Chlorophyll

1 Extraction

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Ш	Acetone (2 mL per sample)
	Centrifuge with rotor 12154
	Ice
	1,000 μ L pipette and tips (2 tips per sample
	Vortexter
	Centrifuge
П	Kimwines

1.2 Method



Chlorophyll is very unstable. Always keep samples on ice and as dark as possible.

- 1. Centrifuge up to 24 sample tubes using rotor 12154 for 5 min at 12,000 rpm at 4° C. Keep centrifuge balanced if less than 24 tubes are used.
- 2. Remove supernatant with 1,000 µL pipette. Carefully remove remaining drops of supernatant with Kimwipes.
- 3. Add 1,000 µL acetone and close tube. Homogenize with pipette and vortex for 15 s. Do not re-use tips.
- 4. Extract chlorophyll for 24 h at 4 °C (Fridge Biomol 2). Keep samples wrapped in aluminium foil.

2 Measurement

2.1 Overview

The chlorophyll concentration ($g mL^{-1}$) is calculated based on the formulas given in (Jeffrey & Humphrey, 1975):

Chlorophyll
$$a = 11.43 \frac{A_{663} - A_{750}}{PL} - 0.64 \frac{A_{630} - A_{750}}{PL}$$

, where A_{663} , A_{630} , and A_{750} is the absorption at 663, 630, and 750 nm, and PL the path length of the cuvette (1 cm).

The photometer can read whole plates (96 samples). However, since acetone is used to extract the chlorophyll, a quartz cuvette has to be used instead of plates made of plastic. The software of the photometer still visualizes the measurements of this cuvette as a plate which can be confusing.

The photometer will automatically measure the absorption at 663, 630, and 750 nm and does the blank-correction automatically. Each measurement is done twice and blanks are measured at the beginning and after every 10 samples, resulting in the following plate outlay:

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	1	2	3	4	5	6	7	8	9	10	11	12
Α	BLK	BLK	SPL1	SPL1	SPL2	SPL2	SPL3	SPL3	SPL4	SPL4	SPL5	SPL5
В	SPL6	SPL6	SPL7	SPL7	SPL8	SPL8	SPL9	SPL9	SPL10	SPL10	BLK	BLK
С	SPL11	SPL11	SPL12	SPL12	SPL13	SPL13	SPL14	SPL14	SPL15	SPL15	SPL16	SPL16
D	SPL17	SPL17	SPL18	SPL18	SPL19	SPL19	SPL20	SPL20	BLK	BLK	SPL21	SPL21
Е	SPL22	SPL22	SPL23	SPL23	SPL24	SPL24	SPL25	SPL25	SPL26	SPL26	SPL27	SPL27
F	SPL28	SPL28	SPL29	SPL29	SPL30	SPL30	BLK	BLK	SPL31	SPL31	SPL32	SPL32
G	SPL33	SPL33	SPL34	SPL34	SPL35	SPL35	SPL36	SPL36	SPL37	SPL37	SPL38	SPL38
Н	SPL39	SPL39	SPL40	SPL40	BLK	BLK	SPL41	SPL41	SPL42	SPL42	SPL43	SPL43

BLK corresponds to blank measurements and SPLXX to samples.

Note

The photometer automatically blank-corrects the measurements which can lead to wrong measurements if the order of samples and blanks differs to the expected order. It is therefore important to note the actual order of the samples in the metadata sheet (described here).

2.2 Materials

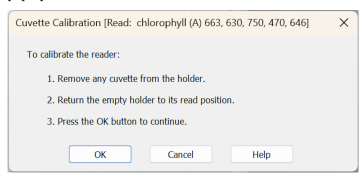
- ☐ Vortexter
- ☐ Centrifuge with rotor 12154
- ☐ Ice
- \square Acetone
- ☐ Kimwipes
- \square 1000 μL pipette & tips
- $\hfill \square$ BioTek Epoch2 microplate reader
- ☐ Quartz cuvette (dark and shattered)

2.3 Method

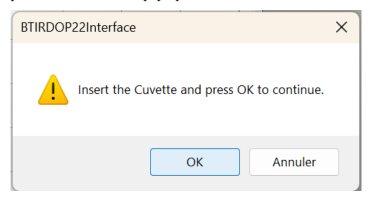
- 1. After the incubation (note down incubation time), vortex samples for 15 s and centrifuge samples using rotor 12154 for 5 min at 12,000 rpm at 4 °C. Keep samples on ice and in darkness. Transport samples carefully to not damage the pellet.
- 2. Turn on BioTek photometer and connect to Dell via USB.
- 3. Double-click on 2024_04_20_chlorophyll+carotinoides_andi.prt to start software (backup stored here).
- 4. Click on Create experiment and read now:



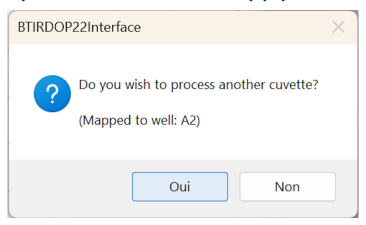
5. At first, the photometer takes a background measurement without any cuvette inserted. Follow the instructions in the pop-up and click OK.



6. Start measuring the blank. Clean quartz cuvette with acetone and Kimwipes, add 500 μL acetone, and insert cuvette into photometer. Click 0K in pop-up window.



7. Repeat blank measurement. Click on Oui in pop-up window.

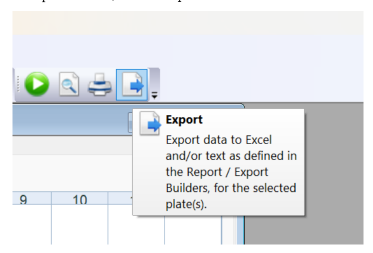


- 8. Measure first sample. Remove cuvette from photometer, clean with acetone and Kimwipes and add $500 \mu L$ sample extract. Click on 0ui in pop-up window (as for second blank measurement).
- 9. Repeat measurement (click Oui).
- 10. Remove cuvette from photometer, clean with acetone and Kimwipes and measure next sample.
- 11. Repeat as described before for 10 samples (i.e. 20 measurements). Then, measure blank as described before (2x) and continue to measure samples. Take care not to damage the pellet.

i Note If you do a mistake in the sequence of measurements, correct it accordingly in the metadata sheet.

- 8. When all samples are measured or after 43 samples, click on Non in pop-up window.
- 9. Save experiment in your folder. Name it as follows: YYYY_MM_DD_chl_adaptome_#.xpt, where YYYY_MM_DD is the date and # the batch number (each day, start with 1), e.g. 2024_11_19_chl_adaptome_1.xpt.

10. To export the data, click on Export.



11. Save data as .txt (default) in your folder and name it YYYY_MM_DD_chl_adaptome_#.txt (as described above).

Caution

Backup your data each day you are measuring. Either upload the folder to Google Drive or save it on a USB key. Keep all data also on the computer.

Jeffrey, S. W., & Humphrey, G. F. (1975). New spectrophotometric equations for determining chlorophylls a, b, c1 and c2 in higher plants, algae and natural phytoplankton. *Biochemie Und Physiologie Der Pflanzen*, 167(2), 191–194. https://doi.org/10.1016/S0015-3796(17)30778-3