REstoring optimal SOil functionality in degraded areas within organic VinEyards

European research project 2015-2018

PROTOCOLS FOR SOIL FUNCTIONALITY ASSESSMENT IN VINEYARDS
ACKNOWLEDGMENTS

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Summary

Summary ............................................................................................................................................. 2
Overview .................................................................................................................................................. 3
About ReSolVe project ............................................................................................................................. 3
How to use this guide ................................................................................................................................. 3
Introduction ............................................................................................................................................... 4
Soil functions and functionality .................................................................................................................. 4
Why optimal soil functionality in vineyard is important .......................................................................... 4
Part I: Soil monitoring .................................................................................................................................. 5
  Methodological sheet n°1: soil profile description and sampling ............................................................. 6
  Methodological sheet n°2: bulk density assessment .................................................................................. 8
  Methodological sheet n°3: water retention curve and available water capacity assessment .................... 10
  Methodological sheet n°4: standard physical and chemical soil analysis ................................................. 12
Part II: Monitoring of soil ecosystem service provision and providers .................................................... 19
  Methodological sheet n°5: organic matter breakdown assessment using Tea-Bag Index ......................... 20
  Methodological sheet n°6: earthworms monitoring .............................................................................. 22
  Methodological sheet n°7: enzymatic activities assessment .................................................................. 23
  Methodological sheet n°8: assessment of soil respiration, microbial biomass and communities ............ 25
  Methodological sheet n°9: soil mesofauna using Berlese-Tullgren extractors ......................................... 28
  Methodological sheet n°10: nematodes monitoring .............................................................................. 30
Part III: Soil rhizosphere monitoring ....................................................................................................... 32
  Methodological sheet n°11: assessment of communities and pure culture isolation of root- or rhizosphere-associated fungi and bacteria ................................................................. 33
  Methodological sheet n°12: direct assessment of mycorrhizal infections in root samples ...................... 36
Part IV: Grapevine monitoring .................................................................................................................. 37
  Methodological sheet n°13: grapevine vegetative growth assessment using machine vision ............... 38
  Methodological sheet n°14: grapevine water status assessment ............................................................. 40
  Methodological sheet n°15: grape yield assessment ............................................................................ 42
  Methodological sheet n°16: grape composition assessment .................................................................. 43
Additional bibliography .............................................................................................................................. 44
Overview

About ReSolVe project

The ReSolVe project aims at testing the effects of selected agronomic strategies for restoring optimal soil functionality in degraded areas within organic vineyard. The term "degraded areas within vineyard" means areas having been reduced vine growth, disease resistance, grape yield and quality. These areas can have lost their soil functionality because of either an improper land preparation, or an excessive loss of soil organic matter and nutrients, erosion and/or compaction, metal accumulation. The project individuated the main causes of the soil functionality loss and tested different organic recovering methods, such as adding compost, adoption of green manure with different species, and dry mulching. The effects of these techniques were evaluated through the monitoring of various components and consequences of soil functionality:

- soil physical, chemical and hydrological properties, such as organic matter content, soil nitrogen and water availability;
- soil and root-zone biodiversity and biological activity: abundance and diversity of microorganisms, microfauna and earthworms, enzymatic activities, organic matter turnover, mycorrhizae;
- grapevine behaviour as a response to soil status: plant water stress, plant phenology, grape yield and quality.

More information about the project can be found on www.resolve-organic.eu.

How to use this guide

The purpose of this guideline is to describe the methods used during ReSolVe project for soil functionality assessment, so they can be implemented in similar studies.

A brief introduction first underlines what are the main functions of soil and why maintaining an optimal soil functionality is particularly of major interest in viticulture.

Then the different protocols selected for ReSolVe project and this guideline are presented according to the following classification:

- Part I: assessment of soil physical and chemical features;
- Part II: assessment of soil biological features (ecosystem service provision and providers);
- Part III: assessment of rhizosphere biological features;
- Part IV: assessment of grapevine quantitative and qualitative indicators reflecting soil functionality.

In each part, global objectives of the monitoring are explained (what is it used for, in which cases...) and the parameters to evaluate are listed with their corresponding methodological sheet.

In these sheets, instructions and information are given about:

- Materials needed to perform the sampling and the measurement
- Sampling procedure
- Analysis procedure
- Possible interpretations and conclusions that can be drawn (value and meaning of the results, indication of reference values when existing, potential limit of the protocol)
- Bibliographic references related to the method described
- Additional helpful information where appropriate (ex: template of sampling sheet)
Introduction

Soil functions and functionality

The soil functions concept emerged during the early 1970’s (Glenk et al., 2012) and was adopted for the development of the EU Soil Framework Directive with seven key soil functions (European Commission, 2006):

- Biomass production, including in agriculture and forestry
- Storing, filtering and transforming nutrients, substances and water
- Biodiversity pool such as habitats, species and genes
- Physical and cultural environment for humans and human activities
- Source of raw materials
- Acting as carbon pool (store and sink)
- Archive of geological and archaeological heritage.

Dominati et al. (2010) suggested the following roles of soils in the provision of services:

- Fertility role
- Filter and reservoir role
- Structural role (i.e. physical support)
- Climate regulation role
- Biodiversity conservation role
- Resource role.

These correspond roughly to the soil functions as presented by the European Commission (2006) and are overlapping with the general concept of an Ecosystem Service. One aspect that might be added is the increasing awareness of cultural services.

Soil functionality is the degree of soil function extend and can be quantitatively or qualitatively evaluated.

Soil function describes what the soil does. Soil functions are: (1) sustaining biological activity, diversity, and productivity; (2) regulating and partitioning water and solute flow; (3) filtering and buffering, degrading, immobilizing, and detoxifying organic and inorganic materials, including industrial and municipal by-products and atmospheric deposition; (4) storing and cycling nutrients and other elements within the earth’s biosphere; and (5) providing support of socioeconomic structures and protection for archeological treasures associated with human habitation. (Seybold et al, 1998).

Why optimal soil functionality in vineyard is important

Of course, these fore-mentioned soil functions apply to vineyards, especially regarding biomass production (grape), water and nutrients regulation, carbon storage and biodiversity. But in viticulture, as an agro-ecosystem looking for high quality production, soil is of particular importance. Combined with other environmental conditions and viticultural practices, it is indeed one of the main factors of terroir, giving to each wine its quality and specificity.
**Part I: Soil monitoring**

**Objectives**

In experimental projects linked to soil, the differences of soil traits between different areas of the fields should be investigated to characterize them and their spatial variability. In this purpose, profile digging allow to obtain detailed information about soil physical, chemical and hydrological properties, grapevine roots description, and to sample for further laboratory analyses *(Methodological sheet n°1: soil profile description and sampling)*.

**Analyses to perform**

The samples collected in the profiles can be analyzed for the following standard laboratory analysis:

<table>
<thead>
<tr>
<th>Analysed parameter</th>
<th>Description of the analysis method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk density assessment</td>
<td>Methodological sheet n°2</td>
</tr>
<tr>
<td>Water retention curve and available water capacity assessment</td>
<td>Methodological sheet n°3</td>
</tr>
<tr>
<td>Standard physical and chemical analysis:</td>
<td></td>
</tr>
<tr>
<td>- Texture</td>
<td></td>
</tr>
<tr>
<td>- pH-water</td>
<td></td>
</tr>
<tr>
<td>- Electrical conductivity (EC)</td>
<td></td>
</tr>
<tr>
<td>- Total carbonates (equivalent CaCO₃)</td>
<td></td>
</tr>
<tr>
<td>- Total organic carbon content (TOC)</td>
<td></td>
</tr>
<tr>
<td>- Total nitrogen content (TN)</td>
<td></td>
</tr>
<tr>
<td>- Cation exchange capacity (CEC) and exchangeable bases (K, Na, Mg, Ca)</td>
<td>Methodological sheet n°4</td>
</tr>
</tbody>
</table>

Some of these parameters can then be monitored throughout the experimentation to evaluate their evolution and the effects of tested modalities (practices, treatments...) where appropriate. During ReSoLVe project, this monitoring was focused on soil moisture, total organic carbon, total nitrogen and pH-water. Once a year, total and exchangeable copper can also be analysed at different soil depths.
Methodological sheet n°1: soil profile description and sampling

Materials needed

- Excavator
- Measuring tape
- Labels to indicate the code of the profile
- Camera
- Small short-handled shovel
- Plastic bags
- Marker pen
- Net with regular squares (15x15cm)

When and where to dig a profile and sample

The soil profile description should be done in spring or early summer, when the soil is slightly humid. To describe the grapevine root distribution, the profile should be dug around 30 cm far from the vine row.

How to proceed

The soil profile might by dug at a minimum depth of 1 m. A vertical surface of the profile must be cleaned and leveled to take a picture. On the picture, a measuring tape and a sign with the code of the profile should be clearly visible. The soil profile has to be described following the methods of FAO (Figure 1): www.fao.org/3/a-a0541e.pdf.

For grapevine roots description, a lateral wall of the profile should be dug at about 10 cm far from the vine row. The surface of the wall must be roughly cleaned from large clods and stones. A net with regular squares (15x15 cm in size) must be fastened to the profile wall under the vine row. For each horizon, fine and coarse roots per square meter should be counted. Fine (feeder) roots: < 2 mm. Coarse roots: > 2 mm.

Soil sampling for laboratory analyses. Undisturbed samples of each soil horizon should be collected in a bag with code. The amount of soil collected for each sample should be around 500 g. In case you have several profiles in the same plot, you can select only few benchmark profiles representative of the different areas of the field where to sample if appropriate. Soil samples should be dry at a maximum temperature of 40 °C.

A database of profile description, soil analyses and soil monitoring should be implemented.
Possible interpretations and conclusions

Since the vineyard soils are usually deep cultivated before vines plantation, the soil horizons are usually mixed and disturbed. For this reason, the individuation of genetic horizons and pedogenetic features are not always easy. The most important features to check in detail during soil profile description in vineyard are the following:

- **Presence of limiting horizons**: one or more horizons that limit grapevine rooting can be observed in the profile. The rooting obstacle could be due to compaction, poor aeration and/or waterlogging, very high coarse fragments fraction (> 50-60%), very high content of calcium carbonate, high salinity, nutrient lack or imbalance. If the limiting horizon is deeper than 80-100 cm, usually it does not have negative effects on grapevine growth, but if it is shallower than 70-80 cm can create serious troubles for grapevine nutrition.

- **Discontinuities**: in vineyard soils, discontinuities are quite common and they are often due to soil truncation and/or accumulation because of the land levelling or erosion.

- **Soil internal drainage**: soil with slow drainage and temporary waterlogging is characterized by redoximorphic mottles, which are spots or blotches of greyish (reducing) and/or yellowish-reddish (oxidizing) colour. Higher is the abundancy of greyish mottles, longer is the waterlogging period during the year. In addition, reducing conditions of the soil can be also characterized by black iron-manganese concretions.

- **Soil structure**: it is the natural organization of soil particles into discrete soil units (aggregates or peds) that result from pedogenic processes. The aggregates are separated from each other by pores or voids. Compaction, scarce organic matter and biological activity can contribute to a scarce structure of the soil. The vine roots can have difficulty to develop in horizons with weak structure or massive.

**References**

**Methodological sheet n°2: bulk density assessment**

**Materials needed**

<table>
<thead>
<tr>
<th>For field work (sampling)</th>
<th>For laboratory work (analysis)</th>
</tr>
</thead>
<tbody>
<tr>
<td>▪ Hammer-driven double-cylinder sampler or metal cylinders with sharpened edge of known volume</td>
<td>▪ Oven (105°C)</td>
</tr>
<tr>
<td>▪ Hammer</td>
<td>▪ Electronic balance, 0.01g sensitivity</td>
</tr>
<tr>
<td>▪ Wood block</td>
<td>▪ 2-mm sieve</td>
</tr>
<tr>
<td>▪ Straight-edged knife</td>
<td></td>
</tr>
<tr>
<td>▪ Small short-handled shovel</td>
<td></td>
</tr>
<tr>
<td>▪ Plastic bags</td>
<td></td>
</tr>
<tr>
<td>▪ Labels</td>
<td></td>
</tr>
<tr>
<td>▪ Marker pen</td>
<td></td>
</tr>
</tbody>
</table>

**When, where and how to sample**

Soil bulk density (Db) is the mass of dry soil per unit of bulk volume, including the void space (Blake and Hartge, 1986). Bulk density is not an invariant characteristic for a given soil. Changes in soil volume due to changes in water content modify bulk density, particularly in swelling soils. Soil mass remains fixed, but the volume of soil may change as water content changes. For this reason, undisturbed soil samples for bulk density determination by the core method (Blake and Hartge, 1986) should be taken when the soil moisture content is near field capacity.

Basically, the core method consists in collecting, drying and weighing a soil sample of known volume. To do this, clean off the pit face or the field surface, and drive the double-cylinder sampler (Figure 2) into the soil; then, carefully remove the inner cylinder to preserve the sample volume. If a hammer-driven double-cylinder sampler is not available, a metal cylinder (ring) with sharpened edge of known volume may be driven into the soil using a hammer and a block of wood.

Extract the cylinder from the sampler, remove excess soil from the bottom using the knife, and transfer the soil sample in a plastic sealable bag. Properly label each sample to show basic information, e.g., project, site, date, plot, depth, and horizon. Collect at least three 100 cm³ volume samples from each soil horizon (or at 0-10 and 10-30-cm depth) to be able to quantify the variability of the system.

Where the soil is too loose (e.g., sandy soil), or skeleton is abundant and preclude the use of core samplers, to allow core sampling it is advisable the adoption of the excavation method (Grossman and Reinsch, 2002).

The excavation method involves digging out a small hole, then oven drying at 105°C and weighing the excavated soil. The volume of the excavation is determined by lining the hole with plastic film and filling it completely with a measured volume of water (or sand, or silicon beads).

![Figure 2. Hammer-driven double-cylinder sampler](image)
How to perform the analysis in laboratory

The dry bulk density (Db) is determined according to the core method (Blake and Hartge, 1986). Undisturbed soil samples are oven-dried at 105°C to constant weight. In order to correct the Db data for the presence of stones, the quantity of material >2 mm in diameter is determined by wet sieving. Each dry sample is washed on the 2-mm sieve; the material that remained on the sieve is collected and oven-dried at 105°C to constant weight. Bulk density is calculated as the mass of dry, coarse fragment-free soil per volume of the excavated soil, where volume is also calculated on a coarse fragment-free basis. Below is the formula used to calculate soil Db, reported to the nearest 0.01 g cm\(^{-3}\).

\[
Db = \frac{\text{Mass of fine material (g)}}{\text{Volume of fine material (cm}^3)}
\]

Possible interpretations and conclusions

High bulk density values are indicators of low porosity and strong compaction of the soil, aspects that adversely affect water retention capacity, root growth (Table 1) and movements of oxygen and water through the soil. As a consequence, soil compaction causes yield and vegetation cover reduction. By impairing water infiltration, compaction also causes the increase of runoff volume and soil erosion in sloping fields, and the occurrence of flooding in flat areas.

References

USDA-NRCS. Soil Bulk Density / Moisture / Aeration – Soil Quality Kit. Guides for Educators. [Link to USDA-NRCS webpage]

Annex

Table 1. General relationship of soil bulk density to root growth based on soil texture

<table>
<thead>
<tr>
<th>Soil texture</th>
<th>Ideal bulk densities for plant growth (g/cm(^3))</th>
<th>Bulk densities that affect root growth (g/cm(^3))</th>
<th>Bulk densities that restrict root growth (g/cm(^3))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sand, Loamy sand</td>
<td>&lt; 1.60</td>
<td>1.69</td>
<td>&gt; 1.80</td>
</tr>
<tr>
<td>Sandy loam, Loam</td>
<td>&lt; 1.40</td>
<td>1.63</td>
<td>&gt; 1.80</td>
</tr>
<tr>
<td>Sandy clay loam, Clay loam</td>
<td>&lt; 1.40</td>
<td>1.60</td>
<td>&gt; 1.75</td>
</tr>
<tr>
<td>Silt, Silt loam</td>
<td>&lt; 1.40</td>
<td>1.60</td>
<td>&gt; 1.75</td>
</tr>
<tr>
<td>Silty clay loam,</td>
<td>&lt; 1.40</td>
<td>1.55</td>
<td>&gt; 1.65</td>
</tr>
<tr>
<td>Sandy clay, Silty clay, Clay loam</td>
<td>&lt; 1.10</td>
<td>1.49</td>
<td>&gt; 1.58</td>
</tr>
<tr>
<td>Clay (&gt; 45% clay)</td>
<td>&lt; 1.10</td>
<td>1.39</td>
<td>&gt; 1.47</td>
</tr>
</tbody>
</table>

Source: USDA-NRCS
Methodological sheet n°3: water retention curve and available water capacity assessment

Materials needed

<table>
<thead>
<tr>
<th>For field work (sampling)</th>
<th>For laboratory work (analysis)</th>
</tr>
</thead>
<tbody>
<tr>
<td>▪ Metal cylinders (rings) with sharpened edge of known volume (70 mm diameter, 30 mm height) and plastic lids</td>
<td>▪ Sand box</td>
</tr>
<tr>
<td>▪ Hammer</td>
<td>▪ Pressure plate extractors</td>
</tr>
<tr>
<td>▪ Wood block</td>
<td>▪ Porous ceramic plates</td>
</tr>
<tr>
<td>▪ Straight-edged knife</td>
<td>▪ Compressor (20 bars)</td>
</tr>
<tr>
<td>▪ Small short-handled shovel</td>
<td>▪ Pressure regulation system</td>
</tr>
<tr>
<td>▪ Plastic bags</td>
<td>▪ Electronic balance, 0.01-g sensitivity</td>
</tr>
<tr>
<td>▪ Marker pen</td>
<td>▪ 2-mm sieve</td>
</tr>
<tr>
<td></td>
<td>▪ Oven (105°C)</td>
</tr>
</tbody>
</table>

When, where and how to sample

Soil samples are taken at 0-10 and 10-30 cm depth at the beginning and the end of the trial period. The samplings are always carried out during the Spring months, with soil moisture content near field capacity. Undisturbed soil samples are taken at each soil horizon during soil profiles description using metal cylinders (70 mm diameter, 30 mm height) with a sharpened edge. Cylinders are hammer-driven into the soil at the 0-10 and 10-30 cm depth and carefully removed with a shovel taking care to preserve soil structure and avoid compaction. The soil extending beyond each end of the cylinder was trimmed with a straight-edged knife, and the ring closed with a plastic lid. Three samples per plot and depth are taken at each sampling date.

How to perform the analysis in laboratory

Soil water retention curve is determined by sand box and pressure plate apparatus (Figure 3) on the undisturbed soil samples collected in the experimental vineyards.

![Figure 3. Sand box (left) and pressure plate extractor (right)](image)

In the pressure plate extractor method, positive pressure is applied to the air phase of soil samples. The water phase is in contact with atmospheric pressure via a fine-pored porous plate, over which the soil samples are drained (Dane and Hopmans, 2002). Place the rings on the ceramic plate and then in the pressure plate extractor. To provide good contact between rings and ceramic plate, cover ceramic plate with a thin layer (2 or 3 mm) of water. Close
container and secure lid. Apply the proper gauged air pressure. When water ceases to discharge from outflow tube, sample is at equilibrium. Remove the rings and record the weight.

Water content is measured at the 0 (saturation), -6, -10, -33, -600 and -1500 kPa matric potential. The water content at 0 (saturation), -6 and -10 kPa matric potential is measured on sand box (Clement, 1966), whereas further three retention measurements at the matric potentials of -33, -600 and -1,500 kPa are determined by means of pressure plate extractors (Klute, 1986). The moisture content at each matric potential is then expressed as percentage by weight of the dry soil ($\theta_d$).

The retention data at field capacity (FC) (-10 kPa) and wilting point (WP) (-1,500 kPa) were used to determine the available water capacity (AWC = FC - WP) (Table 2).

On the same soil samples, at the end of the analysis, soil bulk density was determined according to Blake and Hartge (1986), removing by wet sieving the contribution of skeletal (material >2 mm) and plant roots, which particle density was assumed equal to 2.65 and 0.70 g cm$^{-3}$, respectively. The bulk density values were then used to convert the gravimetric water content ($\theta_d$) data on a volumetric basis ($\theta_v$) by applying the equation (1) (Gardner, 1986).

$$\theta_v = \theta_d \frac{BD}{\rho_w} \quad (1)$$

where $\rho_w$ is the density of water.

Actually, such a procedure was required to run RETC software (van Genuchten, 1980), employed to determine the retention parameters of soil retention curve (Figure 4), by fitting measured water contents.

Table 2. Matching table for evaluation of the Available Water Capacity (AWC).

<table>
<thead>
<tr>
<th>Class</th>
<th>Values (% on volume basis)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Very low</td>
<td>&lt; 5</td>
</tr>
<tr>
<td>Low</td>
<td>5 - 10</td>
</tr>
<tr>
<td>Moderate</td>
<td>10 - 15</td>
</tr>
<tr>
<td>High</td>
<td>15 - 20</td>
</tr>
<tr>
<td>Very high</td>
<td>&gt; 20</td>
</tr>
</tbody>
</table>

References


Methodological sheet n°4: standard physical and chemical soil analysis

Materials needed

<table>
<thead>
<tr>
<th>For field work (sampling)</th>
<th>For sample preparation in laboratory</th>
</tr>
</thead>
<tbody>
<tr>
<td>▪ Shovel</td>
<td>▪ 2-mm sieve</td>
</tr>
<tr>
<td>▪ Auger</td>
<td>▪ 0.250-mm sieve</td>
</tr>
<tr>
<td>▪ Plastic bags</td>
<td>▪ Electronic balance, 0.01-g sensitivity</td>
</tr>
<tr>
<td>▪ Marker pen</td>
<td>▪ Oven (105°C)</td>
</tr>
<tr>
<td>▪ Meter</td>
<td>▪ Distilled water</td>
</tr>
<tr>
<td></td>
<td>▪ Dispersing solution (containing 2 g of sodium hexametaphosphate/L)</td>
</tr>
<tr>
<td></td>
<td>▪ Hydrogen peroxide (H₂O₂) 30% vol.</td>
</tr>
<tr>
<td></td>
<td>▪ Pipettes 25 cm³ capacity</td>
</tr>
<tr>
<td></td>
<td>▪ Cylinders - 500 cm³</td>
</tr>
<tr>
<td></td>
<td>▪ Hand stirrer</td>
</tr>
<tr>
<td></td>
<td>▪ Common laboratory equipment</td>
</tr>
</tbody>
</table>

Additional specific materials are indicated for each parameter hereafter.

When, where and how to sample

Samples can be collected either in soil profiles at each horizon or in the first 30 cm of soil for a more regular monitoring (especially for total organic carbon, total nitrogen and pH).

In this last case, each experimental plot or modality should be monitored during the main vine phenological phases (budburst, berry formation, pre-veraison, post-veraison) and sampled with a shovel in three randomly selected points at two depths 0-10 and 10-30 cm. The samples from the three sampling points are then mixed thoroughly in a bag with a code to provide a single composite sample.

How to perform the analysis in laboratory

Analysis standard procedures will be performed according to the World Reference Base for Soil Resources classification system (IUSS Working Group WRB, 2014). Full descriptions of the procedures can be found in Procedures for soil analysis (Van Reeuwijk, 2002) and the USDA Soil Survey Laboratory Methods Manual (Burt, 2004).

COMMON SAMPLE PREPARATION

Before analysis, soil samples must be air-dried or, alternatively, oven-dried at a maximum of 30°C. The fine earth fraction should be obtained by sieving the dry sample through a 2 mm sieve. Clods not passing through the sieve should be crushed (not ground) and sieved again. To this aim, a variety of manual or mechanical implements may be used. Gravel, rock fragments, etc. not passing through the sieve are collected and set aside.

TEXTURE (PARTICLE-SIZE) ANALYSIS

Soil texture refers to the relative size distribution of the primary particles in a soil. Particle size, using the USDA classification scheme, is divided into three major size classifications: sand (2.0-0.05 mm), silt (0.05-0.002 mm), and clay (<0.002 mm) (Gee and Bauder, 1986). The laboratory method for soil textural analysis accomplished by first dispersing the soil into individual primary particles, followed by fractionation and quantification of each particle-size interval by sieving or sedimentation (Kettler et al., 2001). The hydrometer and pipette methods are sedimentation procedures.
that are widely accepted as standard methods for particle-size analysis (Gee and Bauder, 1986). The procedures are applied to the fine earth (< 2 mm) fraction only.

Materials needed
- 2-mm sieve
- 0.250-mm sieve
- 0.050-mm sieve
- Glass cylinders (500 cm³ volume)
- Hand stirrer
- Pipette
- Calgon (Sodium hexametaphosphate) solution.
- Rubber hammer
- Horizontal agitator
- Oven (105°C)
- Electronic balance, 0.01-g sensitivity

The analysis by the pipette method (Figure 5) followed the standard procedure as described by USDA-NRCS (2004), consisting in the following phases:

i) weighing 10 g of air dried soil sample <2mm;
ii) soil dispersion with 10 cm³ of a Calgon (0.2% vol) solution;
iii) distilled water addition up to the final volume of 250 cm³;
iv) agitating the suspension with horizontal agitator for at least 12 hours (150 rpm); cleaning the suspension at 250 µm with distilled water;
v) topping up the passing fraction to the reference volume of 500 cm³ with distilled water;
vi) analyzing the soil suspension volume (25 cm³).

No pre-treatment for soil organic matter removal has been carried out on ReSolve project samples, since in this study the knowledge of the more "natural" particle-size distribution of the soil was of greater importance from a functional point of view. In that regard, Matthews (1991) assessed that the choice of including removal of organic matter, as well as that of carbonates and/or iron oxides, should correspond to the aim of the investigation and type of material to be analyzed.

The sedimentation analysis was carried out on a soil suspension passed through 250 µm sieve. The wet sieving procedure was employed to determine the sands larger than 250 µm; the wet sieving procedure was employed to determine the sands larger than 250 µm, but also the fine and very fine sand fractions, after that silt and clay analysis was completed.

Once the percentage of sand, silt and clay is known, the identification of the textural class is easily obtained through the textural triangle (Figure 6).
RESOLVE - Protocols for soil functionality assessment in vineyards

Figure 6. Textural triangle for soil texture analysis using the USDA classification scheme (USDA-NRCS, 2004).

References


AGGREGATE STABILITY

Materials needed

- 10.0, 4.75, 2.0, 1.0, 0.25, and 0.050 mm sieves
- Rubber hammer
- Vibrating sieve shaker
- Electronic-controlled sieving machine
- Numbered cans
- Digital balance with readability of 0.01 g
- Oven (105°C)
- Dispersing solution (containing 2 g of sodium hexametaphosphate/L)

To evaluate soil macroaggregate stability, the mean weight diameter (MWD) of water stable aggregates was determined by the procedure described by Kemper and Rosenau (1986). Undisturbed soil samples of the surface (0-10 cm) layer were collected in triplicate down to 0.1 m depth, air dried at room temperature, weighted and separated into different sized fractions (20.0-10.0, 10.0-4.75, 4.75-2.0, 2.0-1.0, 1.0-0.25, 0.25-0.05, <0.05 mm) by a vibrating sieve shaker. After drying, the samples were broken down into smaller mechanical aggregates using a rubber hammer. The sample, passed through a 10-mm mesh and retained at the 4.75-mm mesh, was used for analyses. The MWD was determined on 20 g of dry aggregates of less than -1,500 kPa water potential. The samples were directly soaked for 5 minutes on the top of a nest of sieves of 4.75, 2.0, 1.0, 0.25, and 0.050 mm immersed in water. We preferred this wetting procedure instead of capillary rise because, according to Legout et al. (2005), it should effectively mimic the breakdown mechanisms that aggregates experience under high intensity (>30 mm h-1) rainfall events. The nest of sieves and its content was then vertically oscillated in water by an electronic-controlled machine (Figure 7).
with a stroke of 4 cm per 10 minutes, at a rate of 30 complete oscillations per minute. The mass of oven-dried particles (105°C for 24 hours) in each sieve that resisted breakdown was determined. The mass of the fraction passing through the 0.050-mm sieve was thereby obtained by subtraction. The respective dry masses were used to compute the MWD according to Van Bavel (1949), as follows:

\[ MWD = \sum_{i=1}^{n} W(i) X_i \]

where:
MWD is the mean weight diameter (mm);
\( X_i \) is the arithmetic mean diameter of the \( i \) and \( i-1 \) sieve openings (mm);
\( W(i) \) is the proportion of the total sample mass (corrected for sand and gravel) occurring in the fraction (dimensionless);
\( n \) is the number of size fractions, including the one that passes through the finest sieve (in this case 6).

![Figure 7. The electronic-controlled machine for aggregate stability analysis by wet sieving.](image)

References

PH AND ELECTRICAL CONDUCTIVITY (EC)

Materials needed
- pH-meter
- Conductivity meter
- pH standard solutions
- Common laboratory equipment

The soil pH is measured potentiometrically in the supernatant suspension of a soil:water mixture (fine earth fraction). If not stated otherwise, soil and water are in a ratio of 1:5. Based on the typical procedure, 20 g
fine earth are placed into a 100 ml wide-mouth type bottle with 100 ml water and shaken for 2 hours. The pH is measured by immersing a combination electrode in the upper part of the suspension. For the EC measurement, it is recommended to let the soil:water mixture stand for another 2 hours after pH reading, then to filter it through a hardened low-speed filter paper. If the initial filtrate is turbid, it must be re-filtered using the same filter paper before EC measurement.

**TOTAL CARBONATES**

Materials needed
- Calcimeter (Dietrich-Fruhling equipment or others)
- HCl
- Thermometer for room temperature measurement
- Barometer
- Common laboratory equipment

The total amount of carbonate in the soil is most commonly reported as total CaCO$_3$ equivalent (the analysis is not selective for calcite) and is usually measured by the gas-volumetric method. To this aim, different calcimetry equipments may be used (Dietrich-Fruhling, Scheibler, Bernard, De Astis, etc). Basically, soil samples are treated with a HCl solution; the evolved CO$_2$ is measured manometrically and the amount of CaCO$_3$ equivalent is then calculated. According to the Dietrich-Fruhling calcimeter procedure, 0.5 to 5.0 g of the “fine earth” sample (depending on the presumed carbonate amount) is treated with 10 mL of 1:1 (v:v) HCl solution. The container used for treatment is manually shaken until complete CO$_2$ development. Room temperature and pressure must be recorded during the analysis to allow standardization at 0°C and 760 mm Hg of the CO$_2$ volume developed. The final calculation takes into account the standardized CO$_2$ volume (V0) and the sample weight (P), by the formula:

$$\text{CaCO}_3 \,(\text{g/kg}) = \frac{(V0 \times 0.0044655 \times 1000)}{P}$$

Alternatively, the total CaCO$_3$ equivalent may be determined by the “acid neutralization” method (sample treatment by dilute HCl and titration of the residual acid).

**TOTAL ORGANIC CARBON (TOC) AND TOTAL NITROGEN (TN)**

Materials needed
- Carbon/nitrogen elemental analyzer
- Tin-foil and Ag-foil capsules
- HCl
- Analytical balance (5 decimal points)
- Thermostatically controlled heating plate
- Common laboratory equipment

The recommended procedure for soil TOC and TN determination is based on dry-combustion by CN elemental analyzer. The analysis is applied to finely ground sub-samples (< 0.5 mm), obtained from the fine earth by means of ball mills or other grinding equipment. Particular care must be paid to the sample homogenization, in order to ensure representativeness in the small amounts of soil used for analysis (20-60 mg). In carbonate-containing soils (total C = organic C + mineral C), TOC determination by dry-combustion requires that all mineral C is previously removed from the sample. This is achieved by weighing the sample (20 to 30 mg) in a silver capsule and treating it by a step-wise addition of 10% HCl, until complete removal of carbonates. After the acid treatment, the sample is dried and analyzed for total carbon (total C = TOC). It is furthermore recommended that in calcareous soils, the total N content is determined separately on untreated samples, because the HCl treatment could alter soil N concentration.
Soil samples without carbonates are directly analyzed for TOC and TN by weighing 40-60 mg of soil in tin capsules. The instrument performance as well as calibration must be regularly checked, and a minimum of three aliquots for each sample (replicates) should be analyzed to check the consistency of results. As an alternative to the dry-combustion technique, the Walkley Black and the Kjeldahl methods are suggested for TOC and TN analyses, respectively. Differently from dry-combustion, the organic C oxidation through the Walkley Black reaction is incomplete and an empirical correction factor of 1.3 is applied in the calculation of the result. Compared to dry-combustion, the Walkley-Black and the Kjeldahl procedures are more time-consuming and require extensive use of glassware. They, moreover, involves some health hazards and produces considerable amounts of polluting wastes.

**CATION EXCHANGE CAPACITY (CEC) AND EXCHANGEABLE BASES (K, NA, MG, CA)**

**Materials needed**
- Ammonium acetate, 1 M, pH 7 solution
- Sodium acetate 0.9 M/sodium chloride 0.1 M, pH 7 solution
- Ethanol 96%
- Centrifuge tubes
- Mechanical shaker
- Whatman n. 42 filter paper
- Commercial standard solutions for AAS (K, Na, Mg, Ca)
- Atomic Absorption Spectrophotometer
- Commonly used laboratory equipment

The reference procedure for the analysis of CEC and exchangeable bases is the pH 7.0-buffered ammonium acetate (NH₄OAc) method. Percolation in tubes may be replaced by shaking in centrifuge tubes. As reference for detailed laboratory operations, the NaOAc pH 8.2 Centrifuge Method (5A2a) by USDA (Kellogg Soil Survey Laboratory Methods Manual, Version 5.0, issued 2014) may be used. Summarizing, the sample is shaken 3-4 times in a centrifuge tube with a 1 M pH 7.0 NH₄OAc solution (5 g soil + 33 mL). The supernatants are collected together for the determination of the exchangeable bases. The sample is subsequently washed with ethanol and shaken 3-4 times with a pH 7.0 sodium acetate solution (33 mL). The excess salt is then removed with ethanol and the adsorbed Na is exchanged by repeated shakings with the pH 7.0 NH₄OAc solution (33 mL). The supernatants are pooled together, and the resulting solution is analysed by flame atomic absorption spectrophotometry (FAAS) for Na concentration, which is used for the calculation of soil CEC.

In order to quantify the exchangeable bases, the extract collected from the NH₄OAc sample treatments is made up to volume, filtered and subsequently diluted so that the analyte concentrations meet the calibration ranges (usually, 0-2.5-5-7.5-10 mg/L for both K and Na; 0-5-10-15-20-25 mg/L for Ca; 0-0.5-1.0-1.5-2.0-2.5 mg/L for Mg). The concentrations of K, Na, Mg and Ca in the solutions are measured by FAAS, after calibration of the instrument. For spectrophotometer set-up and operation, it is recommended to refer to the manufacturer's manual.

Calibration standard solutions may be prepared either for a single element or for several elements (e.g., K+Na and Mg+Ca). They should have the same NH₄OAc composition as the sample solutions.

Moreover, in order to overcome common interferences occurring in the flame, the addition of specific suppressors to both calibration and sample solutions is required (Caesium at a final concentration of 1 mg/mL for K and Na measurement; Lanthanum at a final concentration of 10 mg/mL for Ca and Mg measurement). The instrument readings for analyte concentration are in mg L⁻¹. These are then converted to (cmol (+) kg⁻¹) or mg kg⁻¹.

The percentage base saturation (BS%), used as criterion in soil classification, is defined as the fraction CEC occupied by basic cations: BS% = [(K+Na+Mg+Ca)/CEC]*100.
Possible interpretations and conclusions

The pH 7.0 NH₄OAc procedure often leads to overestimation of CEC in acid soils, especially for soils with significant variable charges. Moreover, it easily introduces errors when applied to calcareous soils, due to the fact that high amounts of Ca are readily dissolved from carbonates at pH 7.0 and interfere with the CEC determination.

When CEC is not a diagnostic criterion for soil classification, e.g. saline and alkaline soils, the CEC may be determined at pH 8.2 (IUSS Working Group WRB, 2014). Because of the dissolution of carbonates, the amount of Ca extracted from calcareous soils often greatly exceeds the exchangeable Ca. That’s why in calcareous soils the exchangeable Ca is routinely determined by the difference between the CEC and the sum of the other exchangeable bases (Mg+K+Na), and the base saturation is set to 100%.

Potassium and magnesium are two of the major nutrients for grapevine. Their exchangeable forms are in equilibrium with the soluble forms and contribute predominantly to the available pool for plants. For a correct interpretation of soil K and Mg values, they should be considered not only in absolute terms, but also in terms of the ratio of one to the other. In fact, an unbalanced Mg to K ratio could result in deficiencies in one element or the other, because of antagonism in the root absorption. On average, the Mg/K ratio is considered to be “very low” when less than 0.5 (very likely induced Mg deficiency), “low” to “slightly low” between 0.5 to 2.0, “optimal” between 2.0 and 6.0, “slightly high” between 6 and 10, “high” when higher than 10.0 (very likely induced K deficiency) (Fregoni, 2005; Sbaraglia and Lucci, 1994).

References


IUSS Working Group WRB, World Reference Base for Soil Resources (2014). International Soil Classification System for Naming Soils and Creating Legends for Soil Maps; World Soil Resources Reports No. 106; Food and Agriculture Organization (FAO): Rome, Italy.


Part II: Monitoring of soil ecosystem service provision and providers

Objectives

The level or intensity of soil services can be modified by soil management or restoration treatments. These changes are mainly associated with the quantity and the quality of organic matter in the first soil layers. Therefore, the analytical methods to monitor soil functionality in vineyards mainly focus on soil organic matter turnover and enzymatic activity. A list of selected service providers is proposed below, as indicators of high soil functionality and biodiversity.

Aims of ecosystem services and providers monitoring in soil are mainly to:
- Quantify the major soil functions linked to soil biota (soil organic matter turnover and nutrient recycling).
- Evaluate the abundance and the diversity of main soil service providers (earthworms, microarthropods or mesofauna, nematodes, fungi and bacteria).

Analyses to perform

Different techniques are available from literature:

<table>
<thead>
<tr>
<th>Evaluated parameter</th>
<th>Description of the assessment method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Organic matter breakdown using Tea-Bag Index</td>
<td>Methodological sheet n°5</td>
</tr>
<tr>
<td>Earthworm monitoring</td>
<td>Methodological sheet n°6</td>
</tr>
<tr>
<td>Enzymatic activity</td>
<td>Methodological sheet n°7</td>
</tr>
<tr>
<td>Soil respiration</td>
<td>Methodological sheet n°8</td>
</tr>
<tr>
<td>Soil microbial biomass</td>
<td>Methodological sheet n°8</td>
</tr>
<tr>
<td>Microbial communities through DNA extraction</td>
<td>Methodological sheet n°9</td>
</tr>
<tr>
<td>Soil mesofauna using Berlese extractors</td>
<td>Methodological sheet n°9</td>
</tr>
<tr>
<td>Nematodes monitoring</td>
<td>Methodological sheet n°10</td>
</tr>
</tbody>
</table>
Methodological sheet n°5: organic matter breakdown assessment using Tea-Bag Index

Materials needed

<table>
<thead>
<tr>
<th>For field work</th>
<th>For laboratory work</th>
</tr>
</thead>
<tbody>
<tr>
<td>▪ Lipton Green Tea Sencha</td>
<td>▪ Drying stove (max 60°C)</td>
</tr>
<tr>
<td>▪ Lipton Rooibos and Hibiscus Tea</td>
<td>▪ Scale with 0.001 g precision</td>
</tr>
<tr>
<td>▪ Shovel</td>
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<tr>
<td>▪ Marker pen</td>
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<tr>
<td>▪ Sticks</td>
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</tbody>
</table>

How to proceed

The protocol is based on the methodology proposed in 2013 by Keuskamp et al. and available on http://www.teatime4science.org/method/stepwise-protocol/. This technique consists of burying Lipton pyramid tea-bags for 3 months in the soil, twice per year (1-November to January, 2-April to June). Two types of teas are used, Green tea and Rooibos tea, the main difference being in their contrasting decomposability.

1. Measure the initial weight of the tea bag (.000 g).
2. Open a few bags and measure the bag weight without content (this is approx. 0.283 g).
3. Mark the tea bag codes on the white side of the label with a permanent black marker.
4. Bury the teabags in 8 cm-deep separate holes while keeping the labels visible above the soil (Figure 8), or otherwise tie the tea bags to a marked stick, to be able to find them back. Notice that tea bags could move in the soil due to soil erosion by water, therefore it is necessary to tie them to some hardly removable support.
5. Note the date of burial, geographical position, ecotype and experimental conditions of the site.
6. Recover the tea bags after approximately 90 days
7. Remove adhered soil particles, by gently tapping off the soil on the outside of the bags. Dry in a stove for 48h at 70 °C (not warmer!). Do not use water to remove the soil particles, because that invokes extra loss of material from the bag.
8. Remove what is left of the label but leave the string and weigh the bags (.000 g). Part of the label is paper and decomposes at a different rate as tea. Weight loss of the label may cause unwanted error in the measurement.
Possible interpretations and conclusions

The 2 types of tea show different values of carbon and nitrogen contents (Keuskamp et al. 2013) and decomposition dynamics. The C/N ratio is approximatively 12 for green tea whereas it is higher than 42 for rooibos tea. Rooibos tea decomposition is consequently slower and still in process after 90 days. Based on the weight of tea bags after exposure in the fields, we can calculate a TBI value, i.e., Tea Bag Index, which includes a degradation coefficient k and a stabilization factor S.

Moreover, the rate of decomposition of teas depends on soil parameters, mainly humidity, temperature, structure, compaction, and of course as a consequence, on soil biodiversity.

The comparison of k and S values allows to compare the quality and biological activity between different soils or to evaluate the effects of soil management practices.


References


[http://www.teatime4science.org/method/stepwise-protocol/]
Methodological sheet n°6: earthworms monitoring

Materials needed

<table>
<thead>
<tr>
<th>For field work (sampling)</th>
<th>For laboratory work</th>
</tr>
</thead>
<tbody>
<tr>
<td>▪ Mustard (Amora® Fine et Forte)</td>
<td>▪ Scale with g precision</td>
</tr>
<tr>
<td>▪ 40x40 cm frame</td>
<td></td>
</tr>
<tr>
<td>▪ Shovel</td>
<td></td>
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<tr>
<td>▪ Boxes</td>
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<td>▪ Marker pen</td>
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</tbody>
</table>

When and where to sample

Earthworms should be monitored once per year in spring (April-May), at temperature between 6-15°C, no frost or drought, with presence of rainfall in the week prior to sampling and soil moisture around 10 to 25%.

How to proceed

To assess earthworm populations, mustard extraction method will be used (Lawrence and Bowers 2002):
- Mix 90 g of hot mustard powder (Amora® Fine et Forte) with 100 ml of water in a closed bottle and allowed to sit for at least 1 h. “Amora” can be replaced by other hot mustard powder, but the final allyl isothiocyanate in the water solution should be about 0.025 g/l.
- Immediately prior to sampling, mix the mustard paste in a sprinkling can of 6 l of water.
- Place, at each sample site, a 40x40 cm metal, wooden or plastic frame on the ground and clear all the vegetation and leaves within it.
- Pour half of the mustard solution within the frame and a bit around, repeating the application a second time 10 min later.
- Take only the individuals emerging into the frame during 10 min after each mustard application.
- Place the earthworms in an identified box.
- Dig the soil within the 40x40 cm to a depth of 25 cm and examine for the remaining earthworms.
- Weigh the identified earthworms.

References

**Methodological sheet n°7: enzymatic activities assessment**

### Materials needed

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<tr>
<th>For field work (sampling)</th>
<th>For laboratory work</th>
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</thead>
<tbody>
<tr>
<td>▪ Shovel</td>
<td>▪ 2mm sieve</td>
</tr>
<tr>
<td>▪ Plastic bags or boxes</td>
<td>▪ Microplates</td>
</tr>
<tr>
<td>▪ Marker pen</td>
<td>▪ Flasks</td>
</tr>
</tbody>
</table>

### When, where and how to sample

Enzymatic analysis should be carried out once per year in spring time on soil sampled at two depths: 0-10 and 10-30 cm.

### How to proceed

The samples should be air-dried (max 30°C), sieved at 2 mm and kept at room temperature until analyzed. Enzyme activity is measured according to the methods of Marx et al. (2001) and Vepsäläinen et al. (2001), based on the use of fluorogenic methylumbelliferyl (MUF)-substrates. Soil is analysed for N-acetyl-β-glucosaminidase (NAG), β-glucosidase (BG), butyrate esterase (BUT), acid phosphatase (AP), arylsulphatase (ARYL), β-xilosidase (XYL), cellulose (CELL) and acetate esterase (AC) activity using methylumbelliferyl (MUF) conjugated surrogate substrates (Sigma, St Louis, MO, USA). Briefly, 2 g of soil sample are weighed into a sterile jar and incubated for 24 hours at 20% soil moisture. A homogenous suspension is obtained by homogenizing samples with 50 mL deionized water with UltraTurrax at 9600 rev / min for 3 min. Aliquots of 50 µL are withdrawn and dispensed into a 96 well microplate (3 analytical replicates/sample/substrate). 50 µL of Na-acetate buffer pH 5.5 are added to each well. Finally, 100 µL of 1 mM substrate solution are added giving a final substrate concentration of 500 µM. Fluorescence (excitation 360 nm; emission 450 nm) is measured with an automated fluorimetric plate-reader (Fluoroskan Ascent) after 0, 30, 60, 120, and 180 min of incubation at 28 °C and enzyme activity will be expressed as nmol MUF g⁻¹ h⁻¹.

The order of magnitude of the values obtained for the different enzymatic responses varies considerably depending on the specific activity being determined, thus leading to some enzyme having more weight than others. To resolve this problem, the sum of the percentage of the maximum value found for a specific enzymatic response across all enzymes for a soil, was used for the calculation of the sum of enzymes (SUM). From this percentage of maximum enzyme activities, the Simpson-Yule index was calculated following the equation \( E = 1/\sum p_i^2 \), as indicated by Bending et al. (2004), where \( p_i \) is calculated as a proportion of enzymatic responses summed across all substrates for a soil.

Discriminant function analysis (DFA) is performed using soil enzymatic activities as grouping variables for soil and fractions. Squared Mahalanobis distances between group centroids are determined. Two significant discriminatory roots are derived and the results of DFA are graphically presented in two dimensions.

### Possible interpretations and conclusions

Soil enzyme activities are widely used as sensitive indicators of changes in soil functioning and health, contributing to biogeochemical cycling, organic matter transformations and nutrient availability (Badiane et al., 2001; Vepsäläinen et al., 2001). Measuring the activity of several soil enzymes is useful to understand...
the organic matter turnover and the availability of inorganic nutrients, providing indications on the function and quality of an ecosystem. The studied group of soil enzymes are hydrolases, which are involved in the main biogeochemical cycling of elements and release C compounds as well as N, P and S. Among hydrolytic enzymes, β-glucosidase, α-glucosidase and β-cellobiohydrolase activities play a role in cellulose and starch degradation; phosphatases and arylsulfatase are involved in soil organic P and S mineralization; N-acetyl-β-glucosaminidase is involved in chitin degradation, a major source of mineralizable N in soil; N-acetyl-β-glucosaminidase and arylsulphatase activities are also considered as indirect indicators of the presence of fungal biomass because sulphate esters (substrates of arylsulphatase) are only present in fungal cells and chitin is the main constituent of fungal cell wall tissue. Hydrolytic enzymes have been frequently used as indicators of changes in quantity and quality of soil organic matter.

References

Annex
- Table of substrates

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Storage</th>
<th>Quantity (mg)</th>
<th>Solvent (µl)</th>
<th>Buffer (ml)</th>
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<tr>
<td>MUF-cellobioside</td>
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- Example of microplate

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<thead>
<tr>
<th>Substrate 1</th>
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<td>Buffer (µl)</td>
<td>Substrate (µl)</td>
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Methodological sheet n°8: assessment of soil respiration, microbial biomass and communities

Materials needed

<table>
<thead>
<tr>
<th>For field work (sampling)</th>
<th>For laboratory work</th>
</tr>
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<tbody>
<tr>
<td>▪ Shovel</td>
<td>▪ Beckers</td>
</tr>
<tr>
<td>▪ Plastic bags</td>
<td>▪ NaOH, HCl, K$_2$SO$_4$, ethanol-free CHCl$_3$</td>
</tr>
<tr>
<td>▪ Marker pen</td>
<td>▪ Thermo Flash 2000 CN soil analyser</td>
</tr>
<tr>
<td></td>
<td>▪ Mechanical shaker</td>
</tr>
<tr>
<td></td>
<td>▪ Whatman n. 42 filter paper</td>
</tr>
<tr>
<td></td>
<td>▪ DNeasy PowerLyzer™ PowerSoil® Kit</td>
</tr>
<tr>
<td></td>
<td>▪ Primers for V6-V8 region of bacterial 16S rDNA</td>
</tr>
<tr>
<td></td>
<td>▪ Gel electrophoresis material</td>
</tr>
<tr>
<td></td>
<td>▪ BIONUMERICS software</td>
</tr>
</tbody>
</table>

When, where and how to sample

Samples should be collected in spring (immediately before blooming) at two depths: 0-10 and 10-30 cm. 5 to 10 soil cores (depending also from the block dimension) for a total amount of approximately 2 kg should be collected, pooled together and then air dried and sieved at 2 mm.

How to proceed

Soil Respiration

At least 2 replicates per treatment (three plots per treatment) should be done. The method to be followed is described below.

Place 25 g of sieved soil sample in a 100 ml becker within a 1L stoppered glass jars together with 3 ml of 1M NaOH in a plastic glass. Incubate at 30°C and determine the CO$_2$ evolved after 1, 3, 7, 10, 14, 21, 28 days (replacing NaOH after each determination) by titration of the exceeding NaOH with 0.1 M HCl (Badalucco et al., 1992). The CO$_2$ evolved during the 28th day of incubation will be used as the basal respiration value, while the Basal Respiration Rate is measured as CO$_2$ evolved in a defined time lapse.

Soil Microbial Biomass

At least 2 replicates per treatment (three plots per treatment) should be done. Microbial biomass carbon (MBC) and N (MBN) should be estimated following the Fumigation Extraction (FE) method (Vance et al., 1987): two portions of sieved soil (20 g each) should be incubated at 30°C for 24 hours in two different 50 ml becker; then the first one (not fumigated) should be immediately extracted with 80 ml of 0.5M K$_2$SO$_4$ by oscillating shaking at 200 rpm for 60 min and filtering with Whatman paper n. 42; the second one should be fumigated for 24 h at 25 °C with ethanol-free CHCl$_3$ in a vacuum dryer and then extracted as described above. Organic C and N in the extracts should be determined by dry combustion on a Flash 2000 CN soil analyser (Thermo Scientific) and subtraction of non-fumigated values from fumigated ones.

Microbial communities through DNA extraction

Samples are the same for the other analyses air dried and sieved at 2 mm and then kept at -20°C until analysed; alternatively, a portion of the collected soil samples, if well loose and not particularly clayey and compact, could be stored in a Falcon tube and kept frozen in dry ice during sampling, then stored at -20°C until analysed.
Soil DNA direct extraction is performed on 250 mg of soil with DNeasy PowerLyzer™ PowerSoil® DNA Isolation Kit (QIAGEN) by mean of FastPrep Instrument or MOBIO Power Lyzer or any equivalent soil bead beater instrument. Subsequently DNA is amplified with primers specific for V6-V8 region of bacterial 16S rDNA in the condition outlined in Castaldini et al. 2005 and included below, and the mixture of three amplicons for each DNA are collected and run in a DGGE (Denaturing Gradient Gel Electrophoresis) gel as a single lane. DGGE profiles are digitally processed with BIONUMERICS software version 4.6 (Applied Maths, St-Martens-Latem, Belgium): similarities between the banding patterns are analysed using Dice pairwise coefficients and hierarchical clustering of the Dice similarity matrix is determined using unweighted pair group method with arithmetic averages (UPGMA); finally a cluster diagrams is produced. The band profiles will be used in addition to count the number of bands to estimate richness of the most abundant bacterial populations in each sample and to calculate different biodiversity indexes (Shannon, Simpson; Pielou) (Boon et al., 2002; Lalande et al., 2013).

**Possible interpretations and conclusions**

Soil is a natural ecosystem whose energetic balance is determined by microbial biomass through respiration biogeochemical cycles. In agricultural ecosystems microorganisms play a key role in the transformation of nutritive elements and the maintenance of soil biological fertility. The Basal Respiration Rate is a measure of the microbial essential respiration involved in the total decomposition of organic matter, while the C extracted after fumigation may be used to estimate the soil microbial biomass. Finally, the metabolic quotient $q_{CO_2}$, that is the rate between basal respiration and microbial biomass C, can be calculated to evaluate the ecosystem stability (Anderson, 1994): a low $q_{CO_2}$ value indicate a higher level of stability, while a high level is generally associated with environmental stress.

The amplified DNA bands of each DGGE pattern give information about the bacterial diversity of a sample, as each band represent roughly a bacterial species, or more exactly, a bacterial phylotype or OTU (Operational Taxonomic Unit).

The UPGMA dendrogram of Dice pairwise similarity index of PCR-DGGE profiles illustrates separation of the profiles into clusters according to their percentage of similarity.

The number of bands estimate the species richness i.e the number of the most abundant bacterial populations in each sample.

The different diversity indexes evaluate the samples in terms of species diversity (Shannon Index that take in account the different bands and their relative abundance), presence of dominant species (Simpson Index), evenness of species (Pielou Index).

**References**


Annex

- Protocol for DNA amplification

<table>
<thead>
<tr>
<th>PCR Cycle</th>
<th>Quantity referred to 25 µl reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Step</td>
<td>Temp °C</td>
</tr>
<tr>
<td>1</td>
<td>95</td>
</tr>
<tr>
<td>2</td>
<td>94</td>
</tr>
<tr>
<td>3</td>
<td>55</td>
</tr>
<tr>
<td>4</td>
<td>72</td>
</tr>
<tr>
<td>5</td>
<td>Go to 2</td>
</tr>
<tr>
<td>6</td>
<td>72</td>
</tr>
<tr>
<td>7</td>
<td>4</td>
</tr>
<tr>
<td>8</td>
<td>END</td>
</tr>
</tbody>
</table>

- Conditions for DGGE run after DNA amplification

<table>
<thead>
<tr>
<th>ACRYLAMIDE/Bis 40% 37,5:1 FINAL CONC IN GEL</th>
<th>6%</th>
</tr>
</thead>
<tbody>
<tr>
<td>DENATURATION GRADIENT (7M UREA – 40% FORMAMIDE)</td>
<td>50% - 60%</td>
</tr>
<tr>
<td>BUFFER</td>
<td>TAE 1x</td>
</tr>
<tr>
<td>RUN TEMPERATURE</td>
<td>60°C</td>
</tr>
<tr>
<td>VOLTAGE / AMPERAGE</td>
<td>75 V / 80 mA</td>
</tr>
<tr>
<td>RUN DURATION</td>
<td>17 h 30 min</td>
</tr>
</tbody>
</table>
Methodological sheet n°9: soil mesofauna using Berlese-Tullgren extractors

Materials needed

<table>
<thead>
<tr>
<th>For field work (sampling)</th>
<th>For laboratory work</th>
</tr>
</thead>
<tbody>
<tr>
<td>▪ Shovel</td>
<td>▪ Berlese-Tullgren extractors (funnel, heating lamp, 2 mm sieve, receptacle, alcohol 75%)</td>
</tr>
<tr>
<td>▪ Plastic bags or boxes</td>
<td>▪ Microscope</td>
</tr>
<tr>
<td>▪ Marker pen</td>
<td>▪ Tubes or jars</td>
</tr>
</tbody>
</table>

When, where and how to sample

Sampling should be done to a depth of 10 cm, 1 dm³ (10x10x10cm) soil, in the middle of the row, avoiding the areas compacted by the tractor wheels. 3 sub-samples should be collected per treatment and homogenized in a single sample. Soil samples can be collected at different seasons, particularly Autumn and Spring, when arthropod abundances and diversity are higher.

How to proceed

The extraction of microarthropods should be carried out using Berlese-Tullgren extractors (Figure 9):
- place a sieve (mesh of 2mm to allow the only passage of mesofauna) on the funnel.
- place above the funnel an incandescent lamp of 40 watts (about 20 cm from the sample).
- collect the mesofauna falling under the funnel in a preservative liquid consisting of alcohol 75° and Glycerine 5%.

After the extraction, microarthropods are counted and identified up to the order level, at stereomicroscope. The structure of microarthropod communities is determined by measuring the abundance of the arthropod groups and the respective relative frequencies (%). The biodiversity is evaluated by the following indexes: a) Shannon-Wiener, Simpson, Pielou indexes; b) ecological indicators such as soil biological quality index (QBS) and mites/springtails ratio.
Soil microarthropod communities are studied according to the procedure described by Parisi et al. (2005), who defined the QBS index. Generally, the application of mesofauna-based indicators of soil quality have been limited by the difficulties in classifying organisms to the species level. To overcome this limitation, Parisi et al. (2005) introduced a new approach, based on the use of a simplified Eco-Morphological Index (EMI) for the determination of an QBS of arthropods index. This index is based on the concept that the higher soil quality, the higher will be the number of microarthropod groups adapted to the soil habitat. The degree of microarthropod adaptation is defined by specific morphological characters; in particular, more adapted organisms will typically show reduced pigmentation and visual apparatus, loss or reduction of wings, reduced appendages and streamlined body form (Parisi, 2001). Each biological form (morpho-type) isolated from the soil is classified to the order level and is eco-morphologically scored. The scoring is proportional to organism adaptation degree, ranging from 1 (surface-living organisms) to 20 (deep-living organisms). The sum of all EMI values for a given soil sample provides its QBS index. Once determined, the QBS values can be used to define the QBS class, according to the classification given by Parisi et al. (2005). In particular, each class is identified by a number, ranging from 0 to 7, which increases with increasing complexity and adaptation degree of soil microarthropod communities.

**Possible interpretations and conclusions**

Berlese-Tullgren device combined with the use of morpho-types is a way to evaluate abundance and diversity of soil microarthropods while avoiding the complexity of taxonomy. Indeed, the use of indicators such as QBS index does not require a species-level identification. This index may be then considered as an appropriate tool for large-scale monitoring and based on a great number of samples. Soil microarthropods demonstrated to respond sensitively to management practices and be related to several beneficial soil functions and ecosystem services. It has for the moment to be developed and used in several systems and agrosystems in order to highlight its potential to characterize the effects of practices on biodiversity levels and associated services in soils.

**References**


https://en.wikipedia.org/wiki/Tullgren_funnel
Methodological sheet n°10: nematodes monitoring

Materials needed

<table>
<thead>
<tr>
<th>For field work (sampling)</th>
<th>For laboratory work</th>
</tr>
</thead>
<tbody>
<tr>
<td>▪ Hand auger</td>
<td>▪ Cotton-wood filter extractor</td>
</tr>
<tr>
<td>▪ Plastic bags</td>
<td>▪ Sieve 25 μm</td>
</tr>
<tr>
<td>▪ Marker pen</td>
<td>▪ Stereomicroscope</td>
</tr>
<tr>
<td></td>
<td>▪ Microscope</td>
</tr>
</tbody>
</table>

When, where and how to sample

At each location, 5 cores (0 to 30 cm layer) should be randomly sampled and then mixed to form one composite sample of 500 cc of soil. Each soil sample should be placed in a plastic bag and stored at a 4°C cold chamber until use.

How to proceed

Free living and plant parasitic nematode should be isolated from 100 ml of each soil sample using the cotton-wood filter extraction methods (Oostenbrink, 1960). Nematodes are extracted for 48h at room temperature, approximately 25°C. Each nematode suspension is sieved through a 25 μm sieve and the nematodes are counted at 50x magnification. Nematodes are mounted on temporary slides and identified at higher magnification to genus or family level using keys from Mai et al. (1964), Bongers (1988) and Marinari-Palmisano and Vinciguerra (2014). Taxonomic families are assigned to a trophic grouping based on Yeates et al. (1993).

The characterization of nematode communities should be done using:

1. absolute abundance of individuals
2. richness determined by counting the number of family
3. Shannon-Wiener diversity index (H)
   \[ H = -\sum_{i=1}^{s} p_i \ln p_i \]
   where \( p_i \) represents the relative abundance of one given family;
4. Simpson index (D)
   \[ D = \frac{\sum_{i=1}^{s} n_i (n_i - 1)}{N(N - 1)} \]
   where \( s \) is the number of species, \( N \) is the total number of organisms, \( n_i \) is the number of organisms of a specie
5. the Maturity (MI) and Plant Parasitic (PPI) indices by Bongers (1990), calculated as the sum of the weighted relative abundance of families classified in the \( cp \) scale for free-living and plant parasitic nematodes (c, colonizer nematodes \( r \) strategy; p, persister nematodes \( k \) strategy)
6. the food web indicators (BI, basal index; EI, enrichment index; SI, structure index; CI, channel index), EI is calculate based on the weighted relative abundance of functional guilds that are responsive to nutrient enrichment in \( cp \) groups 1 and 2, while SI is calculated as the weighted relative abundance of functional
guilds responsive to physical disturbance in *cp* groups 3, 4, and 5. The CI is calculated as weighted ratio between fungal to bacterial feeding nematodes in *cp* groups 1 and 2 (Ferris et al., 2001). These indicators add information on functional guilds to develop food web.

**Possible interpretations and conclusions**

Shannon-Weiner and Simpson indices need a determination at species level to be representative. Generally, these indices underestimate their values because the specific identification is not always possible for nematodes.

Maturity and Plant Parasitic indices range from 1 (disturbed soil) to 4 (good soil quality) in a nematode community.

1 – indicator of organic pollution
2 – indicator of stress
3–4 – indicator of good soil quality

**Food web indices**: The combination between Enrichment index and Structure index highlights the soil conditions.

- **EI > 50 and SI < 50** values indicate high soil disturbance enrichment.
- **EI > 50 and SI > 50** values indicate from low to moderate soil disturbance.
- **EI < 50 and SI > 50** values indicate undisturbed soils.
- **EI < 50 and SI < 50** values indicate stressed soils.

**References**


Part III: Soil rhizosphere monitoring

Objectives

Soil rhizosphere contains various micro-organisms. By monitoring them, it is possible to (1) assess microbial diversity in general, (2) isolate culturable microbes among them and (3) evaluate extent of colonization of roots by arbuscular mycorrhizal fungi. These parameters can then be compared and studied in relation with soil functionality status or soil restoration treatments.

Analyses to perform

Analyses on soil rhizosphere micro-organisms can target:

1. Microbial diversity in general, by using a culture independent approach. Principle of analyses:
   - Obtain environmental DNA extracts from any microorganisms present in source sample (here rhizosphere soil or roots).
   - PCR amplify phylogenetic marker loci with universal primers.
   - Separate amplified products of DNA by exposing the mixture of amplicons to a gradient of denaturant in a polyacrylamide gel. Each electrophoretically separated band represents an individual bacterial / fungal species and band profiles represent bacterial / fungal communities.
   - Analyze the obtained number of bands per source sample to calculate diversity indexes.
   - Analyze pattern of obtained band profiles for comparing communities of bacteria / fungi from different source samples, e.g. in Dice cluster analyses.

2. Culturable microbes, by isolating colony forming units from agar media. Principle of analyses: isolate rhizosphere soil or root associated bacteria / fungi by inoculating serial diluted source material on agar medium.


<table>
<thead>
<tr>
<th>Evaluated parameter</th>
<th>Description of the assessment method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Assessment of communities and pure culture isolation of root- or rhizosphere-associated fungi and bacteria</td>
<td></td>
</tr>
<tr>
<td>- Assessing fungi and bacteria communities by generating fingerprints represented by PCR amplified phylogenetic marker genes</td>
<td>Methodological sheet n°11</td>
</tr>
<tr>
<td>- Pure culture isolation of soil rhizosphere and root-associated bacteria and fungi</td>
<td></td>
</tr>
<tr>
<td>Direct assessment of mycorrhizal infections in root samples</td>
<td>Methodological sheet n°12</td>
</tr>
</tbody>
</table>
Methodological sheet n°11: assessment of communities and pure culture isolation of root- or rhizosphere-associated fungi and bacteria

Materials needed

<table>
<thead>
<tr>
<th>For field work (sampling)</th>
<th>For laboratory work (sample preparation)</th>
</tr>
</thead>
<tbody>
<tr>
<td>▪ Shovel or minibagger</td>
<td>▪ 50 ml tubes, sterile</td>
</tr>
<tr>
<td>▪ Clean bags or boxes</td>
<td>▪ Sterile distilled water (SDW)</td>
</tr>
<tr>
<td>▪ Marker pen</td>
<td>▪ Triple layered cheesecloth</td>
</tr>
<tr>
<td>▪ Twig scissors</td>
<td>▪ Centrifuge, e.g., Biofuge Stratos, Heraeus instruments</td>
</tr>
<tr>
<td>▪ 70/96% for ethanol disinfection</td>
<td>▪ 75% ethanol, 1% NaClO (freshly prepared)</td>
</tr>
<tr>
<td>▪ Laboratory gloves</td>
<td>▪ Waring/lab blender, e.g., Ultra-Turrax T25 (IKA Labs)</td>
</tr>
<tr>
<td></td>
<td>▪ Particle separating sieves, e.g., Retsch</td>
</tr>
<tr>
<td></td>
<td>▪ Triple layered cheesecloth</td>
</tr>
<tr>
<td></td>
<td>▪ Sterile laboratory glassware</td>
</tr>
</tbody>
</table>

Additional specific materials are indicated for each analysis hereafter.

When, where and how to sample

Root collection

Collect soil profile exposed roots or roots from shovel-dug soil from 10–30 cm deep layers. Alternatively, roots can be sampled also from a depth of 30–60 cm for allowing comparisons of communities from roots collected from different depths. Digging of excavation holes should target a radius of 30–50 cm away from tree trunks in the inter-row. Fine roots smaller than 3 mm diameter plus systems of attached fine and feeder roots should be targeted. Roots can be cut off with twig scissors. An amount of ca. 100 g per sample is normally sufficient for the amount of required fine roots and rhizosphere soil. Wearing laboratory gloves and cleaning twig scissors etc. with ethanol avoids cross contaminations of samples. Carefully remove bigger soil clumps from roots and collect roots in bags or boxes.

How to proceed

COMMON SAMPLE PREPARATION

Sampling rhizosphere soil

Cut off terminal roots of ca. 1 mm or less and terminal roots in up to 2 cm long pieces collected in sterile 50 ml tubes (Falcon). Carefully/gently wash off rhizosphere soil with sterile distilled water (SDW) through triple layered cheesecloth and collect rhizosphere slurry into new 50 ml tube. Centrifuge at 2200 g for 5 min and discard supernatant.

Preparation of root pieces

Process pieces of roots by following principles described by Collado et al. (2007). Wash collection of 2 cm long root pieces at least 5x exhaustively with SDW. Each washing step is to be terminated by centrifugation, see above, and discarding the supernatant. Additional washing steps to be followed are in 75% ethanol for 1 min, 1% NaClO for 3 min, and 75% ethanol for 30 sec followed by 3 washing steps with SDW. Pulverize clean roots with waring / lab blender, e.g., Ultra-Turrax blender in water and retrieve 100–250 µm big pieces with Retsch sieves. Centrifuge at 2200 g for 5 min to discard supernatant.
ASSESSING ROOT- OR RHIZOSPHERE ASSOCIATED MICROBIAL COMMUNITIES (FUNGI AND BACTERIA) BY GENERATING FINGERPRINTS REPRESENTED BY PCR AMPLIFIED PHYLOGENETIC MARKER GENES

Materials needed: DNA extraction, PCR and DGGE equipped molecular lab facilities

Aliquot soil slurry or root pieces in appropriate amounts and extract DNA by using appropriate spin column-based nucleic acid purification kits such as NucleoSpin® (Macherey-Nagel), PowerSoil DNA isolation kit (MO BIO) by following vendors protocols. For amplifications of phylogenetic marker genes targeting bacteria, see the Methodological sheet n°8 of this document; for amplifications of phylogenetic marker genes targeting fungi, see, Anderson et al. (2003), which is a protocol based on a nested PCR system for amplifying internal transcribed spacer region 1 of the fungal ribosomal DNA gene cluster using primers EF4/ITS4 and GC-ITS1f/ITS2. For exposing PCR amplified mixtures of amplicons to a gradient of denaturant in a polyacrylamide gel (DGGE) and how to analyze DGGE generated community fingerprints, see Methodological sheet n°8; Castaldini et al (2005); Schroers et al (2018).

PURE CULTURE ISOLATION OF SOIL RHIZOSPHERE AND ROOT-ASSOCIATED BACTERIA AND FUNGI

Materials needed

- General microbiological lab facilities (autoclave, clean bench, Bunsen burner, microscopes, pipettes, inoculation needles/loops, etc.)
- Carboxymethyl cellulose (e.g., Sigma)
- Sterile distilled water (SDW)
- Antibiotics (including cycloheximide, Penicillin G, Streptomycin sulphate)
- Agar (e.g., BD Difco)
- Soil extract (DSMZ 12 medium)
- Nutrient broth (e.g., Biolife)
- Yeast extract (e.g., BD Bacto)
- Malt extract (e.g., Oxoid)

Generate inoculum for culture depending inventory as described previously in this methodological sheet. Dilute aliquots of pulverized root material in 0.1% carboxymethyl cellulose to a concentration of 10–20 pieces per 10 µl. Concentration can be checked by counting the number of pieces microscopically in aliquots of 10 µl pipetted on glass slides. Pipett aliquots of 60 µl on agar media in 9 cm Petri-Dishes. Disseminate inocula with drigalski spatel and incubate Petri dishes at room temperature.

Dilute aliquots of rhizosphere soil in SDW to 10⁻⁴ and pipett on agar as described for root pieces.

Agar media for bacteria:
- Soil extract agar (DSMZ 12 medium; http://www.dsmz.de) + 75 mg cycloheximide per liter agar medium
- Diluted nutrient broth (Biolife) (1 g per liter) plus 75 mg cycloheximide per liter agar medium

Agar media for fungi:
- Yeast extract / Malt agar (2 g yeast extract, 10 g malt per liter) plus 0.1 g chloramphenicol diluted in 10 ml absolute ethanol
- Potato dextrose agar diluted to 1/3 strength plus 0.0542 g streptomycin sulphate and 0.0121 g penicillin G diluted in 10 ml SDW

Inspect petri-dishes regularly to subculture individually growing colonies on appropriate media. Further attempts are then required to pure culture retrieved strains including the isolation of single spores (fungi) or streak plate method (bacteria).

Identification of retrieved taxa: Retrieved taxa can be identified based on phenotype or genotype approaches or a combination of both. Genus to species group level (sometimes species level) DNA barcodes are based on sequencing fragments of the ribosomal RNA gene cluster that are universally amplified with
primers in PCRs (Tsoktouridis et al 2014 for bacteria emphasizing 16S rDNA; Schoch et al 2014 for fungi emphasizing nuclear ribosomal internal transcribed spacers). Additional DNA barcodes including protein-encoding housekeeping genes are often required for species level identifications depending on the taxonomic group (EPPO 2016 for selected fungal and bacterial taxa). Unraveling the taxonomic position of organisms showing nucleotide substitutions when compared to sequences of reference material requires determination of patterns of relationships in phylogenetic analyses.

DNA barcode-based identification methodologies (exemplified in EPPO 2016) involve:
- Pure culture isolating of bacteria or fungal isolates
- Biomass generation for the extraction of DNA
- DNA extraction and purification
- PCR for amplification of any targeted DNA barcode
- Visualization of amplified product on agarose gels and product purification
- Sanger sequencing analysis of amplified product
- Assembly of raw sequence data
- Analysis of retrieved consensus sequences using Basic Local Alignment Search Tools (BLAST) and their comparisons with sequences from reference material or
- Phylogenetic analyses of retrieved consensus sequences

Possible interpretations and conclusions

The crucial step in microbial diversity assessments is sampling and initial sampling processions. The strategy described here aims at removing clonally identical microorganisms developing from propagules/spores a single individual representative / strain may produce in a specific environmental situation. For example, a fungal strain may inhabit soil through metabolically/physiologically active mycelium and perhaps a single sporulating structure producing thousands of spores at sampling time. The numerous washing steps implemented in the described procedures thus aim at removing the spores, each of which could produce a colony forming unit, while it is the mycelium colonizing the studied substratum that is measured. By generating “small” (here, 100–250 µm) pieces of plant material, it is purported that each of the pieces can be colonized only by a single microbial organism while a “large” piece (e.g., 2 cm long piece of roots) is colonized by numerous species, of which certain fast growing (measurable) opportunists may suppress the growth of numerous and (overlooked) others.

References


Methodological sheet n°12: direct assessment of mycorrhizal infections in root samples

Materials needed

<table>
<thead>
<tr>
<th>For field work (sampling)</th>
<th>For laboratory work</th>
</tr>
</thead>
<tbody>
<tr>
<td>▪ Shovel or minibagger</td>
<td>▪ Laboratory glassware</td>
</tr>
<tr>
<td>▪ Clean bags or boxes</td>
<td>▪ Tap water</td>
</tr>
<tr>
<td>▪ Marker pen</td>
<td>▪ KOH, 10%, freshly prepared</td>
</tr>
<tr>
<td>▪ Twig scissors</td>
<td>▪ Lactic acid, 5%, freshly prepared and 100%</td>
</tr>
<tr>
<td>▪ 70/96% for ethanol disinfection</td>
<td>▪ Aniline blue, 0.01% in pure lactic acid</td>
</tr>
<tr>
<td>▪ Laboratory gloves</td>
<td>▪ Dissection microscope</td>
</tr>
</tbody>
</table>

When, where and how to sample → For root sampling, see Methodological sheet n°11.

How to proceed

Root preparation includes (i) rinsing of roots in tap water, (ii) clearing of roots in 10% KOH for 24h at room temperature, (iii) acidifying roots in 5% lactic acid for 24h at room temperature and (iv) staining roots in 0.01% aniline blue in pure lactic acid for 24h.

Storing of roots prior to analysis can be done in pure lactic acid.

Mycorrhizal infections are scored following the procedure described by Trouvelot et al (1986), see figure 10.

The procedure involves counting of stained/ mycorrhized roots, what can be done under the dissection microscope. Roots are disseminated in a petridish and root points showing colonization by arbuscular mycorrhizal fungi are counted with a help of a raster scheme.

References

Part IV: Grapevine monitoring

Objectives

Grapevine yield and development is partly depending on soil functionality. That is why monitoring grapevine growth, water status, yield quantity and quality is also a way to highlight soil well or malfunctioning, to assess how soil status influences grapevine development and if restoration practices have an effect.

Analyses to perform

Many parameters can be monitored to evaluate grapevine development. The following ones are particularly interesting to assess for their link with soil status.
In addition, daily weather conditions and main phenological stages (date of budburst, flowering, veraison and harvest) can be registered to check for a possible difference in phenology and help in data interpretation.

<table>
<thead>
<tr>
<th>Evaluated parameter</th>
<th>Description of the assessment method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Assessment of grapevine vegetative growth by machine vision</td>
<td>Methodological sheet n°13</td>
</tr>
<tr>
<td>Assessment of grapevine water status</td>
<td>Methodological sheet n°14</td>
</tr>
<tr>
<td>Assessment of grape yield</td>
<td>Methodological sheet n°15</td>
</tr>
<tr>
<td>Assessment of grape composition</td>
<td>Methodological sheet n°16</td>
</tr>
</tbody>
</table>
**Methodological sheet n°13: grapevine vegetative growth assessment using machine vision**

**Materials needed**

<table>
<thead>
<tr>
<th>For field work (sampling)</th>
<th>For laboratory / office work</th>
</tr>
</thead>
<tbody>
<tr>
<td>▪ Camera of good resolution 10-14MP with a flash light and a tripod</td>
<td>▪ Matlab</td>
</tr>
<tr>
<td>▪ 2 painted canes</td>
<td></td>
</tr>
<tr>
<td>▪ White screen or sheet</td>
<td></td>
</tr>
</tbody>
</table>

**How to proceed**

An adequate and accurate assessment of the canopy status is the first step towards appropriate and effective canopy management, therefore an easy, non-invasive, robust method to evaluate the main features of a grapevine canopy is needed. The reference method for assessing canopy porosity, leaf density and fruit exposure is Point Quadrat Analysis (PQA) (Smart 1987). PQA is based on the insertion of a probe through the canopy of grapevines and counting the number and sections of the vine the probe comes into contact with: leaves, clusters, canes, or gaps. In addition to being subjective, PQA is labour and time consuming, and can potentially damage the fruit. The Televitis group of the University of La Rioja has established a new method to assess canopy status using machine vision (Diago et al 2018). The vines should be photographed under similar natural light conditions, and with diffuse light preferably. A standard digital reflex camera of good resolution 10MP-14MP with a flash light, in order to avoid shadows in the canopy, should be used. The camera should be mounted on a tripod set normal to the canopy 2 m away from row axis and at 1.00 m above the ground. In order to take the same image always, from the same reference point and the same size, the following easy steps could be taken.

1. Two woody or plastic canes could be placed (somehow stuck or driven into the soil) at both left and right end sides of the cordons of the canopy. Regarding the canes, it would be even better if they were painted in two colors (ie. red and white) at 10-15 cm intervals, so the dimensional reference will also exist in all the images. These canes should not be removed until the end of the season or imaging period. This way, the same references are used for the temporal series. Images should be taken always from the same point at the same distance. You can make some sort of "triangle" with two ropes or cotton thread so that the point from the camera to be set (with a tripod) does not change from one date to another across the whole period (see Figure 11).

![Camera shooting point](image)

**Figure 11. Canes or thin stakes at both ends of the vine**

2. To avoid the canopies of adjacent vines of rows behind the vine of interest being also photographed, thus interfering with the image, a white cloth, paper, screen... should be placed behind the canopy of the vine to be photographed. Field of view would be constant during the season. At the very early stages, only the first 50-60 cm of the vertical size of the picture will be filled with canopy but as the season progresses we will eventually fill up the whole vertical axe. All images should have the same number of pixels and field of view, and comparison among them for temporal series would be much easier (Figure 12).

3. Finally, avoid the automatic mode for capturing the pictures. You’d better set a manual configuration and make all pictures under the same set up of the camera.

4. Do not edit the pictures with photoshop or similar software, just rename the pictures indicating the grape variety, grapevine number, treatment and date of acquisition and it will be fine.
RGB images will be processed with Matlab, using a classification algorithm based on the Mahalanobis distance (Mahalanobis, 1936). This algorithm uses a known sample of values to classify an unknown group of pixels into classes based on a specific vector (the RGB color values of each pixel). The process involves two steps: the first step was the delineation of a region of interest (ROI), covering the 50 cm height, from the vine cordons, and delimited by the two conspicuous signs; the second step consists on a supervised manual selection of a representative number of points to be used as reference (also denoted as seed) for each class. The amount of pixels corresponding to exposed leaves, clusters, gaps and shoots will be calculated for each treatment in the defined ROI area of each image. Then, the ratio between the number of pixels of the leaf, cluster and gap classes and total number of pixels in the ROI, constitute the percentage of each feature respectively.

Possible interpretations and conclusions

The vineyard canopy status can be assessed using non-invasive RGB image analysis. Different features of the canopy, such as exposed leaf area, percentage of exposed fruit and canopy porosity can be determined by new non-destructive method. The developed RGB image-based methodology has enabled the assessment of the canopy status of grapevines growing under various soil conditions in an easy and non-invasively way. The methodology is inexpensive and can be adopted by viticulturists and researchers in viticulture to objectively assess the canopy status without damaging the plants, to help them in management decision making and to identify differences among treatments, respectively.

References

Methodological sheet n°14: grapevine water status assessment

Grapevine water stress can be measured through 2 methods: thermography and stem water potential.

Materials needed and prerequisites

<table>
<thead>
<tr>
<th>Leaf water potential assessment</th>
<th>IR-thermography assessment</th>
</tr>
</thead>
<tbody>
<tr>
<td>▪ Pressure chamber</td>
<td>▪ Infrared camera</td>
</tr>
<tr>
<td>▪ Aluminium-plastic bags</td>
<td></td>
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<tr>
<td>▪ Cutter or blade</td>
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</tbody>
</table>

How to proceed

Stem water potential.
Plant water status can be determined by stem water potential (\(\Psi_{\text{stem}}\)) at midday (between 13:00 to 15:00) on fully expanded leaves using a Scholander-type pressure chamber. The best period is during veraison. Measurements should be replicated at least 10 times in each modality. This protocol can be applied in the vineyard:

1. Bag one main leaf per plant of the mid-upper portion of the shoots into an aluminium-plastic bag for minimum 60 minutes prior to taking the readings.
2. Separate the entire leaf (petiole + blade) from the shoot and bring it immediately to the pressure chamber.
3. Cut the petiole tip just before insertion into the chamber.
4. Close the chamber, slightly open the gas (nitrogen) entry and turn it off as soon as you see a droplet of sap emerging from the petiole. Register the corresponding pressure indicated by the chamber and release the gas before opening the chamber for the following measurement.

IR-thermography.
Plant water status can be also assessed by thermal imaging using a handheld thermal camera. Each pixel in the thermal images is considered an effective temperature. The emissivity for vineyard canopy can be set at 0.96 (Jones 2004). Images will be taken at a distance of 1.5 m from the lateral canopy foliage. Minimum, maximum and mean temperatures, coefficient of variation (CV, %) and temperature range (maximum temperature difference) will be calculated for each plant (Pou et al. 2014). Wet and dry reference temperatures can be obtained in each image for the derivation of stress indices. \(T_{\text{wet}}\) and \(T_{\text{dry}}\) reference temperatures can be acquired using an evaposensor having two artificial leaves: a dry one (dry reference) and another one covered with a black cotton wick and receiving continuous water absorption for the wet reference. The average temperature (Tcanopy) is the statistic used to calculate two thermal indices, the conductance index (IG) and the crop water stress index (CWSI), using the reference temperatures (\(T_{\text{wet}}\) and \(T_{\text{dry}}\)). IG and CWSI will be calculated using the following expressions, respectively proposed by Idso et al (1981) and by Jones (2002).

\[
IG = \frac{(T_{\text{dry}} - T_{\text{canopy}})}{(T_{\text{canopy}} - T_{\text{wet}})} \quad (\text{Eq. 1})
\]

\[
CWSI = \frac{(T_{\text{canopy}} - T_{\text{wet}})}{(T_{\text{dry}} - T_{\text{wet}})} \quad (\text{Eq. 2})
\]

where: IG = conductance index; \(T_{\text{dry}}\) = temperature corresponding to the dry surface (\(^{\circ}\)C); Tcanopy = mean canopy temperature (\(^{\circ}\)C); \(T_{\text{wet}}\) = temperature corresponding to the wet surface (\(^{\circ}\)C), CWSI = crop water stress index.
Possible interpretations and conclusions
IR-thermography can be applied in sustainable viticulture for accurate vineyard water assessment and then for improvement of irrigation in viticulture, that is a very needed requirement in the current context of climate change and water scarcity.

References
**Methodological sheet n°15: grape yield assessment**

### Materials needed

- Containers or buckets
- Weighing machine
- Bags and marker pen

### When, where and how to sample

Yield components have to be assessed at harvest.

### How to proceed

Yield weight and number of clusters per vine will be measured and used to calculate cluster weight. In order to determine the berry weight, 10 clusters per plot have to be collected and stored overnight in a cool room (4°C). In the laboratory, a 100-berry sample will be separated from the rachis and weighed to estimate the berry weight. The number of berries per cluster will be then calculated from the cluster and berry weigh.

### Possible interpretations and conclusions

Among all collectable data from a vineyard, grapevine yield estimation outstands for its economical relevance and also for being key to help optimizing plant growth and to improve fruit quality. Yield variability within a vineyard has been proved to be high, here lies the true importance to assess theirs components.

### References


Methodological sheet n°16: grape composition assessment

Materials needed

<table>
<thead>
<tr>
<th>For field work (sampling)</th>
<th>For laboratory work</th>
</tr>
</thead>
<tbody>
<tr>
<td>▪ Bags and marker pen</td>
<td>▪ Pestle</td>
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<tr>
<td>▪</td>
<td>▪ Centrifuge</td>
</tr>
<tr>
<td>▪</td>
<td>▪ Refractometer</td>
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<tr>
<td>▪</td>
<td>▪ Ultra Turrax grind mixer</td>
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</tbody>
</table>

When, where and how to sample
Grape ripening can be monitored from veraison to harvest.

How to proceed
It consists in analyzing the grape composition: sugar content (°Brix), total acidity, pH, polyphenols, and anthocyanins.
200 grapevine berries should be collected in the vineyard and stored at 4°C for fruit composition analysis. Berries will be hand crushed and centrifuged at 4000 rpm for 10 minutes. The total soluble solid concentration (°Brix) can be determined using a temperature-compensating digital refractometer. A must sample of 50 mL will be used to determine pH and titratable acidity according to the methods of the Organisation Internationale de la Vigne et du Vin (OIV 2009). The remaining berry sub-sample can be homogenized using an Ultra Turrax grind mixer (IKA, Germany) at high speed (14,000 rpm for 1 minute). Anthocyanin and phenolic concentrations will be determined according to the method of Iland et al (2004). Total anthocyanins will be expressed as mg per berry and per gram of fresh berry mass, whereas total phenols will be expressed as absorbance units (AU) at 280 nm per berry and per gram of fresh berry mass.

Possible interpretations and conclusions
Wine quality depends on grape composition. Knowledge of the ripening process allow the grapegrowers and winemakers to make decisions about harvest according to the type of wine to be made. The careful control of grape composition parameters during ripening is crucial for the winemaking.

References

Additional bibliography


Dominati, E., Patterson, M., Mackay, A., 2010. A framework for classifying and quantifying the natural capital and ecosystem services of soils. Ecol. Econ. 69, 1858–1868. doi:10.1016/j.ecolecon.2010.05.002