

SLOVENSKI STANDARD **SIST EN ISO 6327:2008** 01-marec-2008

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Gas analysis - Determination of the water dew point of natural gas - Cooled surface condensation hygrometers (ISO 6327:1981)

Gasanalyse - Bestimmung des Wassertaupunktes von Erdgas - Kondensations-Hygrometer mit gekühlter Oberfläche (ISO 6327:1981)

iTeh STANDARD PREVIEW

Analyse des gaz - Détermination du point de rosée des gaz naturels - Hygromètres à condensation à surface refroidie (ISO 6327:1981)

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75.060

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EUROPEAN STANDARD NORME EUROPÉENNE

EUROPÄISCHE NORM

EN ISO 6327

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English Version

Gas analysis - Determination of the water dew point of natural gas - Cooled surface condensation hygrometers (ISO 6327:1981)

Analyse des gaz - Détermination du point de rosée des gaz naturels - Hygromètres à condensation à surface refroidie (ISO 6327:1981) Gasanalyse - Bestimmung des Wassertaupunktes von Erdgas - Kondensations-Hygrometer mit gekühlter Oberfläche (ISO 6327:1981)

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

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EUROPEAN COMMITTEE FOR STANDARDIZATION COMITÉ EUROPÉEN DE NORMALISATION EUROPÄISCHES KOMITEE FÜR NORMUNG

Management Centre: rue de Stassart, 36 B-1050 Brussels

Foreword

The text of ISO 6327:1981 has been prepared by Technical Committee ISO/TC 193 "Natural gas" of the International Organization for Standardization (ISO) and has been taken over as EN ISO 6327:2008 by Technical Committee CEN/SS N21 "Gaseous fuels and combustible gas", the secretariat of which is held by CMC.

This European Standard shall be given the status of a national standard, either by publication of an identical text or by endorsement, at the latest by July 2008, and conflicting national standards shall be withdrawn at the latest by July 2008.

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

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

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Endorsement notice

SIST EN ISO 6327:2008

The text of ISO 6327/1981 has been approved by GEN as EN-ISO 6327:2008 without any modifications. 9628d7278e5d/sist-en-iso-6327-2008

International Standard



6327

INTERNATIONAL ORGANIZATION FOR STANDARDIZATION●MEЖДУНАРОДНАЯ ОРГАНИЗАЦИЯ ПО CTAHДAPTUЗАЦИИ●ORGANISATION INTERNATIONALE DE NORMALISATION

Gas analysis — Determination of the water dew point of natural gas — Cooled surface condensation hygrometers

Analyse des gaz — Détermination du point de rosée des gaz naturels — Hygromètres à condensation à surface refroidie

First edition – 1981-03-15Teh STANDARD PREVIEW (standards.iteh.ai)

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Ref. No. ISO 6327-1981 (E)

Foreword

ISO (the International Organization for Standardization) is a worldwide federation of national standards institutes (ISO member bodies). The work of developing International Standards is carried out through ISO technical committees. Every member body interested in a subject for which a technical committee has been set up 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.

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.

International Standard ISO 6327 was developed by Technical Committee ISO/TC 158, Analysis of gases, and was circulated to the member bodies in September 1979.

It has been approved by the member bodies of the following countries:

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Australia Belgium

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Czechoslovakia

Korea, Rep. of 9628d7278@Romahian-iso-6327-2008

Egypt, Arab Rep. of

South Africa, Rep. of Libyan Arab Jamahiriya

United Kingdom

Mexico Netherlands

USSR

France Germany, F.R.

Philippines

No member body expressed disapproval of the document.

Gas analysis - Determination of the water dew point of natural gas — Cooled surface condensation hygrometers

Scope

This International Standard describes hygrometers used for the determination of the water dew point of natural gases by detecting water vapour condensation occurring on a cooled surface or by checking the stability of the condensation on this surface.

Field of application

The water dew point of processed natural gases in transmission lines normally lies between - 25 °C and + 5 °C which cor responds to water concentrations of 50 to 200 ppm (V/V), according to the pressure of the gas.

/standards.iteh The hygrometers considered in this International Standard may be used for determining water vapour pressure, without requiring calibration, in a system operating under total pressures greater than or equal to atmospheric pressure. The relationship between water vapour partial pressure and the observed dew point confers on the method the quality of absolute measurement.

If the test atmosphere contains gases which condense at a temperature in the region of, or above, that of the water dew point, it is very difficult to detect the condensed water vapour.

Principle

3.1 Principle of the apparatus

With this type of apparatus, which determines the water content of a gas by measuring the corresponding dew point, a surface (generally a metallic mirror), the temperature of which may be artificially lowered and accurately measured, is exposed to a sample of the gas being tested. The surface is then cooled to a temperature at which condensation occurs and is observed as dew.

Below this temperature, condensation increases with time, whilst above it, condensation decreases or does not occur. This surface temperature is then (for practical applications) taken as the dew point of the gas flowing through the apparatus.

3.2 Determination of water vapour pressure

The partial water vapour pressure in the gas samples is the saturated vapour pressure corresponding to the observed dew point, provided that the gas in the hygrometer is at the same pressure as the gas at the time of sampling.

Published documents are available giving the relationship between saturated vapour pressure and temperature.

It should be noted that/if methanol is present, this method determines methanol in addition to water. However, if the methanol content is known, the annex gives, for information, correction factors allowing determination of the actual water dew point.

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It is essential that all sample lines be as short as possible and be sized to produce a negligible pressure drop during measurement. The sample lines and the hygrometer, apart from the mirror, shall be above the water dew point temperature.

Characteristics of the apparatus

4.1 General

Condensation apparatus may be designed in various ways. The differences lie mainly in the nature of the condensation surface, the methods used for cooling the surface and for controlling its temperature, the methods used for measuring the surface temperature and the method of detecting the condensation. The mirror and its associated components are normally placed in a small cell through which a sample of the gas flows; at high pressures, the mechanical strength and leak tightness of the cell have to be suitable.

It is recommended that the mirror should be easily removable for cleaning.

Adequate precautions shall be taken if measurements are to be made in the presence of condensable hydrocarbons.

Measurements can be carried out manually or automatically.

Caution: Manufacturers' instructions should be carried out before gas at high pressure is admitted to the cell.

4.2 Automatic and manual types

Devices for measuring dew point can be designed to make isolated measurements at different times or to make more or less continuous measurements. For isolated measurements, methods of mirror cooling may be chosen which require continuous attention by the operator responding to changes in the condensed deposit which is observed by the naked eye. If there is less moisture in the gas sample, i.e. if the gas has a lower dew point, the rate at which water vapour flows through the apparatus per unit time decreases so that condensation forms more slowly, and it becomes more difficult to judge whether condensation is increasing or diminishing. Observation of the deposit can be made easier by using a photoelectric cell or any other device which is sensitive to light, if a simple indicator is needed, while maintaining manual control of the cooling device.

With certain types of manually operated instruments, it is very difficult to observe the water dew point in the presence of condensed hydrocarbons. In such cases, a liquid paraffin bubbler may be used to assist such observations. It is very important, however, that the principles involved and the limitations in the use of such a bubbler are understood.

An equilibrium is established between the gas passing through the bubbler and the liquid paraffin oil contained in it, at the temperature and pressure of the bubbler. This involves the following reactions: following reactions:

- water to the paraffin until equilibrium is achieved, at which the inlet gas. Therefore, the temperature of the pubbler shows the compressed air or other suitable pressurized gas. must be above that of the water dew point of the gas to be tested and sufficient gas must be passed into the bubbler for equilibrium to be established before observations can be made.
- b) Until equilibrium is established, heavy hydrocarbon components pass from the gas into the liquid paraffin. It is this exchange that reduces the volume of potentially condensable hydrocarbons in the gas, thereby reducing the masking effect of the condensed hydrocarbon liquid. As there is a continuing exchange of components, the liquid paraffin becomes saturated with condensible hydrocarbons the content of which increases in the gas. The liquid paraffin must then be replaced and the bubbler conditioned before further observations can be made.

The device can be fully automated by using the output signal of the photoelectric cell to stabilize the mirror at the required condensation temperature. Automatic operation is indispensable for continuous reading or recording.

4.3 Mirror illumination

Manual devices can involve observation of condensation with the naked eye; if a photoelectric cell is used, the mirror is illuminated by a light source built into the test cell. The lamp and photoelectric cell can be arranged in various ways, provided that diffusion in the direction of the light source from the mirror is reduced by the polishing of the mirror. In any case, the mirror must be clean before use.

In the absence of any condensation, the diffused light falling on the photocell must be reduced. The effects of light diffused from internal surfaces of the cell can be reduced by blackening these surfaces and this precaution can be supplemented by an arrangement of the optical system so that only the mirror is illuminated and the photocell views only the mirror.

4.4 Methods for cooling the mirror and controlling its temperature

The following methods are used for reducing and adjusting the mirror temperature. The methods described in 4.4.1 and 4.4.2 require constant attention from the operator and are not suitable for automatic devices. For automatic devices, two cooling methods are used: indirect contact with a coolant or cooling by the thermoelectric (Peltier) effect as described in 4.4.3 and 4.4.4. In any case, the rate of cooling of the mirror shall not exceed 1 °C per minute.

4.4.1 Solvent evaporation

A volatile liquid in contact with the rear face of the mirror can be evaporated and cooled by an air flow. Hand bellows are generally used for this purpose/but an adjustable source of low pressure compressed air or any other suitable pressurized gas is preferable. The liquid used can be ethylene oxide, a very efficient liquid giving cooling of the mirror of approximately 30 °C, without effort, when hand bellows are used. However, a) The first gas passing through fresh liquid paraffingoses IN IS (1 toxicity) (18 a risk, acetone can be used to obtain cooling of stand approximately 20 °6 with hand bellows or even greater cooling

4.4.2 Gas cooling by adiabatic expansion

The mirror can be cooled by discharging onto its rear face a gas which has just expanded through a nozzle. Compressed carbon dioxide, available from small cylinders, is often used for this purpose, but other gases such as compressed air, compressed nitrogen, propane or halogenated hydrocarbons can also be used. Mirror temperatures of at least 40 °C below the gas sample temperature can be obtained.

4.4.3 Indirect contact with a coolant

The mirror is connected to a cooler through a thermal resistor. Normally, a solid copper rod is plunged into the cooler and connected to the mirror by a small piece of insulating material forming the thermal resistor. The mirror is heated by an electric element. Current intensity should be controlled so that the mirror temperature can be adjusted easily and accurately. Using liquid nitrogen as coolant, temperatures of - 70 °C to - 80 °C may be obtained; for temperatures down to approximately - 50 °C (according to the apparatus design), a mixture of solid carbon dioxide plus acetone may be used, and for temperatures around - 30 °C liquefied propane can be used.

4.4.4 Cooling by thermoelectric (Peltier) effect

A single stage Peltier effect element normally allows maximum cooling of approximately 50 °C. With two stages, cooling of approximately 70 °C can be obtained.

The mirror temperature can be adjusted by varying the current in the Peltier effect elements, but thermal inertia tends to be high, and more rapid adjustment is achievable by maintaining a constant cooling current, connecting the mirror to a thermal resistance, and heating the mirror with an adjustable electric heating device.

4.5 Temperature measurement

It is essential that the temperature of the mirror on which the deposit is formed is measured as accurately as possible. To avoid temperature differences on the surface, a mirror of high thermal conductivity is preferred. Manual devices generally incorporate a mercury thermometer, and with automatic devices a thermoelectric probe is used (for example a resistance thermometer, a thermistor or a thermocouple).

Sources of error — General precautions for operation

5.1 Interfering substances

5.1.1 General

Substances other than the gas or water vapour can enter the device and affect its operational characteristics. Such substances may be solid particles, dust, etc., which can be deposited on the mirror. Vapour other than water vapour can condense on the mirror. Gases soluble in water, voluntarily or 632 5.308 Equilibrium temperature approach accidentally introduced into the test cell, can also induce an observed dew point different from the dew point which would correspond to the actual water vapour content.

5.1.2 Solid impurities

If solid impurities are absolutely insoluble in water, they do not modify the observed condensation temperature, but can hinder condensation observation. In an automatic device, without a compensation device for such impurities, these tend to obstruct the operation of the device if the amount of condensate is low. Defects resulting from an excess of solid impurities on the mirror generally result in an unexpected increase of the mirror temperature for a few minutes and call for dismantling of the device and cleaning of the mirror. (It is essential for this purpose that the hygrometric cell can be rapidly dismounted.) It may be desirable to remove solid impurities by using a nonhygroscopic filter¹⁾ to avoid such difficulties.

To prevent the influence of dust particles, some automatic devices are fitted with a "calibration" sequence. This consists of an optional superheating of the mirror, so as to remove all condensate, water and hydrocarbons, followed by a rebalancing of the measuring bridge.

5.1.3 Impurities in vapour form

Hydrocarbons can condense on the mirror. In principle these do not interfere because hydrocarbon surface tension is very different from that of water. They spread on the mirror and form a continuous layer which does not diffuse light. Manual detection of condensates is, nevertheless, not easy because although the dew point is very much lower than the condensation temperature of hydrocarbons, only a few water droplets can be detected in a large hydrocarbon droplet (see clause 6).

The presence of a hydrocarbon condensate does not modify the water dew point since the condensates are not miscible.

If the gas contains methanol, this will be deposited with the water, and a dew point for the mixture water and methanol will be obtained. If hydrocarbons are also present, then two condensates are formed, one aqueous, the other oily. In this case, the condensation temperature of the aqueous condensate is not due solely to the water content.

5.2 Cold wall error

It is essential that the parts of the pipes and device other than the mirror are at a temperature greater than the condensation temperature; if not, water vapour will condense at the coldest points and modify the moisture content of the gas sample.

7e898cd-c248-4743-b1bf-If the quantity of water conveyed to the mirror per unit time is small, the mirror shall be cooled as slowly as possible because of the risk of greatly exceeding the actual condensation temperature without observing the first deposit.

The quantity of dew which can normally be observed with the naked eye is about 10⁻⁵ g/cm². Automatic devices, if very sensitive, can detect considerably lower quantities of water.

If a manual device is necessary, and especially for lower dew points, the following precautions shall always be taken:

- a) The mirror cooling rate shall be as small as possible in the condensation temperature range. (It is good practice to carry out a rapid test to determine the approximate condensation temperature before an accurate measurement is made.)
- b) The mean value of the temperature measured at the time of first dew appearance, while the mirror temperature slowly decreases, and of the temperature at which dew disappears, while the mirror temperature slowly increases, may be considered as the approximate dew point.

¹⁾ If a filter is used, even if it is stated to be non-hygroscopic, it should be in equilibrium with the water vapour content of the gas; this is obtained by allowing gas to flow through it for a period of time before the test at a rate considerably higher than that to be used during the test.