TC

INTERNATIONAL STANDARD

Surface active agents – Evaluation of certain effects of laundering – Methods of analysis and test for unsoiled cotton control cloth

INTERNATIONAL ORGANIZATION FOR STANDARDIZATIONOMEXDYHAPODHAR OPFAHM3AUMR ПО СТАНДАРТИЗАЦИИФОRGANISATION INTERNATIONALE DE NORMALISATION

Agents de surface – Contrôle de certains effets de blanchissage – Méthodes d'analyse et d'essai d'un tissu de coton témoin non souille

(standards.iteh.ai)

First edition - 1979-02-01

<u>ISO 4312:1979</u> https://standards.iteh.ai/catalog/standards/sist/b62059b5-34aa-45bf-8f70e8d9d4bd926b/iso-4312-1979

UDC 661.185 : 620.16

Ref. No. ISO 4312-1979 (E)

Descriptors : surfactants, tests, washing tests, wear tests, cotton fabrics, test specimens.

с	CONTENTS			e
0		ntroduction		1
1	S	cope and field of application	. '	1
2	F	References	. 1	1
3	S	election of specimens and samples from test pieces	. 2	2
4	E	Determination of intrinsic greying (greying measured without ultraviolet		-
А	1	Scope and field of application		<u>'</u>
4.	· · ·			2
iTeh S	Í	ANDARD PREVIEW		2
4.	.3 St	andards.iteh.ai)		3
4	.4	Apparatus.	. :	3
4.	.5	Procedure <u>180 4312:1979</u>		3
https://standards.4	¢6.8	a Expressionadi resultis b6205965-34aa-45bt-81/0- e8d9d4bd926b/iso-4312-1979		3
5	C V	Determination of intrinsic yellowing (yellowing measured without ultra- iolet radiation)	4	ł
5.	.1	Scope and field of application	. 4	1
5.	.2	Definitions	4	ł
5.	.3	Principle	. 4	ł
5.	.4	Apparatus	4	ł
5.	.5	Procedure	. 4	ł
5.	.6	Expression of results	. 5	5
6	D	Determination of increase in organic depositions content	. 5	5
6.	.1	Scope and field of application	. 5	5
6.	.2	Principle	. 5	5
6.	.3	Reagent	. 5	5
6.	.4	Apparatus	5	5
6.	.5	Procedure	5	;
6.	.6	Expression of results	6	5

ANNEXES

Α	Determination on increase (or decrease) in degree of whiteness	15
В	Determination of chemical damage based on variation in average degree of viscometric polymerization in an ammonium copper(II) hydroxide solution	17
С	Determination of chemical damage based on variation in average degree of viscometric polymerization in a copper(II) diethylenediamine (Cuen) solution	19
F	IGURES	
1	Viscometer	22
2	Supporting tube for viscometer	22
3	Mean degree of viscometric polymerization as a function of fluidity value	23
4	Examples of vessels for dissolution of cellulose (C.5.1)	24
5	Curve showing the variation in the viscosity relative increment of cellulose solutions in the Cuen solution as a function of the product of the limiting viscosity number and the cellulose concentration in the solutions.	25

iTeh STANDARD PREVIEW (standards.iteh.ai)

<u>ISO 4312:1979</u>

https://standards.iteh.ai/catalog/standards/sist/b62059b5-34aa-45bf-8f70e8d9d4bd926b/iso-4312-1979

Surface active agents — Evaluation of certain effects of laundering — Methods of analysis and test for unsoiled cotton control cloth

0 INTRODUCTION

It will be recalled that some effects produced by laundering on textiles can be evaluated by means of test pieces of unsoiled cotton control cloth which are washed together with normally soiled textile articles. It is impossible to effect this evaluation with normally soiled textile articles themselves since these are subject to damage and to modification due to actual use between successive launderings and which cannot be controlled.

This damage and these modifications are not part of the effects of laundering and, furthermore, their extreme variability would make the measurements very difficult to reproduce and even harder to compare between one.itch.ai) laboratory and another. In addition, it is hardly ever possible to operate with normally soiled textiles having 1070

The use of the test pieces of strictly defined unsolided-43 cotton control cloth enables the causes of variations which are extraneous to the laundering process itself to be largely eliminated. For this reason, the use of test pieces of cotton control cloth has been recognized as essential for verifying the laundering processes employed by industrial laundries.

Conclusions based on the behaviour of the unsoiled control cloth cannot be used to predict that of other textiles laundered in the same way, if these differ too much from the control cloth with regard to the nature of the fibres, the yarn linear density, the mass per unit surface, the presence of finishes or the initial degree of damage. In such cases, the results obtained on the control cloth can at most show qualitative differences between different laundering processes or variations of a process.

When the unsoiled cotton control cloth is used to determine the influence of a single factor (for example the type of washing machine, the nature or concentration of the detergent, the nature or concentration of the bleaching agent) from the point of view of the effects produced on the textiles, the comparisons are only valid, of course, so long as all the other factors are kept constant. In particular, care should be taken to see that the test pieces of cotton control cloth to be used come from the same consignment, that the normally soiled textiles are always of the same kind and have a uniform degree of soiling, and that the hardness of the water is always the same (so long as it is not precisely the influence of the hardness that is to be studied).

It follows from this that it is advisable in practice to compare the results obtained in different laboratories with one another only after a preliminary study has provided an assurance that all the conditions for comparison have been met.

In particular cases, it may be decided to determine only certain of the characteristics from among those the determination of which is described in this International Standard.

This International Standard should be read in conjunction with ISO 2267.

1 SCOPE AND FIELD OF APPLICATION

used to determine, under strictly controlled conditions, some characteristics of unsoiled cotton control cloth, namely intrinsic greying and yellowing, increase in organic depositions content and incineration residue, determination of overall loss of breaking strength, of loss of breaking strength resulting from chemical degradation of cellulose and of loss of breaking strength resulting from mechanical factors in laundering both before and after processing, so that certain effects of laundering can be evaluated.

> For the purpose of the routine assessment of the effects of processing on cotton textile articles in commercial laundries. the application of some of the methods only may be appropriate. Furthermore, while the methods given in this International Standard evaluate damage caused by excessive mechanical action, they do not distinguish between the effects of smaller differences in mechanical action on cleansing.

2 REFERENCES

ISO 139, Textiles - Standard atmospheres for conditioning and testing.

ISO/R 1628, Plastics - Directives for the standardization of methods for the determination of the dilute solution viscosity of polymers.

ISO 2267, Surface active agents - Evaluation of certain effects of laundering - Preparation and use of unsoiled cotton control cloth.

4 The degree of visible whiteness of white textiles is, of course, one of the factors that determine the quality of the laundering.

However, standardization of the measurement of the degree of visible whiteness is not sufficiently advanced to permit the inclusion of a method in this International Standard.

Annex A describes an empirical method which satisfies the majority of cases met with in practice.

4.3 Principle

Measurement of the reflectometer value (R_g) of the cotton control cloth before and after the 25 (or 50) laundering cycles, by means of a reflectometer (photoelectric colorimeter) fitted with a Y (green) tristimulus filter and under operating conditions selected so as to eliminate any fluorescent effect.

The decrease in the reflectometer value (perfect diffuser = 100), which is identical in this case to the decrease in the percentage luminance, is a measure of the intrinsic greying.

4.4 Apparatus

4.4.1 Reflectometer (photoelectric colorimeter), capable of being fitted with a Y (green) tristimulus filter which in conjunction with the photoelectric cell and light source, affords a spectral sensitivity corresponding to the CIE

colour-matching function \overline{y} for the C source and enabling 1979 Repeat the measurements on each test piece. any fluorescent effects to be excluded (see note 2/follow s/sist/b62059b5-34aa-45bf-8f70ing 4.2.5). e8d9d4bd926b/iso-43124.6)7Expression of results

NOTES

1 There are two possible methods that may be used to eliminate fluorescence. The first consists in placing the Y (green) tristimulus filter between the light source and the cloth to be examined. This prevents the ultra-violet and violet light required to excite fluorescence from reaching the cloth. The second consists in illuminating the cloth with white light, but using a filter which blocks the ultra-violet rays between the light source and the cloth; the Y (green) tristimulus filter is placed between the cloth and the photoelectric cell. The U.V. filter should block all radiations below 450 nm.

2 In theory, it is possible to obtain an identical result by calculations based on the spectral reflectance curve determined by means of a spectrophotometer. In this case, the data obtained from the spectrophotometric curve must be multiplied by the tristimulus values $(\overline{\nu} H)\lambda$ for standard source C and the products integrated over the whole of the spectrum. In this case also, any fluorescence must, of course, be eliminated.

4.4.2 Two calibration plates, for calibrating the reflectometer (4.4.1), one of neutral white with a percentage luminance between 85 and 90 and the other of neutral grey with a percentage luminance between 60 and 75. The calibration of these plates should be based on the perfect diffuser (= 100) for the Y (green) and Z (blue) tristimulus filters, under the measuring conditions of the reflectometer used.

Calibration plates in various materials are obtainable from various national calibration bodies. Bright materials are easier to maintain but dull materials give reflectometer values which are less dependent on the geometrical conditions of lighting and observation.

 ${\sf NOTE}$ — The use of a single calibration plate around 100 % leads to errors if the reflectometer response is not strictly linear over the whole of the measuring range. It is thus essential to use at least two calibrated calibration plates.

4.5 Procedure

The procedure is based on the apparatus used. It should enable a correct measurement of the percentage luminance of the cloth to be obtained with the Y (green) tristimulus filter specified, excluding any fluorescence phenomena.

Carry out the measurements on the various test pieces of cotton control cloth at least 12 h and at most 7 days after final ironing (or pressing). During this period, keep the test piece in conditions of the strictest cleanliness and protected from the light.

For each individual measurement, fold the test piece under test in such a way as to give a thickness of eight layers with the uppermost layer, on which the measurement is made, corresponding to the surface of the cloth which has been in contact with the polished part of the press or ironing machines. If it is not possible to distinguish the two surfaces, for example on a tumbler-dried test piece, carry out half the measurements on one surface and half on the other. Carry out ten individual measurements in this manner at different points on each test piece.

The intrinsic greying $\Delta {\pmb G}$ is given, as a percentage, by the formula

$$\Delta G = R_{g_0} - R_g$$

where

 R_{g_0} is the arithmetic mean, expressed to one decimal place, of the reflectometer values for the Y (green) tristimulus filter (equal to the percentage luminance) recorded on the cotton control cloth;

 R_g is the arithmetic mean, expressed to one decimal place, of 3×10 reflectometer values for the Y (green) tristimulus filter (equal to the percentage luminance) recorded on the three test pieces that have undergone the 25 (or 50) laundering cycles.

Express the result to one decimal place.

NOTE – The R_{g_0} value for the cotton control cloth is not necessarily the highest value that might be obtained for it as the laundering operations that will subsequently be applied to it could, in some cases, perfect the results of the preparatory treatment operations alone.

In certain cases, therefore, it is possible to record negative ΔG values.

There is no disadvantage in this so far as comparative tests are concerned.

If, on the other hand, it is desired to use ΔG as an absolute index for the quality of laundering, for example in checking the quality

5.6 Expression of results

The degree of intrinsic yellow J of the laundered test pieces is conventionally obtained from the formula

 $J = R_{\rm g} - R_{\rm b}$

and the degree of intrinsic yellow J_0 of the cotton control cloth from the formula

 $J_0 = R_{g_0} - R_{b_0}$

where

 R_{g} and $R_{g_{0}}$ are as defined in 4.6;

 $R_{\rm b}$ is the arithmetic mean, expressed to one decimal place, of 3×10 reflectometer values for the Z (blue) tristimulus filter as recorded on the three test pieces that have undergone 25 (or 50) laundering cycles;

 R_{b_0} is the arithmetic mean, expressed to one decimal place, of the reflectometer values for the Z (blue) tristimulus filter recorded on the cotton control cloth.

The intrinsic yellowing ΔJ , caused by the 25 (or 50) laundering cycles on the control cloth, is given by the formula

```
\Delta J = J_{25(50)} - J_0 iTeh STANDARD Pre
```

Express the result to one decimal place. standards.it

When $\Delta J > 0$, the laundering has caused intrinsic yellowing. the ingress of humidity for $\Delta J = 0$, the laundering has not caused intrinsic yellow: 1979 ing the following items: ing. Finally, if $\Delta J < 0$, the laundering has is ither reduced state between the degree of intrinsic yellow in the cotton control cloth or -43126.4

NOTE — The R_{b0} value for the cotton control cloth is not necessarily the highest value that might be obtained for it as the laundering operations to which it will later be subjected could, in some cases, perfect the results given by the preparatory treatment operations alone.

In certain cases, it is therefore possible to record negative values for ΔJ .

There is no disadvantage in this so far as comparative tests are concerned.

If, on the other hand, it is desired to use ΔJ as an absolute index for the quality of laundering, for example in checking the quality achieved by commercial laundries, it would be preferable to establish once and for all a constant value for R_{b_0} equivalent to the highest values obtained for the cotton control cloth in the course of the various tests.

As this maximum value depends on the working conditions and the apparatus used in each laboratory, it is impossible to propose a universally acceptable value. For guidance, it may be stated that it usually lies between 86 and 90 %.

6 DETERMINATION OF INCREASE IN ORGANIC DEPOSITIONS CONTENT

6.1 Scope and field of application

This clause specifies a method for the determination of the increase in the content of organic matter insoluble in water in the unsoiled cotton control cloth, caused by the 25 (or 50) laundering cycles specified in ISO 2267. These deposits, arising mainly from calcium and magnesium soaps, can cause various undesirable conditions : greying, yellowing, a greasy feel, more difficult wetting, an unpleasant smell, calendering difficulties.

6.2 Principle

Treatment of a known mass of cotton control cloth with a benzene/ethanol mixture before and after the 25 (or 50) laundering cycles. After extraction, evaporation of the solution of organic matter extracted and weighing of the dry residue.

6.3 Reagent

During the analysis, use only reagents of recognized analytical grade and only distilled water or water of equivalent purity.

Extraction solvent, consisting of a mixture of 9 volumes of benzene and 1 volume of 96 % (V/V) ethanol.

6.4 Apparatus

Ordinary laboratory apparatus and

6.4.1 Soxhlet type extraction apparatus, having standardized ground joints, capable of being protected against the ingress of humidity from the atmosphere, and comprising the following items:

z/ 5t/b62059b5-34aa-45bf-8f70-

6.4.1.2 Extractor, of capacity between 150 and 250 ml.

6.4.1.3 Condenser, fitting both the receiving flask (6.4.1.1) and the extractor (6.4.1.2).

6.4.2 Weighing bottle.

6.4.3 Oven, capable of being controlled at 103 ± 2 °C.

6.4.4 Drying tube.

6.4.5 Water bath or flameless heater.

6.4.6 Vacuum pump.

6.5 Procedure

6.5.1 Test portion

Take a test portion of mass approximately 3 g, weighed to the nearest 0,000 1 g, from that remaining part of the test piece which is intended for measurement of loss of tensile strength and has been prepared in accordance with clause 6 of ISO 2267 (see also clause 3).

At the same time, take a second test portion for determining the water content of the cloth. In the first case, the ash consists of depositions of mineral salts as such or having undergone certain chemical modifications as a result of calcination, and of the residue of incineration of organic salts. As this residue comprises only a quite small fraction of the organic depositions, in the majority of cases, the amount of total ash depends above all on the magnitude of the mineral depositions.

However, the amount of mineral ash is a more accurate index of the presence of mineral depositions. The latter derive from salts contained in the washing water or from detergents or from the reaction of both.

Depositions of mineral products on cloth indicate a defect in the detergent action; they may contribute to the premature rejection of the cloth by modifying its organoleptic qualities (harsh feel, dull appearance, and greying or yellowing) or by promoting chemical or mechanical damage.

7.2 Principle

After free combustion of the cotton control cloth in a tared crucible, incineration of the resulting ash in a muffle furnace at 800 $^{\circ}$ C, followed by weighing. Determination of the ash before and after the 25 (or 50) laundering cycles.

The determination may be carried out on the cloth as it is, the result being the total amount of ash. It may also be carried out on the cloth after the extraction of organic depositions (clause 6) and, in this case, it will give the amount of mineral ash.

7.3 Apparatus

IANDAKD P to the dry state of the cloth;

Ordinary laboratory apparatus and

(standards.itemais)the mass, in grams, of the empty crucible;

 m_4 is the mass, in grams, of the crucible and ash after calcination.

Provise the mass, in grams, of the test portion, referred

catches fire. Withdraw the flame and leave the cloth to

burn freely; when the flame goes out, allow the ignition of

the combustion points to continue. Repeat the operation

Transfer the crucible to the muffle furnace (7.3.3), con-

Transfer the crucible to a dessiccator, allow to cool to

2 To obtain the ash of cotton control cloth before laundering, the

The ash of the original dry cloth is given, as a percentage

procedure specified in note 4 to 6.5.3 may be followed.

until complete combustion is achieved.

ambient temperature and reweigh.

NOTES

where

1 See note 3 to 6.5.3.

7.5 Expression of results

by mass, by the formula

 $A = \frac{m_4 - m_3}{m_0} \times 100$

trolled at 800 \pm 25 $^{\circ}\text{C}$, and calcinate for 1 h.

7.3.1 Silica, porcelain of platinum crucible in accordancels/sist/E2059b534aa-45b58f70with ISO 1772. e8d9d4bd926b/iso-4312-1979

ISO 4312:1979

7.3.2 Mecker or Bunsen burner.

7.3.3 Muffle furnace, capable of being controlled at 800 \pm 25 °C.

7.4 Procedure

7.4.1 Test portion

Weigh the crucible (7.3.1) after heating it in the muffle furnace (7.3.3), controlled at 800 ± 25 °C, and allowing it to cool to ambient temperature in a dessiccator.

To determine the amount of total ash, take a test portion of mass approximately 3 g, weighed to the nearest $0,000 \ 1 \ g$, and place it in the tared crucible (7.3.1). Determine the water content of a second test portion as indicated in 6.5.2.

To determine the amount of mineral ash, it is only necessary to take the whole of the test portion in the Soxhlet extractor (6.4.1), the dry mass of which is known. Place the test portion in the tared crucible and expel the solvent by evaporation under a ventilated hood.

7.4.2 Determination

Place the crucible with its contents on a tripod and heat it over a small flame from the burner (7.3.2) until the cloth

Calculate the result of the test by taking the arithmetic mean of the values for three test pieces that have undergone 25 (or 50) laundering cycles $(A_{25(50)})$ and three test pieces intended for the determination of the initial characteristics of the cotton control cloth (A_0) .

The increase ΔA in the ash due to the 25 (or 50) laundering cycles is given by the formula

$$\Delta A = A_{25(50)} - A_0$$

8 DETERMINATION OF OVERALL LOSS OF BREAKING STRENGTH (TOTAL DAMAGE)

8.0 Introduction

After repeated washings, the cotton control cloth generally exhibits less breaking strength than it did originally, due to the combined action of the mechanical and chemical factors operating during the 25 (or 50) laundering cycles.

Any loss of breaking strength, expressed as a percentage of the initial breaking strength, is determined from the variation in the breaking strength measured in the direction of the warp of the control cloth before and after laundering.

The measurement is effected on cloth stabilized in the standard conditioning atmosphere specified in ISO 139. For particular purposes, however, measurements may be made on water-wet cloth (wet strength).

contains coloured or thicker threads demarcating the width of the specimens (see ISO 2267, sub-clause 5.1.1, note 2).

Place the specimens, after removal of deposits, where necessary, in the conditioning enclosure (8.4.3) and leave them there for 24 h before measuring.

8.5.2 Measurement of the breaking strength

Carry out the measurement in the direction of the warp on the conditioned cloth in the atmosphere specified in 8.4.3 and complying with the procedure of ISO 5081 for a specified period for the average time-to-break of 30 ± 5 s.

Examine the laundered control cloth for evidence of localized damage.

8.6 Expression of results

The total wear U_t of the cotton control cloth, expressed as a percentage, is given by the formula

$$U_{t} = \frac{F_{0} - F_{25(50)}}{F_{0}} \times 100$$

where

 $F_{25(50)}$ is the breaking strength, expressed in newtons, of cotton control cloth subjected to the 25 (or 50) laundering cycles;

 F_0 is the breaking strength, expressed in newtons, of cotton control cloth measured under the same conditions.

Express the result to one decimal place.

ttps://standards.itch.ai/catalog/standards The breaking strength is calculated by taking the arithmetic mean of the individual values determined for each specimen that has been subjected to measurement, i.e., in principle, 30 measurements for all three test pieces representing a test.

NOTES

1 F_0 can be determined by proceeding as specified in 6.1 of ISO 2267. In view of the inherent variability of textiles, it is always preferable to make the 30 measurements specified for each test and to calculate the F_0 value by taking the arithmetic mean of these measurements.

 $2\,$ The conditioning of the specimen may be omitted when measuring the breaking strength of the piece in the wet state and refer to prescriptions of ISO 5081.

If any localized damage is evident, describe it fully.

9 DETERMINATION OF LOSS OF BREAKING STRENGTH RESULTING FROM CHEMICAL DEGRADATION OF CELLULOSE DUE TO LAUNDERING (CHEMICAL DAMAGE)

9.0 Introduction

The chemical degradation of the cellulose in cloth, caused by the action of chemical agents during the laundering operation, usually accompanies a variation in the intrinsic mechanical properties of the fibres and, correlatively, a loss of breaking strength in the cloth. This chemical damage, which is a function of the chemical agressiveness of the laundering process, is characterized by a reduction in the degree of polymerization of the cellulose constituent. Knowing the degree of polymerization - or a related parameter - of the cotton control cloth before and after the 25 (or 50) laundering cycles, and the link between the variation in this parameter and the corresponding loss of tensile strength, it is possible to calculate the part of the loss of breaking strength in the cotton control cloth which is due solely to chemical degradation.

Chemical damage is caused in the first place by the chemical agressiveness of washing products, mainly oxidizing agents such as hypochlorite and peroxides.

9.1 Scope and field of application

This clause specifies a method for the determination of the loss of breaking strength corresponding to the chemical degradation of the unsoiled cotton control cloth, due to the 25 (or 50) laundering cycles specified in ISO 2267.

9.2 Principle

ISO 4312:1979

Measurement of the fluidity value of a solution [in ammonium copper(II) hydroxide] of cotton provided by warp threads from the cotton control cloth before and after the 25 (or 50) laundering cycles.

The loss of breaking strength due to chemical degradation, expressed as a percentage of the initial breaking strength, is an approximate linear function of the increase in the fluidity value.

NOTE -- There are many methods of measuring viscosity or its reciprocal, the fluidity, of cellulose (see 9.6).

bthe present method has been chosen because it is the only one for which the relation with the loss of breaking strength in cotton control cloth has been established experimentally.

9.3 Preparation of test samples

Ravel out warp threads from each of the specimens intended for the determination of total damage and take enough to give a sample of mass about 4 g for each test piece to be examined.

If it has been necessary to remove deposits from the cloth (see 8.5.1.1), take the threads from the specimens after the treatment to remove the deposits.

Cut the threads into strands from 1 to 2 mm long. Homogenize them. One part of this test sample is intended for the measurement of the fluidity value (or of the viscosity or degree of polymerization), the other for the determination of the water content.

9.4 Determination of fluidity value in ammonium copper(II) hydroxide solution

WARNING — Appropriate safety precautions should be observed when carrying out the operations described in this sub-clause.

9.4.1 Definition

fluidity of a solution : The reciprocal of the viscosity of the solution expressed in pascal seconds (Pa·s); fluidity is thus expressed in reciprocal pascal seconds $[(Pa-s)^{-1}]$. For

 V_4 is the volume, in millilitres, of the standard volumetric *diammonium* iron bis-sulphate solution used for the titration of the ammonium copper(II) hydroxide solution.

9.4.4 Apparatus

9.4.4.1 Viscometers, of the shape shown in figure 1 and the following dimensions :

- inside diameter of the capillary E : 0,88 ± 0,01 mm
- outside diameter of the capillary E : 6 \pm 2 mm
- length of the capillary $E: 25 \pm 0.5$ mm
- inside diameter of the tube A : 10 ± 0.05 mm

 $-\,$ distance from the reference mark B to the end of the capillary E : 242 \pm 0,5 mm

- distance from the reference mark C to the end of the capillary E : 122 ± 0.5 mm

 $-\,$ distance from the reference mark D to the end of the capillary E : 62 \pm 0,5 mm

The end of the capillary should be flat. The viscometer may be closed at the top by means of a plastic stopper pierced by a capillary tube; the latter may be closed by a small cap. The lower capillary may be closed by means of a rubber tube and a clamp. For each viscometer, the internal volume must be determined for the closed viscometer, subtracting a volume of 0,7 ml corresponding to the stirring device (see 9.4.4.4).

9.4.4.2 Supporting tubes for viscometers (see figure 2 for -43 an example of a suitable shape).

The tubes should support the viscometers (9.4.4.1) in the thermostatic bath (9.4.4.3) for the period required to reach a temperature of 20 °C and while the flow time is being measured. In the latter case, the viscometers must be kept absolutely vertical.

9.4.4.3 Glass-walled thermostatic bath, capable of being controlled at 20 ± 0.2 °C and large enough to enable the tubes containing the viscometers to be kept immersed up to the upper rim by means of suitable fixtures.

9.4.4. Rotary stirrer, enabling the viscometers (9.4.4.1) to be rotated in such a way that the mercury (or the stirrer) slides from one end to the other of the tube A during each demi-rotation. The rotational frequency may vary between 4 and 10 min⁻¹.

9.4.4.5 Filling device, allowing the ammonium copper(11) hydroxide solution (9.4.3) to be maintained under nitrogen pressure and the viscometers (9.4.4.1) to be refilled by causing the liquid to rise through the capillary E (see figure 1).

9.4.5 Procedure

Weigh a portion of the test sample (9.3), to the nearest 0,0001 g, such that a solution containing 0,5 g of the pure,

dry cotton in 100 ml can be placed in the viscometer (9.4.4.1). To achieve this, the water content of the test sample, determined separately in accordance with 6.5.2 (see also note 3 to 6.5.2), and the organic and mineral deposits present in the test sample, determined in accordance with clauses 6 and 7, must be taken into account.

Introduce the test portion into the viscometer, which must be held upright. Using the short tube fitted with a clamp, connect the lower capillary tube of the viscometer to the vessel containing the ammonium copper(II) hydroxide solution (9.4.3) and make the solution rise into the viscometer under nitrogen pressure. When the viscometer is almost full, pinch the clamp of the lower tube and detach the viscometer. Add 0,7 ml of mercury by means of a short burette (alternatively, a stainless steel spiral with a volume of 0,7 ml may be used). Close the viscometer by means of a plastic stopper so that the liquid fills the viscometer completely and overflows through the capillary tube in the stopper, close this tube with a cap.

It is important to ensure that there is not a single air bubble left in the viscometer. In order to achieve this, the stopper must be inserted carefully but quickly. For the same reason, it may be necessary to tap the viscometer lightly before it is closed in order to detach air bubbles that might cling to the cotton fibres.

Place the viscometer, which shall not be exposed to light, on the stirring device (9.4.4.4). Allow it to rotate overnight at a frequency of 4 min^{-1} or for 4 h at 10 min^{-1} (see

https://standards.iteh.ai/catalog/standards/sist/lnote51/b5-34aa-45bf-8f70-

Withdraw the viscometer from the stirring apparatus, remove the rubber tube from the lower capillary tube and allow the mercury to flow by opening the capillary in the upper stopper for a moment (this is not necessary when a spiral is used).

Place the viscometer in a supporting tube (9.4.4.2) immersed in the thermostatic bath (9.4.4.3), maintained at 20 ± 0.2 °C. Wait until the temperature has stabilized (at least 1 h). Check that the viscometer is in a vertical position. Remove the stopper and leave the solution to flow freely into the bulb of the support tube. Measure the time t (in seconds to the nearest 0.2 s) required for the meniscus to fall from the reference mark B to the reference mark D (see note 2).

NOTES

1 Since the control cloth is bleached so that the fluidity value of the cotton always exceeds, by definition, 40 (Pa·s)⁻¹, there is no difficulty in dissolving it in the ammonium copper(II) hydroxide solution and a stirring period of 4 h at 10 min⁻¹ (or 1 night at 4 min⁻¹) is always adequate.

2 To check that the solution is flowing freely through the capillary, it may be useful to measure both the time t_1 taken by the meniscus to fall from B to C and the time t_2 taken by it to fall from C to D. The ratio t_1/t_2 should be greater than 0,85 for a fluidity value of 40 (Pa·s)⁻¹ and should tend towards unity for fluidity values higher than 40 (Pa·s)⁻¹.

9.4.6 Expression of results

The fluidity value F of a solution containing 0,5 g of pure