<table>
<thead>
<tr>
<th>Standard No.</th>
<th>SASO-ISO-29581-2</th>
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<tr>
<td>Reference No.</td>
<td>ISO 29581-2</td>
</tr>
<tr>
<td>Issue Year</td>
<td>2010</td>
</tr>
<tr>
<td>Adoption Year</td>
<td>2014</td>
</tr>
<tr>
<td>Numeric Coding</td>
<td>922-29581-2</td>
</tr>
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Cement -- Test methods -- Part 2: Chemical analysis by X-ray fluorescence
The Saudi Standards, Metrology and Quality (SASO) has adopted without any technical changes the International Standard:

ISO 29581-2 Issue Year 2010

Cement -- Test methods -- Part 2: Chemical analysis by X-ray fluorescence
Cement — Test methods —
Part 2: Chemical analysis by X-ray fluorescence

Ciments — Méthodes d’essais —
Partie 2: Analyse chimique par spectrométrie de fluorescence X
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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.

International Standards are drafted in accordance with the rules given in the ISO/IEC Directives, Part 2.

The main task of technical committees is to prepare International Standards. 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 a vote.

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

ISO 29581-2 was prepared by Technical Committee ISO/TC 74, Cement and lime.

ISO 29581 consists of the following parts, under the general title Cement — Test methods:

— Part 1: Analysis by wet chemistry

— Part 2: Chemical analysis by X-ray fluorescence
Introduction

This part of ISO 29581 incorporates the following technical principles based on comments received by the Secretariat.

a) It provides an analytical method based on X-ray fluorescence (XRF) for use as the alternative method for the analysis of cement. When correctly calibrated according to the specified procedures and reference materials, it provides a method of suitable precision for conformity and information purposes.

b) It introduces a reference method for TiO₂, P₂O₅, SrO and Br analysis.

c) Traceability of the method relies upon reference materials and “pure” chemicals so that the ultimate traceability to basic international chemical standards relies upon classical analytical methods that are outside of the scope of this part of ISO 29581.

XRF and other instrumental methods, such as differential thermal analysis for determination of carbon dioxide, atomic absorption spectroscopy, etc., can be used as alternative methods, provided they are calibrated against the reference methods, or against internationally accepted reference materials.
Cement — Test methods —

Part 2:
Chemical analysis by X-ray fluorescence

1 Scope

This part of ISO 29581 describes a performance-based method for the chemical analysis of cement for SiO$_2$, Al$_2$O$_3$, Fe$_2$O$_3$, CaO, MgO, SO$_3$, K$_2$O, Na$_2$O, TiO$_2$, P$_2$O$_5$, Mn$_2$O$_3$, SrO, Cl and Br using X-ray fluorescence (XRF). It can be applied to other relevant elements when adequate calibrations have been established.

This part of ISO 29581 describes an alternative method for analyses of cement for conformity and information purposes, based on beads of fused sample and analytical validation using certified reference materials, together with performance criteria.

A method based on pressed pellets of unfused sample can be considered as equivalent, providing that the analytical performance satisfies the same criteria.

NOTE 1 The use of fused beads generally improves the accuracy of analysis for non-volatile elements, since it eliminates variability arising from differences in mineralogical forms or oxidation states. Pressed pellets generally improve the accuracy of analysis for volatile elements and can give adequate accuracy for the routine analysis of non-volatile elements.

NOTE 2 The presence of sulfide in a sample also leads to restrictions on the scope of the analysis that can be undertaken using the XRF technique based upon fused beads. In particular, sulfate (SO$_3$) cannot be determined directly from such a fused bead because of the contribution to the analysis from the unknown amount of sulfide. In addition, sulfide cannot be determined directly (or accurately, indirectly) because of the contribution of the unknown amount of sulfate to the analysis and because of the possibility that some sulfide can be lost by volatilization during fusion. Consequently, the method of ISO 29581-1, included as Annex D to this part of ISO 29581, is the reference method for determining the sulfate content of samples containing sulfide species.

Other methods can be used, provided they are calibrated, either against the reference method or against internationally accepted reference materials, in order to demonstrate their equivalence.

In the case of dispute, unless otherwise agreed by all parties, only the reference method in ISO 29581-1 can be used.

This part of ISO 29581 describes methods that apply principally to cements, but which can also be applied to their constituent materials and to other materials, the standards for which call up these methods.

International Standard specifications state which methods can be used.
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.

ISO Guide 30, Terms and definitions used in connection with reference materials

ISO Guide 31, Reference materials — Contents of certificates and labels

ISO 29581-1, Cement — Test methods — Part 1: Analysis by wet chemistry

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

EN 197-1, Cement — Part 1: Composition, specifications and conformity criteria for common cements

3 Terms and definitions

For the purposes of this document, the following terms and definitions apply.

3.1 beads
glassy discs of fused sample for analysis by X-rays in the spectrometer

3.2 pellets
compressed discs of finely ground sample for analysis by X-rays in the spectrometer

3.3 calibration beads or pellets
beads or pellets used for establishing the calibration equation

3.4 analysis beads or pellets
beads or pellets containing the sample being analysed

3.5 accuracy
closeness of agreement between a test result and the certified value for a reference material

3.6 repeatability
closeness of agreement among independent test results obtained with the same method on identical test items in the same laboratory by the same operator using the same equipment within short intervals of time

3.7 reproducibility
closeness of agreement between independent test results obtained with the same method on identical test items in different laboratories with different operators using different equipment

3.8 expert laboratory
laboratory capable of consistently meeting the expert performance criteria set out in Clause 14

3.9 normal laboratory
laboratory capable of consistently meeting the normal performance criteria set out in Clause 14
4 General requirements for testing

4.1 Number of tests

Analysis of a cement can require the determination of a number of its chemical elements. For each determination, one or more tests shall be carried out in which the number of measurements taken shall be as specified in the relevant clause of this part of ISO 29581.

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

Where the analysis method (including preparation and measurement) is checked at least once a week, as in accordance with 12.5.1, a determination of each chemical element by a single test shall be the minimum required. In the other cases, the number of tests for the determination of each chemical element shall be two; see also Clause 13.

4.2 Accuracy and precision limits

4.2.1 Accuracy limit

The accuracy performance criterion in this part of ISO 29581 is measured as a limit on the closeness of agreement between a test result and an accepted reference value for a certified reference material. The limits for accuracy, expressed in percent absolute, are set out in Table 2; one set is appropriate to the performance it is expected that an “expert” laboratory can achieve, whereas the other is appropriate for a “normal” laboratory.

4.2.2 Repeatability limit

The repeatability performance criterion in this part of ISO 29581 is measured as a limit on the repeatability where independent test results are obtained with the same method on identical test items (material) in the same laboratory by the same operator using the same equipment within a short interval of time. The limits for repeatability, expressed in percent absolute, are set out in Table 1; one set is appropriate to the performance it is expected that an “expert” laboratory can achieve, whereas the other is appropriate to a “normal” laboratory.

4.2.3 Reproducibility limit

The reproducibility performance criterion in this part of ISO 29581 is measured as a limit on the reproducibility where test results are obtained with the same method on identical test items (material) in different laboratories with different operators using different equipment. The limits for reproducibility, expressed in percent absolute, are set out in Table 3; one set is appropriate to the performance it is expected that an “expert” laboratory can achieve, whereas the other is appropriate to a “normal” laboratory.

4.2.4 Laboratory competence

The laboratory shall demonstrate that it can achieve the required performance in accordance with 12.3.3 and 12.3.4.

4.3 Expression of mass

Express mass in grams to the nearest 0,000 5.

4.4 Other methods

Other methods may be used, provided they are calibrated, either against the reference method or against internationally accepted reference materials, in order to demonstrate their equivalence.
5 Reagents and reference materials

5.1 Pure reagents

Reagents shall be of analytical quality and, wherever possible, pure oxides or carbonates, except for the calibration of such elements as sulfur, chlorine, bromine or phosphorus, which do not form stable oxides or carbonates, where some guarantee of stoichiometry is required.

Reagents shall be free of (or corrected for) the presence of water (and, in the case of oxides, carbon dioxide) when weighed out for fusion. Also, the reagents shall be in a known oxidation state. The specified procedure ensures that the correct oxidation state is obtained.

The reagents used to prepare the standard beads for cations shall be pure oxides or carbonates of at least 99.95 % purity (excluding moisture or CO₂).

Reagents shall be used in a known stoichiometry in terms of content. In order to achieve this, they can be treated before use as follows.

a) Determine the loss on ignition for silicon dioxide (SiO₂), aluminium oxide (Al₂O₃) and magnesium oxide (MgO) as follows.

1) Ignite the reagent at, for example, (1 175 ± 25) °C and maintain at this temperature for 30 min.
2) Cool in a desiccator to room temperature and reweigh.
3) After allowing for this loss, weigh the appropriate amount of the unignited material to prepare the bead.

b) Dry manganese oxide (Mn₂O₃) and titanium(IV) oxide (TiO₂) as follows.

1) Ignite the reagent at, for example, (1 000 ± 25) °C and maintain at this temperature for 30 min.
2) Cool in a desiccator to room temperature before use.

c) Dry iron (III) oxide (Fe₂O₃) as follows.

1) Ignite the reagent at, for example, (700 ± 25) °C and maintain at this temperature for 30 min.
2) Cool in a desiccator to room temperature before use.

d) Dry calcium carbonate (CaCO₃), strontium carbonate (SrCO₃), potassium carbonate (K₂CO₃) and sodium carbonate (Na₂CO₃).

1) Heat the reagent at, for example, (250 ± 10) °C and maintain at this temperature for 2 h.
2) Cool in a desiccator to room temperature before use.

5.2 Reference materials

5.2.1 Certified reference materials

Certified reference materials (CRMs) are materials, e.g. cement, supplied by an organization conforming to the requirements for the competence of reference material producers in accordance with ISO Guide 30.

CRMs shall be supplied with a certificate of analysis giving information on the average value and standard deviation in accordance with ISO Guide 31.
5.2.2 Industrial reference materials

Industrial reference materials (IRMs) are materials, e.g. cement, prepared and homogenized by a laboratory. The reference analysis of an IRM shall be the average result from inter-laboratory co-operative testing involving at least four laboratories able to meet the performance criteria given in 12.3.

5.3 Calibration standards

Calibration standards are prepared in the laboratory from pure, analytical-grade reagents, IRMs, CRMs or a combination of these. They shall be formulated to provide a series of calibration standards covering the range of maximum to minimum values for each element being analysed and shall be evenly distributed between those limits. The variation in concentrations of the elements shall be independent of each other. There shall be a minimum of seven calibration standards in a series.

5.4 Binding agent

A binding agent, e.g. wax, whose influence on the elements being analysed has been determined, is used in the grinding of samples during the preparation of pressed pellets. Carry out a pellet-preparation monitoring check (see 12.5) whenever the batch of binding agent is changed.

6 Apparatus

6.1 Balance, capable of weighing to an accuracy of ± 0,000 5 g.

6.2 Fusion vessels and casting moulds, of a non-wetted platinum alloy, such as Pt/5 % Au or Pt/Rh.

Vessels that serve both as a fusion vessel and as a casting mould (i.e. a combined fusion mould) may be used. If moulds become distorted in use, then they shall be reshaped by pressing in a suitable former. If the bottom (flat) surface of the bead is used for analysis, it is necessary that the internal base of the mould also be kept flat and free from blemishes.

NOTE Cleanliness of fusion vessels is important in achieving accurate analyses. This can be achieved, for example, by boiling in dilute hydrochloric acid, 1:10 by volume or citric acid, 100 g/l.

6.3 Lids, optional, of a platinum alloy (not necessarily non-wetted).

6.4 Furnace, e.g. an electric resistance, muffle or high-frequency induction furnace, capable of operating at (250 ± 10) °C, (700 ± 25) °C, (950 ± 25) °C, (1 000 ± 25) °C and (1 175 ± 25) °C.

6.5 Automatic fusion apparatus, for use in automatic bead preparation (see 11.4).

An automatic fusion apparatus may be used, provided that the performance criteria in 12.3 can be met.

6.6 Cooling apparatus, consisting of any device, such as a narrow jet of air that can be directed to the centre of the base of the casting mould (for example, by the base of a bunsen burner without a barrel) or a water-cooled metal plate.

NOTE Normally, cooling in air is sufficient but some difficult samples can require a cooling apparatus in order to cool the melt rapidly. This is necessary to obtain a homogeneous bead and to free the melt from the casting mould.

6.7 Heat reservoir, for the casting mould, required in special circumstances when using moulds of small sizes, so that the mould does not cool too rapidly when removed from the furnace.

6.8 Spectrometer, X-ray fluorescence, capable of meeting the performance criteria given in 12.3.

NOTE It is required to set appropriate measuring conditions to satisfy the performance criteria based on the type of samples, the type of apparatus, elements being analysed and their content, etc.
6.9 Flow gas, maintained at as constant a room temperature as possible.

The temperature of the flow gas cylinder and of the connecting pipework is critical in order to prevent drift in sensitivity of the flow proportional counters. Pipework shall be as short as practical and run, whenever possible, within the temperature-controlled room housing the spectrometer. Where this is not possible, the cylinder shall be kept in a temperature-controlled cabinet (room temperature ± 2 °C) or otherwise maintained at a constant room temperature. For the same reason, new cylinders shall be allowed to equilibrate for at least 2 h to room temperature before use.

NOTE 1 The flow gas is used in the gas flow proportional counter of the XRF spectrometer.

NOTE 2 The composition of gas can change as the cylinder becomes exhausted. Cylinders should be replaced before they become completely empty.

6.10 Grinding equipment, capable of grinding the sample, with binding agent if necessary, to a suitable fineness.

6.11 Pellet press, capable of applying a pressure suitable for production of pellets with a consistent, consolidated surface to meet the performance requirements given in 12.3.

6.12 Mould, usually of steel, of suitable strength to withstand the press without distortion and of suitable size to produce a pellet to fit the spectrometer.

7 Preparation of a test sample of cement

Before chemical analysis, treat the laboratory sample, taken in accordance with EN 196-7, as follows to obtain a homogeneous test sample.

a) Take approximately 100 g of the laboratory sample by means of a sample divider or by quartering.

b) Sieve this portion on a 150 µm or 125 µm sieve until the residue remains constant.

c) Remove metallic iron from the material retained on the sieve by means of a magnet (see Note 1).

d) Grind the iron-free fraction of the retained material so that it completely passes the 150 µm or 125 µm sieve.

e) Transfer the sample to a clean, dry container with an airtight closure and shake vigorously to mix it thoroughly.

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

NOTE 1 Where the analysis is one of a series subject to statistical control and the level of the metallic iron content has been shown to be insignificant in relation to the chemical properties being determined, it is not necessary to remove metallic iron. Where the level of metallic iron is significant, it is required to record and report the amount in the results.

NOTE 2 Where the sample contains quartz, it can be necessary to grind the sample to pass a 90 µm sieve in order to obtain a satisfactory fusion (see Clause 11). The time and temperature required to obtain a satisfactory fusion is affected by the fineness of the sample.

NOTE 3 Where pressed pellets are used, accuracy can be improved by grinding the sample more finely.
8 Flux

8.1 Choice of flux

8.1.1 General

One of the advantages of the XRF fused cast bead method is that a wide variety of fluxes may be chosen. For a given calibration, the same flux shall be used throughout. The conditions given in 8.1.2 to 8.1.4 shall be met for any flux used.

NOTE 1 Fluxes used with success in the analysis of cement are given in Annex A. Pre-fused fluxes have the advantage of a lower moisture content.

NOTE 2 Reducing the particle size of the flux has been shown to improve fusion at a given temperature.

8.1.2 Dissolution

Under the conditions of preparation used, the sample shall be totally dissolved by the flux and shall not come out of solution in the casting procedure.

8.1.3 Heavy-element absorber

A heavy-element absorber, such as lanthanum or vanadium oxide, may be incorporated into the flux, provided

— it does not reduce sensitivities to the point that the performance criteria given in 12.3 cannot be met;
— the heavy element does not have a line overlap with any of the elements being determined.

NOTE Lanthanum oxide assists the formation and stabilization of glass but reduces the intensity of the emitted X-rays.

WARNING — There are restrictions on the use of heavy-metal chemicals in some countries. Care should be taken in the handling of these and national safety rules observed.

8.1.4 Flux purity

The flux shall be pure with respect to the elements being determined.

Most reagents sold as “flux” grade quality by reputable manufacturers meet this requirement, but an analysis shall be obtained for each batch of flux supplied. Carry out a bead-preparation monitoring check (see 12.5) when a batch of flux is changed.

8.2 Moisture in flux

Pre-molten fluxes having a loss on ignition not exceeding 0.50 % mass fraction are preferred. However, if the flux contains moisture, it shall be dried at a suitable temperature.

8.3 Flux-to-sample ratio

The flux-to-sample ratio selected shall be such that the performance criteria given in 12.3 are met. The mass fraction ratio, \( R \), of flux to sample used for the calibration shall be used for subsequent analyses.

NOTE As the flux-to-sample mass fraction ratio is greater than one, impurities in the flux can have a negative influence on the measured result. The greater the ratio of the flux to sample, the greater the influence.

The total mass of sample and flux shall be chosen for the particular casting mould type used, and this mass shall always remain the same.
8.4 Anti-wetting agent

A small amount of anti-wetting agent may be used, if necessary. An anti-wetting agent, such as lithium bromide, ammonium bromide, lithium iodide, lithium iodate or ammonium iodide, may be added to the melt to assist in preventing the cracking of the fused beads on cooling and to aid in the release from the mould. Where an anti-wetting agent is used, all beads shall be prepared using the same anti-wetting agent added in the same quantity and at the same stage of bead preparation.

NOTE Bromine or iodine in the anti-wetting agent can remain in the bead under some fusing conditions. It is required to check for residual bromine or iodine, since these elements can cause line overlapping, such as Br Lα on Al Kα or I Lβ₂ on Ti Kα.

9 Determination of loss on ignition and the change in mass on fusion of the cement

9.1 Principle

In order to be able to total (to 100 % mass fraction) any oxide analysis of cement, a loss on ignition, i.e. the amount of any combined water and carbon dioxide, is required. In addition, in order to be able to convert an oxide analysis obtained on the fused-basis using fused-bead XRF, to an oxide analysis on the as-received basis, an “observed” loss on ignition is also required.

NOTE Where the sample contains no oxidizable species, the loss on ignition and the “observed” loss on ignition are the same.

The “observed” loss on ignition (see 9.3.1) is a very close approximation to the “change in mass on fusion” that occurs when a sample is prepared as a fused bead for analysis by XRF. This “observed” loss on ignition is used in this method to calculate a factor, $f$, (see Clause 10) to convert test results obtained on the fused basis to the as-received basis.

The traditional loss-on-ignition determination carried out in an oxidizing atmosphere by igniting in air can be used to determine both the loss on ignition and the “observed” loss on ignition. Where any oxidizable species are present, in particular sulfide or sulfur-containing species, a correction can be applied to the “observed” loss in order to derive a “corrected” loss on ignition for use in totalling the oxide analysis. However, any error resulting from the oxidation of any metallic iron, bivalent iron or bivalent manganese is usually considered to be negligible and only the correction for the extent of oxidation of sulfides is applied in the correction.

An alternative method, e.g. automatic equipment, may be used, provided that it can be demonstrated that the performance criteria given in 12.3 are satisfied.

9.2 Procedure

Weigh, to ± 0,000 5 g, (1,00 ± 0,05) g of cement into a crucible which has been previously ignited and tared. Record this mass as $m_1$. Place the covered crucible in the furnace (6.4) controlled at (950 ± 25) °C. After heating for 5 min, remove the lid and leave the crucible in the furnace for a further 10 min. Allow the crucible to cool to room temperature in a desiccator. Determine constant mass by making successive 15 min ignitions followed each time by cooling and then weighing, and record as $m_2$. Constant mass is reached when the difference between two successive weighings is less than 0,000 5 g.

NOTE For cements containing sulfides, a more accurate determination of the loss on ignition can be obtained by determining the sulfate content before and after ignition. The correction applicable to these cements is given in 9.3.2.

When vacuum desiccators are used, the appropriate desiccator inlet trap should be used when any vacuum is released. Phosphorus pentoxide should be avoided where surface-active materials are being stored, since P₂O₅ can be absorbed by the sample, particularly under vacuum conditions.
9.3 Calculation and expression of results

9.3.1 Observed loss on ignition

Calculate the observed loss on ignition (i.e. the change in mass on fusion), \( L \), expressed as a percentage mass fraction, as given in Equation (1):

\[
L = \frac{m_1 - m_2}{m_1} \times 100
\]  

where

- \( m_1 \) is the mass of the test portion, expressed in grams;
- \( m_2 \) is the mass of the ignited test portion, expressed in grams.

9.3.2 Correction to the observed loss on ignition for the oxidation of sulfides

Calculate the correction to the “observed” loss on ignition for the extent of oxidation of sulfides, by determining the sulfate present before and after ignition. The gravimetric method of ISO 29581-1, reproduced in Annex D of this part of ISO 29581, is the reference method for these sulfate determinations.

The amount of \( \text{SO}_3 \), \( w_{S,up} \), resulting from the oxidation of sulfides, expressed as a percentage mass fraction, is calculated as given in Equation (2):

\[
w_{S,up} = w_{S,f} - w_{S,i}
\]  

where

- \( w_{S,f} \) is the final \( \text{SO}_3 \) value, expressed as a percentage mass fraction;
- \( w_{S,i} \) is the initial \( \text{SO}_3 \) value, expressed as a percentage mass fraction.

A correction, \( w_{O,up} \), for the oxygen taken up by the test portion, expressed in percent, is given by Equation (3):

\[
w_{O,up} = 0.8 \times w_{S,up}
\]  

The corrected loss on ignition, \( L_{cor} \), is calculated as given in Equation (4):

\[
L_{cor} = L + w_{O,up}
\]  

where \( L \) is the observed loss on ignition, expressed as a percentage mass fraction.

Indicate any corrections applied in the test report.

9.4 Repeatability and reproducibility for loss on ignition

The standard deviation for repeatability is 0.04 % mass fraction.

The standard deviation for reproducibility is 0.08 % mass fraction.
10 Factoring test results and correcting total analyses for presence of sulfides and halides

10.1 General

Analyses may be carried out on a fused-bead or pressed-pellet test portion. In the case of analysis using fused beads, it is necessary to determine the change in mass that occurs during fusion (see 9.3.1) in order to derive a multiplying factor for converting test results obtained on the fused basis to analytical results reported on the as-received basis.

In addition, where sulfide, chloride or bromide are present, it is also necessary to correct the total oxide analysis of a cement for the fact that, traditionally, these species are also expressed as if they were oxygen (combined with calcium) in the analytical result reported for calcium oxide. This latter correction also applies to analyses carried out using pressed pellets.

10.2 Factoring test results obtained from fused-bead analysis

10.2.1 Principle

Where fusion is employed, it is necessary to determine the change in mass that occurs during fusion in order to derive a multiplying factor for converting test results obtained on the fused basis to analytical results on the as-received basis.

The change in mass that occurs during fusion is determined by igniting, in air, a weighed test portion to constant mass at (950 ± 25) °C and calculating the "observed" loss on ignition; see 9.3.

NOTE 1 The change in mass occurring during fusion in air is normally a loss as, generally, water and carbon dioxide are released. Where, however, the sample contains oxidizable species, there is an increase in mass of these species during fusion resulting from oxygen uptake. This might or might not lead to an overall increase in mass of the test portion depending on the balance of mass lost versus mass gained. The oxidizable species that can lead to an overall gain in mass of a test portion is sulfur, generally present as sulfide (typically calcium sulfide) in test portions that contain a blast-furnace slag constituent. Other oxidizable species, where present, such as metallic iron, bivalent iron or bivalent manganese, are generally present in small amounts and make only a minor contribution to mass change.

NOTE 2 The presence of sulfide in a sample also leads to restrictions on the scope of the analysis that can be undertaken using the XRF technique based upon fused beads. In particular, sulfate (SO₃) cannot be determined directly from such a fused bead because of the contribution to the concentration from the unknown amount of sulfide. In addition, sulfide cannot be determined directly (or accurately, indirectly) because of the contribution of the unknown amount of sulfate to the analysis and from the possibility that some sulfide could have been lost by volatilization during fusion. Consequently, the method of ISO 29581-1, included as Annex D to this part of ISO 29581, is the reference method for determining the sulfate content of samples containing sulfide species.

10.2.2 Calculating the factor for converting the results obtained using fused beads

Calculate the factor, \( f_{\text{LOI}} \), for converting test results obtained on the fused basis to analytical results on the as-received basis, from the observed loss on ignition, \( L \), (see 9.3.1) as given in Equation (5):

\[
 f_{\text{LOI}} = \left( \frac{100 - L}{100} \right)
\]  

(5)

where \( L \) is expressed as a percentage mass fraction.

When \( L \) is a negative number, which occurs when the test portion has undergone an overall gain in mass, preserve the sign together with \( L \) in Equation (5).
10.2.3 Use of the conversion factor, $f_{\text{LOI}}$

Convert all test results obtained on the fused basis to analytical results on the as-received basis by multiplying each result by the factor, $f_{\text{LOI}}$.

10.3 Correcting total oxide analysis for sulfides and halides

10.3.1 Correcting total oxide analysis for sulfides

Irrespective of whether the analysis was carried out on a fused (bead) or pressed (pellet) test portion, determine the sulfide content of the cement, on the as-received basis, in accordance with the method for sulfide determination in ISO 29581-1.

The factor, $f_{\text{S}}$, to correct the total analysis of cement containing sulfide species, is calculated as given in Equation (6):

$$f_{\text{S}} = w_{s^{2-}} + L_{\text{cor}} + w_{O,\text{tot}} - w_{O,s^{2-}}$$

where

- $w_{s^{2-}}$ is the sulfide content, expressed as a percentage mass fraction;
- $L_{\text{cor}}$ is the “corrected” loss on ignition, expressed as a percentage mass fraction; see 9.3.2;
- $w_{O,\text{tot}}$ is the sum of all oxides, expressed as a percentage mass fraction; see Clause 1;
- $w_{O,s^{2-}}$ is the oxygen equivalent of sulfide, expressed as a percentage mass fraction, which is calculated as given in Equation (7):

$$w_{O,s^{2-}} = 0.5w_{s^{2-}}$$

10.3.2 Correcting total oxide analysis for halides

10.3.2.1 Correcting total oxide analysis for chloride

Irrespective of whether the analysis has been carried out on a fused (bead) or pressed (pellet) test portion, calculate the factor, $f_{\text{Cl}}$, to correct the total analysis of cement containing chloride as given in Equation (8):

$$f_{\text{Cl}} = w_{\text{Cl}} + L_{\text{cor}} + w_{O,\text{tot}} - w_{O,\text{Cl}}$$

where

- $w_{\text{Cl}}$ is the chloride content, expressed as a percentage mass fraction;
- $w_{O,\text{Cl}}$ is the oxygen equivalent of chloride, expressed as a percentage mass fraction, which is calculated as given in Equation (9):

$$w_{O,\text{Cl}} = 0.2w_{\text{Cl}}$$

NOTE Any correction for the presence of chloride is generally very small and is usually ignored.
10.3.2.2 Correcting total oxide analysis for bromide

Irrespective of whether the analysis has been carried out on a fused (bead) or pressed (pellet) test portion, calculate the factor, \( f_{\text{Br}} \), to correct the total analysis of cement containing bromide as given in Equation (10):

\[
f_{\text{Br}} = w_{\text{Br}} + L_{\text{cor}} + w_{\text{O,tot}} - w_{\text{O,Br}}
\]

where \( w_{\text{Br}} \) is the bromide content, expressed as a percentage mass fraction, which is calculated as given in Equation (11):

\[
w_{\text{Br}} = \frac{10}{w_{\text{O,Br}}}
\]

NOTE Any correction for the presence of bromide is generally very small and is usually ignored.

11 Preparation of fused beads and pressed pellets

11.1 General

At several of the stages, a choice of procedures is given. Once a choice has been made, the procedure shall be adhered to throughout calibration and analysis, unless a new calibration is carried out in accordance with Clause 12.

The bead or pellet preparation conditions shall be set such that the criteria given in 12.3 are met.

11.2 Conversion of the sample to bead form

11.2.1 Weighing of the sample

11.2.1.1 General

Weigh the sample, the flux and any anti-wetting agent to the nearest 0,000 1 g and to the ratio, \( R \), selected. Add the anti-wetting agent, when used as a solution, using a micro-pipette. The sample being used shall be weighed as given in either 11.2.1.2 or 11.2.1.3.

11.2.1.2 Unignited samples

The required mass, \( m_3 \), expressed in grams, of the unignited sample is calculated as given in Equation (12):

\[
m_3 = \frac{m_4}{1 - \frac{L}{100}}
\]

where

\( m_4 \) is the mass of fused sample (see 8.3) required to prepare a bead, expressed in grams;

\( L \) is the observed loss on ignition, expressed in percent (see 9.3.1).

11.2.1.3 Ignited samples

Prepare the sample in accordance with 9.2 but, in this case, the appropriate amount of the sample may be used. A constant mass is reached when the difference between two successive weighings is less than 0,05 % mass fraction. Store ignited samples in a desiccator.

When preparing a bead using the ignited sample, carry out weighings as quickly as possible to avoid contamination.
Where the sample contains significant amounts of carbonate that can result in loss of material through spurting, or where it contains carbides, iron or other metals that can interact with platinum and damage the crucible, the ignited sample should be used in the preparation of beads.

The amount of the sample should not be more than 5 g, because it is difficult with a larger sample to reach a constant mass without repeated ignitions. However, if a large amount of sample is being used, lumps in the sample should be broken up after the first ignition.

11.2.2 Fusion procedure

Thoroughly mix the sample, the flux and any anti-wetting agent before fusion. Where the anti-wetting agent is added as a solution, it shall be added after placing the mixture of sample and flux into the fusion vessel and evaporating all moisture at a low temperature before the fusion. Fuse the mixture for the prescribed time, e.g. 10 min, at the selected, controlled temperature with occasional swirling until the sample dissolves and the melt is homogeneous.

NOTE The volatilization of sulfur trioxide becomes significant for samples fused at temperatures in excess of 1 100 °C. Where sulfur trioxide is being determined, it is essential to keep the temperature below this level.

The fusion temperature may be specified according to cement or element type. For more volatile elements, e.g. sulfate, sulfide, chloride or alkalis, fusion at reduced temperature or the use of pressed pellets can be necessary to achieve the required precision.

Increasing the temperature in two stages has been shown to improve precision.

11.3 Casting of the beads

11.3.1 General

Cast the beads using one of the following methods. Beads produced shall be free from blemishes on the analytical measuring surface.

a) Outside the furnace: after 5 min, remove the casting mould (and the heat reservoir) from the furnace and place on a horizontal surface. Remove the lid from the fusion dish, remove the dish from the furnace and immediately pour the melt into the casting mould.

a) In the furnace: after 5 min, remove the lid from the fusion dish and pour the melt into the casting mould inside the furnace, ensuring that as much of the melt is transferred to the casting mould as possible. Remove the casting mould from the furnace and place on a horizontal surface.

b) Combined fusion mould: after 5 min, remove the fusion mould from the furnace and by swirling ensure the transfer of the whole of the melt into the mould part of the dish.

c) Mould heated over burner: after preparation of the melt, pour the melt into the preheated casting mould and turn the burner off.

d) Inside the fusion mould: after completion of the fusion, leave the melt inside the fusion dish and remove it from the furnace.

11.3.2 Cooling of the beads

Allow the casting mould to cool on a horizontal surface.

If rapid cooling is required, when the melt has cooled from red heat, transfer the casting mould to an air jet (6.6) (or alternatively, a water-cooled plate). The melt can be molten or solid at this stage. Hold the dish in a horizontal position above the air jet so that the air is directed onto the centre of the base of the mould. When the bead has solidified and released itself, turn off the air jet.

NOTE It can be necessary to encourage the release of the beads at this stage by gently tapping the casting mould on a solid surface.
11.4 Automatic bead preparation

An automatic fusion apparatus (6.5) may be used to prepare beads.

11.5 Storage

Protect beads from deterioration caused by adverse temperature and humidity conditions, e.g. by storage in a polyethylene self-seal bag. If the laboratory environment is suitably controlled (e.g. air-conditioned), then the bag shall be stored in a desiccator. Alternatively, if the environment is not controlled, the bags shall be stored in a temperature-controlled oven at 25 °C to 30 °C.

Where pressed pellets are being stored for a limited period, they may be placed unwrapped in a desiccator.

The bags themselves can cause surface contamination due to the use of "anti-blocking agents" (the effect being more apparent for the lighter elements). After long-term storage, thoroughly clean the measuring surfaces of beads before use, e.g. by washing with ethanol or acetone or by polishing.

NOTE Reported sources of contamination are as follows:

— sulfur from vacuum oil in the spectrometer or from the laboratory atmosphere;
— sodium and chlorine from the atmosphere if the laboratory is near the sea;
— sodium from manual contact;
— potassium from cigarette smoke.

11.6 Pressed pellets

Weigh, to the nearest 0.1 g, a quantity of sample suitable to fill the mould to excess. Grind to a suitable fineness, with a binding agent (5.4) if necessary. Place the ground sample into the pellet mould (6.12) and spread evenly. Apply pressure at a rate and for a time shown to be suitable to ensure that the criteria given in 12.3 are met. Store in accordance with 11.5 until required.

12 Calibration and validation

12.1 Principles

The calibration equations and inter-element corrections are established using calibration standards produced using beads or pellets made from pure reagents, CRMs, IRMs or any combination of these (see Clause 5). The validity of the calibration is confirmed by analysing one or more CRMs, representative of the range of analyses being undertaken. The ongoing calibration monitoring for intensity, bead preparation and calibration drift is determined immediately after calibration and at intervals during the analytical process. Typical ranges for calibration standards for analysis of CEM I cements are given in Annex C. Examples of sources of CRMs are shown in Annex B. Performance criteria are set out in Clause 14.

12.2 Calibration and validation samples

12.2.1 Calibration standards

Prepare a series of beads or pellets from pure reagents, IRMs, CRMs or combinations of these (see Clause 5) as calibration standards. The series shall cover the range from the minimum to the maximum values for each element being analysed and shall be evenly distributed between those limits. The variations in the concentration of each element in the calibration standard shall be independent of each other. The minimum number of standards is seven.
An example of a series of calibration standards suitable for the analysis of cements of type CEM I, in accordance with EN 197-1, is given in Annex C. Other reagents may be added to cover the scope of the elements being determined.

Where ten standards or fewer are used, duplicate beads or pellets should be prepared. Where a larger number of standards is used, single preparation can be sufficient.

For pellet analysis, a calibration made from IRMs or CRMs can provide unsatisfactory results due to mineralogical effects. In this case, the calibration may be done with secondary standards, made from samples routinely analysed by the laboratory, that have been characterized with at least one analysis by reference methods or by X-ray fluorescence using fused beads.

### 12.2.2 Calibration validation of a CRM or IRM

Prepare one or more CRMs, not used in the calibration standard (12.2.1) and having a composition within the calibration range for each element being analysed.

When only one validation CRM is being used, select a sample in the middle of the concentration ranges. Where several validation CRMs are used, select samples covering high and low values.

Examples of sources of CRMs are given in Annex B.

If no suitable CRM is available, choose an IRM that satisfies the above criteria. It is recommended that each IRM be validated by at least four laboratories.

**NOTE** In the case of pressed pellets, and especially for composite cements, variations in mineralogy can affect the accuracy. In such cases, it is necessary to use an IRM matching closely each type of sample being analysed, and preferably made from the samples routinely analysed by the laboratory.

### 12.2.3 Intensity correction samples

**NOTE 1** These are sometimes referred to as “monitor” samples.

Use one or more samples, (glass beads or other stable material) each having a composition giving an intensity level similar to the calibration range for each element being analysed. If several samples are used, select high and low values for each element. These samples shall be different from the calibration standards (see 12.2.1). Store these samples for future reference. Excessive exposure to the X-ray beam shall be avoided to prevent ageing.

**NOTE 2** Samples stored as pellets are subject to rapid ageing and, therefore, it is recommended that fused beads or glass be used for intensity control, even where the analysis predominantly uses pellet calibrations.

The samples shall be used to calculate a correction factor to apply to the raw intensity measurements, to take account of the ageing of the apparatus (X-ray tube, detectors, etc.) (see 12.4.3).

### 12.2.4 Re-calibration standards

Select two of the calibration standards (12.2.1), having high and low values for each element of interest, and store for future reference.

**NOTE** Depending on the equipment, it can be required to define re-calibration standards at the time of the initial calibration.

They shall be used to adjust the equation of the calibration curve in the event that the drift of the apparatus is not sufficiently corrected by the intensity correction procedure (see 12.4.4).
12.2.5 Spectrometer control samples

12.2.5.1 Control for bead analysis

Prepare and store one or more beads within, or close to, the calibration range. Calibration validation samples (see 12.2.2) are suitable.

A replacement bead may be prepared if ageing is suspected.

12.2.5.2 Control for pellet analysis

Due to the more rapid ageing of pellets, it is not possible to ensure control of the spectrometer using pellets. Therefore, the spectrometer control for pellet analysis shall be made using a fused bead or glass sample.

12.2.6 Preparation procedure control sample

Prepare a new bead or pellet of the calibration validation samples (see 12.2.2) as a control for the preparation procedure.

12.3 Initial calibration

12.3.1 Calibration procedure

Establish the calibration relationship between the intensity of measured X-rays and the element concentration, including corrections for any mass absorption and line overlap effects.

At a reasonable counting time (e.g. 200 s), measure and record the intensities for each element being determined in all of the calibration beads or pellets. Using regression analysis, establish the calibration curve for each element being determined.

NOTE Increasing the counting time on the measurement of the standards can improve the accuracy of the calibration.

12.3.2 Calibration for inter-element effects

Where inter-element effects significantly influence the accuracy of the calibration, e.g. the influence of potassium on calcium, it can be necessary to establish a correction. For each inter-element correction applied, prepare at least one additional calibration standard bead or pellet.

12.3.3 Validation of calibration

Validate the accuracy of the calibration for the elements being determined by analysing, in duplicate, at least one accuracy validation sample [see 12.2.2 (CRM or IRM)].

NOTE When selecting a CRM for validation, it is recommended that the series of analyses used to produce the certified analysis be examined in relation to the criteria for repeatability, reproducibility and accuracy (see Clause 14).

The calibration is valid if all the elements being determined satisfy the accuracy limits set down in Table 2.

If the calibration is not valid:

— consider the need for a calibration adjustment for inter-element effects (see 12.3.2);
— consider whether the set of standards used is adequate;
— determine other cause(s), and take appropriate corrective action; or,
— repeat initial calibration in accordance with 12.3, as appropriate.
12.3.4 Validation of repeatability

12.3.4.1 General

Validate the repeatability of analysis for a new spectrometer or when there has been any major modification of the preparation procedure or the spectrometer.
12.3.4.2 Repeatability of the spectrometer

Measure, over a period of not more than one week, the same bead of the spectrometer control sample (see 12.2.5) at least 10 times. Record the average and calculate the standard deviation of these measurements. To validate the spectrometer repeatability, the standard deviation shall be less than one-third of the repeatability limits defined in Table 1.

The limits in Table 1 allow for the overall procedure repeatability. The repeatability of the spectrometer itself should be lower, to ensure the required repeatability of the method.

If the repeatability of the spectrometer is not validated, implement one of the following and repeat the repeatability validation:

— check the stability of the apparatus (gas flow, temperature regulation, etc.);
— increase the counting time;
— increase the sample-to-flux ratio for the bead to improve sensitivity.

12.3.4.3 Repeatability of the method

Prepare, over an appropriate period of time (e.g. at least two weeks), at least 10 different preparations of the preparation procedure control sample (see 12.2.6). Record the average and the calculated standard deviation.

The repeatability of the method is valid if the difference in results for each new preparation, compared with the previous example, is within the repeatability limits defined in Table 1.

If the repeatability of the method is not validated:

— check the stability of the apparatus (gas flow, temperature regulation, etc.);
— increase the counting time;
— increase the sample-to-flux ratio for the bead to improve sensitivity;
— increase the fineness of the sample;
— increase the temperature and/or duration of the fusion for beads;
— consider the use of a grinding agent or binding wax for pellets;
— increase the duration and/or pressure on the press for pellets.

12.3.5 Initial values of the intensity correction samples

At the time of the initial calibration, record the initial values of the intensity correction samples (“monitors”; see 12.2.3).

NOTE It is generally possible to store these values in the software.

Store the corresponding samples for future reference.

12.3.6 Initial values of the re-calibration standards

At the time of the initial calibration, record the initial values of the re-calibration standards (see 12.2.4).

NOTE It is generally sufficient in the software to define two of the calibration standards (see 12.2.1) as the re-calibration standards.
12.3.7 Initial values of the spectrometer control sample

At the time of the initial calibration, record the average value, calculated in accordance with 12.3.4.2, of the spectrometer control sample (see tolerance limits in 12.4.2).

NOTE If ageing of the spectrometer control sample is suspected, prepare a new bead, and establish new initial values.

12.3.8 Initial values of the preparation procedure control sample

At the time of the initial calibration, record the average calculated in 12.3.4.3 as the initial value of the preparation procedure control sample (see 12.2.6 and preparation repeatability validation limits in 12.5.1).

12.4 Spectrometer follow-up

12.4.1 Principle

The main bias in X-ray fluorescence is the intensity drift due to ageing. The consistency of the results is monitored by measuring the spectrometer control sample (see 12.2.5), before carrying out each series of sample analyses.

In the first instance, this intensity drift is corrected using the intensity correction samples (12.2.3). Where this is not sufficient, correction factors may be applied to the calibration equation using the re-calibration standards. Alternatively, a full re-calibration can be necessary.

After each action that modifies the result provided by the spectrometer (e.g. drift correction or re-calibration), the accuracy shall be assessed again in accordance with 12.3.3.

12.4.2 Spectrometer follow-up validation

12.4.2.1 General

If the spectrometer control sample is within the limits set out below, or if the calibration has been re-validated in accordance with 12.3.3, then the spectrometer is ready for analysis. If the analysis is not valid, apply the decision diagram in Figure 2.

12.4.2.2 Re-validation of bead calibration

The bead calibration remains valid if the results, compared with the initial values (see 12.3.7), are equal to or less than three times the standard deviation for the spectrometer determined in 12.3.4.2.

12.4.2.3 Re-validation of pellet calibration

Depending on the capabilities of the software used, record the measurements

— as a corrected intensity, taking into account the intensity drift correction (see 12.4.4);
— as a concentration.

The pellet calibration remains valid if the results, compared with the initial values (see 12.3.7), are equal to or less than three times the standard deviation for the spectrometer determined in 12.3.4.2.
Figure 2 — Analysis validation decision scheme

Instrument check

| Measure preparation procedure control sample according to 12.2.6 with appropriate frequency |
| Measure spectrometer control sample according to 12.2.5 |
| Validated according to 12.4.2? |
| Check instrument according to 12.4.3 |
| Spectrometer OK? |
| Apply remediation |
| Check that calibration is valid according to 12.3.3 |
| Correction for intensity drift according to 12.4.4 |
| Re-calibration 12.4.5 |
| Renew initial calibration according to 12.3 |
| Ready for analysis |

Carry out the following checks, as necessary:

— whether any automatic fine adjustment, present in software on some spectrometers, has been made;
— gas flow;
— temperature regulation;
— resolution counter problems;
— 2-theta angle settings;
— high-voltage settings of the detector.
12.4.3 Correction for intensity drift

Where necessary (see Figure 2), a correction factor is calculated for each element to compensate for the drift in intensity due to ageing of the tube, detector, etc. Subsequently, the corrected intensities are then used in the calibration equation to derive the concentrations.

NOTE In most software applications, this correction is applied to all calibrations for each element.

The intensity drift should be small and regular with time. A sudden drop in intensity indicates an instrumental problem, and correction by this intensity drift method is not appropriate.

12.4.4 Re-calibration

Where necessary (see Figure 2), if the intensity correction (see 12.4.4) is insufficient to achieve accurate results for the analysis validation (see 12.4.2), it is possible to apply correction factors to the calibration curves using the re-calibration standards.

The modifications in the calibration equation due to instrumental drift should be small and regular with time. Any rapid or major bias in the analytical results can indicate an instrumental or analytical problem, and correction by this re-calibration method is not appropriate.

12.5 Preparation procedure follow-up

12.5.1 Bead and pellet preparation validation

On a regular basis, the whole analysis method shall be checked, including preparation of the bead or pellets and the measurement on the spectrometer.

Prepare a new preparation procedure control sample (see 12.2.6). The preparation is valid if

— the results of the new analysis, compared with the initial values (see 12.3.8), are within the limits for repeatability set down in Table 1;

— the ongoing average values for new analyses, compared with the certified values of the CRM or IRM, are within the limits for accuracy set down in Table 2.

12.5.2 Remedial action for preparation bias

If the analysis fails to satisfy the repeatability or accuracy limits set down in 12.5.1, check whether

— the flux is loss-on-ignition-free;

— drift can have occurred due to the preparation apparatus (check temperature, fineness, quality of sample surface, etc.);

— a new batch of flux has been introduced;

— a new bead or pellet preparation apparatus has been installed;

— the sample preparation procedure has been changed or modified, e.g. from manual to automatic;

— a new type of flux or new sample-to-flux ratio has been used;

— other procedural or apparatus changes have been introduced.

In the first two cases, take remedial action and re-check the analysis.

In all other cases, carry out a full re-calibration, including new preparation of all of the samples in accordance with 12.2.
Table 1 — Repeatability limits for analysis validation

<table>
<thead>
<tr>
<th>Element mean value</th>
<th>Repeatability limit for “normal” performance</th>
<th>Repeatability limit for “expert” performance</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>% mass fraction absolute</td>
<td>% mass fraction absolute</td>
</tr>
<tr>
<td>0 to 0,49</td>
<td>0,057</td>
<td>0,023</td>
</tr>
<tr>
<td>0,50 to 0,99</td>
<td>0,080</td>
<td>0,032</td>
</tr>
<tr>
<td>1,00 to 1,99</td>
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<td>0,044</td>
</tr>
<tr>
<td>2,00 to 3,99</td>
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<td>4,00 to 4,99</td>
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<td>80,00 to 100</td>
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<td>0,260</td>
</tr>
</tbody>
</table>

NOTE  All values of repeatability are expressed on the basis of fused, i.e. loss-on-ignition-free, samples.

Table 2 — Accuracy limits for validation

<table>
<thead>
<tr>
<th>Content of element species</th>
<th>Accuracy limit for “normal” performance</th>
<th>Accuracy limit for “expert” performance</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>% mass fraction absolute</td>
<td>% mass fraction absolute</td>
</tr>
<tr>
<td>0 to 0,49</td>
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<td>80,00 to 100</td>
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</table>

NOTE  All values of accuracy are expressed on the basis of fused, i.e. loss-on-ignition-free, samples.
Table 3 — Reproducibility limits achieved using certified reference materials

<table>
<thead>
<tr>
<th>Element mean value % mass fraction</th>
<th>Reproducibility limit for “normal” performance % mass fraction</th>
<th>Reproducibility limit for “expert” performance % mass fraction</th>
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<td>60</td>
<td>1,125</td>
<td>0,450</td>
</tr>
<tr>
<td>65</td>
<td>1,162</td>
<td>0,465</td>
</tr>
<tr>
<td>70</td>
<td>1,237</td>
<td>0,495</td>
</tr>
<tr>
<td>75</td>
<td>1,275</td>
<td>0,510</td>
</tr>
<tr>
<td>80</td>
<td>1,312</td>
<td>0,525</td>
</tr>
</tbody>
</table>

NOTE All values of reproducibility are expressed on the basis of fused, i.e. loss-on-ignition-free, samples.

13 Calculation and expression of results

Calculate the concentrations on a fused basis from the calibration curve (see 12.3.1), corrected, where necessary, for any inter-element effect (see 12.3.2).

Calculate the results, where a single test result has been obtained, as a percentage mass fraction to four significant figures when the value exceeds 1,00 % mass fraction, or to three decimal places for values below 1,00 % mass fraction.

Calculate the results, where two test results have been obtained, as the mean of the results, as a percentage mass fraction to four significant figures when the value exceeds 1,00 % mass fraction, or to three decimal places for values below 1,00 % mass fraction.

If the two test results differ by more than the appropriate repeatability limits set out in Table 1, repeat the test and take the mean of the two closest test results.
Convert test results on a fused basis to results on the as-received basis using the factor, $f_{\text{LOI}}$, in accordance with 10.2.3.

Where the sample contains oxidizable sulfur species, or any chloride or bromide, see 10.3 with regard to totalling the oxide analysis.

The results of all individual tests shall be recorded.

Express the results, generally as the oxide, to two decimal places.

14 Performance criteria (repeatability, accuracy and reproducibility limits)

Performance criteria are given in Tables 1, 2 and 3.
Annex A  
(informative)

Examples of fluxes

There are three examples of fluxes that have proved suitable:

a) 66 % lithium tetraborate + 34 % lithium metaborate; purity 99,95 % (minimum);

b) 100 % lithium tetraborate; nominal 100 %;

c) lithium tetraborate/lithium metaborate, in ratios of 100/0 to 0/100 %; purity 99,98 %.
Annex B
(informative)

Sources of certified reference materials

Examples of sources of certified reference materials include the following:

- BCR Community Bureau of Reference (Belgium);
- BCS British Chemical Standards (UK);
- CANMET Canada Centre for Mineral and Energy Technology (Canada);
- DL Dillinger Laboratory (Germany);
- ECRM Euronorm Certified Reference Materials (EU);
- GBW State Bureau of Technical Supervision (China);
- JSS Iron and Steel Institute of Japan (Japan);
- NCB National Council for Cement and Building Materials (INDIA);
- NIST National Institute of Standards and Technology (USA);
- SARM South African Bureau of Standards (South Africa); and
- Glass beads BREITLÄNDER Eichproben + Labormaterial GmbH (Germany)\(^1\).

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1) This is an example of a suitable commercial supplier. This information is given for the convenience of users of this part of ISO 29581 and does not constitute an endorsement by ISO.
### Annex C
(informative)

Examples of calibration standards and monitor beads and pellets

Table C.1 — Calibration standards for the calibration of analysis of CEM I cements

<table>
<thead>
<tr>
<th>Reagent</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiO₂</td>
<td>0.240</td>
<td>0.195</td>
<td>0.204</td>
<td>0.223</td>
<td>0.180</td>
<td>0.187</td>
<td>0.214</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>0.072</td>
<td>0.081</td>
<td>0.026</td>
<td>0.054</td>
<td>0.045</td>
<td>0.063</td>
<td>0.036</td>
</tr>
<tr>
<td>Fe₂O₃</td>
<td>0.015</td>
<td>0.061</td>
<td>0.068</td>
<td>0.042</td>
<td>0.053</td>
<td>0.023</td>
<td>0.034</td>
</tr>
<tr>
<td>CaCO₃</td>
<td>1.042</td>
<td>1.091</td>
<td>1.096</td>
<td>1.148</td>
<td>1.135</td>
<td>1.188</td>
<td>1.199</td>
</tr>
<tr>
<td>MgO</td>
<td>0.011</td>
<td>0.017</td>
<td>0.040</td>
<td>0.026</td>
<td>0.022</td>
<td>0.032</td>
<td>0.005</td>
</tr>
<tr>
<td>K₂CO₃</td>
<td>0.014</td>
<td>0.005</td>
<td>0.002</td>
<td>0.011</td>
<td>0.008</td>
<td>0.017</td>
<td>—</td>
</tr>
<tr>
<td>CaSO₄</td>
<td>0.068</td>
<td>0.030</td>
<td>0.064</td>
<td>0.003</td>
<td>0.057</td>
<td>0.017</td>
<td>0.039</td>
</tr>
<tr>
<td>Totals</td>
<td>1.462</td>
<td>1.481</td>
<td>1.482</td>
<td>1.508</td>
<td>1.502</td>
<td>1.528</td>
<td>1.527</td>
</tr>
</tbody>
</table>

Table C.2 — Composition of the calibration standards listed in Table C.1

<table>
<thead>
<tr>
<th>Oxide</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiO₂</td>
<td>24.0</td>
<td>19.5</td>
<td>20.4</td>
<td>22.3</td>
<td>18.0</td>
<td>18.7</td>
<td>21.4</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>7.2</td>
<td>8.1</td>
<td>2.6</td>
<td>5.4</td>
<td>4.5</td>
<td>6.3</td>
<td>3.6</td>
</tr>
<tr>
<td>Fe₂O₃</td>
<td>1.5</td>
<td>6.1</td>
<td>6.8</td>
<td>4.2</td>
<td>5.3</td>
<td>2.3</td>
<td>3.4</td>
</tr>
<tr>
<td>CaO</td>
<td>61.2</td>
<td>62.4</td>
<td>63.3</td>
<td>64.5</td>
<td>66.0</td>
<td>67.3</td>
<td>68.8</td>
</tr>
<tr>
<td>MgO</td>
<td>1.1</td>
<td>1.7</td>
<td>4.0</td>
<td>2.6</td>
<td>2.2</td>
<td>3.2</td>
<td>0.5</td>
</tr>
<tr>
<td>K₂O</td>
<td>1.0</td>
<td>0.4</td>
<td>0.2</td>
<td>0.8</td>
<td>0.6</td>
<td>1.2</td>
<td>0.0</td>
</tr>
<tr>
<td>SO₃</td>
<td>4.0</td>
<td>1.8</td>
<td>2.7</td>
<td>0.2</td>
<td>3.4</td>
<td>1.0</td>
<td>2.3</td>
</tr>
<tr>
<td>Totals</td>
<td>100.0</td>
<td>100.0</td>
<td>100.0</td>
<td>100.0</td>
<td>100.0</td>
<td>100.0</td>
<td>100.0</td>
</tr>
</tbody>
</table>
Annex D
(informative)

Determination of the sulfate content
of samples containing sulfide species

D.1 Principle

Sulfate ions, produced by the decomposition of cement with hydrochloric acid, are precipitated at a pH between 1.0 and 1.5 by a solution of barium chloride. The precipitation of barium sulfate is carried out at boiling point.

The determination is then completed gravimetrically and sulfate expressed as SO_3.

D.2 Reagents and equipment

D.2.1 Hydrochloric acid (HCl), concentrated, density 1.18 to 1.19.

D.2.2 Hydrochloric acid, dilute, 1 + 11.

D.2.3 Ammonium hydroxide, dilute, 1 + 16.

D.2.4 Barium chloride, solution, prepared by dissolving 120 g of barium chloride (BaCl_2·2H_2O) in water and making up to 1 000 ml.

D.2.5 Silver nitrate, solution, prepared by dissolving 5 g of silver nitrate (AgNO_3) in water, adding 10 ml of concentrated nitric acid (HNO_3) and making up to 1 000 ml with water.

D.2.6 Filter paper, ashless.

NOTE Filter papers with a mean pore diameter of around 2 µm are termed fine, those with a mean pore diameter of around 7 µm are termed medium and those with a mean pore diameter of around 20 µm are termed coarse.

D.3 Procedure

Weigh, to ±0.0005 g, (1.00 ± 0.05) g of cement and record this mass as m_5. Place the cement sample in a 250 ml beaker, and add 90 ml of water. While stirring the mixture vigorously, add 10 ml of concentrated hydrochloric acid (D.2.1). Heat the solution gently and crush the sample with the flattened end of a glass stirring rod until decomposition is complete. Allow the solution to digest for 15 min at a temperature just below boiling.

Filter the residue on a medium filter paper (D.2.6) into a 400 ml beaker. Wash thoroughly with hot water until free from chloride ions, as determined by the silver nitrate test (D.7).

Adjust the volume to about 250 ml; if necessary, adjust the pH of the solution to between 1.0 and 1.5 with dilute hydrochloric acid 1 + 11 (D.2.2) or ammonium hydroxide 1 + 16 (D.2.3).

Bring to a boil and boil for 5 min. Check that the solution is clear; if not, start the determination again using a new test portion. While stirring vigorously, maintain the solution at the boiling point and add, drop by drop, 10 ml of the barium chloride solution (D.2.4) heated to just below boiling. Maintain the solution at just below boiling point for at least 30 min, ensuring that the volume is kept between 225 ml and 250 ml, and then allow the covered beaker to stand at room temperature for 12 h to 24 h before filtration.
Filter the precipitate on a fine filter paper (D.2.6) and wash with boiling water until free from chloride ions, as determined by the silver nitrate test (D.7).

Ignite at (950 ± 25) °C to constant mass (D.6) and record this mass as $m_6$.

NOTE In general, an ignition period of 15 min is sufficient to achieve constant mass.

D.4 Calculation and expression of results

Calculate the sulfate content, expressed as SO$_3$, in percent, using the equation:

$$w_{SO_3} = \frac{m_6 \times 0.343 \times 100}{m_5} = 34.3 \times \frac{m_6}{m_5}$$  \hspace{1cm} (D.1)

where

- $m_5$ is the mass of the test portion, expressed in grams;
- $m_6$ is the mass of barium sulfate, expressed in grams.

D.5 Repeatability and reproducibility

The standard deviation for repeatability is 0.07 % mass fraction.

The standard deviation for reproducibility is 0.08 % mass fraction.

D.6 Determination of constant mass

Determine constant mass by 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.

D.7 Check for absence of chloride ions (silver nitrate test)

After five to six washes of a precipitate, rinse the base of the filter stem with a few drops of water. Wash the filter paper and its contents with several millilitres of water and collect this in a test tube. Add several drops of silver nitrate solution (D.2.5). Check the absence of turbidity or precipitate in the solution. If present, continue washing while carrying out periodic checks until the silver nitrate test is negative.
Bibliography

[1] ISO 5725-1, Accuracy (trueness and precision) of measurement methods and results — Part 1: General principles and definitions