

dpxi5pke

PlanktoScope protocol for plankton imaging V.(dpxi5pke)



Adélaïde PERRUCHON¹. Lombard Fabien^{1,2}

¹Sorbonne Université, Centre National de la Recherche Scientifique, Laboratoire d'Océanographie de Villefranche (LOV), Villefranche-sur-Mer, France;

²Institut Universitaire de France, 75231 Paris, France



Lombard Fabien

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External link: https://www.planktoscope.org/

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Disclaimer

This protocol applies to the version 2.5 of the PlanktoScope and the 2.3 version of software. It is optimized to image 20µm-200µm organisms using the 25mm lens (as tube lens) and 12mm one as objective one and may be inaccurate with other configurations or light. Please note that the segmenter is currently also optimized for this and may need to be recoded (or adjusted) for other configurations, notably the size threshold but also the intensity threshold.

Abstract

This protocol is for using PlanktoScope and collecting usable results for quantitative imaging of plankton.

This project has received funding from the European Union's Horizon 2020 research and innovation programme "Atlantic Ecosystems Assessment, Forecasting and Sustainability" (AtlantECO GA#862923)

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See also https://www.planktoscope.org/

Image Attribution

Fabien Lombard, Thibaut Pollina, Karine Leblanc, Will Major, Pierre Kostyrka

Guidelines

Planktoscope is an optical instrument. As it optical elements (camera, lenses, flowcell) are highly sensible to dust and dirt. we recommend that you never touch any of those component with fingers and store the planktoscope in a dust free and humidity free area (or in a box when not used)

complete manual of assembly and software could be found at https://planktonscope.readthedocs.io/en/latest/



Materials

- Plankton net or other kind of microplankton collector
- 200µm sieve
- Squizing bottle
- Micrometer slide (or millimetric ruller)
- Optical paper
- Dry gas dispenser
- PlanktoScope box
- Fine forceps
- A computer

Softwares:

- ImageJ (last version, needs to compute RGB images)
- FileZilla
- BalenaEtcher

Safety warnings



- Planktoscope is an electronic device, powered with electricity. It is therefore sensible to water.
 - Place it in an environment where water can not enter in contact with the instrument and secure its electrical part.
 - Be careful when manipulating samples, take care of having the exhaust tube in a "trash" contained to avoid spillage
 - glass parts are present (flowcell) and should be manipulated with caution (can break and injure you), but also should be kept clean (avoid touching it with fingers)
 - For an easiest navigation, you can see the table of contents by clicking on "Show the table of contents" on the top left.
 - Figures are numbered by section:
 - If a figure is cited in its corresponding section, only the number will be cited (ex: fig.1).
 - If a figure is cited in a different section, the section and the number will be cited (ex: S3-fig.1 for figure 1 of section 3).

Before start

- -Test the protocol before acquisition of your first sample
- -Calibrate your instruments to ensure coherent measures
- -Create an Ecotaxa account and request the right to create project way before
- -Collect a plankton sample using a net



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Introduction

2 The **PlanktoScope** is a frugal, microfluidic microscope designed with an open-hardware and open-software approach. It was conceived with the idea of providing the thousands of scientists and sailors exploring the oceans with a high-quality instrument suitable for deepening our knowledge of the sea around us. In this manual you will learn how to operate the PlanktoScope and take images of plankton.

Quick usage version

3 This part is a quick version of the protocol, where you will find the essential steps to follow. If you are using your PlanktoScope for the first time or need to calibrate it, please read the full



protocol. Do not wait too long between the sampling and the processing of your sample through the PlanktoScope to avoid sedimentation and aggregation.

3.1 Preparation

- 1. Assemble and plug in the PlanktoScope 5 go to step #4
- 2. Connect to PlanktoScope's Wi-Fi go to step #5.1
- 3. Go to http://planktoscope.local > Node-RED Dashboard)
- 4. Fill ALL the sample metadata (in "Sample") [Critical] = go to step #10.1
- 5. Turn on the light (in "Optic Configuration")
- 6. Check the WB parameters (you should have already done the calibration during the initial connection) = go to step #6
- 7. Check Flow Cell alignment and check the focus **5** go to step #10.2

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5 go to step #10.3
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- 8. Put 20 mL sample go to step #10.4
- 9. Turn on the bubbler
- 10. Check the focus (in "Optic Configuration") go to step #10.3

3.2 Acquisition

- 1. Fill acquisition parameters (in "Fluidic Acquisition") = go to step #10.7
- 2. Pump to drain sedimented organisms before acquisition (in "Optic Configuration")
- 3. Start the acquisition (in "Fluidic Acquisition")

3.3 Segmentation and data export

- 1. Fill in segmentation parameters (in "Segmentation") = go to step #11
- 2. Start segmentation
- 3. Back-up data (in "Gallery" or by using FileZilla) go to step #14
- 4. Import on EcoTaxa 5 go to step #14

3.4 Cleaning ≡5 go to step #13

- 1. Drain the syringe (disconnect your system)
- 2. Drain the content
- 3. Replace with distilled water and drain several times (pushing with the syringe may helps)
- 4. Replace the system and drain first with distilled water and then air
- 5. Empty waste tube

If not used immediately

- 1. Put 20 mL diluted bleach
- 2. Leave 15 minutes
- 3. Drain the content (high pump speed)
- 4. Put 10 mL distilled water
- 5. Drain the content (high pump speed)



3.5 Shut down

- 1. Turn off virtually (in "Home") and wait one minute
- 2. Unplug the PlanktoScope

Assemble the PlanktoScope

4 The PlanktoScope kit

In this part you will learn:

- 1. What is inside the PlanktoScope kit
- 2. How to connect the fluidic system with the Flow Cell
- 3. How to assemble the bubbler
- 4. How to put all the components together

Open the PlanktoScope box and check that everything is in it. The PlanktoScope kit includes a PalnktoScope, a bubbler, power cable, waste tube, sample tube, syringe, tube holder, Flow Cell holder and Flow Cell (Fig.1; Fig.2; Fig.3).



Figure 1: The PlanktoScope Kit. (A) The main box with the pump A1, the USB cover A2, the pi camera A3 and the screws A4; (B) Tube holder; (C) Bubbler; (D) Power cable; (E) Flow Cell holder; (F) Supplementary materials box; (G) IFIXIT kit; (H) Flow Cell



Figure 2: Material inside the box F: (F1) 200 μ m filter; (F2) plastic dropper; (F3) cleaning kit; (F4) fluidic system; (F5) SD card adaptor; (F6) syringe; (F7) waste tube; (F8) sample tube; (F9) cleaning blower



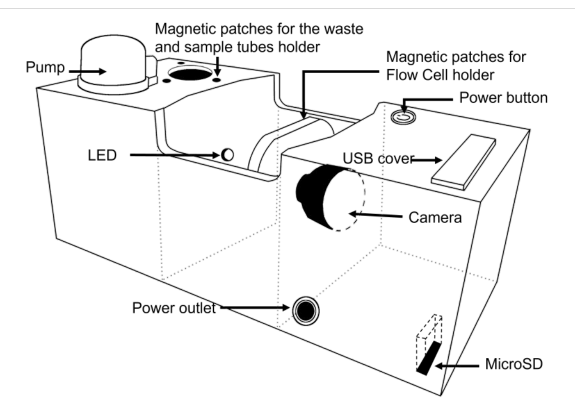


Figure 3: Main components of the PlanktoScope (A in Fig.1)



Safety information

If it is present, do not forget to remove the transparent cover from the camera (Fig.4).



Figure 4: The transparent cover of the camera

4.1 The Flow Cell

Safety information

The Flow Cell can break easily. It is a part to handle with care. There is multiple Flow Cell in the box to replace broken ones.

- Do not touch it with your fingers or leave it on a surface.
- If a Flow Cell is dirty, you can clean it softly with the cleaning kit.
- The Flow Cell receptacle should be placed with caution on the lens to not break the Flow Cell. It will clips with the magnets.
- Do not stretch the tubes attached to the Flow Cell, as this may damage the Flow Cell.
- 1. Take a Flow Cell and plug the short part to the syringe with the help of a male adaptor.
- 2. Install it like in the image on the left of the Fig.5 and press the tube where the arrows point.
- 3. Screw softly the two parts together.
- 4. Place the receptacle on the magnetic Flow Cell holder on the PlanktoScope. You can put some tape on the magnetic parts to reduce the impact between the Flow Cell holder and the receptacle.





Figure 5: How to install the Flow Cell. You will need the syringe (F6), the Flow Cell holder (E) and one Flow Cell (H).

4.2 The fluidic system

Now that you have fixed the Flow Cell, assemble all the fluidic system (Fig.6). Do not forget to install the flux stopper on the matching tube (Fig.6). It will allow you to stop the flow when it is necessary.



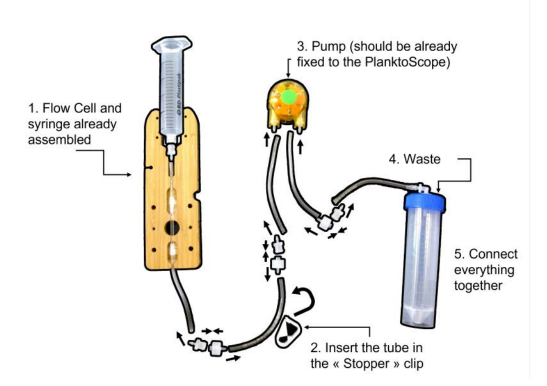


Figure 6: How to assemble the fluidic system

4.3 The bubbler

A bubbler is needed to prevent sedimentation, as the PlanktoScope takes images of fluids at low speed.

Safety information

Not agitating your sample will let plankton to sediment and could even block the fluidic system. More importantly, the organisms concentration will be inhomogeneous, and because you will first get the sinking plankton, will lead your measurements to overestimate true concentrations.

- 1. Assemble the bubbler like in the image below (Fig.7). It is recommended to use a cut glass pipette or something similar instead of the needle provided because the needle can be clogged and does not allow a good control on the air flux.
- 2. Plug the bubbler into one of the USB ports on the PlanktoScope.
- 3. Place the tubing into the syringe so that it reaches the bottom. **Do not put the tip of the bubbler in the middle of the syringe or it will inject bubbles into the Flow Cell.**
- 4. You can secure the tubing to the syringe using a rubber band, string or similar.
- 5. Switch on the bubbler: the flow of air in the water needs to be adjusted to approximately 1 bubble/sec.



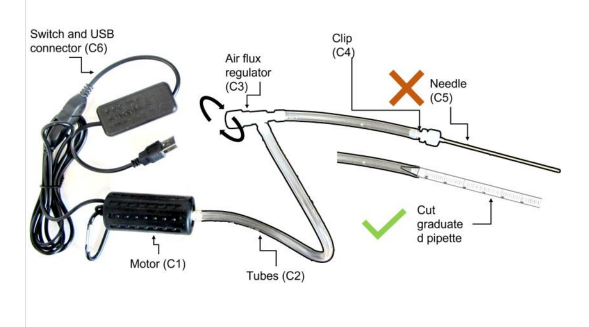


Figure 7: The bubbler

4.4 Assemble the PlanktoScope

The fluidic system, the Flow Cell and the bubbler should be all assembled now. Now you just need to (Fig.8):

- Place the tube holder (B in Fig.1) on the magnetic patches in front of the pump
- Place the waste tube in the outer hole of the tube holder and the sample tube in the inner hole
- Place the Flow Cell receptacle on the magnetic Flow Cell holder in front of the camera
- Connect all the tubes of the fluidic system together with tube adaptors
- Plug the power cable



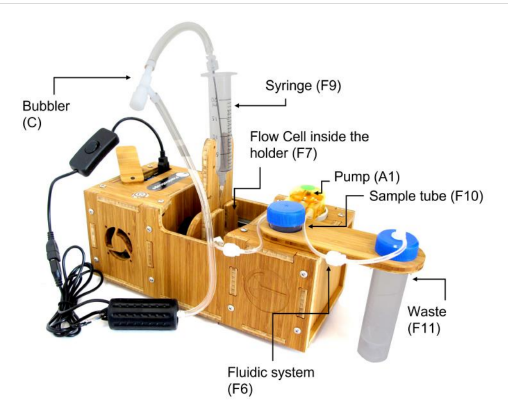


Figure 8: Assembled PlanktoScope

User Interface and initial connection

5 Initial connection

You can access to the user interface of the PlanktoScope by connecting to the Wi-Fi generated by the PlanktoScope. Your computer will be only a projection of the software of the PlanktoScope. It means that even if you disconnect from the Wi-Fi of the PlanktoScope, it will still be running.

- 1. Power your PlanktoScope by connecting the power cable to an appropriate electrical outlet
- 2. Within 1 minute of turning on your PlanktoScope, you should see the LED flash once. This means that the PlanktoScope is ready to be connected by Wi-Fi.
- 3. You should see a new option for Wi-fi appearing on your computer. Connect to it with the password: "copepode"

For more information and alternative methods of connection, see the designer's Connectivity Tutorial here: PlanktoScope - Connectivity Tutorial ...

Note

Note that when you are connected to the Wi-Fi of the PlanktoScope, you cannot access to the internet. If you want to use internet at the same time, you should use your smartphone's hotspot or an ethernet cable.



5.1 The User Interface (Node-RED dashboard, UI)

Open the PlanktoScope's User Interface (UI) on your web browser (Chrome, Firefox, Edge etc.) using the following webpage link and go to the Node-RED dashboard:

http://192.168.4.1:1880/ui/ (v.2022)

<u>http://home.pkscope/</u> > Node-RED dashboard (v.2023)

http://planktoscope.local > Node-RED Dashboard (v.2024)

There are several tabs on the User Interface (UI, Fig.1) that can be used to adjust setting, run samples and take images. To navigate around the UI, all tabs are available from the Home tab, including the Shutdown button. We can also use the **Hamburger Menu**, situated in the top-left corner of the UI, to navigate between these tabs.

Note

v.2023:

In addition to the Node-RED Dashboard, you can also access to the data file manager, the protocol for the PlanktoScope, or the log in case of errors.

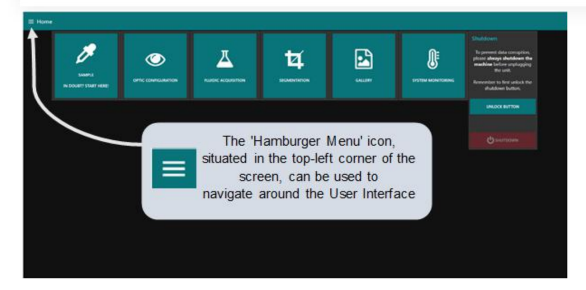


Figure 1: The "Home" tab of PlanktoScope's User Interface

You will find:

- the "Sample" tab to fill all the metadata of your sample
- the "Optic Configuration" tab to control the various features of PlanktoScope (focus, LED and pump)
- the "Fluidic acquisition" tab to launch an acquisition and edit its parameters
- the "Segmentation" tab to start the segmentation of the images taken during acquisition
- the "Gallery" tab with files including the exports for EcoTaxa, the original images and the extracted images
- the "System Monitoring" tab to check the correct operation of the device (not use in standard use)
- the "WIFI" tab for the characteristics of the wifi generated by the PlanktoScope, to which you will connect in order to control the device. You will not have to modify anything on this page.



 the "Hardware Settings" tab (not needed for processing samples and strongly advised to not change anything)

Note

If all the tabs are not visible, you can adjust the zoom on your browser (usually Ctrl + scroll UP or DOWN on Windows or command + scroll UP or DOWN on Mac).

5.2 The "Optic Configuration" tab

Once the UI has loaded on your browser, navigate to the **"Optic Configuration"** tab and we will make sure the PlanktoScope is operating correctly (Fig.2). This tab can be used to adjust the camera settings. If only "Preview" is visible on your screen, the other options should be available below by scrolling down.

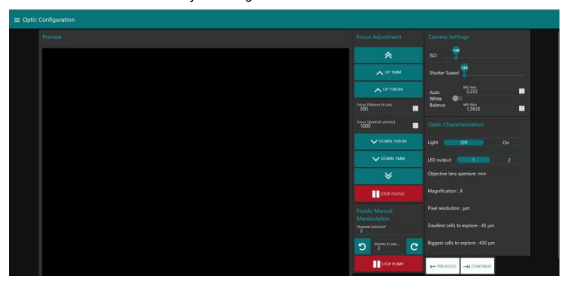


Figure 2: The "Optic Configuration" tab

• Under "Optic Characterisation", switch on the "Light" (Fig.3). You should see the "Preview" image turning from dark to light. The Preview image could be any color so do not worry if yours show blue, red, green, etc.; it will be adjusted later during the White Balance (WB) calibration. You will need to switch on the light every time you use your PlanktoScope.





Figure 3: The red box highlights the location for turning on the LED

Under "Focus Adjustment" (Fig.4), click "UP 1MM" and "DOWN 1MM" to ensure focus buttons turn the focus motor (same for the 100 UM version of the button). You should see the frame moving further from (UP) or closer to (DOWN) the camera. You can also use the ">>" button to move your focus according to a personalized value ("Focus distance").

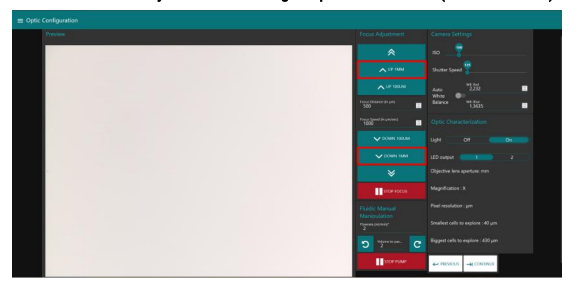


Figure 4: Red boxes highlight "UP 1MM" and "DOWN 1MM"

• Under "Fluidic Manual Manipulation" (Fig.5), click clockwise arrow to check that the peristaltic pump is working. You should see the pump rotating in an clockwise direction. You can choose to pass a volume, but keep in mind that this parameter will not impact the acquisition. On the other hand, the flowrate parameter will also be effective during an acquisition.





Figure 5: The red square highlights the location of the clockwise arrow that will rotate your peristaltic pump

• Under "Camera Settings", you can see a button to change the ISO value (Fig.6). Set the ISO to 150. You should see your Preview image change color when you adjust this setting. The ISO is the light sensitivity of the camera. A low ISO will be less sensitive to light and then darker. A low ISO tends to provide images of better quality.

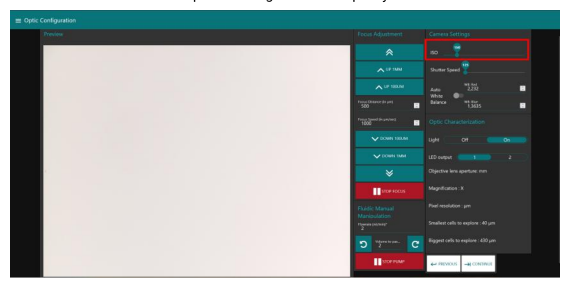


Figure 6: The red box highlights the location of the ISO setting

Safety information

Keep the shutter speed at the minimal value. The shutter speed determine how long the light will enter the camera to take one image. A low shutter speed provides better quality, but it also requires the objects to not move too fast.

White Balance calibration



6 White Balance (WB) calibration

Stay in the "Optic Configuration" tab. For this step, you will need the software ImageJ: https://imagej.net/. To check the WB, do not put the Flow Cell or anything between the camera and the LED.

Note

To calibrate the PlanktoScope or if you just need to do a test, you can use the **"Test"** mode in the "Sample" tab. It allows you to not fill the date or the location of the sample.

Safety information

PlanktoScope are normally cross-calibrated for white balance initially, this information could be recovered from the provider. **We strongly encourage you to note the initial values** before trying to change those and this procedure should not be done without reasons (incorrect image with initial calibration; reboot or update of the software.

Note your initial calibration here:

WB Red:

WB Blue:

Try pressing the Auto White Balance (AWB) button to its "on" and "off" positions on the "Optic Configuration" tab; you will likely see the Preview image changing colour (Fig.1). In this example (Fig.1), the correct setting was WB Red = 4 and WB Blue = 1.21. The Auto WB button should be set to "off" once you have completed this step.

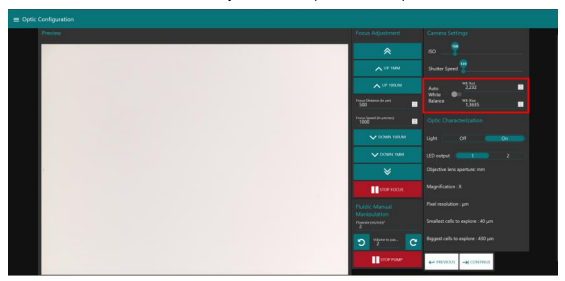


Figure 1: The red box highlights how to adjust the white balance of the Preview image

The Auto White Balance feature is currently not optimised. In addition, not using Auto White Balance enhances the performance of the PlanktoScope over time (the camera will try to adjust it in between every image). It is then recommended to manually adjust the white balance of your PlanktoScope. Set the AWB button to "off". You will need to adjust WB Red and WB Blue until it looks white/grey. The WB should never be perfect white but grey, as it imply over-exposition during the segmentation step.



To manually set to the White Balance:

- 1. Turn off the "Auto White Balance" button. Make sure the ISO value is 150. Never set it to 0 or less, or it will create a bug in the software.
- 2. Take one image in the "Fluidic Acquisition" tab. You should not use the preview image as it will not show the real colours taken by the camera.
- 3. Download the raw image in the "Gallery" tab or by using FileZilla.
- 4. Open the software ImageJ.
- 5. Click on "File > open" and open your raw image.
- 6. Click on "Plugins > analyze > RGB". If you do not have the plugin or if it is not working go to "Analyze > Histogram > Click on RGB" until you see the mean red, green and blue values.
- 7. Once the values are extracted, calculate the ratio G/R and G/B like in the example below.
- 8. Multiply it by the old ones to get the new ones.
- 9. Change the configuration and make sure the ISO value is still on 150.

Example:

If the default configuration is:

$$egin{aligned} WB^{old}_{red} &= 1 \ WB^{old}_{blue} &= 2 \end{aligned}$$

With this actual configuration, we measure the RGB (Red, Green, Blue) values:

$$R = 245, G = 240, B = 230$$

 $G/R = 0.98$
 $G/B = 1.04$

The corrected WB Red or Blue is the ratio between the green value (G) on the corresponding colour value (Red = R; Blue = B) multiplied by the old configuration. The new configuration should be:

$$\begin{array}{l} WB_{red}^{new} = WB_{red}^{old} * G/R = 1 * 0.98 = 0.98 \\ WB_{blue}^{new} = WB_{blue}^{old} * G/B = 2 * 1.04 = 2.08 \end{array}$$

Note

The green colour is a fixed value. It is then a way to check the exposure of your image. Because the camera changes the exposure between the preview and the acquisition, it is recommended to **check the green value with an acquisition** and not with the preview. It should never be at 255.

If the light exposure is too important (>250), put some tape on the LED to decrease it, and do the WB calibration again.

Also, if the exposure is too important, it will be impossible to correct the WB.

Pump calibration

7 **Pump calibration**



Safety information

Peristaltic pump tubes flexibility varies with age, care and type of liquid used (e.g. lugol may age it quicker), calibrating the pump regularly could be needed but is not highly important to get good quantitative count since it is the number of images (therefore the volume imaged) which is important (not the pumped volume).

7.1 Calculate pump step

- 1. Prepare a large volume of distilled water and put a total volume A of 20mL in the syringe.
- 2. On the "Optic Configuration" tab: choose to pass a volume Y = 10 mL and record the exact volume X it finally ends to pass (eg. by looking on the graduation B of the syringe).
 - $X = final \ volume \ passed for a 10 \ mL \ instruction = A B$
- 3. If the pump calibration is correct, X should be equal to Y = 10 mL
- 4. If it is not correct, update the pump step (see next step)

For example, in Fig.1: X = A - B = 20 - 15 = 5 mL. There is then an error in the calibration and we need to update the pump step according to this error.

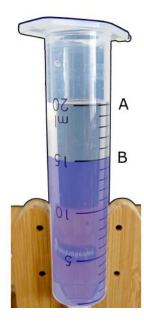


Figure 1: Volume pumped

7.2 Update pump step

- 1. Go to the "Hardware Settings" tab (Fig.2)
- 2. Note the initial calibration "Pump: step per mL" here:
- 3. Calculate the calibrated "Pump: step per mL" such as New Step = (Y/ X)*old step
- 4. Replace the "Pump: step per mL" parameter in "Hardware Settings" with the value calculated in the previous step



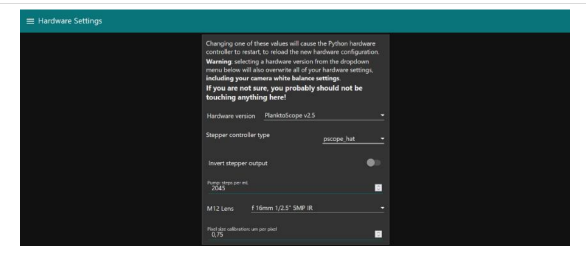


Figure 2: Example of Hardware Settings

Size calibration

8 Size calibration

Safety information

Size calibration is an important process to get good data and should be absolutely done and noted. It will allows to know the real size of the plankton.

8.1 Set up the scale

- 1. Remove the Flow Cell holder and tilt the PlanktoScope on the side with the camera on the bottom (Fig.1)
- 2. Place a micrometric ruler (or a millimetric one) in front of the camera, at sample level, such that the ruler is either vertical or horizontal but not in diagonal (Fig.1)
- 3. Make the focus on the scale of the micrometric ruler (Fig.2)





Figure 1: PlanktoScope on the side with the micrometric ruler

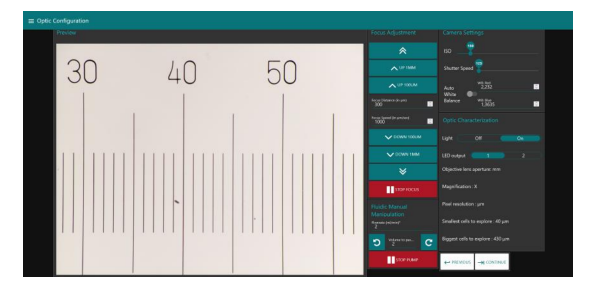


Figure 2: Focusing on the scale of the micrometric ruler

8.2 Take images of the scale

- 1. Fill metadata in the "Test" mode in the sample tab (Fig.3)
- 2. Take 1 or 2 images in "Fluidic Acquisition" (pumped volume does not matter, Fig.4). You should only use acquired images to measure the real size of the pixels, and not the preview image as it will give you wrong results.



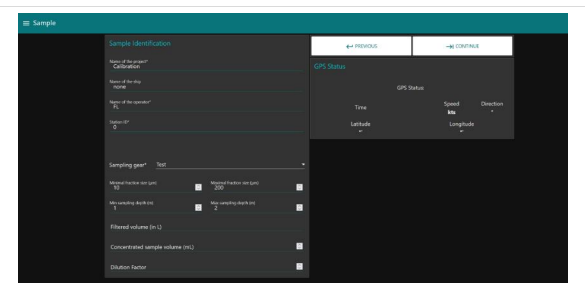


Figure 3: Example of metadata entered in the "Sample" tab

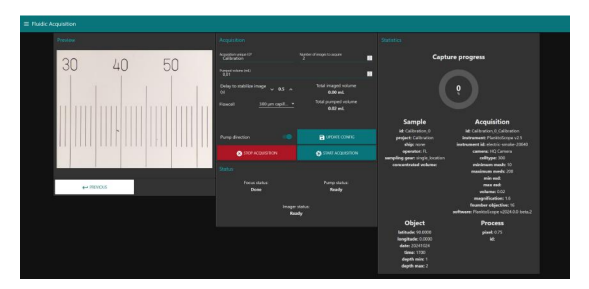


Figure 4: Here two images are acquired

8.3 Measure the number of pixels

- 1. Download the images on a computer using FileZilla or the "Gallery" tab according to your software version
- 2. Open ImageJ
- 3. Click on "File > Open" to open your image
- 4. Click on the line button (see Fig.5) and trace a line
- 5. Click on "Analyze > Measure"
- 6. Check the length value (in pixel)



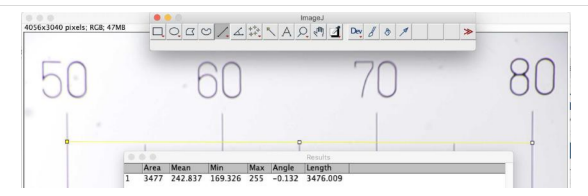


Figure 5: Line drawn using ImageJ on the picture taken with the PlanktoScope

In this example (Fig.5), the line is 3476 pixel length for 3mm, which means that here one pixel is 0.86, which differ from the expected value of 0.75 for this camera.

8.4 Calculate the micron/pixel ratio

Calculate how much microns are represented by each pixel. The expected value for the default camera is expected to be around 0.75. You will need to do **Length(micron)/Length(pixels)** (see the example in Fig.5).

8.5 Update the micron/pixel ratio

Enter the calibrated pixel size value in the "Hardware Settings" (Fig.6).

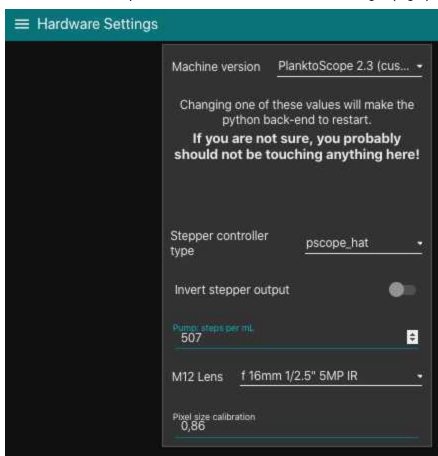


Figure 6: Example of pixel size calibration settings



Get your sample

- 9 This protocol considers that a plankton net is used to collect the sample.
 - In this section, you will:
 - 1. Collect your sample
 - 2. Fill a log sheet
 - 3. Filter your sample with a 200 µm sieve

9.1 Collect your sample and the metadata

Safety information

Using logsheets:

- -Record Latitude Longitude (taking photos of the GPS when launching and recovering the net could serve, if UTC time is on the GPS this could also be interesting)
- -if vertical net, record min and max depth
- -if horizontal records initial/final positions, speed and length (min) of deployment
- -if you have flowmeter, record the initial/final digits of the flowmeter and calculate the filtered volume

in all cases the diameter of the net opening will be needed

Those are critical information to get to quantitative sampling.

You can find here a logsheet example (based on the datas needed in the SAMPLE Tab).

Operator:

Station ID:

Sampling gear:

Process time:

Net throw Lat:

Net throw Lon:

Net throw date/time:

If it a horizontal sampling:

Net retrieval Lat:

Net retrieval Lon:

Net retrieval date/time:

Minimal fraction (µm):

Maximal fraction (µm):

Min depth (m):

Max depth (m):

/!\ Critical information

Filtered Volume (L):

Concentrated sample volume (mL):

Dilution factor (<1 if there has been a dilution):

Net opening dimension (mm):

Speed Through Water (kts):

9.2 Filter your sample



Safety information

Larger organisms may clog the Flow Cell. It is then necessary to filter the volume through a 200 µm sieve. Rinse the sieve using seawater and a squeezing wash bottle (helps to pass small objects).



Figure 1: How to filter the sample with the 200 µm sieve.

Pass the sample on PlanktoScope

10 Assemble and start the PlanktoScope before (see 5 go to step #4)

In this section you will:

- 1. Fill the metadata of the logsheet in the "Sample" tab
- 2. Check the focus and the Flow Cell
- 3. Switch on the light and the bubbler
- 4. Put your sample and check the concentration of it
- 5. Dilute your sample if necessary
- 6. Pass sedimented organisms
- 7. Launch an acquisition

10.1 Fill the metadata

Go to the "Sample" tab and fill the metadata. This step is critical because those data are the ones that will make your sample usable or not.



Safety information

If the PlanktoScope has already been used before, the old metadata are keept. It can be useful, but do not forget to change them if it is necessary.

• Fill the sample identification (project, name, ship used, your name and the station number; Fig.1)

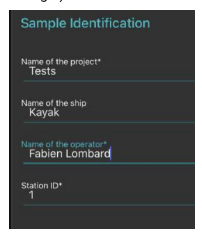


Figure 1: Sample ID parameters.

- Note how you sampled the plankton recording mesh size with "minimal fraction size" (will be used afterwards in the segmentation process, objects smaller than this will not be segmented; Fig.2)
- "Maximal fraction size" is the size of the mesh used (to remove oversized organisms; Fig.2)
- Filtered volume is important if you recorded it but could be calculated from other parameters. Make sure to either have filled it or to have filled either min and max depth if using a vertical net and initial/final coordinates, speed and length (min) of deployment if using a horizontal towed net.

Safety information

Known bug: If filtered volume is provided but also initial/final latitude and longitude, calculation from this latter may replace the measured filtered volume.

In all cases the diameter of the net opening will be needed to calculate the filtered volume (only if you used a plankton net).

- Note the mesh size used for collection in "minimal fraction size" (it will be used afterwards in the segmentation process, object smaller than this will not be segmented);
- The "Maximal fraction size" is the mesh size used to filter the sample during preparation (It must have been done at 200 µm so as not to block the fluidic system);
- The "Filtered volume" is the volume passed through the net during sampling. It is better if you recorded it but could be calculated from other parameters. So, make sure to either have filled it or to have filled either min and max depth if using a vertical net; initial/final positions, speed and length (min) of deployment if using an horizontal towed net; and in all cases the diameter of the net opening (to be able to calculate the volume afterwards).



- "Concentrated sample volume" is the volume recovered by the net (Fig.2).
- If a **dilution** has been done, note the "dilution factor" (if not, write "1"). <1 if it is diluted, >1 if it has been concentrated.

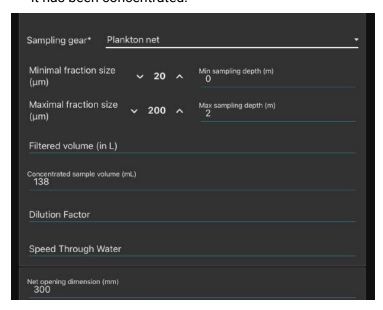


Figure 2: Example of sampling parameters

• Fill the net initial and final position (if towed horizontally) remember to validate both of them (Fig.3). Note that values disappear once validated but are saved.

Note

1. When you click on "VALIDATE", the coordinates will disappear in any case. You can check that they are correctly registered in the "Fluidic Acquisition" tab. If not, you will have a warning.



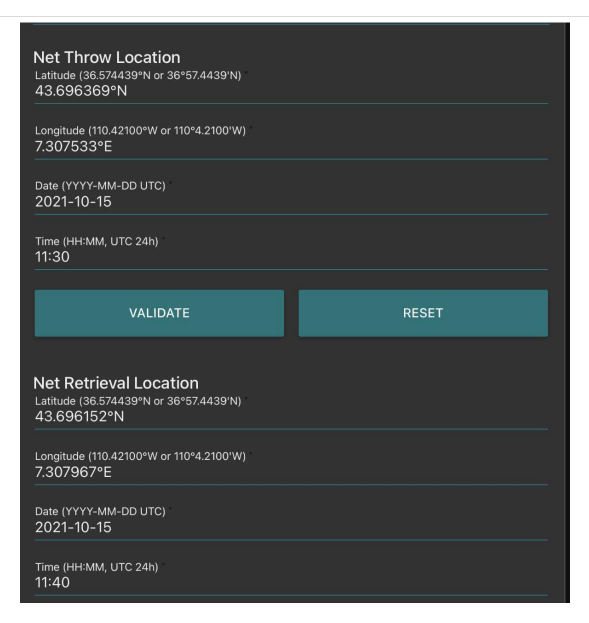


Figure 3: Example of date and time of the sample

10.2 **Check Flow Cell alignment**

- 1. Turn on the light
- 2. Check for lens alignment with the Flow Cell. Move slowly the Flow Cell receptacle until there is no black background like in Fig.4.



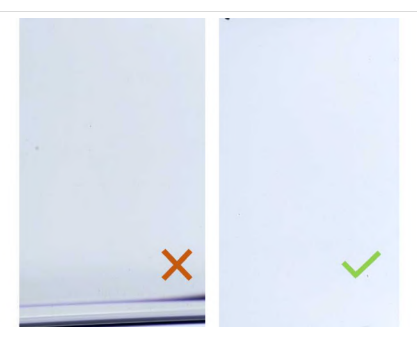


Figure 4: The figure on the left shows a misaligned Flow Cell. The background should be homogeneous like in the figure on the right.

10.3 Do the focus

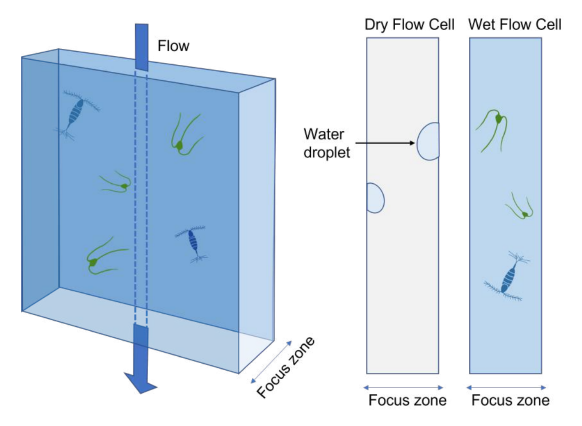


Figure 5: Focus zone of the Flow Cell

Go to the "Optic Configuration" tab:

1. Switch on the light if you have not already done so



2. Check focus on the two sides of the Flow Cell, and try to have a focus between the two sides (Fig.5). In the Fig.6, you can see that on a dry Flow Cell that has been exposed to water, there are still water droplets on both sides. Use them as an indicator to do the focus.

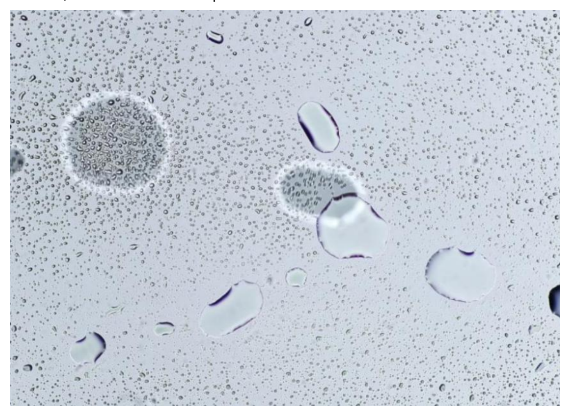


Figure 6: Water droplets

On Fig.6, there is a good focus on the front face, and you can see two other water droplets badly focused on the back side of the Flow Cell.

Note

tip#1: start using the "1 MM" buttons, then the 100 μ m buttons and finish by typing 25 or 50 μ m adjustments in the middle box and pressing external arrows of focus; **tip#2**: you can connect your phone or a tablet to the PlanktoScope to have controls on the focus while checking a zoomed portion on the streamed image on another device.

This step also allows you to check that your Flow Cell is properly levelled. If bubbles in the same layer do not all have the same focus, your Flow Cell is not straight. You are going to have to reposition it by taking it out of its holder, so it's important to check this before putting your sample in the PlanktoScope. **Be very careful not to break the Flow Cell.**

10.4 Put your sample in the PlanktoScope

- 1. Close the flux stopper
- 2. Fill the sample in the syringe. For this you can just remove the full sample holder and fill it on top of a sink (to not risk spills on-top of the PlanktoScope; Fig.7)
- 3. Reconnect the syringe to the pump
- 4. Open the flux stopper
- 5. Place the bubbler and adjust the flow to 1 bubble/second



Safety information

Never forget to open the flux stopper. If the flux stopper is closed for a long time, it can deteriorate the fluidic system.

Safety information

Not agitating your sample will let plankton sediment and could even block the fluidic system. More importantly, the organisms concentration will be inhomogeneous, and because you will first get the sinking plankton, will lead your measurements to overestimate true concentrations. You should agitate your sample using bubbling and use a sufficient pumping rate to avoid sinking/clogging of sample.

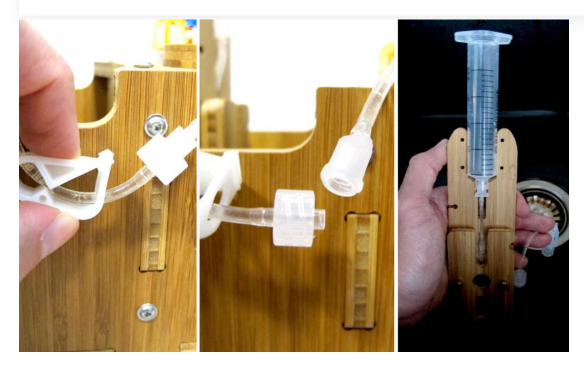


Figure 7: How to remove the part of the fluidic system containing the Flow Cell to fill the syringe with water

10.5 Dilute your sample if necessary

Adjust the concentration of the sample: ideally not more than 20-30 objects should be present per frame. If the sample is over-concentrated, dilute it by a factor 2 (add in a jar 1/2 of the sample -after agitating it- and 1/2 of seawater). Note the dilution factor in the metadata!

Safety information

Having too many objects per frame will:

- increase the probability to aggregate objects (making them impossible to count or identify)
- increase the probability of clogging the fluidic system
- create artefacts during the segmentation step



10.6 Pump sedimented particles

In the "Optic Configuration" tab, pump with high flow rate a good amount of water to remove plankton that have sunk in the fluidic system. You do not need to pump a large amount of your sample, 1 mL is sufficient (Fig.8).



Figure 8: How to pump a specific volume without image acquisition.

Safety information

Be careful, even if the volume chosen here will not impact the future acquisition (it is completely independent), the flowrate will be the one used during the acquisition. It is recommended to have a low flowrate for an acquisition (around 2 mL/min). **Do not change the flowrate during an acquisition and stay around 2 mL/min after you pumped sedimented organisms.**

If your fluidic system is not optimised to avoid plankton sedimentation, some plankton could accumulate in the fluidic system. This can be checked by pinching the tube halfway in between the Flow Cell and the pump during 1-2 seconds (to accumulate suction pressure) and releasing it (Fig.9). If a large quantity of plankton passes suddenly this means that plankton have sedimented between the syringe and the Flow Cell.





Figure 9: How to unclog the fluidic system

10.7 Acquisition

Go to "Fluidic Acquisition" and set parameters (Fig.11).

• "Number of images to acquire" (to be chosen depending on the desired final object number and the observed concentration on images)



Safety information

Pump significantly between two images will help to:

- Avoid plankton sedimentation in the fluidic system
- Avoid imaging two times the same plankton

Target a sample size (by setting the number of images to acquire) that finally have something like 1000-2000 final objects or more (e.g. if you have 10 objects per image, imaging 100-200 frames would be enough; Fig.10). Lower numbers of objects would be statistically problematic.

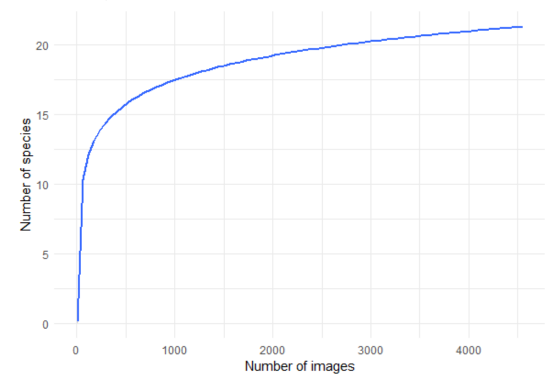


Figure 10: Number of species per number of images acquired for a random sample. Note that the number of species starts to stabilise around 2000 images here.



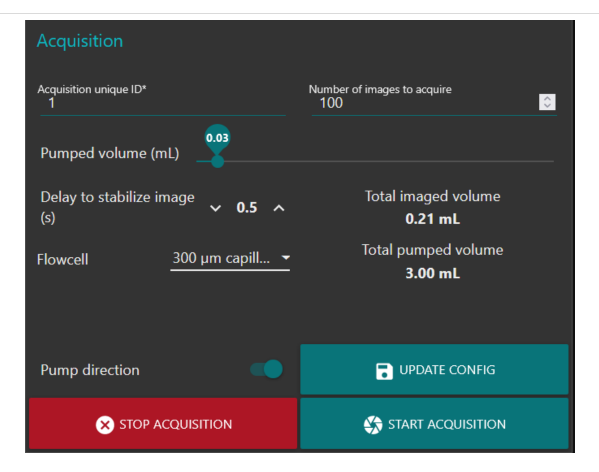


Figure 11: Acquisition parameters. **This figure is not a recommendation**, as it depends on the number of objects you want to acquire and the type of Flow Cell you use.

- "Volume pumped" is the volume to pump in between two images: it should be large enough to: avoid taking twice the same object in picture; avoid large sedimentation in the fluidic system; avoid objects to stick on the Flow Cell. It is recommended testing it in order that the volume passed between two images correspond at least to 5-10 times to the volume imaged (see here the discrepancy between imaged volume/pumped volume).
- "Delay to stabilise image" is the time lag in between the stop of the pump and the
 acquisition of the image. It should be large enough to avoid objects moving while imaged.
- 1. Go to "Fluidic Acquisition" and start the acquisition
- 2. Take two or three images to check if the focus is good. If it is not, try to do it again directly on the plankton.
- 3. Change the acquisition ID, the old one is used by the focus test
- 4. Launch the real acquisition
- 5. Wait for the acquisition to be done
- 6. Results can be consulted by consulting the "Gallery" tab or the file manager (data > export)



Safety information

The PlanktoScope is using a rolling shutter camera which means that there is a small delay in between the first line of pixel imaged and the last line of pixel imaged. To overcome this, it uses a "stop and go" strategy where the imaging only takes place when the flow of the pump is stopped. Not setting this properly will generate artefacts, swimming organisms will also suffer from this (Fig.12).



Figure 12: Copepod nauplii moving while imaged.

Segment the acquisition

11 Segmentation

- 1. Go on segmentation and click on the "UPDATE ACQUISITION'S FOLDER LIST"
- 2. Select the samples you wish to segment, either the parent folder or the file itself
- 3. Setup the different options of the segmenter (Fig.1)



Safety information

Even if the segmentation process can take a lot of time, it is not recommended to start the acquisition of a new sample during this part in case the results are corrupted. You should do the cleaning of the fluidic system and process a new sample only if you are sure that all your images are not corrupted.

- Recursive folder: it will segment all samples within a selected sample
- EcoTaxa archive: it will create a zip file containing all files needed for a easy importation within EcoTaxa
- Force rework: if yes it will re-segment samples already segmented
- Keep objects: it will keep the final segmented images visible in the PlanktoScope. It could be accessed by the "Gallery" in the objects folder

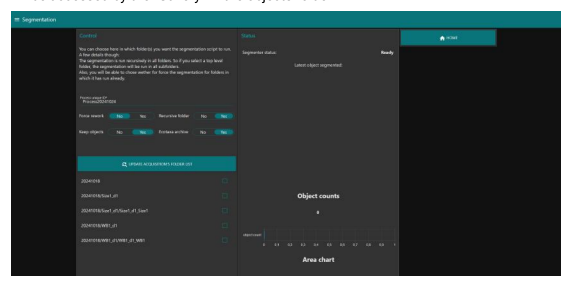


Figure 1: Where to find the folders containing the images to do the segmentation. **Do not forget to update the folder list.**

- 1. Scroll down and click on start segmentation
- 2. Wait for the segmenter status to turn to "Done" (Fig.2)
- 3. Export your data on your computer



Figure 2: End of the segmentation.

How to export data

12

Safety information

In v.2023, you can now use the datafile manager (data > export) to download your files directly. In both versions, you can also use the "Gallery" and directly download the images. It is then possible to no longer use FileZilla.

With FileZilla

You will need a computer connected to the PlanktoScope together with the free software FileZilla (https://filezilla-project.org/).

- 1. Open FileZilla
- 2. Either click on the top left to create a new connection or use the quick-connection fields below (it will not save the ftp site for later, Fig.1)
- 3. To create a new connection "file > site manager > new site"

Enter the following information:

Host: sftp://192.168.4.1 (note images were taken with a previous version, the address does not correspond to i

Username: pi

Password: copepode

Port: 22

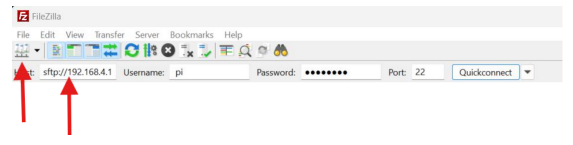


Figure 1: How to log to a ftp host

- 1. Click on "Quickconnect".
- 2. On the bottom panels you have, on the left, the access to what is in your computer, and on the right, the access to what is in the PlanktoScope (Fig.2). Click and slide to transfer data in between both.



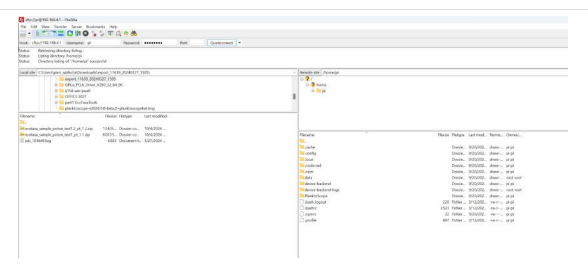


Figure 2: Interface of FileZilla. Personal files are on the left, files of the ftp host are on the right

- Exports file for EcoTaxa are in /home/pi/data/export/ecotaxa.
- Raw images files are in /home/pi/data/img.
- Different control files to check the segmentation process (images after background subtraction, masks of the different objects etc) are in /home/pi/data/clean.
- Final images are in /home/pi/data/objects.

With the "Gallery"

Go to the "Gallery" tab (Fig.3):

- For raw images: img
- For segmented images used in EcoTaxa: export > ecotaxa
- For detailed process: clean
- For segmented images without metadata: objects

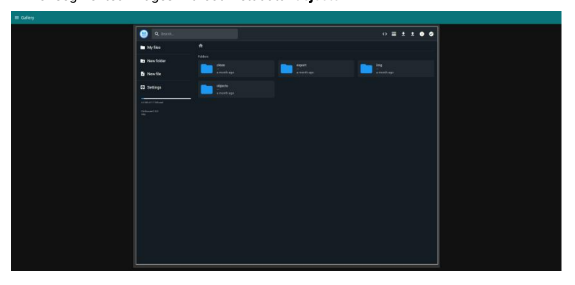


Figure 3: User interface of the "Gallery" tab

Clean the PlanktoScope

13 **Cleaning**



- 1. Drain the sample out of the syringe
- 2. Disconnect the syringe and clean it with distilled water
- 3. Pump (at high speed!) the full content of the fluidic system to remove any liquid
- 4. Reconnect the syringe
- 5. Fill it with distilled water
- 6. Pump (at high speed!) while regularly pinch the tubing to detach any plankton in the system (see **5** go to step #10.6
- 7. Drain again the syringe (repeat steps 2-7 at least 2 more times until no plankton is visible on the camera)
- 8. Finally drain the system

If not used again immediately afterwards

- 1. Put 20 mL diluted bleach
- 2. Leave 15 minutes
- 3. Drain the content (high pump speed)
- 4. Put 10 mL distilled water
- 5. Drain the content (high pump speed)
- 6. If there are traces of calcification, use a diluted acid solution (like HCl diluted by 6).

Upload your images on EcoTaxa

- 14 In this section you will learn to:
 - 1. Do your first connection on EcoTaxa
 - 2. Create a new project
 - 3. Connect to EcoTaxa with FileZilla
 - 4. Import data of the PlanktoScope to EcoTaxa

First connection

• Create an account on EcoTaxa (https://ecotaxa.obs-vlfr.fr/) by clicking on the top right "Log in/Register" then on "Create your EcoTaxa account" (Fig.1).



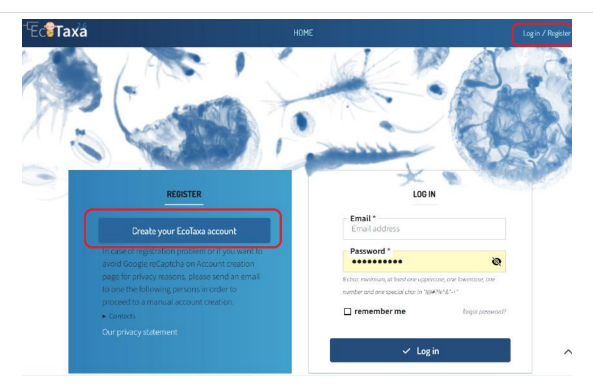


Figure 1: Log in/Register interface.

Put your real name and a valid mail so that you can be contacted (Fig.2)

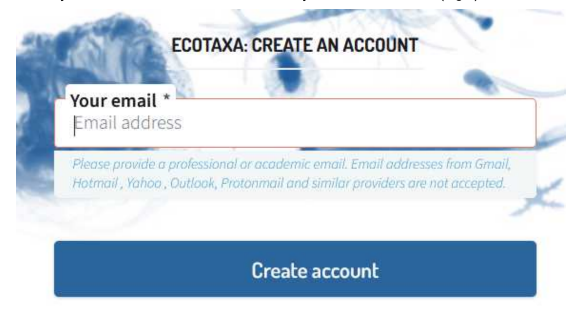


Figure 2: EcoTaxa account creation interface.

14.1 **Projects**

Once logged in, you can consult the project on which you are registered (e.g. your own projects and the ones you have been invited by the different data owners) by clicking on "Contribute to a project" on the main page (Fig.3).





Figure 3: Accessible buttons on the home page.

14.2 Create a project

Go to "Contribute to a project > Create a new project". You can create your own project on which you will be able to import, visualise and classify images.

In the creation panel, you can (Fig.4):

- Add the title of your project.
- Describe your project.
- Comment your project.
- Define the instrument used (here, the PlanktoScope).
- Choose if you want to annotate (define taxonomy) or only explore images, etc. and who can see your project.
- Define what pre-trained Deep Learning features to use on your project (it is recommended to use «Planktoscope_2022-09 » unless you see a more recently trained model on PlanktoScope image).
- Choose a license for your images (it is recommended to use one of the CC-BY one or CC-0 if you want data to have a future use for science).
- Define a list of taxa to help you classify your sample (in the "TAXONOMY" tab).
- Add useful sorting variables in "SORTING TOOLS" that will be added to the top bar filters:
 - area=area
 - meanhue=meanhue
 - meansaturation=meansaturation
 - meanvalue=meanvalue
- Invite new contributors/viewer/manager.

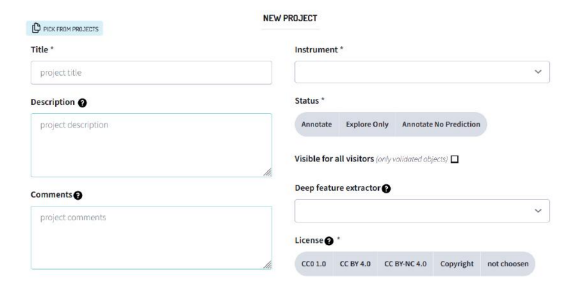


Figure 4: New project interface.



If you want to edit the settings later, go to "Edit project settings" (Fig.4).

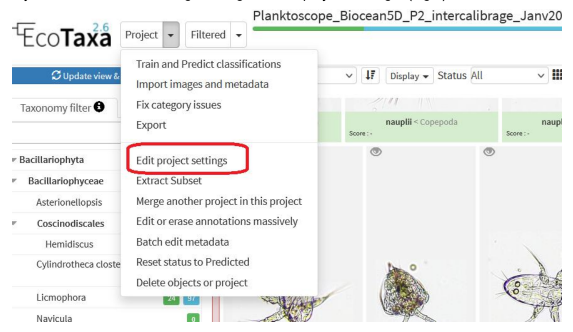


Figure 5: "Edit project settings" button.

14.3 Connect to EcoTaxa ftp

Safety information

In v.2023, you can also download the zip file directly from your computer, without using FileZilla. Otherwise, FileZilla is recommended for huge amounts of data.

Upload the EcoTaxa archives (see step 6-7) on the EcoTaxa ftp

Select File > Site Manager...

Create a New Site called: Ecotaxa_VLFR

In General tag:

Host: plankton.obs-vlfr.fr

Protocol: FTP – File Transfer Protocol Encryption: Only use plain FTP (insecure)

Logon Type: Normal User: ftp_plankton

Password: Pl@nkt0n4Ecotaxa

Once this is done you could use FileZilla to load the Zip files downloaded from the PlanktoScope onto the EcoTaxa ftp server (e.g.

/Ecotaxa_Data_to_import/PLANKTOSCOPE).

Safety information

- Please eventually create your own folder to try to keep it clean and tidy.
- Please think to regularly remove those temporary files from the ftp, at this point they
 are not secured at all and everybody can access them (and disk space is not free).



14.4 Import data in your project

- 1. In your project: on your project options button, select import images and metadata (Fig.6)
- 2. Locate your file on the EcoTaxa ftp folders and import it (only works for one zip file at a time for now; Fig.7).
- 3. Check the quality of your images and the quality of the segmentation once the images are imported

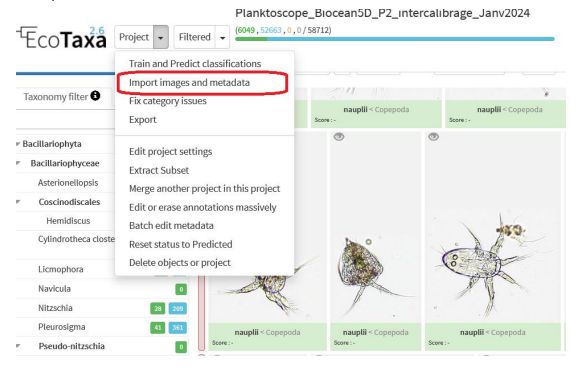


Figure 6: How to import images and metadata.

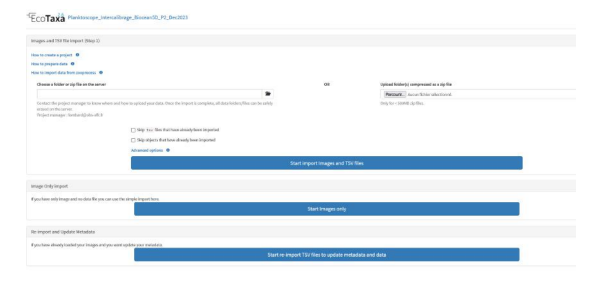


Figure 7: Data import interface

How to use efficiently EcoTaxa



- 15 In this section, you will learn to:
 - 1. Use filters
 - 2. Validate taxonomy
 - 3. Do a prediction
 - 4. Export the results

Note

For more details, please check the online manuals https://sites.google.com/view/piqv/piqv-manuals/ecotaxaecopart-manuals?authuser=0

15.1 Use filters wisely

There are three layers of filters in EcoTaxa (Fig.1):

The taxonomic filter tab that allows to filter by taxonomic groups either from the list of taxa that you defined when creating the project (they will be underlined in the suggestions when you start typing) or from all the taxonomic categories registered on EcoTaxa.

The other filter tabs allows you to filter by sample and by all the parameters of the samples (depth, location, time, annotator, and all the information entered in your metadata).

The top bar is for the status and features filters. They are very useful because they allow you to sort objects according to descriptive values specific to each image (eg. mean saturation in Fig.3, to quickly observe objects that have lots of chlorophyll). You can revert the sorting order of those filters by ascending or descending order. You can also choose to display images according to their status (validated, predicted, dubious, etc) as well as the number of images you want to see per page.

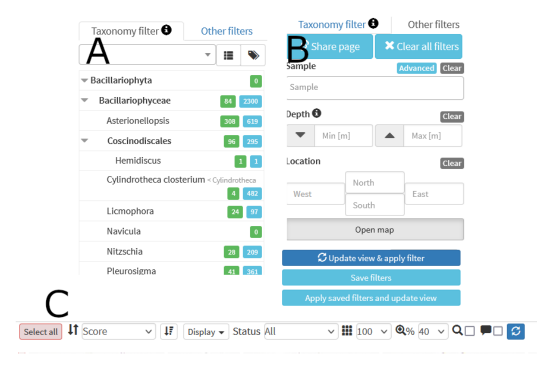


Figure 1: (A) Taxonomy filters; (B) Sample filters; (C) Status and feature filters.



Filters are additive (Fig.2), so you can add filters on location, date, annotator, taxonomic group and every metadata fields entered in EcoTaxa to search for specific things. You can also get rid of them easily by clicking on the cross in the grey fields that you can see at the top of the Fig.2.

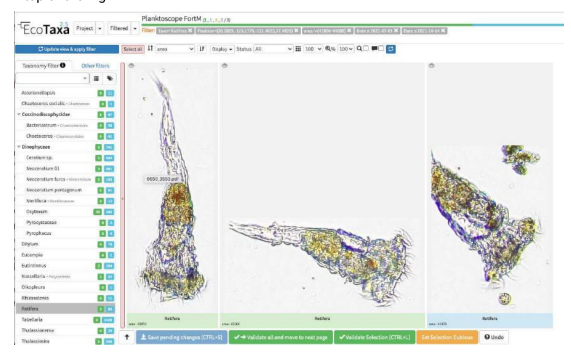


Figure 2: Multiple filters applied in a project. At the top of the figure you can see that the filters allows to see only the taxon "Rotifera" of a specific size (area of the image), at a specific time and location.

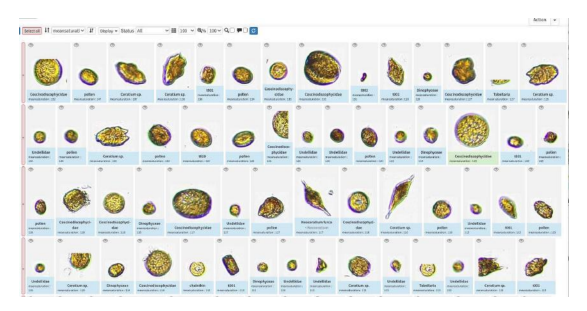


Figure 3: Objects sorted by the mean saturation.

15.2 The different validation status in EcoTaxa and how to validate

Image imported in EcoTaxa have the status "Unclassified" (grey surrounding of the image, Fig.4).



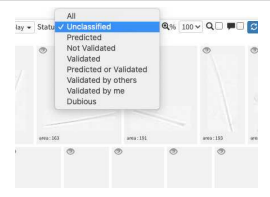


Figure 4: Unclassified images.

However they could be also set as "Predicted" (blue surrounding; classified automatically by taking as example one pre-existing project), "Validated" (green surrounding; checked and annotated by a human), or "Dubious" (orange surrounding; checked and annotated as dubious by a human) (Fig.5).



Figure 5: Types of images status: blue predicted, green validated, orange classified as dubious and grey unclassified.

You can validate an image either by dragging it into a taxonomy filter among the taxa defined in the preset (Fig.6), or by typing the name of the taxon directly into the search bar of the "Taxonomy filter" tab (Fig.7). Once validated the name appears in red below the images and they appear surrounded by green (Fig.). For the validation to be taken into account, it is important to always save either with ctrl + S or with the "Save pending changes" button (Fig.9).



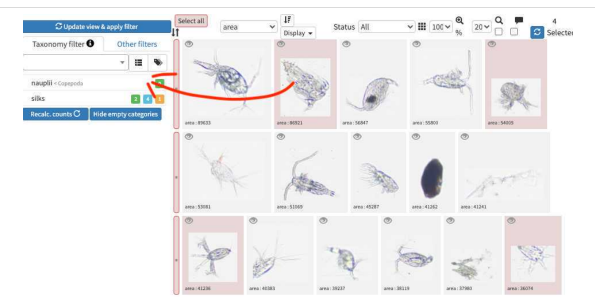


Figure 6: Dragging image to validate

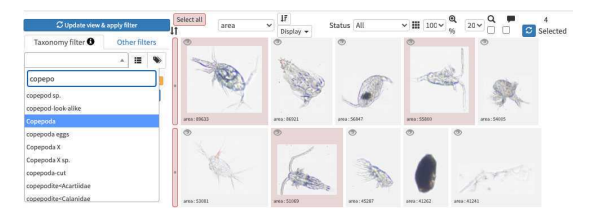


Figure 7: Typing in the search bar of the "Taxonomy filter" tab to validate

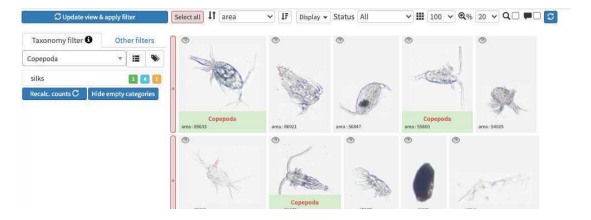


Figure 8: Image configuration after validation



Figure 9: Do not forget to save your validations

When you have a lot of images and/or are dealing with unfamiliar taxonomic categories, the validation process can take a lot of time and energy. To speed up this process, you can use prediction tools. They allow you to validate thousands of images with a single click (Fig.10).



Figure 10: Example of well predicted objects which would be easily validated.

15.3 **Prediction**

Once you have validated at least 20 images per taxonomic category, you can use the prediction tools to speed up the validation process. To predict the images of your taxa in the taxonomy filter, you can use your project or another pre-existing PlanktoScope project, preferably in the same location and with similar plankton community as a reference (Fig.12). You can run a prediction directly after importing your images with another project as reference, but be aware that the quality of the prediction will not be ideal. The more images you validate, the more reliable the prediction will be. 20 images is the smallest number of validations per taxonomic category necessary to ensure the quality of the prediction if you are using your own project as a reference.

Safety information

You can use the PlanktoScope machine learning algorithm in your project. *Go to Project > Edit project settings* and chose the SCN Network "planktoscope_2022-09". Do not forget to save changes.

 In "Project" (or "Filtered", if only the filtered images need to be predicted), select "Train and Predict classifications" (Fig.11).



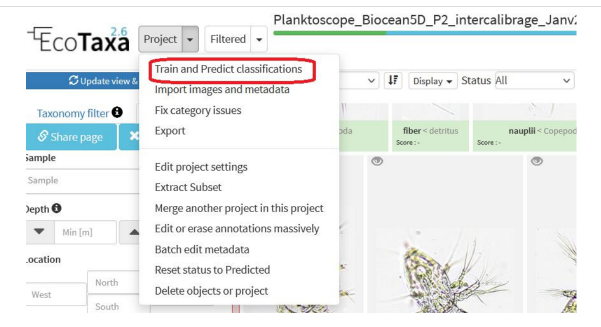


Figure 11: Click on the button "Train and Predict classifications" to start a prediction

Launching a prediction directly after importing the images into EcoTaxa will allow you to quickly validate around twenty images on your own project. However, until you have validated the images, you should use a pre-existing project as a reference for the prediction (Fig.12). Note that currently, only few PlanktoScope projects acquired with the same segmentation procedure than we do exist, we therefore strongly encourage you after a first trial of prediction to quickly validate to then predict on your own project.

Projects that could be used for first prediction:

- #6818 MOOSE-GE-2022_tests_ID_vignettes (Med sea; Processed with current segmenter;
 Fully validated) https://ecotaxa.obs-vlfr.fr/prj/6818
- #9621 APERO_PP&THA_Phytonet_35mu_Planktoscope https://ecotaxa.obs-vlfr.fr/prj/9621
- #10 056 AtlantECO_P2_Ada_2023 (PlanktoScope; Fully validated) https://ecotaxa.obs-vlfr.fr/prj/10056
- #6765 Planktoscope Reference (Processed with other segmenter, works only with adding Deep Learning features into play); Fully validated) https://ecotaxa.obs-vlfr.fr/prj/6765



Figure 12: Prediction learning sets.

Click on "Next: Choose objects in selected projects". You then have the possibility to select what types and quantity of objects to consider. It is recommended to avoid selecting too many objects in a category, in order to partially correct the usual strong imbalance between categories (Fig.13, limited to 500 objects per group).



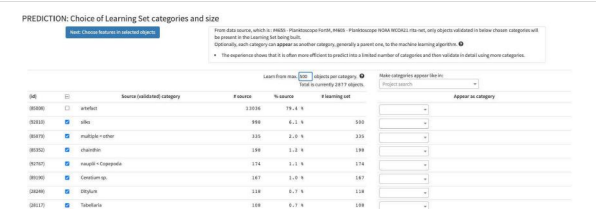


Figure 13: Choice of objects to train the machine learning algorithm.

Click on "Next: Choose features in selected objects" to activate the pre-trained deep learning features (if not available see step 10). You can uncheck variables that are not relevant for prediction and relate to position of the vignette in the initial images such as bx, by, depth min/max, label, local centroid col/row, x, y (Fig.14).

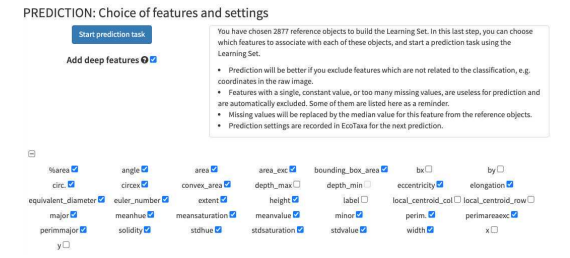


Figure 14: Prediction features and settings.

Click on "Start prediction task". Once done, images have the status "Predicted". Each image has a "Score" that represent the reliability of the prediction (Fig.15). It is therefore a good filter to use when you want to validate a large number of images quickly to then launch a prediction with your own project. Do not hesitate to launch a prediction as soon as you have validated most of the images with a high score.



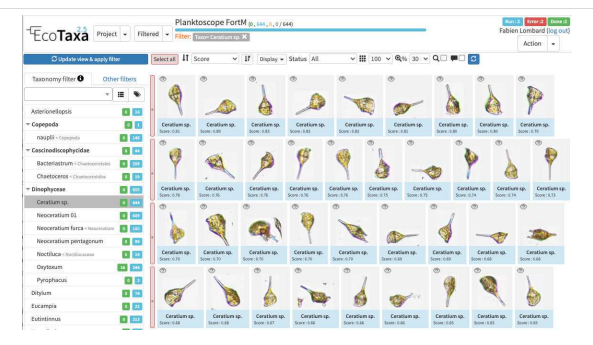


Figure 15: Example of images sorted by score of prediction.

Safety information

Making repeated predictions on your own project is always better than doing so on preexisting random projects.

Keep validating and making predictions until your project is fully validated.

15.4 Export your results from EcoTaxa

Once fully validated, export your results (Fig.16). Different solutions exist, general export for configurable objects export (Fig.17), back up export for restoring or archiving and summary export for synthetic taxon-oriented export (Fig.18).

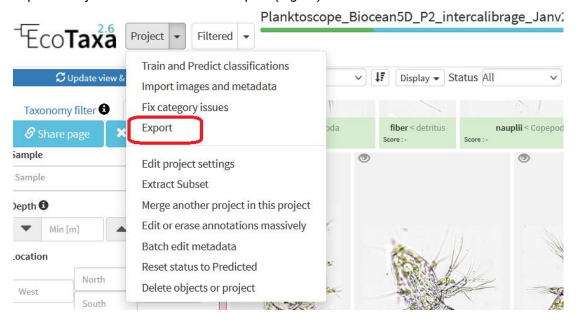


Figure 16: How to export data.



A general export will give you all the metadata of your project. You can chose to separate (in ZIP sub-directories) output by sample, acquisition, taxon or to not separate the output. We recommend to chose to separate by sample (Fig.17).

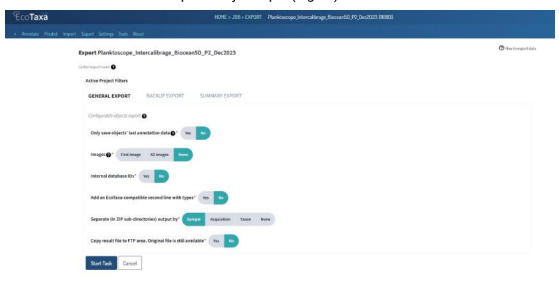


Figure 17: General export interface

With the summary export, you will not see all the metadata and thus the potential errors lying in it. It is very important to check for errors and what is missing from the metadata to ensure that we have a full quantitative signal and not just a relative abundance signal. This type of export allow you to chose what you want to compute, either the abundance, the concentration or the biovolume (Fig.18). Depending on what you choose as the to compute parameter, you need to enter a specific formula to extract them. For the abundance, this is "just" counts of vignettes, you do not need everything. For the concentration and the biovolume it depends a lot on the method you used to sampled and on the metadata.

Concentration:

total_water_volume: **sam.total_volume/1000/1000** (%from mL to m³) subsample_coeff:

sam.concentrated_sample_volume/(sam.total_volume*ssm.imaged_volume*sam.dilution_factor)

Biovolume: (plain area)

individual_volume: 4.0/3.0*math.pi*(math.sqrt(obj.area/math.pi)*ssm.pixel)**3

With:

sam: sample part of the data

ssm: process and acquisition part of the data

Safety information

These formula only work when all volume have the same units: either \mathbf{mL} everywhere or $\mathbf{m^3}$ everywhere.



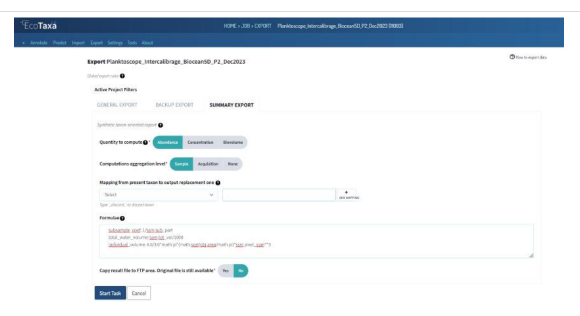


Figure 18: Summary export interface

Safety information

Before starting the data analysis, it is very important to check for errors and what is missing from the metadata to ensure that we have a full quantitative signal and not just a relative abundance signal.

How to compute biovolumes

16 Calibrate your data

EcoTaxa generate a lot of interesting variables to analyse your sample. Only a few of them are depicted here, but do not hesitate to explore the other ones.

Before any analysis, it is important to relate all the parameters computed by EcoTaxa with the acquisition parameters. You will need:

Variable	Unité EcoTaxa	Unité Script	Calcul
Sample_concentrated_sample_volume	mL	m ³	X/1 000 000
(V _{concentrated})			20
Acq_imaged_volume	mL	m ³	X/1 000 000
(V _{imaged})			
Sample_total_volume	L	m ³	X/1 000
(V _{total})			(=1 si NA)
Pixelsize	μm	mm	X/1 000
Major	рх	mm	X * pixelsize
Minor	рх	mm	X * pixelsize
Area_exc	рх	mm ²	X * pixelsize ²
Area	рх	mm ²	X * pixelsize ²

Table 1: Transformations to apply to essential variables.



It will allow you to compute the conversion factor "conver". The dilution factor should be < 1 if a dilution has been made and > 1 if the sample has been concentrated (it is not the sample concentrated volume, which is the original volume taken from the collector):

$$conver = rac{V_{concentrated}}{V_{imaged}*V_{total}*Dilution}$$

Safety information

Do not forget to convert all the parameters like above. Otherwise, it will produce wrong results.

Compute Biovolume

There is three methods to calculate the biovolume (BV) of an object.

Ellipsoid:

$$AR = \pi * rac{major}{2} * rac{minor}{2} \ BV = rac{4}{3} * rac{minor}{2} * AR$$

Plain:

$$ESD = 2*\sqrt{rac{area}{\pi}} \ R^3 = (rac{ESD}{2})^3 \ BV = rac{4}{3}*\pi*R^3$$

Riddled:

$$ESD = 2 * \sqrt{rac{area_exc}{\pi}}$$
 $R3 = (rac{ESD}{2})^3$
 $BV = rac{4}{3} * \pi * R^3$

Maintenance of your PlanktoScope

17 Clean tubing and Flow Cell from inside

Imaging plankton will lead to have a lot of organic material and seawater in the fluidic system. Some may clog or accumulates in some parts of the fluidic system.

- 1. Do not let it dry and try to get rid of it as soon as possible. If it occurs during sample acquisition, abort it, take care of the clog. You may need to dilute the sample, note the dilution in the metadata and restart acquisition.
- 2. Pump distilled water with high pumping rates helps to unclog the system. Make sure no plankton organisms remain in the fluidic system and especially on the internal walls of the Flow Cell. If it is the case do not hesitate to pinch (during 1-2 second) and release the tubing between the Flow Cell and the pump while pumping to create a sudden variation of pressure.
- 3. Over time, wet conditions and organic matter may create favourable condition for the growth of a bacterial film. The Flow Cell and tubing will look dirty from the inside. You can avoid this by pumping diluted bleach, let it in for 1-2 hours and carefully rinse the whole system.
- 4. Water, bacteria, and bleach together may favour the apparition of a calcium carbonate film inside the tubing and Flow Cell. It may either appear as dispersed crystals attached inside the Flow Cell or a white coating inside the tubing. To remove and clean this, pump some



acidic solution (vinegar, citrus juice or other kind of other acids), let it rest for a few hours and rinse the system.

Clean Flow Cell outside:

The Flow Cell is an optical critical component, keeping it clean is an absolute necessity. Do not touch it with fingers or other kind of dirty material. If dirty:

- 1. If only dry dusts are present, gently blow the Flow Cell with the cleaning blower.
- 2. If the dirt is not only dry dusts it could be cleaned with optical paper and ethanol.

DO NOT USE CLASSICAL PAPER TOWELS!

They are usually enriched in silica fibres for solidity and may create scratches on the Flow Cell. If optical paper is not available, paper tissues are a better alternative.

Clean optical lenses

As for the Flow Cell, optical lenses are critical elements of your PlanktoScope and should be kept as clean as possible. It starts by never touching them with fingers! Cleaning those would require a lot of patience, efforts and may even lead to unexpected disappointments.

- 1. Dry dust: dry gas with even more caution than previously.
- 2. Others: only used optical paper.

Clean the camera sensor

Critical part! **NEVER** touch it, only use dry gas.

Regularly calibrate the pump and the WB.

Update the software or reset the PlanktoScope

- 18 If you need to update the software, or if there are a lot of bugs and you want to reset it, follow this procedure.
 - 1. Withdraw the micro SD card at the bottom of the PlanktoScope with a fine forceps (Fig.1).
 - 2. Connect the SD card to your computer with a SD card adapter if needed (you can find one in the supplementary materials box).





Figure 1: How to withdraw the micro SD card

- 1. Open BalenaEtcher > Flash from file (Fig.2)
- 2. Choose the desired version of the software (.img file). You can find the actual one on the github of the PlanktoScope in the "External links" section of the protocol.
- 3. Choose the SD card you want to flash
- 4. Click on "Flash from file" and wait until it is flashed
- 5. Eject it from your computer and put it back in the PlanktoScope with a fine forceps



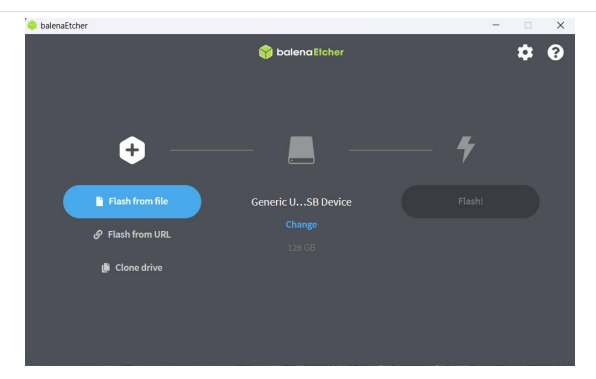


Figure 2: BalenaEtcher interface

Troubleshooting

19 The Flow Cell is clogged with plankton Why this happens:

- First this may happen if your sample does not have been pre-filtered. It is recommended to do a pre-filtration to 200 µm.
- It may also happen if your sample is too concentrated. If you got more than 20 plankton objects per image this may already be the case, dilute your sample and fill the dilution factor in the sample metadata.
- If you forget to use a bubbler to agitate your sample, or if you let your sample stagnate too long in the fluidic system.

Unclogging the Flow Cell:

- Try to pinch the tube in between the Flow Cell and the pump while the pump is running.
- Try to do the same while pumping in the reverse direction eventually at high speed.
- Dismount the Flow Cell but keeping the tube adapter on it. On the side which was connected with the pump, connect a syringe and pass air/water to chase the blocked plankton.

19.1 The image is partly blurred

Why this happens:

• The focus is correctly done, but the Flow Cell is not well positioned.

How to correct it:

• Try to adjust the position of the Flow Cell with tape (Fig.1).



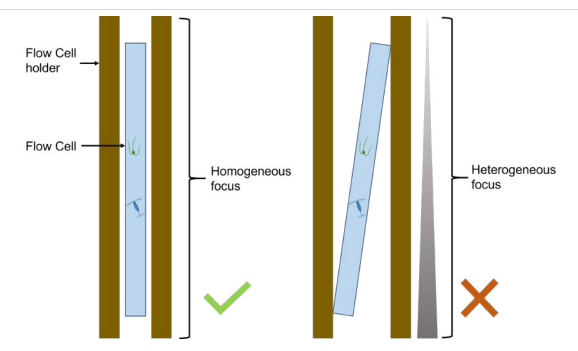


Figure 1: On the left, a Flow Cell correctly positioned. On the right, the Flow Cell do not fit correctly between the fixations of the Flow Cell receptacle, creating a gradient blur-sharp.

19.2 The software is not working

Why this happens:

- The python code encountered a bug.
- There is a segmentation error because the number of objects is too important.
- The optical configuration tab does not work (black screen, impossible to change the WB, etc...).

Solutions:

- Try to restart the PlanktoScope.
- Try to change values of the WB, check if you use commas or dots and restart the PlanktoScope.
- Try to stand by a number of final objects around 2000~3000 per sample.
- Ask guestions on the PlanktoScope Slack (see "External links" section).
- If you do not find any solution, flash the SD card of the PlanktoScope with the software by using BalenaEtcher.

19.3 The pump is not working

If there is any problem with the pump, check that it is properly positioned (Fig.2). If it is not the problem, remove it and check that the pump tube is correctly positioned.

- 1. If you need to change the internal pump tube, you can take the pump off by turning it counter-clockwise.
- 2. Position the tube correctly
- 3. To install it again, place it in the position shown in the image below until it clips, push it in and turn it clockwise until it snaps into the white socket (Fig.3).
- 4. Be careful to clean the grease afterwards



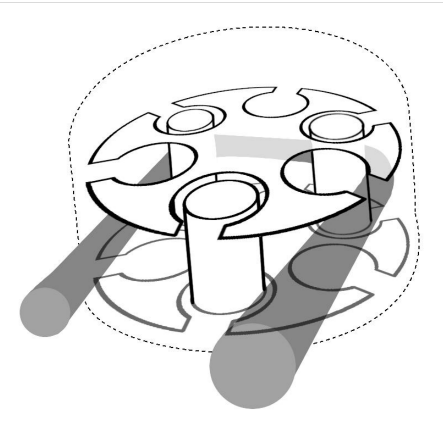


Figure 2: How the tube should be installed inside the pump.

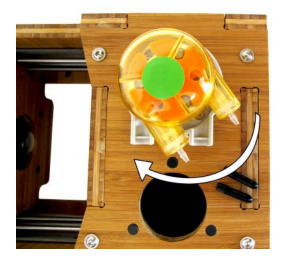




Figure 3: How to install the pump.

External links

20 PlanktoScope website

https://www.planktoscope.org/

PlanktoScope github

https://github.com/PlanktonPlanet/PlanktoScope



PlanktoScope complete assembly guide and complete documentations

https://planktoscope.curious.bio/ (v2.5)

PlanktoScope Slack channel (to exchange ideas/protocols/solutions/questions)

https://forms.gle/qvh5jwuMvmyBKMQC7

Plankton Planet website

https://planktonplanet.org/

EcoTaxa tutorials:

https://sites.google.com/view/piqv/piqv-manuals/ecotaxaecopart-manuals?authuser=0

https://www.youtube.com/watch?v=PSO6ZS765tk

https://www.youtube.com/watch?v=RaWUqloKk0E