (63X oil immersion objective; 2048 dpi; 8-bit). **(B)** P4 cochlear image after applying a 50-pixel rolling ball radius background subtraction with sliding paraboloid.

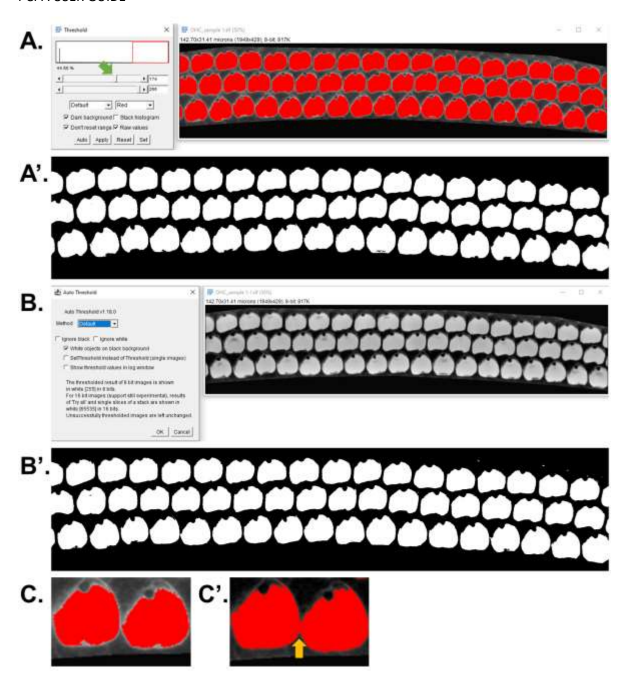


Figure 3. Threshold image to make binary. (A) To use the Manual Threshold function, users should use the top bar (green arrow) to adjust the amount of thresholding applied (appearing as red pseudostaining in the open image). Users should set the threshold to maximize inclusion of cell mass pixels while minimizing inclusion of background staining. (A') Binary image created by Manual Threshold function.
(B) Fiji ImageJ features an Auto Threshold option with various in-built thresholding methods. Auto Threshold may be useful on well imaged samples with minimum background staining. (B') Binary image created by Auto Threshold function (default method). (C) Whatever approach chosen, users should endeavor to prevent threshold levels that cause cells to dilate to the point of (C') touching surrounding cells.

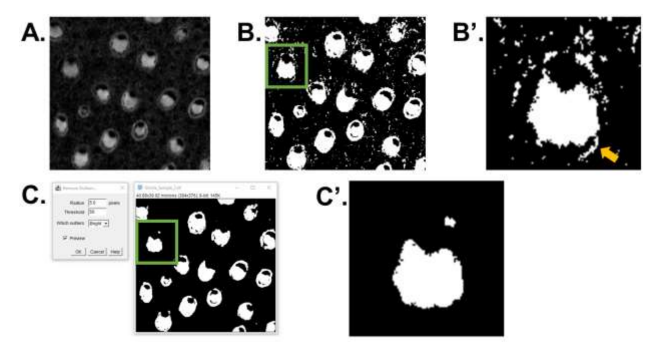


Figure 4. ImageJ's Remove Outliers function. **(A)** E17.5 utricle (40X oil immersion objective; 2048 dpi; 8-bit) after background subtraction. **(B)** Image after manual thresholding, with accidental inclusions of background staining. **(B')** Subset of B, showing pixels near the bottom of the cell (yellow arrow) that may cause PCPA's plastic wrap function to create an incidental cave that could interfere with correct angle calculations. **(C)** Fiji ImageJ's inbuilt remove outliers function allows users to remove outlier pixels via by a user defined pixel radius. **(C')** Subset of C, showing cell after removing outlier pixels (radius: 5 pixels; threshold 50) from the image.

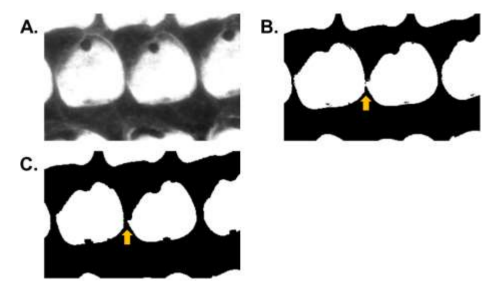


Figure 5. ImageJ's erode function. **(A)** P4 cochlear hair cells (63X oil immersion objective; 2048 dpi; 8-bit) original image. **(B)** After manual thresholding of the image, two cells touch (yellow arrow). Left unaddressed, this will cause PCPA to categorize the two cells as one long cell and subsequently produce incorrect angle measurements for this doublet. **(C)** After applying 2 iterations of ImageJ's erode function the cells no longer touch.

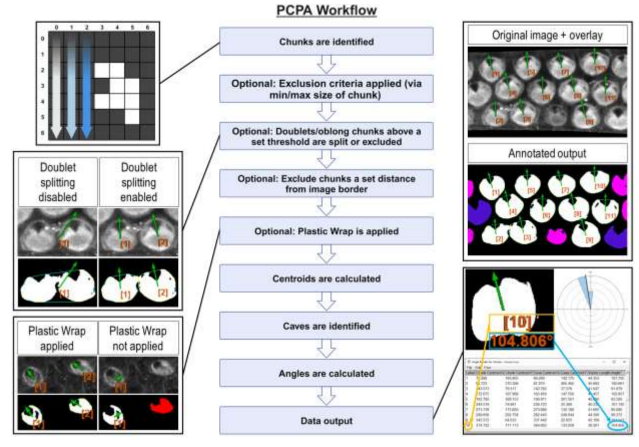


Figure 6. Overview of PCPA's algorithmic flow. (A) PCPA identifies aggregates of white pixels in a binary image, termed chunks. (B) Chunks can be discarded from subsequent operations via an optional pixel size minimum and/or maximum. (C) PCPA has limited (maximum of two adjacent cells) capacity to separate (or exclude) chunks that touch as a result of the thresholding process during pre-processing. (D) Cells that fell partially outside of the imaging frame can be discarded by setting an exclusion zone of a set number of pixels away from the edge of the image frame (cells pseudo-colored blue). (E) In the absence of Plastic Wrap, fonticuli that are not completely enclosed in β-spectrin staining cannot be recognized by PCPA. Applying Plastic Wrap will add pixels to bridge the gap in the cell's perimeter where the unenclosed fonticulus would be. (F) The center of the chunk's mass is calculated. (G) The cave best meeting selection criteria is identified as the directional cave of interest. (H) planar cell polarity is calculated using the chunk centroid as the vertex and the cave centroid as the ray node indicating directionality. (I) PCPA outputs include annotated images specifying chunks excluded due to violating chunk size limits (cells pseudo-colored pink) or touching the border of the image (cells pseudo-colored blue). Cells are labeled in the annotated output and correspond with the angle results table, which proves a detailed output of summary statistics for each chunk that was quantified. Rose diagrams and summary statistics tables (not shown) are also produced.

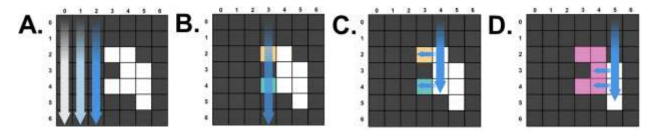


Figure 7. Example image demonstrating how PCPA identifies pixels of interest. **(A)** PCPA scans down column rows to find white pixels. At the end of each column, PCPA recurses and begins again at the top of the next column. **(B)** If PCPA encounters a white pixel it will determine if the white pixel abuts any other white pixels in its current column or the column to its left. If it does not, a new pixel aggregate called a chunk is recorded. **(C-D)** If white pixels in the current column abut any white pixels to their left, all of these pixels will be assigned to the same chunk.

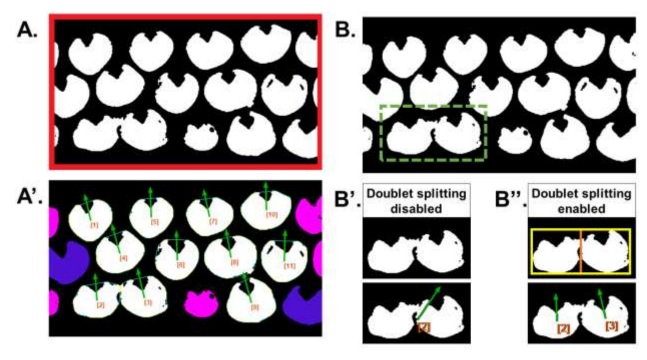


Figure 8. (A) Example of a border exclusion zone (red). (A') Any chunks that meet size exclusion criteria are excluded from subsequent analysis (pink). Any pixels that touch or fall within the exclusion zone will be excluded from subsequent analysis (blue). Note that because the size exclusion step occurs before the border exclusion step, some chunks that would otherwise violate the border exclusion step have already been disregarded from analysis by the size exclusion step, and are thus categorized under size excluded chunks instead of border excluded chunks. (B) Thresholded image featuring two cells that touch. (B') Subset of touching cells from A. If no doublet splitting options are applied, PCPA will treat this chunk as one mass and the angle measurements will incorrectly calculate based on the total chunk mass and dominant cave of the chunk. (B") Doublet splitting settings recognize this chunk as a doublet, draw a bounding box (yellow), then split the bounding box in half (orange). The two resulting halves will be treated as independent chunks with chunk and cave centroid calculations being based on the mass contained within each respective box. (B) Example of a border exclusion zone (red). (B') Any chunks that meet size exclusion criteria are excluded from subsequent analysis (pink). Any pixels that touch or fall within the exclusion zone will be excluded from subsequent analysis (blue). Note that because the size exclusion step occurs before the border exclusion step, some chunks that would otherwise violate the border exclusion step have already been disregarded from analysis by the size exclusion step, and are thus categorized under size excluded chunks instead of border excluded chunks.

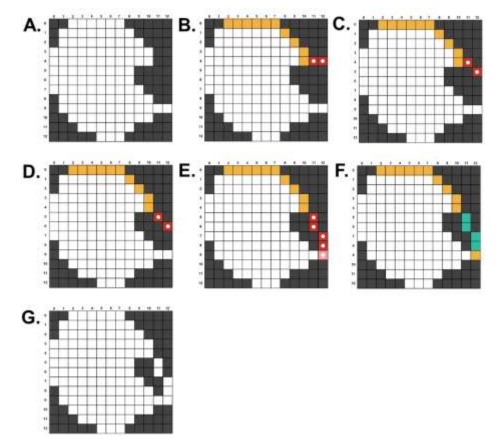


Figure 9. Demonstration of PCPA's plastic wrap feature. **(A)** Example chunk with no enclosed caves. PCPA will draw a bounding box encompassing all pixels of the chunk. **(B-D)** Example showing plastic wrapped pixels (yellow) projecting an exploratory line (red with white dot) from an orphaned pixel (coordinates 10, 3). When PCPA finds no chunk pixels that fall within the exploratory line, the next exploratory line is drawn from the orphan pixel to the next row of the last column. **(E)** PCPA will continue in the same manner until the exploratory line encounters a white pixel that falls within the exploratory line (pink with white dot). **(F-G)** All pixels of the exploratory line that connect the two pixels of the chunk (teal) will be added to the chunk mass for subsequent analyses.

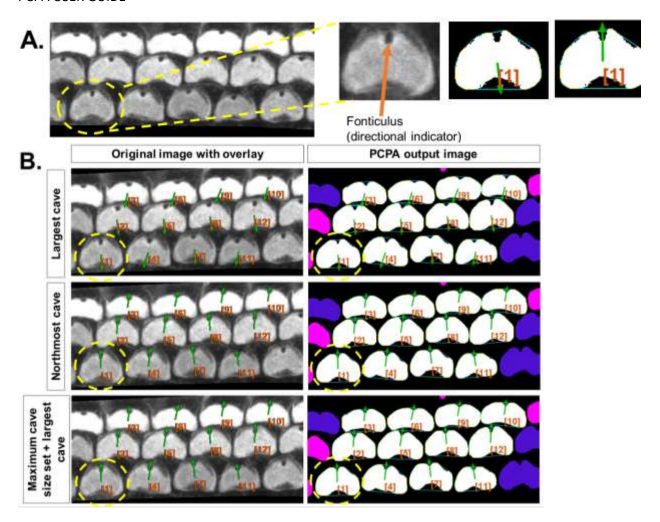


Figure 10. (A) When using PCPA on cells lacking a uniform oval shape, using Plastic Wrap (blue outline) may cause large caves to form that are the result of the shape distortion and not of inherent cell characteristics. **(B)** Cave selection criteria can be used to select for the user's cave of interest. If the largest cave is selected, PCPA selects the large inclusion created by Plastic Wrap. This can be addressed by changing the cave selection to northmost cave, which will disqualify the largest inclusion at the bottom of the cell. Alternatively, users can combine the largest cave setting with cave maximum and/or minimum size restrictions—here, a maximum size has been applied to exclude the large inclusion on the bottom caused by Plastic Wrap.

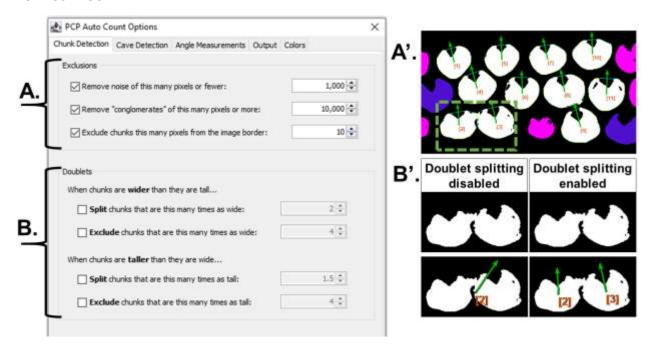


Figure 11. User option menu for PCPA chunk detection. **(A)** The exclusion criteria box allows users to disregard chunks smaller and/or larger than a specified pixel size, and to exclude cells a set pixel distance from the image border. **(A')** Example output where size excluded chunks are shown as pink pseudocoloring and border excluded chunks are shown as blue pseudocoloring. **(B)** To address instances were two chunks touch after applying thresholding, users can set options to split chunks meeting a set ratio of wider-than-tall or taller-than-wide, or to exclude these chunks from further analyses. **(B')** Subset of A' (green box). If no doublet splitting options are applied, PCPA will treat this chunk as one mass and the angle measurements will incorrectly calculate based on the total chunk mass and dominant cave of the chunk. When doublet splitting options are applied, PCPA will split the doublet in half. The two resulting halves will be treated as independent chunks with chunk and cave centroid calculations being based on the mass contained within each respective box.

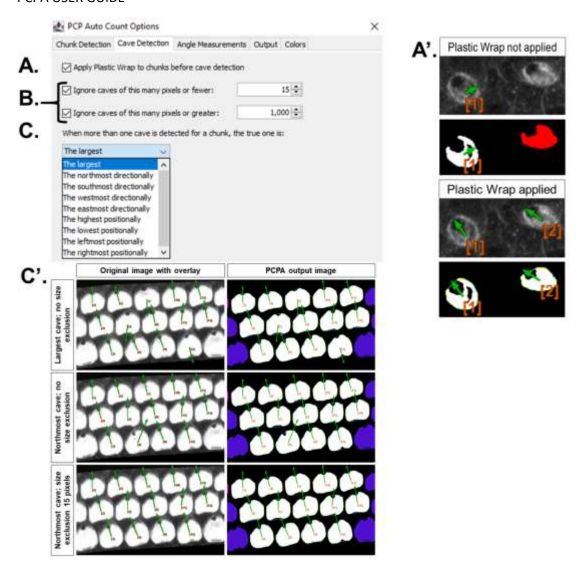


Figure 12. User menu for PCPA cave detection. **(A)** Users are recommended to apply the Plastic Wrap function to their analyses. Plastic Wrap allows PCPA to group pixels that are proximal, but not abutting, into the same chunk. **(A')** Example images show that when plastic wrap is not applied, chunk 1 selects an incorrect cave, and chunk 2 is excluded from analysis for having no cave. When plastic wrap is applied, PCPA artificially closes off the open cave and selects the correct cave as the cave of interest. **(B)** Users can set minimum and maximum size requirements for cave of interest selection. **(C)** Users can set a directional requirement for cave of interest selection. Selecting a directional characteristic (e.g. northmost) will instruct PCPA to designate the cell's cave as the largest northmost inclusion. **(C')** Users can combine directional characteristics with a lower size limit for caves, in order to filter out incidental inclusions created by Plastic Wrap, pitting caused by the thresholding process, or large inclusions caused by irregular cell shapes.

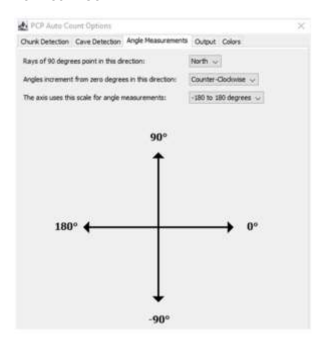


Figure 13. User options for angle measurement calculations.

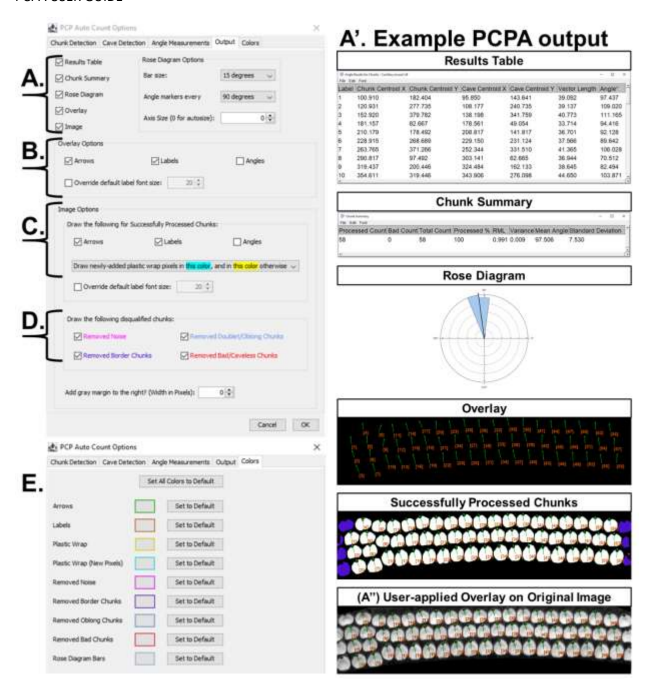


Figure 14. PCPA data output tab. **(A)** Options for data metrics produced by PCPA. **(A')** Example of data metrics in A. **(A'')** Example of overlay superimposed on original preprocessed image. **(B)** The overlay (and selected overlay information) can be superimposed over the original image for data visualization. **(C)** Optional labels can be viewed on the post-PCPA image. **(D)** Color coding of the post-PCPA image can indicate to the investigator what chunks may have been excluded and why. **(E)** The Colors tab allows users to customize color coding options from D.



Figure 15. Cell counting when PCP measurements are not needed. **(A)** Example of recommended output settings when a user only desires to view total cell count. **(B)** Example of the PCPA cell labels overlaid on original image of cochlear IHC. **(B')** Annotated PCPA output showing counted cells with labels, noncounted cells (i.e. lacking a cave; red Pseudocolor), and border excluded cells (blue). **(C)** The Chunk Summary table shows how many cells were analyzed ("Processed Count"), how many cells were rejected because they lacked a cave ("Bad Count"), and the "Total Count" value of all recognized chunks.

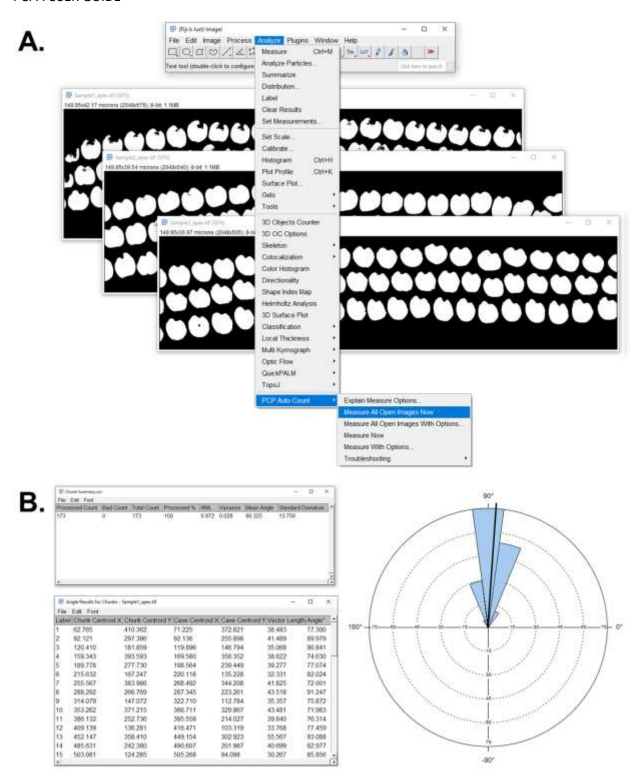


Figure 16. Batch process multiple images as one data set. **(A)** To batch process multiple images, open each image of the batch in ImageJ then select "Measure All Open Images Now" or "Measure All Open Images With Options" under the PCPA menu. **(B)** All output metrics will aggregate the data collected from each open image into a combined data set. Summary statistics and a rose diagram for the combined data set can be viewed

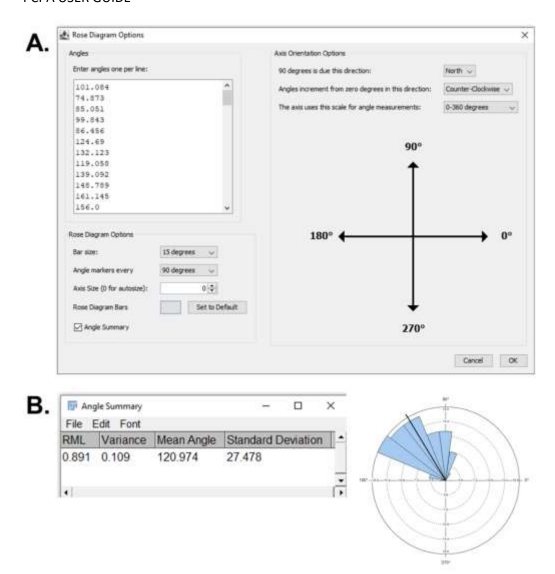


Figure 17. Stand-alone rose diagram function. **(A)** Users will input their data into the data box (one angle per line) then set axis orientation options. Users can customize the rose diagram output and optionally view summary statistics by selecting the Angle Summary box. **(B)** Example output from the standalone rose diagram function.