

Department of Geomorphology and Quaternary Geology – Procedure

Bulk pigment extraction from lake sediments

Consumables and equipment

Chemical reagents

- Acetone 100% (grade HPLC): solvent.
- Acetone **technical**: cleaning.
- Redistilled water (MilliQ).
- Technical nitrogen.
- Neodisher Laboclean.

Neodisher LaboClean FLA: liquid, highly-alkaline cleaning, dispergator agent.

Equipment

• Laboratory fume hood.



Acetone

Acetone (C_3H_6O) is an organic solvent requiring work under the **running fume hood**. Acetone is highly **flammable** and is an **irritant**.

- Ultrasound bath.
- Centrifuge.
- Vortex shaker.
- Laboratory furnace for glassware baking.
- Laboratory oven.
- Freeze drier.
- Glass beakers for acetone pippeting.
- Acetone pipette $1000 \mu l$ and tips (for acetone).
- Heating plate.
- Nitrogen evaporator.

- Needles for \mathcal{N}_2 evaporator.
- Metal rack for amber vials on the heating plate.

For example *Turbovap* instrument.

- Racks for centrifuge tubes and amber vials.
- Analytical balance, antistatic device and stainless steel spatula.
- Ice bath.

Plastic tray with cooling packs.

• Crimper (for closing HPLC vials).

Other

• Polypropylene (PP) centrifuge tubes (falcons) **25 ml** or **50 ml** (CorningTM).

Other plastics will melt in contact with acetone.

One-time use.

• Amber vials with caps 25 ml or 50 ml.

Tape the labels.

• PTFE hydrophobic filters of **0.2** m (**13** mm).

One-time use.

• Polypropylene (PP) small syringes with Luer Lock for filtering (2 ml).

One-time use.

Important

Always check size and fit compatibility.

- Aluminum foil.
- Glass Pasteur pipettes.

One-time use.

• Silicone nipples for pipettes.

Cleaning

Amber vials

- Put into soap water with Neodisher Laboclean for few hours.
- Wash in the washing machine.
 - Glassware program and Neodisher Laboclean.
- Dry in the drying oven.
- Close with aluminum foil (cleaned with technical acetone).
- Bake in the furnace in 500 °C.

Minimum 3 h.

Amber vials caps

- Rinse with tap water.
- Put into beaker filled with technical acetone and redistilled water (MilliQ) mixture (1:1).
 - Sonicate for **10 minutes**.
- Put into beaker filled with technical acetone.
 - Sonicate for **10 minutes**.
- Dry in the drying oven.

Washing solvents can be used again. Store in the glass bottles.

Evaporator needles

- Put into beaker filled with technical acetone.
- Sonicate 3×10 minutes.

Disposal

- Acetone accordingly.
- PP falcons and glass Pasteur pipettes need to be washed and recycled.

Sediment preparation

• Freeze dry.

High temperature speeds up pigment degradation.

• Loss on ignition 550 °C.

Paleoenvironmental Research Laboratory: calculate organic matter from CNS.

• Weigh **0.5** g homogenized sediments into polypropylene falcons (PP) **25** ml.

Clean tools with technical acetone, use antistatic device.

If there is not enough material, use no less than **0.25** g.

• Label falcon and cap with sample name.

Pigment extraction

- Prepare workplace, cover surfaces under the fume hood with aluminum foil cleaned with technical acetone.
- Switch off the lights whenever possible.
- Fill glass beaker with 100% acetone.

Don't pipette from the bottle.

• Rinse pipette with acetone.

Pipette into waste beaker.

• Add 5 ml of acetone to the falcon.

pipette right, there should be no air bubbles in the tip.

- Shake with vortex for 1 minute.
- Sonicate for 1 minute.
- Centrifuge:
 - 10 minutes.
 - 5000 rpm.

This step needs to be adjusted to the centrifuge. Lower rpm requires longer centrifugation.

• Check if supernatant is clear after centrifugation.

If it is not clear centrifugate again.

• Remove (transfer) supernatant using glass Pasteur pipette into amber vial.

Label amber vial accordingly.

Be sure all supernatant is transferred.

- Repeat two times using 5 ml acetone each time.
- Hold labelled Pasteur pipettes in a glass beaker between the extraction steps.

Every amber vial and falcon has its own Pasteur pipette.

• Amber vials between extraction steps should be covered from light and stored in the ice bath.

After three extraction cycles, if supernatant still has a strong color continue with additional extraction steps.

It is important that all samples are prepared the same way.

Extracts can be stored for a few days in the -20 °C temperature.

Concentration and reconstitution

Evaporation

- Set heating plate and nitrogen evaporator.
- Evaporate samples to dry residue.

Before use be sure that needles are clean (Section ??).

- Put amber vials in the rack on the heating plate set to 35 °C.
- Open nitrogen flow.

Set flow so small flexion of the liquid surface is visible.

• Evaporate to dry residue (around 4 to 6 h).

Reconstitution and filtering

• Add 2 ml 100% acetone to the amber vial using 1000 µl pipette.

Be very precise, check for air bubbles in the pipette tip.

- Homogenize solution with the glass Pasteur pipette.
- Rinse the vial walls and *vortex*.

All pigments need to redissolve in the acetone.

- Take a 2 mL syringe, attach the 0.2 μm PTFE hydrophobic filter on.
- Pipette directly with a pateur pipette into the syringe.
- Pass the solvent through the filter into the small HPLC glass vials with the piston.
- Loose the filter from the syringe to get all solvent out.
- Close the HPLC vials with a crimper and store your samples in a labelled sample box at -20 °C.

Credits

Andrea Sanchini, Giulia Wienhues and Paul Zander – University of Bern, Switzerland.

Changelog

- 09.12.2022, MZ initial version.
- 09.02.2023, MZ full version.
- $\bullet~17.03.2023,~\mathrm{MZ}-\mathrm{update}$ with filtering and call outs.

maurycy.zarczynski@ug.edu.pl

http://geomorfologia.ug.edu.pl