

## APPENDIX C. SPECIFIC METHODOLOGY

### C.1 Map unit determination

Map units were delineated according to geology and slope category (McDonald *et al.* 1984) using geological mapping, topographical mapping, aerial photography and field survey techniques.

### C.2 Field observations

Most field descriptions are based on McDonald *et al.* (1984), Northcote (1979) and Isbell (1994). The definition for soil horizon boundaries is listed below.

S	Sharp	< 5 mm
A	Abrupt	5 - 20 mm
C	Clear	20 - 50 mm
G	Gradual	50 - 100 mm
D	Diffuse	> 100 mm
+	Continuing	

### C.3 Field tests

#### C.3.1 Saturated hydraulic conductivity

Site selection:

Considerable time and effort is required to obtain meaningful permeability ( $K_{sat}$ ) values. It is imperative that sites are chosen carefully prior to the day of measurement. The sites should have nil, or at most, minimal disturbance.

Procedure:

- i) Insert five small (35 cm diameter) and five large (40 cm diameter) infiltration rings with the small rings placed inside the large rings, so that each ring is approximately 100 mm into the main clay horizon. Remove some topsoil if necessary but care should be taken to cause minimal soil disturbance.
- ii) Rings need to be at least two metres apart and located at random. Relocate rings if obstacles such as stones or roots prevent an even downward movement of the ring into the soil.
- iii) Fill rings with water and set up reservoir tanks so that water is added when the level drops below the outlet tube. Record the time and date on field sheets.
- iv) Place lids on rings to minimise evaporation and interference.

v) Check that all containers are full and will last overnight to allow soil to saturate and conductivity rate to equilibrate.

vi) Record water levels at various times during the day (depending upon infiltration rate), and leave for 24 hour period without any interruptions to the water flow, if possible .

vii) Next day dig out each ring taking care not to disturb the soil contained within the ring. Up-end the ring and record the proportion of soil area that has been transmitting water for each ring and record if water movement has been evenly distributed or confined to root/worm holes or structural cracks. Note any other differences, i.e. rocks, sand, clay patches.

### C.4 Laboratory analysis

Samples collected for each soil horizon were air dried, ground with a mortar and pestle and separated with 4.75 and 2 mm sieves into a gravel fraction (4.75 - 2 mm), and soil. The gravel fraction was reported as a percentage of the air dried field sample and discarded, while all subsequent tests were carried out on the soil samples and reported in terms of oven dried (105 °C) samples (except for EC, pH and Cl).

#### C.4.1 Physical properties

##### 1. Particle size analysis

The method used for particle size analysis is based upon that of Hutton (1956), which divides the soil sample into the following four principal size groups:

Coarse sand 2.0 - 0.20 mm  
Fine sand 0.20 - 0.02 mm  
Silt 0.02 - 0.002 mm  
Clay < 0.002 mm

In this method the soil sample is mechanically dispersed using pentasodium triphosphate (sodium tripolyphosphate), shaken in a sedimentation cylinder, and silt and clay percentages determined on a 2% soil water mixture using a plummet balance. After hand decanting the silt and clay suspension, the sand fractions are determined by sieving and weighing the oven dried (105 °C) sand fractions.

Due to the presence of both organic material and solutes in the soil and also due to the limitations of the technique used, the sum of the four fractions does not always equal 100%. Limits of 4% variation for surface horizons and 2% variation for lower horizons are regarded as acceptable. The determination is repeated for samples outside these limits. If repeat samples still remain outside these limits, then the closest result is accepted.

## 2. Emerson class

Soil dispersion is tested using the method of Emerson, (1967), and based upon the Australian Standard AS1289, C8.1, (1980). This gives eight dispersion classes from E1 to E8, where E1 is the most dispersive class and E8 the least dispersive class. Class E5 was further divided into four sub-classes E5(A), E5(B), E5(C) and E5(D), where E5(A) is more dispersive than E5(D). Also, classes E2 and E3 were each divided into four sub-classes according to the modification of Loveday and Pyle (1973), as quoted in Craze and Hamilton (1991). In this classification E2(1) is less dispersive than E2(4) and E3(1) is less dispersive than E3(4).

The order of soil dispersion from most dispersive to least dispersive is therefore:

E1  
E2(4), E2(3), E2(2), E2(1)  
E3(4), E3(3), E3(2), E3(1)  
E4  
E5(A), E5(B), E5(C), E5(D)  
E6  
E7  
E8

## 3. Atterberg limits

Atterberg investigated the behaviour of fine grained soil with varying water content. He used the following definitions, quoted in Hicks (1991):

(a) The liquid limit is the water content at which a trapezoidal groove of specified shape, cut in moist soil held in a special cup, is closed after 25 taps on a hard rubber plate.

(b) The plastic limit is the water content at which the soil begins to break apart and crumble when rolled by hand into threads three mm in diameter.

(c) The shrinkage limit is the water content at which the soil reaches its theoretical minimum volume, as it dries out from a saturated condition.

The plasticity index is the difference between the liquid and plastic limits, and represents the range of water contents that the soil remains in the plastic state.

Atterberg limits are determined on a sieved soil fraction with particles < 0.425 mm in size. The methods are based upon the Australian Standard 1289 (1977), as follows:

Liquid limit AS1289. C1.1  
Plastic limit AS1289. C2.1  
Plasticity index AS1289. C3.1  
Linear shrinkage AS1289. C4.1

## C.4.2 Chemical properties

Soil chemical analyses were carried out by the State Chemistry Laboratory, South Road, Werribee, Vic., 3030.

### 1. EC, pH, and Cl determinations

These determinations are carried out on a 1:5 soil water suspension shaken for one hour, and allowed to equilibrate.

#### (a) Electrical conductivity

This test is used to estimate the concentration of soluble salts in the soil. Measurements are made on the soil water suspension using a dip cell and direct reading meter. Values are determined at 25 °C.

State Chemistry Laboratory, Method 004, July 1986.

#### (b) pH in H<sub>2</sub>O at 20 °C

The pH of the above suspension is determined using a calomel electrode and digital pH meter. State Chemistry Laboratory, Method 009 (1986).

#### (c) pH in CaCl<sub>2</sub>

This is carried out on the soil water suspension after the pH in H<sub>2</sub>O determination. One mL of 1M calcium chloride solution is added to the soil water suspension, and the mixture stirred. The pH is then measured again.

State Chemistry Laboratory, Method 009 (1986).

#### (d) Chloride

A fresh 1:5 soil water suspension is titrated with a silver nitrate solution, using an electrical circuit to determine the end point of the titration. Note that this determination may be omitted if the EC determination is < 0.1 dS/m.

State Chemistry Laboratory, Method 003 (1982).

## 2. Oxidizable organic carbon

In this determination the soil sample is oxidised by chromic acid in the presence of excess sulphuric acid, without the application of external heat (Walkley and Black, 1934). The colour produced is measured with a spectrophotometer.

State Chemistry Laboratory, Method 014 (1987).

### 3. Total nitrogen

Total nitrogen is determined by a Kjeldahl method, where the sample is digested with a sulphuric acid/selenious acid mixture. The resulting solution is analysed for nitrogen colorimetrically. State Chemistry Laboratory, Method 021 (1985).

### 4. Available potassium

The Skene method is used where soil potassium is extracted with 0.05M hydrochloric acid, and the potassium determined with an atomic absorption spectrophotometer (Skene 1956).

State Chemistry Laboratory, Method 011 (1987).

### 5. Available phosphorus

Phosphorus is determined by the Olsen method in which the soil phosphorus is extracted with a 0.5M sodium bicarbonate solution at pH 8.5, (Olsen *et al.* 1954). The phosphorus is then measured colorimetrically after reduction with ascorbic acid.

State Chemistry Laboratory, Method 010 (1982).

### 6. Exchangeable aluminium and manganese

The soil sample is extracted with a 1M potassium chloride solution, and both determinations are made on the one extract. Aluminium is determined colorimetrically using pyrocatechol violet. Manganese is determined by atomic absorption spectrophotometry.

State Chemistry Laboratory, Method 001 (1985).

### 7. Extractable bases, calcium, magnesium, potassium and sodium

The bases are extracted from the soil with a 1M ammonium acetate solution at pH 7, and the bases are then analysed by atomic absorption spectroscopy.

State Chemistry Laboratory (1993) - draft procedure.

### 8. Total exchangeable bases

This is a calculated value consisting of the sum of the exchangeable bases calcium, magnesium, potassium and sodium, as determined in method 7 (above).

### 9. Exchangeable hydrogen

The exchangeable hydrogen is extracted from the soil using 0.053N triethanolamine and back titrated with 0.2M hydrochloric acid. This is a method modified by Peech *et al.* (1962).

State Chemistry Laboratory, Method 005 (1984).

### 10. Cation exchange capacity

This is a calculated value consisting of the sum of the exchangeable bases calcium, magnesium, potassium and sodium plus exchangeable hydrogen, as determined in methods 7 and 9 (above).