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Surface active agents – Determination of flow properties using a rotational viscometer

Agents de surface — Détermination des propriétés d'écoulement au moyen d'un viscosimètre rotatif

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<u>ISO 6388:1989</u>

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Foreword

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Draft International Standards adopted by the technical committees are circulated to the member bodies for approval before their acceptance as International Standards by the ISO Council. They are approved in accordance with ISO procedures requiring at least 75 % approval by the member bodies voting.

International Standard ISO 6388 was prepared by Technical Committee ISO/TC 91, COM 2019 Surface active agents.

This second edition cancels and replaces the first edition (ISO 6388 : 1983), of which it constitutes a minor revision.

<u>ISO 6388:1989</u>

Annex A of this International Standard is for information only. 0e1a8-5a2b-40eb-8acb-5891d6ad68c2/iso-6388-1989

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International Organization for Standardization

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Introduction

The rheological behaviour of surface active agents in general is determined using a rotational viscometer, which is suitable for studying substances having a complex rheological behaviour, e.g. plastics, paints.

In developing this International Standard, reference has been made to ISO 3219, which deals with plastics, for the definitions and the apparatus. The method specified in this International Standard, however, is specific to surface active agents.

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Surface active agents — Determination of flow properties using a rotational viscometer

1 Scope

This International Standard specifies a method for the characterization of the flow properties of non-solid surface active agents, either alone or in the form of mixtures, and of products consisting essentially of surface active agents, using a co-axial cylinder, cone and plane, or double cone, etc., rotational viscometer.

NOTE — The rheological behaviour of a system containing surface active agents is frequently marked by anomalies. These are mostly due to the tendency of the molecules of surface active agents to associate. The rheological behaviour varies essentially as a function of the nature and concentration of the surface active agents. Small variations in temperature, in the concentration of mineral salts, and the presence of any other substance, may also cause variations in the rheological behaviour of surface active agents; even the rheological pattern itself may sometimes be changed. The method specified in this International Standard attempts to take account of all these factors. In the case of certain very special surface active agents, other methods of determination may be used. For newtonian systems, for example, reference should be made to ISO 3104 and ISO 1652, which are much more precise.

When the newtonian character of a system is in question, the method specified in this International Standard will make it possible to select measuring equipment which will allow a determination to be carried out.

2 Normative references

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

ISO 607 : 1980, Surface active agents and detergents – Methods of sample division.

ISO 862 : 1984, Surface active agents - Vocabulary.

3 Definitions, symbols and units

3.1 General

The viscosity of a fluid sheared between two parallel planes, one of which moves in its own plane with linear and uniform motion relative to the other, is defined by Newton's equation:

$$\eta = \frac{\tau}{D}$$

where

 η is the (dynamic) viscosity;

 τ is the shear stress;

$$D = \frac{\mathrm{d}v}{\mathrm{d}z}$$
 is the rate of shear,

v being the velocity of one plane relative to the other, and

9 z being the coordinate perpendicular to both planes.

NOTE – Products for which the viscosity is independent of the rate of shear at which the measurement is taken are regarded as having newtonian behaviour and are called "newtonian". Others are regarded as having non-newtonian behaviour and are called "non-newtonian".

The apparent viscosity of a non-newtonian product is the ratio of the produced shear stress to the applied rate of shear.

The value of the apparent viscosity, a function of the rate of shear, can depend on thermal and rheological hysteresis of the sample in the apparatus.

The dimensions of viscosity are $ML^{-1}T^{-1}$, and in the International System of Units (SI), the unit is the newton second per square metre (N·s/m²) or the pascal second (Pa·s).¹⁾

3.2 Rheological phenomena (see figures 1 and 2 and ISO 862).

NOTE — Rheological phenomena are described for a defined shear stress.

¹⁾ $1 \text{ N} \cdot \text{s/m}^2 = 1 \text{ Pa} \cdot \text{s} = 10^3 \text{ cP}$ (centipoise)

 $^{1 \}text{ mN} \cdot \text{s/m}^2 = 1 \text{ mPa} \cdot \text{s} = 1 \text{ cP}$ (centipoise)

3.2.1 shear thinning: The reduction under isothermal and reversible conditions and without hysteresis of the apparent viscosity with increasing rate of shear.

3.2.2 dilatancy: The increase under isothermal and reversible conditions and without hysteresis of the apparent viscosity with increasing rate of shear.

3.2.3 time-dependent viscosity: Under reversible and isothermal conditions, a change of the apparent viscosity during flow with a constant magnitude of the shear rate.

3.2.3.1 thixotropy: The reduction of viscosity or consistency, under isothermal and reversible conditions, by the application of shear from the viscosity or consistency at rest (immediately after beginning to shear) to a final value (dependent on the magnitude of the shear rate).

When the shear is discontinued the viscosity or consistency at rest must be reestablished within a certain time, the "time of thixotropic recovery".

3.2.3.2 rheopexy: The phenomenon in which the time of thixotropic recovery, after discontinuation of a relatively high shear rate, is shortened by applying a small shear rate.

3.2.3.3 anti-thixotropy: The increase of viscosity or consistency under isothermal and reversible conditions, by the application of shear from the viscosity or consistency at rest (immediately after beginning to shear) to a final value (dependent on the magnitude of the shear rate).

^{*h*i} ≥ 1,5 When the shear is discontinued, the viscosity at rest must be re-established within a certain time, the "time of thixotropic recovery".

3.2.3.4 rheological hysteresis: If the shear rate under isothermal and reversible conditions is increased linearly with respect to time from zero to a maximum value (up-branch) and then decreased in the same manner (down-branch), the rate of shear-diagram shows a hysteresis loop, which is used to detect and to characterize thixotropy or anti-thixotropy.

3.2.4 plasticity: A plastic body behaves as an elastic body when it is subjected to a stress less than a critical value, τ_{0} , the "yield stress". Above this limiting value, a flow takes place. When the function $D = f(\tau)$ (*D* is the rate of shear) for $\tau \ge \tau_0$ is represented by a straight line, the substance is said to follow the Bingham model.

Principle

Determination of flow properties of a newtonian or nonnewtonian test portion by means of a specified rotational viscometer1), which will allow, for the newtonian products, the simultaneous measurements of the shear rate at which the determination is carried out, but for the non-newtonian products, possible measurement of various apparent rates of shear used for the determination.

Reference substances 5

Newtonian liquids shall be used as reference substances.

Apparatus

Viscometers, with co-axial cylinders, a cone and plate, a 6.1 double cone or a combination of two of these types, with the following specifications:

6.1.1 In the case of viscometers with co-axial cylinders, the external and internal diameters of the cylinders (stator and rotor), d_{e} and d_{i} respectively, shall be such that the value of the ratio $d_{\rm e}/d_{\rm i}$ is as small as possible and, preferably, less than or equal to 1,10, and in no case greater than 1,50. If the ratio has a value greater than 1,10, this shall be indicated in the test report and a correction shall, if possible, be made, which shall also be reported in the test report.2)

In addition, if the apparatus does not include a geometric device (conical base and guard ring above it) for correcting the end effect, the following additional requirement shall be respected:

where h_i is the height of the inner cylinder.

6.1.2 In the case of viscometers with a cone and plate, or a double cone, the value of the angle α , formed by the generating line of the cone and the plate or by the generating lines of the two cones shall be as small as possible and, preferably, less than or equal to 1°, and in no case greater than 4°. If the angle α has a value greater than 1°, this shall be indicated in the test report and a correction shall, if possible, be made, which shall also be reported in the test report.2)

6.1.3 In the case of viscometers combining two of the preceding principles, each of the specifications shall be observed, unless one of the devices contributes at a very low percentage and can be regarded as being a correction.

The first type is most commonly used, and it is the use of this type which is described in this International Standard.

¹⁾ Two types of viscometer exist: one where the shear stress is determined at a constant shear rate (constant frequency of rotation), in which case the results of the determination are presented in the form of a plot of τ as a function of D (see figures 1 and 2), the other where the shear rate is determined at a constant shear stress, in which case the results of the determination are presented in the form of a plot of D as a function of τ (see figures 1 and 2).

²⁾ The correction formulae should usually be supplied by the manufacturer of the apparatus.