## INTERNATIONAL STANDARD

ISO 10693

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# Soil quality — Determination of carbonate content — Volumetric method

iTeh Soualité du sol R Détermination de la téneur en carbonate — Méthode volumétrique (standards.iteh.ai)

<u>ISO 10693:1995</u> https://standards.iteh.ai/catalog/standards/sist/22eeba20-922e-4d6e-b660-27fe3d3731d9/iso-10693-1995



### Foreword

ISO (the International Organization for Standardization) is a worldwide federation of national standards bodies (ISO member bodies). The work of preparing International Standards is normally carried out through ISO technical committees. Each member body interested in a subject for which a technical committee has been established has the right to be represented on that committee. International organizations, governmental and non-governmental, in liaison with ISO, also take part in the work. ISO collaborates closely with the International Electrotechnical Commission (IEC) on all matters of electrotechnical standardization.

Draft International Standards adopted by the technical committees are circulated to the member bodies for voting. Publication as an International Standard requires approval by at least 75 % of the member bodies casting VIEW a vote.

International Standard ISO 10693 was prepared by Technical Committee ISO/TC 190, Soil quality, Subcommittee SC 3, Chemical methods and soil characteristics. https://standards.iteh.ai/catalog/standards/sist/22eeba20-922e-4d6e-b660-

Annexes A and B of this International Standard7are1for information3 only5

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## Soil quality — Determination of carbonate content — Volumetric method

## iTeh STANDARD PREVIEW (standards.iteh.ai)

#### 1 Scope

## ISO 10693:1995 Principle

This International Standard specifies a method for the 2/16303/sist/22eeba20-922e-4d6e-b660determination of carbonate content in soil samples. compose any carbonates present. The reaction in simplified form reads as follows (Me means metal):

It is applicable to all types of air-dried soil samples.

#### Normative references 2

The following standards contain provisions which, through reference in this text, constitute provisions of this International Standard. At the time of publication, the editions indicated were valid. All standards are subject to revision, and parties to agreements based on this International Standard are encouraged to investigate the possibility of applying the most recent editions of the standards indicated below. Members of IEC and ISO maintain registers of currently valid International Standards.

ISO 3696:1987, Water for analytical laboratory use ---Specification and test methods.

ISO 11464:1994, Soil quality - Pretreatment of samples for physico-chemical analyses.

ISO 11465:1993, Soil quality — Determination of dry matter and water content on a mass basis ----Gravimetric method.

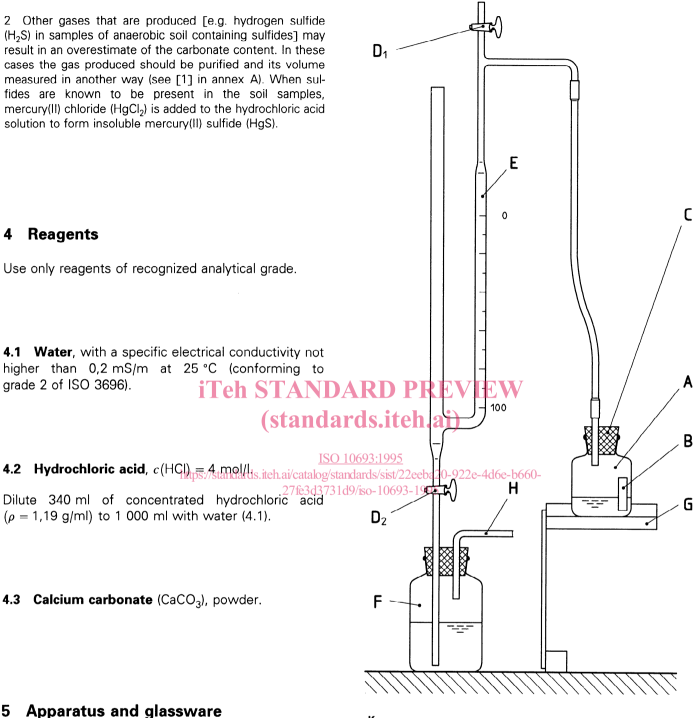
$$MeCO_3 + 2H^+ \rightarrow Me^{2+} + H_2CO_3$$

$$H_2CO_3 \rightarrow H_2O + CO_2$$
 (gaseous state)

The volume of the carbon dioxide produced is measured by using a Scheibler apparatus (5.1), and is compared with the volume of carbon dioxide produced by pure calcium carbonate. To avoid making corrections for differences in temperature and pressure, all determinations are carried out under the same conditions. The determination should be carried out in a temperature-controlled room.

#### NOTES

1 The carbonate content is expressed as an equivalent concentration of calcium carbonate (CaCO<sub>3</sub>). In fact all carbonates and bicarbonates present in the sample are measured. Many carbonates appear in the form of calcite and aragonite (CaCO<sub>3</sub>), dolomite [CaMg(CO<sub>3</sub>)<sub>2</sub>], siderite (FeCO<sub>3</sub>) and rhodochrosite (MnCO3). In soils in dry (arid) regions, soda (Na<sub>2</sub>CO<sub>3</sub> 10H<sub>2</sub>O) may be present. When it is known that a certain form of carbonate, other than calcium carbonate, is mainly present in the soil under study, the final concentration of this form can be used.



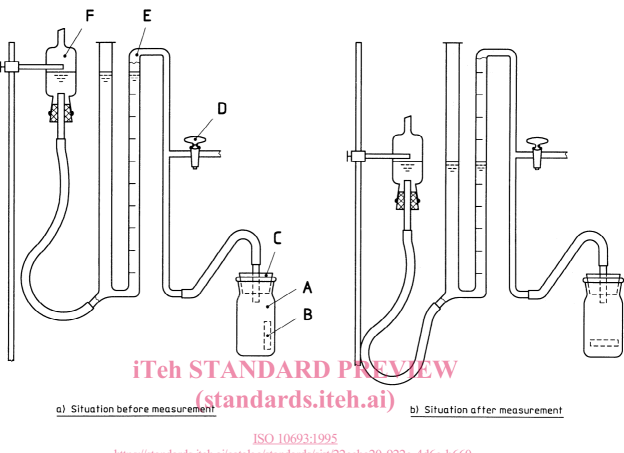
**5.1 Scheibler apparatus**, adapted for carrying out a single sample determination (see figure 1). An example of a single unit of apparatus, together with an indication of the water level before and after the

measurement, is given in figure 2.

#### Key

- A Reaction vessel (5.3)
- B Plastics cup (5.4) with hydrochloric acid (4.2)
- C Rubber stopper
- D<sub>1</sub> Stop-cock
- D₂ Stop-cock E U-shaped c
- E U-shaped calibration tube E Storage vessel with water
- F Storage vessel with water G Shaking beam
- H Air-inlet tube

## Figure 1 — Example of a Scheibler unit for a single sample determination



- Key https://standards.iteh.ai/catalog/standards/sist/22eeba20-922e-4d6e-b660-Reaction vessel (5.3) 27fe3d3731d9/iso-10693-1995
- Plastics cup (5.4) with hydrochloric acid (4.2) R
- С Rubber stopper
- D Stop-cock
- Е U-shaped calibration tube
- F Storage vessel with water

#### Figure 2 — Example of a single Scheibler unit

#### NOTES

Δ

3 For the determination specified in this International Standard, a Scheibler apparatus with two glass U-shaped calibration tubes each having a volume of 100 ml is used. It is also possible to use tubes with a different volume (i.e. 50 ml or 200 ml). In this case the mass of the test portion used for the measurement has to be adapted.

4 The temperature of the water in the system should be equal to the ambient temperature.

5 When only a few soil samples have to be analysed, a minimum of five Scheibler units is sufficient. In this case the reaction vessels can be shaken by hand. However, it is then necessary to take into account the temperature variations in the air inside the vessels.

6 By lowering the water level in the tube on the left (see figure 2) with the stop-cock closed, an underpressure is created in the tube on the right. By creating such an underpressure for some time, the unit can be checked for leaks.

5.2 Analytical balance, with an accuracy of at least 0,1 mg.

5.3 Reaction vessels, of capacity 150 ml and with a wide neck.

5.4 Plastics cup, of capacity about 10 ml, which can pass through the neck of the reaction vessel (5.3).

#### **5.5 Tongs**, acid-proof.

#### 5.6 Watch glass.

#### 6 Laboratory sample

Use the < 2 mm fraction of air-dried soil samples pretreated according to ISO 11464. Use part of the laboratory sample to determine the water content according to ISO 11465.

#### 7 Procedure

#### 7.1 Preparation

For a preliminary test, add some hydrochloric acid (4.2) to a portion of the soil on a watch glass (5.6). The carbonate content of the sample can be estimated on the basis of the intensity and time of effervescence; see table 1. Determine from table 1 the mass of test portion for the determination according to 7.2.

Table 1 — Relationship between intensity and time of effervescence, estimated carbonate	ľ
time of effervescence, estimated carbonate	L
content of the soil and mass of test portion	

Intensity of effervescence	ttps://standard Carbonate content	testead portion	g/s1 73
	g/kg	g	
None or only limited	< 20	10	
Clear, but for a short time	20 to 80	5	
Strong, for a long time	80 to 160	2,5	
Very strong, for a long time	> 160	≼ 1	

#### NOTES

7 If the mass of the test portion is less than 2 g, it should be taken from a representative portion of the laboratory sample that is crushed to a particle size of less than  $250 \ \mu m$  (according to ISO 11464).

8 Soils with a  $pH-H_2O$  less than 6,5 contain carbonates only in exceptional circumstances (e.g. recent liming or the presence of carbonates with a low solubility such as shells).

#### 7.2 Measurement

Determinations of samples, blanks and the calcium carbonate used as standard material, shall be performed simultaneously in a room where temperature and pressure do not vary too much during the measurement. Weigh a representative test portion of the laboratory sample (see 7.1 and table 1). Transfer this amount quantitatively into the reaction vessels (5.3) and add 20 ml of water (4.1). Also weigh the standards of 0,200 g and 0,400 g of calcium carbonate, transfer these amounts quantitatively into the reaction vessels (5.3) and add 20 ml of water (4.1). For the blank determinations, use reaction vessels containing 20 ml of water.

Place the reaction vessels in front of the Scheibler units (5.1, see figure 1) in such a way that blanks and standards are randomly distributed over the units. Open the stop-cocks  $D_1$  and  $D_2$  and bring the water level in the tube H to 3 ml for the samples and standards, and to 20 ml and 80 ml for the respective blanks. Close stop-cock  $D_2$ .

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3d3731d9/iso-10693-1995

Moisten the rubber stoppers of the apparatus with water and connect the reaction vessels. Close stopcock D<sub>1</sub> and once again check the water level in the tubes. Carefully add the hydrochloric acid from the cup (5.4) to the soil by tilting the reaction vessel at an angle. The gas produced will lower the water level in the tube on the right and at the same time will raise the water level in the tube on the left. Shake for 5 min and note the volume when it no longer varies. If it still varies, continue shaking until the volume is stable, but not longer than 1 h. Note the volume. During shaking, take care that the difference between the water levels in both tubes does not exceed 3 ml. This can be controlled by opening the stop-cock D<sub>2</sub>. At the end of the shaking period, bring the water level in both tubes to the same height and measure the volume of gas in the calibrated tube with an accuracy of 0,1 ml.

#### NOTES

9 The changes in volume of the blanks should not exceed 1,0 ml.

10 More water should be added to soils with a high organic matter content.

#### Calculation 8

Calculate the changes in volume of the gas produced by the sample  $(V_1)$ , the calcium carbonate standards  $(V_2)$  and the blanks  $(V_3)$  by subtracting the initial value in the calibrated tube from the final value.

Calculate the carbonate content of the sample using the following equation:

$$w(CaCO_3) = 1\ 000 \times \frac{m_2(V_1 - V_3)}{m_1(V_2 - V_3)} \times \frac{100 + w(H_2O)}{100}$$

where

- $w(CaCO_3)$  is the carbonate content, in grams per kilogram, of the oven-dried soil;
- is the mass, in grams, of the test  $m_1$ portion:
- is the mean mass, in grams, of the  $m_2$ calcium carbonate standards;
- $V_1$ is the volume, in millilitres, of carbon dioxide produced by the reaction of the test portion;
- is the mean volume, in millilitres, of S.1  $V_2$ carbon dioxide produced by the calcium carbonate standards; ISO 10693:1995
- $V_3$ the blank determinations (this value can be negative);
- $w(H_2O)$ is the water content, expressed as a percentage by mass, of the dried sample, determined according to ISO 11465.

#### 9 Repeatability

The repeatability of the determination of carbonate content carried out in two separate, consecutive measurements shall satisfy the conditions given in table 2.

Table 2 — Repeat	ability
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<b>Carbonate content</b> g/kg	Acceptable variation	
0 to 50	3 g/kg	
> 50 to 150	6 % of the value	
> 150 to 180	9 g/kg	
> 180	5 % of the value	

#### 10 Test report

The test report shall contain the following information:

a) a reference to this International Standard; KE

b) all information necessary for the complete identi**tenfication** of the sample;

the results of the determination in whole numhttps://standards.iteh.ai/catalog/standards/sist/22eebers.jmgrams.perokilogram, calculated on the bais the volume change, in millilitres dinso-10693-19is of oven-dried soil expressed as calcium carbonate or as another specified form of carbonate, when it is known to be the main form present;

> d) details of any operation not specified in this International Standard or regarded as optional, as well as any factor which may have affected the results.

## **Annex A** (informative)

# Results of an interlaboratory trial for the determination of the carbonate content of soils

An interlaboratory trial was organized in 1993 by the Wageningen Agricultural University (Netherlands), to verify the procedure specified in this International Standard.

For this interlaboratory trial, the determination of the carbonate content of five soils was carried out by nine laboratories.

The summary of the results of the interlaboratory trial is presented in table A.1.

The repeatabilities (*r*) and the reproducibilities (*R*) listed in table A.1 were calculated according to ISO 5725:1986, *Precision of test methods* — *Determination of repeatability and reproducibility for a standard test method by inter-laboratory tests.* 

Soil No.	Origin	<b>Mean value r</b> g/kg	ds.iteh.	$(ai)_R$	п	
1	Great Britain	662 <u>ISO 1(</u>	<u>693:B198</u>	94,8	9	
2	Frances://standards.	teh.ai/c5t4bg/stan	lards/2i8t/22eeb	a20-933,04d6e	-b660-9	
3	France	158	7,6	18,6	8	
4	Netherlands	66	3,8	19,2	9	
5	Argentina	1	2,0	3,3	9	
n = number of laboratories retained after eliminating outliers.						

## Table A.1 — Results of the interlaboratory trial for the determination of the carbonate content

### Annex B

(informative)

### Bibliography

[1] ALLISON, L.E. Wet combustion apparatus and procedure for organic and inorganic carbon in soils. *Soil Science Society of America Proc.* **24** (1960), pp. 36-40.

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