



SLOVENSKI STANDARD

SIST EN 196-5:2011

01-junij-2011

Nadomešča:
SIST EN 196-5:2005

Metode preskušanja cementa - 5. del: Določanje pucolanske aktivnosti za pucolanske cemente

Methods of testing cement - Part 5: Pozzolanicity test for pozzolanic cement

Prüfverfahren für Zement - Teil 5: Prüfung der Pozzolunität von Pozzolanzementen

Méthodes d'essais des ciments - Partie 5: Essai de pouzzolanité des ciments pouzzolaniques

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Ta slovenski standard je istoveten z: EN 196-5:2011

ICS:

91.100.10 Cement. Mavec. Apno. Malta Cement. Gypsum. Lime. Mortar

SIST EN 196-5:2011

en,fr,de

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EUROPEAN STANDARD
NORME EUROPÉENNE
EUROPÄISCHE NORM

EN 196-5

March 2011

ICS 91.100.10

Supersedes EN 196-5:2005

English Version

Methods of testing cement - Part 5: Pozzolanicity test for pozzolanic cement

Méthodes d'essais des ciments - Partie 5: Essai de pouzzolanité des ciments pouzzolaniques

Prüfverfahren für Zement - Teil 5: Prüfung der Pozzolanzität von Pozzolanzementen

This European Standard was approved by CEN on 20 February 2011.

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This European Standard exists in three official versions (English, French, German). A version in any other language made by translation under the responsibility of a CEN member into its own language and notified to the CEN-CENELEC Management Centre has the same status as the official versions.

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Foreword

This document (EN 196-5:2011) has been prepared by Technical Committee CEN/TC 51 “Cement and building limes”, the secretariat of which is held by NBN.

This European Standard shall be given the status of a national standard, either by publication of an identical text or by endorsement, at the latest by September 2011, and conflicting national standards shall be withdrawn at the latest by September 2011.

Attention is drawn to the possibility that some of the elements of this document may be the subject of patent rights. CEN [and/or CENELEC] shall not be held responsible for identifying any or all such patent rights.

This document supersedes EN 196-5:2005.

This European Standard on the methods of testing cement comprises the following Parts:

- EN 196-1, *Methods of testing cement — Part 1: Determination of strength*;
- EN 196-2, *Methods of testing cement — Part 2: Chemical analysis of cement*;
- EN 196-3, *Methods of testing cement — Part 3: Determination of setting times and soundness*;
- CEN/TR 196-4, *Methods of testing cement — Part 4: Quantitative determination of constituents*;
- EN 196-5, *Methods of testing cement — Part 5: Pozzolanicity test for pozzolanic cement*;
- EN 196-6, *Methods of testing cement — Part 6: Determination of fineness*;
- EN 196-7, *Methods of testing cement — Part 7: Methods of taking and preparing samples of cement*;
- EN 196-8, *Methods of testing cement — Part 8: Heat of hydration — Solution method*;
- EN 196-9, *Methods of testing cement — Part 9: Heat of hydration — Semi-adiabatic method*;
- EN 196-10, *Methods of testing cement — Part 10: Determination of the water-soluble chromium (VI) content of cement*.

NOTE A previous part, EN 196-21: *Methods of testing cement — Part 21: Determination of the chloride, carbon dioxide and alkali content of cement*, has been revised and incorporated into EN 196-2.

This edition introduces the following technical changes based on comments received by the secretariat:

- a) the procedure, reagents and layout of the standard have been aligned with the relevant clauses of EN 196-2;
- b) the procedure for preparation of a test sample has been clarified;

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- c) Patton and Reeders reagent has been included as an additional, optional indicator for visual determination of EDTA titrations;
- d) the specification for apparatus has been extended to include a balance of specified accuracy; apparatus for measuring the absorbance of a solution whilst being stirred and a pH meter of specified accuracy.

According to the CEN/CENELEC Internal Regulations, the national standards organizations of the following countries are bound to implement this European Standard: Austria, Belgium, Bulgaria, Croatia, Cyprus, Czech Republic, Denmark, Estonia, Finland, France, Germany, Greece, Hungary, Iceland, Ireland, Italy, Latvia, Lithuania, Luxembourg, Malta, Netherlands, Norway, Poland, Portugal, Romania, Slovakia, Slovenia, Spain, Sweden, Switzerland and the United Kingdom.

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1 Scope

This European Standard specifies the method of measuring the pozzolanicity of pozzolanic cements conforming to [1] EN 197-1. This standard does not apply to Portland pozzolana cements or to pozzolanas.

This method constitutes the reference procedure.

2 Normative references

The following referenced documents are indispensable for the application of this document. For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.

EN 196-7, *Methods of testing cement — Part 7: Methods of taking and preparing samples of cement*

EN ISO 385:2005, *Laboratory glassware — Burettes (ISO 385:2005)*

EN ISO 835:2007, *Laboratory glassware — Graduated pipettes (ISO 835:2007)*

3 Principle

The pozzolanicity is assessed by comparing the concentration of calcium ion, expressed as calcium oxide, present in the aqueous solution in contact with the hydrated cement, after a fixed period of time, with the quantity of calcium ion capable of saturating a solution of the same alkalinity. The cement is considered to satisfy the test, i.e. gives a positive result, if the concentration of calcium ion in the solution is lower than the saturation concentration.

NOTE Experiment has shown that a mixture of 20 g of cement and 100 ml of water at 40 °C achieves equilibrium after a period of between 8 d and 15 d. If the cement satisfies the test at 8 d (see 10.2) it is not necessary to continue to 15 d.

4 General requirements for testing

4.1 Number of tests

Where the determination is one of a series subject to statistical control, determination by a single test shall be the minimum required.

Where the determination is not part of a series subject to statistical control, the number of tests shall be two (see also 10.1).

In the case of dispute, the number of tests shall be two.

4.2 Repeatability and reproducibility

Repeatability and reproducibility in this document are expressed as repeatability standard deviation(s) and reproducibility standard deviation(s).

EN 196-5:2011 (E)**4.3 Expression of masses, volumes and factors**

Express masses in grams to the nearest 0,000 1 g and volumes from the burette in millilitres to the nearest 0,05 ml. Express the factors of solutions, given by the mean of three determinations, to three decimal places.

4.4 Determination of constant mass

Determine constant mass by drying for successive periods at the stated temperature, or making successive 15 min ignitions, followed each time by cooling and then weighing. Constant mass is reached when the difference between two successive weighings is less than 0,000 5 g.

5 Preparation of a test sample of cement

Before starting the determinations, treat the laboratory sample, taken in accordance with EN 196-7, as follows to obtain a homogenous test sample.

Take approximately 100 g of the sample using a sample divider or by quartering. Sieve this portion on a 150 μm or 125 μm sieve until the residue remains constant. Grind the retained material so that it completely passes the 150 μm or 125 μm sieve. Transfer the sample to a clean dry container with an airtight closure and shake vigorously to mix it thoroughly.

Carry out all operations as quickly as possible to ensure that the sample is exposed to ambient air only for the minimum time.

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6 Reagents

Use only reagents of analytical quality. References to water mean distilled or de-ionised water having an electrical conductivity $\leq 0,5$ mS/m. The quantities of reagents listed are to indicate concentrations; actual quantities to be prepared shall be adjusted according to the amounts required.

Unless otherwise stated (%) means percent by mass.

6.1 Concentrated hydrochloric acid (HCl), ($\rho = 1,18$ g/cm³ to 1,19 g/cm³).

6.2 Hydrochloric acid, about 0,1 mol/l, prepared by measuring with a graduated cylinder (7.16) 8,5 ml of concentrated hydrochloric acid (6.1) to a litre volumetric flask (7.10) containing about 500 ml of water and make up to 1 000ml with water. Determine the factor of normality of the solution as indicated in 8.2.

6.3 Dilute hydrochloric acid (1 + 2), prepared by adding 250 ml of concentrated hydrochloric acid (6.1) to 500 ml water.

6.4 Methyl orange, (dimethylaminoazobenzene p-sodium sulfonate).

6.5 Methyl orange indicator, prepared by dissolving (0,020 \pm 0,002) g of methyl orange (6.4) in water and making up to 1 000 ml.

6.6 Sodium hydroxide, (NaOH).

6.7 Sodium hydroxide solution, prepared by dissolving (100 \pm 1) g of sodium hydroxide (6.6) in water and making up to 1 000 ml.

6.8 Calcium carbonate, (CaCO₃), dried to constant mass at (200 \pm 10) °C (purity greater than 99,9 %).

- 6.9 Sodium chloride**, (NaCl), dried to constant mass at $(110 \pm 5) ^\circ\text{C}$.
- 6.10 Murexide**, (ammonium purpurate).
- 6.11 Murexide indicator**, prepared by grinding $(1,0 \pm 0,1)$ g of murexide with (100 ± 1) g of dry sodium chloride (NaCl).
- 6.12 EDTA**, (dihydrated disodium salt of ethylenediaminetetra-acetic acid).
- 6.13 EDTA solution, about 0,03 mol/l**, prepared by dissolving $(11,17 \pm 0,01)$ g of EDTA in water and making up to 1 000 ml. Store in an air-tight polyethylene container. Determine the factor of molarity of the solution as indicated in 8.1.
- 6.14 Sodium carbonate**, (Na_2CO_3), dried to constant mass at $(250 \pm 10) ^\circ\text{C}$.
- 6.15 Mixed calcein and methylthymol blue indicator**, prepared by grinding $(0,20 \pm 0,02)$ g calcein (bis [bis (carboxymethyl)-amino-methyl] -2', 7'-fluorescein (fluorescein, Flurorescein di-(methylimino diacetic acid) sodium salt) and $(0,10 \pm 0,01)$ g methylthymol blue, sodium salt of 3', 3''-bis- [bis (carboxy-methyl)-aminomethyl]-thymolsulfophthalein, ($\text{C}_{37}\text{H}_{41}\text{N}_2\text{O}_{13}\text{SNa}_3$) with (100 ± 1) g of potassium nitrate (KNO_3).
- 6.16 Calcon indicator**, prepared by grinding $(1,0 \pm 0,1)$ g of calcon, sodium 2-hydroxy-4-(2-hydroxy-1-naphthylazo) naphthalene-1-sulfonate, (Eriochrome Blue-Black R) with (100 ± 1) g of anhydrous sodium sulfate (Na_2SO_4).
- 6.17 Patton and Reeders reagent**, prepared by mixing $(1,0 \pm 0,1)$ g of Calcon carboxylic acid, (2-hydroxy-1-(2-hydroxy-4-sulfo-1-naphthylazo)-3-napthoic acid, ($\text{C}_{21}\text{H}_{14}\text{N}_2\text{O}_7\text{S}$), with (100 ± 1) g of anhydrous sodium sulfate (Na_2SO_4).

7 Apparatus

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- 7.1 500 ml cylindrical polyethylene container**, of about 70 mm diameter with a pressure seal-plug locked by a screw plug, capable of preventing evaporation during storage.
- 7.2 Wide stem funnel**.
- 7.3 Porcelain Büchner funnel**, of 60 mm inner diameter.
- 7.4 Filter paper**, with low porosity (mean pore diameter of about $2 \mu\text{m}$).
- 7.5 250 ml vacuum flask**.
- 7.6 250 ml and 400 ml beakers**.
- 7.7 50 ml and 100 ml pipettes**, class A of EN ISO 835:2007.
- 7.8 50 ml burette**, class A of EN ISO 385:2005.
- 7.9 Uniform temperature enclosure**, controlled thermostatically at $(40 \pm 1) ^\circ\text{C}$.
- 7.10 500 ml and 1 000 ml volumetric flasks**.
- 7.11 250 ml conical flask**.
- 7.12 Balance**, capable of weighing to an accuracy of $\pm 0,0005$ g.