INTERNATIONAL STANDARD

Second edition 1990-12-15

Corrected and reprinted 1991-03-15

Lead chromate pigments and lead chromate-molybdate pigments — Specifications and methods of test

iTeh STANDARD PREVIEW

Rigments à base de chromate et de chromomolybdate de plomb — Spécifications et méthodes d'essai

ISO 3711:1990 https://standards.iteh.ai/catalog/standards/sist/366d4c1e-6731-433e-adfefb4ec57861f2/iso-3711-1990



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 EVIEW casting a vote.

International Standard ISO 3711 was prepared by Technical Committee 1) ISO/TC 35, Paints and varnishes.

This second edition cancels and replaces the <u>37</u> first <u>290</u> edition (ISO 3711:1976), of which it constitutes a technical advection devision and sist <u>366d4c1e-6731-433e-adfe-fb4ec57861f2</u> iso-3711-1990

© ISO 1990

Case Postale 56 ● CH-1211 Genève 20 ● Switzerland

Printed in Switzerland

All rights reserved. No part of this publication may be reproduced or utilized in any form or by any means, electronic or mechanical, including photocopying and microfilm, without permission in writing from the publisher.

International Organization for Standardization

Lead chromate pigments and lead chromate-molybdate pigments — Specifications and methods of test

Scope 1

This International Standard specifies the requirements and the corresponding methods of test for the following pigments identified by Colour Index numbers¹): orange 21, yellow 34 and red 104. These pigments are suitable for general use.

ISO 787-4:1981, General methods of test for pigments and extenders — Part 4: Determination of acidity or alkalinity of the aqueous extract.

ISO 787-5:1980, General methods of test for pigments and extenders - Part 5: Determination of oil absorption value.

ISO 787-7:1981, General methods of test for pigments The chemical identity of these pigments is NOTE 1 and extenders ____ Part 7: Determination of residue on given in table 1. iTeh STANDARI sieve - Water method -- Manual procedure.

Normative references 2

(standards.itso 7878) 1979, General methods of test for pigments and extenders — Part 8: Determination of matter The following standards contain provisions which 711:1995 oluble in water - Cold extraction method.

through reference in this text a constitute provisions and sist of this International Standard. At the time of public/iso-37 cation, 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 385-1:1984, Laboratory glassware – Burettes – Part 1: General requirements.

ISO 648:1977, Laboratory glassware – One-mark pipettes.

ISO 787-1:1982, General methods of test for pigments and extenders - Part 1: Comparison of colour of pigments.

ISO 787-2:1981, General methods of test for pigments and extenders — Part 2: Determination of matter volatile at 105 °C. 150,787-9:1981, General methods of test for pigments and extenders — Part 9: Determination of pH value of an aqueous suspension.

ISO 787-15:1986, General methods of test for pigments and extenders - Part 15: Comparison of resistance to light of coloured pigments of similar types.

ISO 787-16:1986, General methods of test for pigments and extenders - Part 16: Determination of relative tinting strength (or equivalent colouring value) and colour on reduction of coloured pigments - Visual comparison method.

ISO 787-20:1975, General methods of test for pigments — Part 20: Comparison of ease of dispersion (Oscillatory shaking method).

ISO 842:1984, Raw materials for paints and varnishes - Sampling.

and the

¹⁾ The Colour Index is published by The Society of Dyers and Colourists, PO Box 244, Perkin House, 82 Grattan Road, Bradford, West Yorkshire BD1 2JB, United Kingdom

American Association of Textile Chemists and Colorists, National Headquarters, Box 12215, Research Triangle Park, NC 27709, USA.

ISO 1042:1983, Laboratory glassware — One-mark volumetric flasks.

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

ISO 3856-1:1984, Paints and varnishes — Determination of "soluble" metal content — Part 1: Determination of lead content — Flame atomic absorption spectrometric method and dithizone spectrophotometric method.

ISO 4793:1980, Laboratory sintered (fritted) filters — Porosity grading, classification and designation.

3 Classification

In this International Standard, lead chromate pigments and lead chromate-molybdate pigments are classified as one of the following two types:

Standard type (type 1): Yellow to red pigments consisting of normal or basic lead chromate with or without lead sulfate and/or molybdate or other coprecipitated water-insoluble compounds of lead. Such pigments shall be free from organic colouring matter and shall not contain extenders. Pigments corresponding to pigment yellow Colour Index No. 34 and pigment red Colour Index No. 104 may con-

solely for the control of the crystal structure of the pigment.

Stabilized type (type 2): Yellow to red pigments consisting of normal or basic lead chromate with or without lead sulfate and/or molybdate or other coprecipitated water-insoluble compounds of lead. These pigments contain other materials introduced during manufacture specifically to improve certain pigmentary properties. They shall be free from organic colouring matter and shall not contain extenders. If type 2 is specified, the purchaser may require the vendor to state the nature of the improved properties which are claimed to result from the additions and to declare the minimum content of total lead.

4 Required characteristics and associated tolerances

4.1 For lead chromate pigments and lead chromate-molybdate pigments complying with this International Standard, the essential requirements are specified in table 2 and the conditional requirements are listed in table 3. The reference pigment and the conditional requirements listed in table 3 shall be specified by agreement between the interested parties.

tain co-precipitated compounds of, for example, al- **4.2** The agreed reference pigment shall comply uminium and/or silicon when these are required ISO 3 with the requirements of table 2.

https://standards.iteh.ai/catalog/standards/sist/366d4c1e-6731-433e-adfefb4ec57861f2/iso-3711-1990

Type of pigment	Shade of pigment	Colour Index No.	Chemical identity	
	Primrose and lemon	Pigment yellow No. 34, Part 2, Ref. 77 603	Lead sulfochromate	
Lead chromate	Yellow	Pigment yellow No. 34, Part 2, Ref. 77 600	Lead chromate	
	Orange	Pigment orange No. 21, Part 2, Ref. 77 601	Basic lead chromate	
Lead chromate- molybdate	Orange to red	Pigment red No. 104, Part 2, Ref. 77 605	Lead sulfochromate-molybdate	

Table 1 — Classification of lead chromate pigments and lead ch	romate-molybdate pigments
--	---------------------------

Characteristic	Unit	Requirement	Method of test
Matter volatile at 105 °C	% (m/m)	max. 1	ISO 787-2
Matter soluble in water (cold extraction method)	% (<i>m/m</i>)	max. 2	ISO 787-8, taking a test portion of 20 g
Acidity or alkalinity of the aqueous extract	ml of 0,1 mol/l solution per 100 g of pigment	max. 20	ISO 787-4, taking a test portion of 20 g
pH value of aqueous suspension		4 to 8	ISO 787-9
Residue on sieve (45 µm)	% (m/m)	max. 0,3	ISO 787-7

Table 2 — Essential requirements

Table 3 — Conditional requirements

Characteristic	Unit	Requirement	Method of test	
Colour		Equal to that of the agreed refer-	ISO 787-1	
Colour on reduction	STAN	ence pigment (see 4.2) to within a tolerance agreed between the in- terested parties	ISO 787-16	
Relative tinting strength	(stand	lards.iteh.ai)		
Ease of dispersion https://standard		Shall not be inferior to that of the agreed reference pigment (see 4.2) gstandards/sist/300d4c1e-6/31-433e-adfe- 7861f2/iso-3711-1990	ISO 787-20, measuring fineness of grind after 2,5 min, 5 min and thereafter every 5 min.	
Resistance to light		Shall not be inferior to that of the agreed reference pigment (see 4.2)	ISO 787-15	
Oil absorption value		Shall not differ by more than 15 % from the agreed value	ISO 787-5	
Total lead content as Pb	% (m/m)	Shall not differ by more than 3% (<i>m</i> / <i>m</i>) from the agreed value	See clause 6	
Soluble lead content as Pb in 0,07 mol/l HCl	% (m/m)	If required, to be agreed between the interested parties	See clause 7	

5 Sampling

Take a representative sample of the product to be tested, as described in ISO 842.

6 Determination of total lead content

For the determination of the total lead content, two methods (one gravimetric, the other titrimetric) are specified. The method to be used shall be agreed on between the interested parties. The gravimetric method (6.1) shall be used as the referee method in cases of dispute.

6.1 Gravimetric method

6.1.1 Principle

Dissolution of the test portion in hydrochloric acid. Separation of the lead as lead sulfide and gravimetric determination of the lead after precipitation as lead sulfate and subsequent extraction with ammonium acetate.

6.1.2 Reagents

During the analysis, use only reagents of recognized analytical grade and only water of at least grade 3 purity as defined in ISO 3696. WARNING --- Use the reagents in accordance with the appropriate health and safety regulations.

6.1.2.1 Ammonium acetate, crystals,

6.1.2.2 Hydrogen sulfide.

6.1.2.3 Tartaric acid.

6.1.2.4 Nitric acid/bromine reagent.

Saturate nitric acid, $c(HNO_3) = 4 \text{ mol/l}$, with bromine.

6.1.2.5 Hydrochloric acid, concentrated, approximately 37 % (m/m), $\rho \approx 1.19$ g/mI.

6.1.2.6 Sulfuric acid, concentrated, approximately 96 % (*m/m*), $\rho \approx$ 1,84 g/ml.

6.1.2.7 Sulfuric acid, dilute.

Dilute 5 ml of the concentrated sulfuric acid (6.1.2.6) by adding it to water, with cooling, and making up to 100 ml with water.

6.1.2.8 Ethanol, 95 % (V/V).

6.1.2.9 Sodium sulfide, 50 g/l solution, freshly prepared.

iTeh STANDA

(standai

6.1.2.10 Ammonium acetate, saturated solution fractional transfer it to the crucible with a jet of the same water

6.1.2.11 Ammonia solution. approximately 25 % (*m/m*), $\rho \approx 0.91$ g/ml.

6.1.3 Apparatus

Use ordinary laboratory apparatus and glassware complying with the requirements of the relevant International Standards (see clause 2), together with the following.

6.1.3.1 Sintered-glass crucibles, grade P16, complying with the requirements of ISO 4793.

6.1.3.2 Sintered-silica crucibles, grade P16, complying with the requirements of ISO 4793.

6.1.3.3 Drying oven, capable of being maintained at 105 °C ± 2 °C.

6.1.3.4 Muffle furnace, capable of being maintained at 500 °C ± 25 °C.

6.1.4 Procedure

Carry out the determination in duplicate.

6.1.4.1 Test portion

Weigh out, to the nearest 0,1 mg, about 0,5 g of the sample (see clause 5).

6.1.4.2 Determination

Place the test portion (6.1.4.1) in a 600 ml beaker and add, as a reducing agent, 2 ml of ethanol (6.1.2.8), followed by 100 ml of water and 15 ml of hydrochloric acid (6.1.2.5). Cover the beaker with a watch-glass and heat to boiling. Boil gently until all odour of aldehyde has disappeared. Dilute to 200 ml with hot water. Filter the solution whilst hot through a fine filter paper and wash the filter and residue well with hot water until a few drops of the filtrate give no coloration with sodium sulfide solution (6.1.2.9). Combine the filtrate and washings and allow to cool to room temperature.

Slowly add ammonia solution (6.1.2.11), whilst stirring, until a faint permanent precipitate forms. Then add 0,5 g of tartaric acid (6.1.2.3) and hydrochloric acid (6.1.2.5), until the pH-value of the solution is between 1 and 2. Pass hydrogen sulfide (6.1.2.2) through the solution in a fume cupboard until it is saturated. Dilute the solution to 400 ml and again saturate with hydrogen sulfide. Allow the precipitated lead sulfide to settle, preferably overnight, and pour off the clear supernatant liquor through a sintered-glass crucible (6.1.3.1), using gentle

ISO 3suction Wash the precipitate once by decantation https://standards.iteh.ai/catalog/stanWilth/swaterd4saturated13with16hydrogen sulfide and using a rubber-tipped glass rod to aid the transfer. Wash the precipitate on the crucible five times with water saturated with hydrogen sulfide and discard the filtrate and washings.

> Place the sintered-glass crucible in the original 600 ml beaker and add nitric acid/bromine reagent (6.1.2.4) to dissolve the lead sulfide precipitate. Wash the crucible with hot water, collecting the washings in the same beaker. Cover the beaker. heat the contents in a fume cupboard, and filter through another sintered-glass crucible (6.1.3.1).

> Wash the beaker, the cover and the crucible five times with hot water, transferring the filtrate and washings to a second 600 ml beaker. Cool the solution, add 15 ml of concentrated sulfuric acid (6.1.2.6), and evaporate the solution carefully until dense white fumes are evolved. Cool the beaker and contents, wash down the sides with water and reevaporate the contents until white fumes are again evolved. Cool the beaker again, add 40 ml of ethanol (6.1.2.8) and 75 ml of water and allow the whole to stand overnight.

> Pour off the clear liquor through a tared sinteredsilica crucible (6.1.3.2), wash the precipitate once by decantation with a mixture of equal parts of ethanol and dilute sulfuric acid (6.1.2.7) and transfer to the

crucible by a jet of the same wash fluid, using a rubber-tipped glass rod to aid the transfer. Wash the precipitate with ethanol until the washings are neutral, dry the crucible and precipitate in the drying oven (6.1.3.3) and heat the crucible at dull red heat $(500 \degree C \pm 25 \degree C)$ to constant mass in the muffle furnace (6.1.3.4). Allow to cool in a desiccator and weigh.

Fill the crucible with ammonium acetate crystals (6.1.2.1) and slowly pour 50 ml of boiling ammonium acetate solution (6.1.2.10) through it. Wash very thoroughly with hot water until a few drops of the filtrate give no coloration with sodium sulfide solution, dry, ignite, cool and reweigh as before.

6.1.5 Expression of results

6.1.5.1 Calculation

Calculate the total lead content w(Pb) of the pigment, expressed as a percentage by mass, using the equation

$w(Pb) = \frac{0,6832 \times m_1}{m_0} \times \frac{100}{100} eh \text{ STANDARD } 70\% (m/m), p \approx 142 \text{ g/ml}.$

where

(standards, if 6.2.2.2, Hydrogen peroxide, 30 % (m/m) solution.

m_0	is the mass, in grams, of thester 1:19	0.2.2.3 990	Urea.
	portion; https://standards.iteh.ai/catalog/standards/		Copper nitrate
m_1	is the mass, in grams, of the precip- itate (PbSO ₄), i.e. the difference be-	711-1990 6.2.2.5	Acetone.
	tween the two weighings before and after extraction with ammonium acetate;		EDTA, standa) = 0,025 mol/l.

is the factor for the conversion of 0.6832 grams of PbSO₄ to grams of Pb.

If it is required to express the total lead content as a percentage by mass of PbO, use the equation

$$w(PbO) = \frac{0.736 \times m_1}{m_0} \times 100$$

where

is the total lead content of the w(PbO) pigment, expressed as a percentage by mass of PbO;

 m_0 and m_1 are as defined above;

0,736 is the factor for the conversion of grams of
$$PbSO_4$$
 to grams of PbO .

If the two determinations differ by more than 0.5 % (m/m) Pb, repeat the procedure.

Calculate the mean of two valid determinations and report the result to the nearest 0,1 % (m/m).

6.1.5.2 Precision

No precision data are currently available.

6.2 Titrimetric method (using electrolytic separation)

6.2.1 Principle

Dissolution of the test portion in nitric acid. Separation of the lead by electrolysis as lead dioxide. Dissolution of the lead dioxide deposit in nitric acid and hydrogen peroxide, and complexometric titration of the lead with EDTA solution.

6.2.2 Reagents

During the analysis, use only reagents of recognized analytical grade and only water of at least grade 3 purity as defined in ISO 3696.

WARNING — Use the reagents in accordance with the appropriate health and safety regulations.

6.2.2.1 Nitric acid, concentrated, approximately

volumetric solution, Idard 1

Take an ordinary standard solution or dissolve 9.306.3 g of disodium ethylenediamine tetraacetate dhydrate (EDTA disodium salt) in water in a 1000 ml one-mark volumetric flask, dilute to the mark and mix well.

6.2.2.7 Xylenol orange mixture.

Add 1 g of xylenol orange to 100 g of sodium chloride (or potassium nitrate) and mix well.

6.2.2.8 Hexamethylene tetramine.

6.2.3 Apparatus

Use ordinary laboratory apparatus and glassware complying with the requirements of the relevant International Standards (see clause 2), together with the following.

6.2.3.1 Electrolysis equipment.

6.2.3.2 Platinum electrode.

6.2.4 Procedure

Carry out the determination in duplicate.

6.2.4.1 Test portion

Weigh out, to the nearest 0,1 mg, about 0,25 g of the sample (see clause 5).

6.2.4.2 Preparation of the test solution

Place the test portion (6.2.4.1) in a 250 ml tall-form beaker and add 10 ml of nitric acid (6.2.2.1), 10 ml of water and a small amount of copper nitrate (6.2.2.4).

Heat until the test portion is dissolved. Add 10 drops of hydrogen peroxide solution (6.2.2.2) and boil gently for 10 min. Dilute with about 50 ml of water and add 5 ml of nitric acid and a small amount of urea (6.2.2.3). Heat to boiling.

NOTE 2 A small insoluble residue may be left at this stage, but it is not relevant to the determination.

6.2.4.3 Separation of lead by electrolysis, as PbO₂

NOTE 3 The optimum voltages for the electrolysis are dependent on the equipment used. They may be between dal 1,6 V and 2,8 V. Thus, in order to determine the optimum voltage to be applied for the equipment used, a prelimi-

voltage to be applied for the equipment used, a preliminary analysis should be carried out, using lead chromate $\frac{150.3711:1990}{1000}$ b) = $\frac{V \times 0.00518}{4 \times 100} \times 100$ with known lead content. The optimum/voltage is that at log/standards/sist/36644c1e- $\frac{1200}{100}$ and $\frac{1200}{100} \times 100$ which the deposition of lead dioxide is total, without the 5786112/iso-3711-1990 occurrence of significant gassing.

The voltages given in the following description should be regarded as examples only.

Before the electrolysis, dip the platinum electrode (anode) (6.2.3.2) in acetone (6.2.2.5) in order to remove any grease. Start the electrolysis, without any stirring, at 2 V (see note 3). After several seconds, when a deposit has appeared, reduce the voltage to 1,6 V to 1,7 V (see note 3). Continue the electrolysis, with gentle stirring, at about 80 °C for 30 min. After 30 min, raise the beaker by about 1 cm to 2 cm and continue the electrolysis for a further 15 min. Then lower the beaker to its original level and visually examine the exposed electrode for deposition of lead dioxide. If deposition has occurred, continue the electolysis for an additional 15 min.

After completion of the electrolysis, slowly raise the electrode out of the solution without changing the applied voltage. Simultaneously, rinse the electrode with a jet of water from a wash-bottle.

6.2.4.4 Determination of lead by complexometric titration

NOTE 4 When present, molybdenum and antimony are partly deposited as oxides during the electrolysis. How-

ever, these elements have no influence on the titrimetric determination of lead.

Immerse the platinum electrode with the lead dioxide deposit (see 6.2.4.3) in a 250 ml beaker containing 150 ml of water, 3 ml of nitric acid (6.2.2.1) and 2 ml of hydrogen peroxide solution (6.2.2.2), for 15 min. The deposit will dissolve rapidly. Remove the platinum electrode from the solution, at the same time rinsing it with water. Decompose any remaining hydrogen peroxide by evaporating the solution to about 50 ml. Then add about 100 ml of water.

After addition of 0,1 g of xylenol orange mixture (6.2.2.7), add small portions of hexamethylene tetramine (6.2.2.8) until the colour changes from yellow to pink. Then add an excess of 0,4 g to 0,5 g of hexamethylene tetramine. Afterwards, whilst stirring, titrate with EDTA solution (6.2.2.6) until the colour just changes to yellow.

6.2.5 Expression of results

6.2.5.1 Calculation

Calculate the total lead content w(Pb) of the pigment expressed as a percentage by mass, using the equation

- *m*₂ is the mass, in grams, of the test portion;
- *V* is the volume, in millilitres, of EDTA solution (6.2.2.6) used;
- 0,00518 is the factor for the conversion of millilitres of EDTA solution, c(EDTA) = 0,025 mol/I, to grams of Pb.

If it is required to express the total lead content as a percentage by mass of PbO, use the equation

$$w(PbO) = \frac{V \times 0,00558}{m_2} \times 100$$

where

w(PbO) is the total lead content of the pigment, expressed as a percentage by mass of PbO;

 m_2 and V are as defined above;

0,00558 is the factor for the conversion of millilitres of EDTA solution, c(EDTA) = 0,025 mol/l, to grams of PbO.

If the two determinations differ by more than 0.8 % (m/m) Pb, repeat the procedure.

Calculate the mean of two valid determinations and report the result to the nearest 0.1 % (m/m).

6.2.5.2 Precision

a) Repeatability (r)

The value below which the difference between two test results obtained on identical material within a short interval of time by one operator in one laboratory using the same equipment and the standardized test method may be expected with 95 % probability to lie is 0,85 %.

b) **Reproducibility** (R)

The value below which the difference between two test results obtained on identical material by operators in different laboratories using the standardized test method may be expected with 95% probability to lie is 2,0%.

7 Determination of soluble lead content

KD The extraction method described in 7.1 has NOTE 5 been derived from ISO 6713:1984, Paints and varnishes - Is. iteh Preparation of acid extracts from paints in liquid or powder form.

ISO 3711:1990

where

7.1 Extraction of the two luble read and a design a design and a design and a design and a design a de lead chrome pigments, expressed as a fb4ec57861f2/iso-3711-1990

7.1.1 Reagents

During the analysis, use only reagents of recognized analytical grade and only water of at least grade 3 purity as defined in ISO 3696.

WARNING - Use the reagents in accordance with the appropriate health and safety regulations.

7.1.1.1 Hydrochloric acid. dilute, c(HCI) =0.07 mol/l.

7.1.1.2 Hydrochloric acid, diluted 1 + 1.

Add 1 part by volume of concentrated hydrochloric acid [approximately 37 % (m/m), $\rho \approx$ 1,19 g/ml] to 1 part by volume of water.

7.1.2 Apparatus

Use ordinary laboratory apparatus and glassware complying with the requirements of the relevant International Standards (see clause 2), together with the following.

7.1.2.1 Suitable mechanical stirrer (see 7.1.3.2, third paragraph).

7.1.2.2 pH meter with electrodes.

7.1.2.3 Membrane filter, pore diameter 0.15 um, or other suitable filter capable of giving a clear filtrate in 7.1.3.2.

7.1.2.4 Filtration apparatus, for the membrane filter (7.1.2.3).

7.1.2.5 Water bath, capable of being maintained at $23 \degree C \pm 2 \degree C$.

7.1.3 Procedure

7.1.3.1 Test portion

Determine the total lead content of the pigment as described in clause 6.

From the total lead content of the pigment, calculate the mass m_3 , in grams, of the test portion for the determination of the soluble lead content, using the equation

$$m_3 = \frac{60}{w(Pb)} \times 0.5$$

w(Pb) is the total lead content of the pigment. expressed as a percentage by mass;

percentage by mass.

7.1.3.2 Extraction

Carry out the extraction of the pigment in duplicate.

Weigh out the test portion (m_3) to the nearest 1 mg and place it into a clean, dry 1000 ml beaker. Fit the stirrer (7.1.2.1) and add 500 ml of dilute hydrochloric acid (7.1.1.1), previously brought to $23 \degree C \pm 2 \degree C$ by means of the water bath (7.1.2.5). Place the beaker into the water bath and immediately commence stirring the mixture (see next paragraph). Insert the electrodes of the pH meter (7.1.2.2) into the mixture and, if necessary, adjust the pH to that of dilute hydrochloric acid (7.1.1.1) by adding 1 + 1hydrochloric acid (7.1.1.2).

During the whole period of extraction, keep the speed of the stirrer adjusted so that the pigment is kept in continuous suspension whilst taking care to avoid splashing.

Continue stirring for 60 min \pm 1 min, checking that the mixture is maintained at 23 °C \pm 2 °C throughout the test period. Maintain the pH of the mixture constant by carefully adding 1 + 1 hydrochloric acid (7.1.1.2). At the end of the period of stirring, allow the mixture to stand for a further 60 min \pm 1 min at