
**Steel — Measurement method for the
evaluation of hydrogen embrittlement
resistance of high strength steels —**

**Part 1:
Constant load test**

*Acier — Méthode de mesure pour l'évaluation de la résistance à la
fragilisation par l'hydrogène des aciers à haute résistance —*

Partie 1: Essai de charge constante

STANDARDSISO.COM : Click to view the full PDF of ISO 16573-1:2020



STANDARDSISO.COM : Click to view the full PDF of ISO 16573-1:2020



COPYRIGHT PROTECTED DOCUMENT

© ISO 2020

All rights reserved. Unless otherwise specified, or required in the context of its implementation, no part of this publication may be reproduced or utilized otherwise in any form or by any means, electronic or mechanical, including photocopying, or posting on the internet or an intranet, without prior written permission. Permission can be requested from either ISO at the address below or ISO's member body in the country of the requester.

ISO copyright office
CP 401 • Ch. de Blandonnet 8
CH-1214 Vernier, Geneva
Phone: +41 22 749 01 11
Email: copyright@iso.org
Website: www.iso.org

Published in Switzerland

Contents

	Page
Foreword.....	iv
Introduction.....	v
1 Scope.....	1
2 Normative references.....	1
3 Terms and definitions.....	1
4 Principle.....	1
5 Specimen preparation.....	2
6 Hydrogen charging methods.....	3
6.1 General.....	3
6.2 Cathodic charge method.....	3
6.2.1 Hydrogen charging solution.....	3
6.2.2 Hydrogen charging conditions.....	4
6.3 Hydrogen absorption in aqueous solution at free corrosion potential.....	4
6.4 Hydrogen absorption in atmospheric corrosion environments.....	4
6.5 Hydrogen absorption in high pressure hydrogen gas.....	4
7 Preparation of electroplating solution and electroplating condition.....	5
7.1 General.....	5
7.2 Electroplating solution.....	5
7.3 Electroplating conditions.....	5
8 Constant loading test.....	5
8.1 Constant loading test procedures.....	5
8.2 Presentation of the results.....	6
9 Post-test specimen treatment.....	8
10 Hydrogen thermal desorption analysis.....	9
10.1 General.....	9
10.2 Experimental apparatus (gas chromatograph).....	10
10.3 Experimental apparatus (mass spectrometry).....	10
11 Test report.....	10
Bibliography.....	11

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.

The procedures used to develop this document and those intended for its further maintenance are described in the ISO/IEC Directives, Part 1. In particular, the different approval criteria needed for the different types of ISO documents should be noted. This document was drafted in accordance with the editorial rules of the ISO/IEC Directives, Part 2 (see www.iso.org/directives).

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. Details of any patent rights identified during the development of the document will be in the Introduction and/or on the ISO list of patent declarations received (see www.iso.org/patents).

Any trade name used in this document is information given for the convenience of users and does not constitute an endorsement.

For an explanation of the voluntary nature of standards, the meaning of ISO specific terms and expressions related to conformity assessment, as well as information about ISO's adherence to the World Trade Organization (WTO) principles in the Technical Barriers to Trade (TBT) see www.iso.org/iso/foreword.html.

This document was prepared by Technical Committee ISO/TC 17, *Steel*, Subcommittee SC 7, *Methods of testing (other than mechanical tests and chemical analysis)*.

This edition cancels and replaces the first edition (ISO 16573:2015), which has been technically revised. The main changes compared to the previous edition are as follows:

- the addition of a note to provide the definition of ρ as the radius of the notch bottom. The definition of r was unclear and was used in a different way in 2b).
- the temperature in [6.1](#) and [Clause 7](#) where different, the temperature below -50 °C is used;
- the addition of Figures of unbroken notched specimen and unbroken smooth specimen;
- the addition of research papers in Bibliography.

Any feedback or questions on this document should be directed to the user's national standards body. A complete listing of these bodies can be found at www.iso.org/members.html.

Introduction

The mechanical properties of high strength steels, such as tensile strength, elongation and reduction of area, would be degraded by the effect of hydrogen, known as hydrogen embrittlement, and the susceptibility of hydrogen embrittlement becomes greater with increasing the strength level of steels. This document suggests a standardized test method for the evaluation of hydrogen embrittlement resistance of high strength steels.

STANDARDSISO.COM : Click to view the full PDF of ISO 16573-1:2020

[STANDARDSISO.COM](https://standardsiso.com) : Click to view the full PDF of ISO 16573-1:2020

Steel — Measurement method for the evaluation of hydrogen embrittlement resistance of high strength steels —

Part 1: Constant load test

1 Scope

This document provides a method for the evaluation of the resistance to hydrogen embrittlement (i.e. hydrogen delayed fracture) using constant loading test with hydrogen pre-charged specimens. The amount of hydrogen content absorbed in the specimens is analysed quantitatively by thermal desorption analysis such as gas chromatography, mass spectrometry and so on. In the case of hydrogen continuous charging such as hydrogen absorption in aqueous solution at free corrosion potential, hydrogen absorption in atmospheric corrosion environments and hydrogen absorption in high pressure hydrogen gas, the evaluation method is also briefly described. This method is mainly applicable to the evaluation of hydrogen embrittlement resistance of high strength steel bolts.

2 Normative references

There are no normative references in this document.

3 Terms and definitions

No terms and definitions are listed in this document.

ISO and IEC maintain terminological databases for use in standardization at the following addresses:

- ISO Online browsing platform: available at <https://www.iso.org/obp>
- IEC Electropedia: available at <http://www.electropedia.org/>

4 Principle

This test method is used to evaluate material resistance to hydrogen embrittlement. [Figure 1](#) shows schematic sequences for a) hydrogen pre-charging method and b) hydrogen continuous charging method. For the hydrogen pre-charging method [see [Figure 1](#) a)], prepare a test specimen which has a higher hydrogen level by forcibly charging hydrogen into the specimen. Apply constant load to the hydrogen charged test specimen and measure the time to failure. By testing specimens containing various contents of diffusible hydrogen, which is mainly responsible for hydrogen embrittlement, the relationship between diffusible hydrogen content and times to failure can be obtained. Diffusible hydrogen content can be measured by thermal desorption analysis using the test specimen after failure. This method can provide at least qualitative comparison of the resistance to hydrogen embrittlement among several high strength steels having different microstructures or compositions. For the hydrogen continuous charging method [see [Figure 1](#) b)], a test specimen is loaded in one of the following three conditions:

- a) in aqueous solution at free corrosion potential;
- b) in atmospheric corrosion environments;

c) in high pressure hydrogen gas.

Then, hydrogen analysis is carried out after failure of the specimen. If specimens do not fail within 100 h (up to 200 h if so formerly agreed), qualitative comparison of the resistance to hydrogen embrittlement can be made by hydrogen analysis of unbroken specimens.

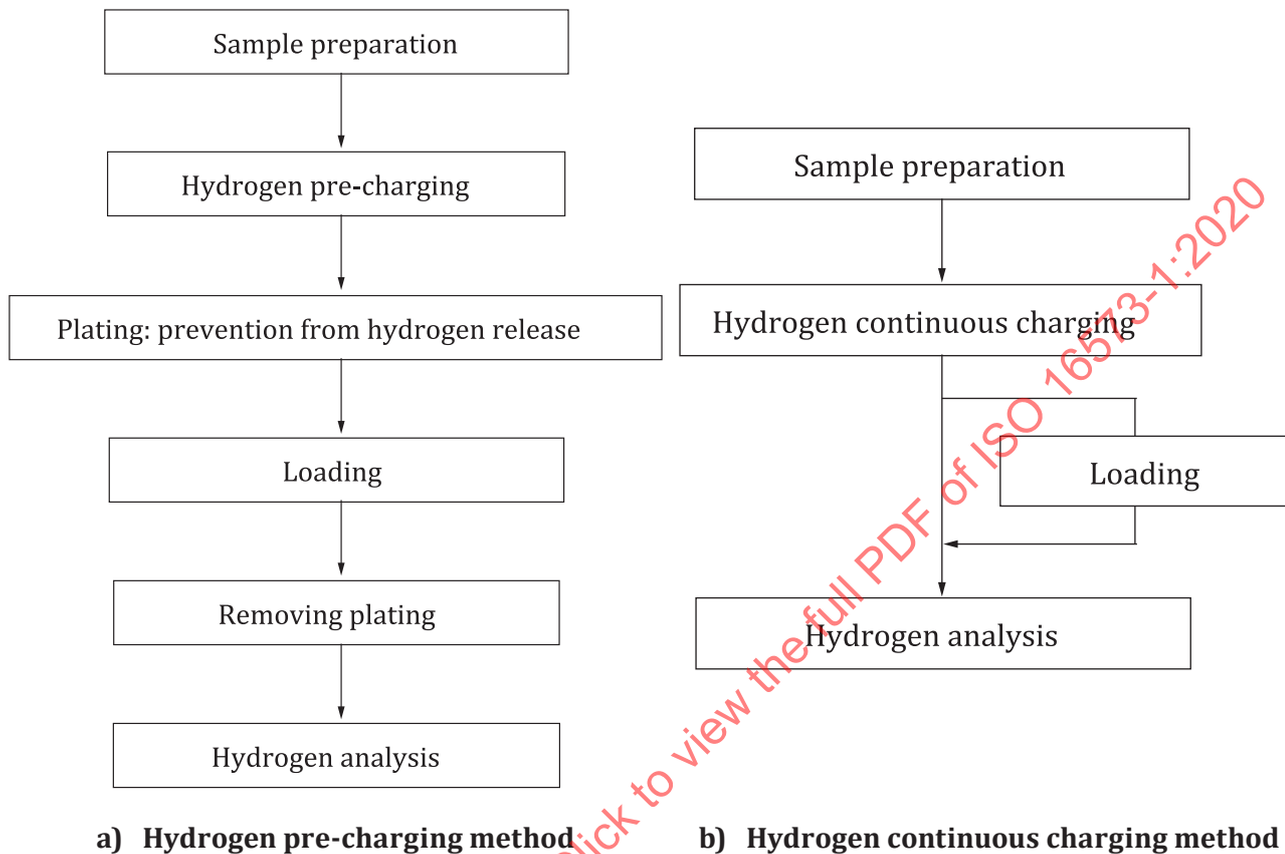
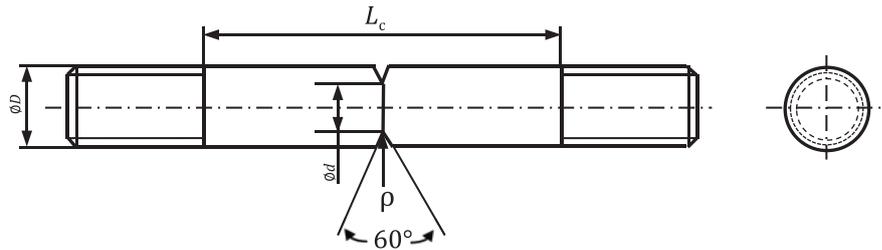


Figure 1 — Flow chart illustrating the test methods

5 Specimen preparation

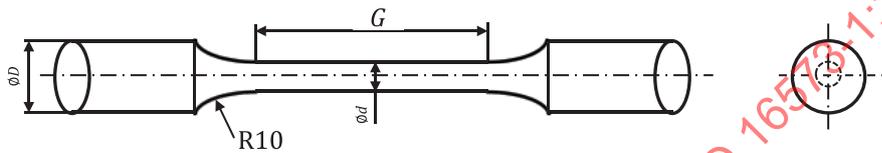
The dimensions of the specimens shall be in accordance with [Figure 2](#), and other configurations of the test specimen may be applied. Unless otherwise specified, diameter of the specimen shall be 10 mm. For samples with smaller diameter (i.e. $D = 5$ mm), $\rho/D = 0,02$ may be applied^{[2][3]}.



NOTE 1 ρ is radius of the notch bottom.

NOTE 2 Some types of specimen don't have thread.

a) Notched specimen



b) Smooth specimen

Key

d/D 0,6

ρ/D 0,01 or 0,02

L_c/D 7

G/D 5

NOTE 1 ρ is radius of the notch bottom.

NOTE 2 Some types of specimen don't have thread.

Figure 2 — Dimensions and shape of specimens

6 Hydrogen charging methods

6.1 General

There are four hydrogen charging methods, such as cathodic charging, hydrogen absorption in aqueous solution at free corrosion potential, hydrogen absorption in atmospheric corrosion environments and hydrogen absorption in high pressure hydrogen gas. The examples of the condition of each method are as follows.

6.2 Cathodic charge method

6.2.1 Hydrogen charging solution

To estimate the effect of hydrogen on the mechanical properties of steels, the hydrogen is forced to diffuse into the specimens by the cathodic charging method. For hydrogen pre-charging, the charging solution should be prepared and the chemical compositions of the solutions are listed in [Table 1](#).

Two kinds of solutions may be used for hydrogen pre-charging. Solution 1 may be used for introducing a relatively large amount of hydrogen to the specimens and Solution 2 may be used for introducing a small amount of hydrogen.

Table 1 — Chemical composition of the solutions for hydrogen charging

Charging solution	Element	Content g/l	Mark
Solution 1	NaCl	30	Large amount of hydrogen
	NH ₄ SCN	3	
Solution 2	NaOH	4	Small amount of hydrogen

6.2.2 Hydrogen charging conditions

The electro-chemical cell for hydrogen pre-charging may be placed in a 200 ml to 1 000 ml beaker. It is recommended that the anode of the electrochemical cell be made of platinum wire of spiral type of 0,5 mm in diameter and 2 m in length (counter electrode), and the specimen works as the cathode (working electrode). After the Pt wire and the specimen are placed in the cell, apply the constant current of its current density in the range of 0 A/m² ~ 20 A/m² by using potentiometer/galvanostat for 48 h. A charging time of 48 h is recommended, but other charging times are also optional as long as a total time of 72 h is reached for hydrogen charging and the homogenization treatment by room temperature exposure after cadmium (Cd) plating. For materials with low hydrogen diffusion coefficient, the hydrogen charging time and the total time may be increased. The specimen's surface area shall be calculated for proper current supply. The pre-charged hydrogen amount may be changed by varying the current density or pre-charging time. However, it is recommended to use fixed pre-charging time and current density to get reproducible test results.

6.3 Hydrogen absorption in aqueous solution at free corrosion potential

For hydrogen charging by corrosion in acid, HCl solutions or HCl with CH₃COOH/CH₃COONa buffered solutions are often used. For example, the specimen shall be immersed in 5 % HCl solution at room temperature. Immersion time shall be determined based on the specimen size and hydrogen diffusion coefficient of the tested material.

6.4 Hydrogen absorption in atmospheric corrosion environments

For hydrogen charging by atmospheric corrosion, the salt spray test (SST) or cyclic corrosion tests (CCT) including salt spraying, drying and humidifying are carried out. As an example, desirable CCT processes are listed in [Table 2](#).

Table 2 — Example of CCT processes

Process	Conditions	Time
Salt spray	5 % NaCl, 35 °C	2 h
Dry	20 % to 30 % humidity, 60 °C	4 h
Wet	≥95 % humidity, 50 °C	2 h

6.5 Hydrogen absorption in high pressure hydrogen gas

For hydrogen charging by hydrogen gas, the specimens shall be exposed directly to gaseous hydrogen up to 140 MPa at above room temperature. Exposure time shall be determined based on the specimen size and hydrogen diffusion coefficient of the tested material. However, extreme care is necessary, and it is not recommended due to the danger of experiment.

7 Preparation of electroplating solution and electroplating condition

7.1 General

The plating process is applied for the hydrogen pre-charging method in order to prevent hydrogen release during the loading test.

7.2 Electroplating solution

After hydrogen pre-charging, plating shall be conducted to prevent the release of hydrogen from the specimens during constant loading test. Cd is a well-known toxic material but the hydrogen diffusivity in Cd is known to be nearly zero. Instead of using toxic Cd, other appropriate plating materials, for example, Zn may be applied. The elements for Cd plating solution and the amounts of each element are listed in Table 3. The pH of the solution shall be 3 pH to 5 pH. To adjust the pH of the solution, one can add H_3BO_4 or ammonium solution (see Table 3).

IMPORTANT — When Cd coating is applied, solution should be treated with special care including regulatory compliance.

Table 3 — Elements for Cd and Zn plating solutions and the amounts of each element

Solution	Element	Type	Amount
Cd plating solution	$Cd(BF_4)_2$	solution [(C) = 50 %]	427 g
	NH_4BF_4	solute	48 g
	H_3BO_4	solute	21,6 g
	H_2O (distilled)	solvent	460 ml
	pH	—	3 to 5
Zn plating solution	$ZnCl_2$	solute	45 g
	KCl	solute	234 g
	H_3BO_4	solute	21,6 g
	Sodium Dodecyl Sulfate	solute	0,5 g
	H_2O (distilled)	solvent	700 ml
	pH	—	5 to 6

7.3 Electroplating conditions

An electrochemical reaction is used for Cd plating. Immerse the metal platinum (Pt) anode into the prepared Cd solution, and immerse the specimen in order for it to work as a cathode. After hydrogen pre-charging, clean the specimens by mechanical grinding (polishing) or with 0,1 N HCl and 0,1 N NaOH by ultrasonic cleaner for 30 s, respectively. Then, immerse the specimens in Cd plating solution and apply a constant current of 0,5 A for 300 s. Other conditions of current and time may also be used for Cd plating. Using the potential difference, Cd is adhered to the surface of the specimen.

The thickness of Cd plating should be at least 15 μm or higher to prevent hydrogen release. Extreme care is needed to handle the solutions containing Cd during the entire Cd plating procedures.

8 Constant loading test

8.1 Constant loading test procedures

The suggested methods for the constant loading test are the following.

- Use adequate jig to apply uniaxial loading to the specimens.

- b) Specimens containing various diffusible hydrogen contents shall be used to measure the time to failure. In the case of hydrogen continuous charging, various current densities shall be applied to measure the time to failure of a specimen.
- c) Unless otherwise specified, stress ratio (applied stress/tensile stress) shall be 0,8. Other stress ratio may be agreed between manufacturer and purchaser.

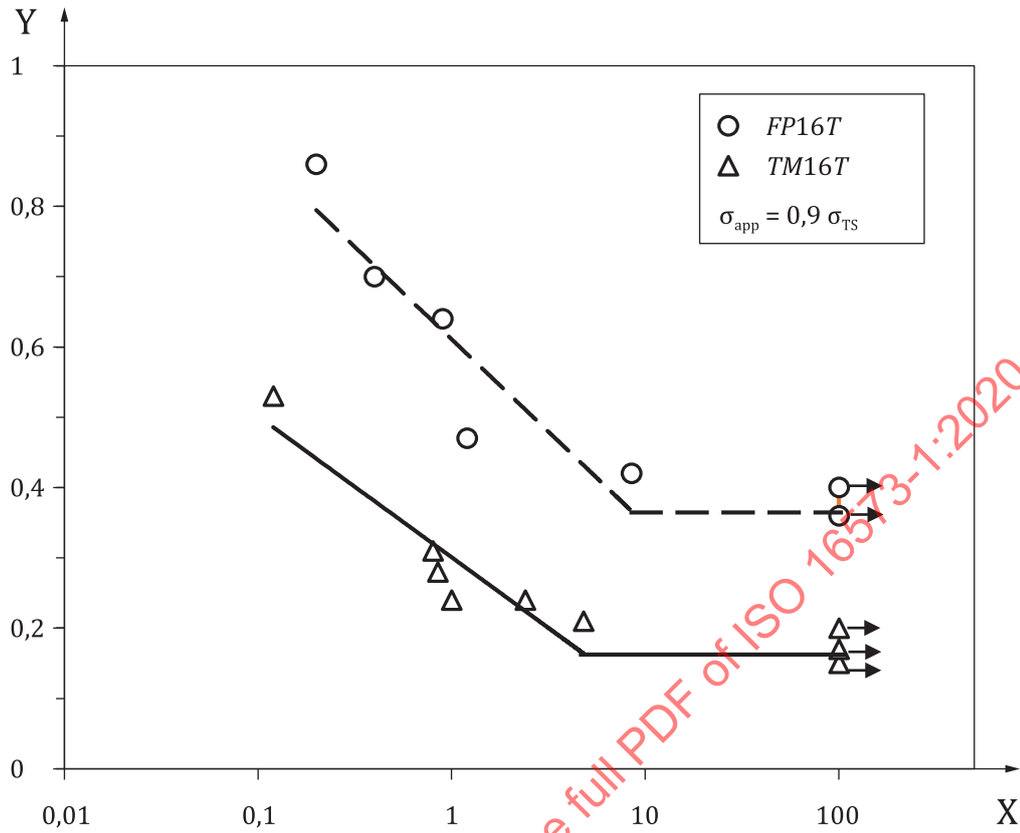
NOTE Tensile strength is measured in evaluated type of specimen without hydrogen charge in advance.

- d) If the specimens are not fractured after up to 100 h (up to 200 h, if necessary), one can conclude that these specimens do not fracture under these conditions and discontinue the test.
- e) To ensure reliability, it is recommended that at least three specimens be tested at the same stress ratio.
- f) After the constant loading test, diffusible hydrogen contents of the specimen should be immediately measured. If it is impossible to measure diffusible hydrogen content immediately following the test, specimens shall be preserved below $-50\text{ }^{\circ}\text{C}$, preferably in liquid nitrogen, to prevent hydrogen release from the fractured surface.

8.2 Presentation of the results

- a) [Figure 3](#) shows an example of the results of the constant loading test.
- b) The abscissa indicates the time to failure of the specimens and the ordinate represents the diffusible hydrogen content or the applied stress ratio (applied stress/notch tensile stress).
- c) The critical hydrogen content below which the fracture does not occur is defined as the amount of hydrogen of the specimen which is not fractured up to 100 h (see [Figure 3](#)).
- d) It may be concluded that the susceptibility to hydrogen embrittlement is greater if time to failure is shorter when tested at the same stress ratio and comparable hydrogen content.

As an additional evaluation method, local hydrogen concentration and local stress may be calculated by numerical method for comparing the results of specimens with different stress concentration factors^[5].



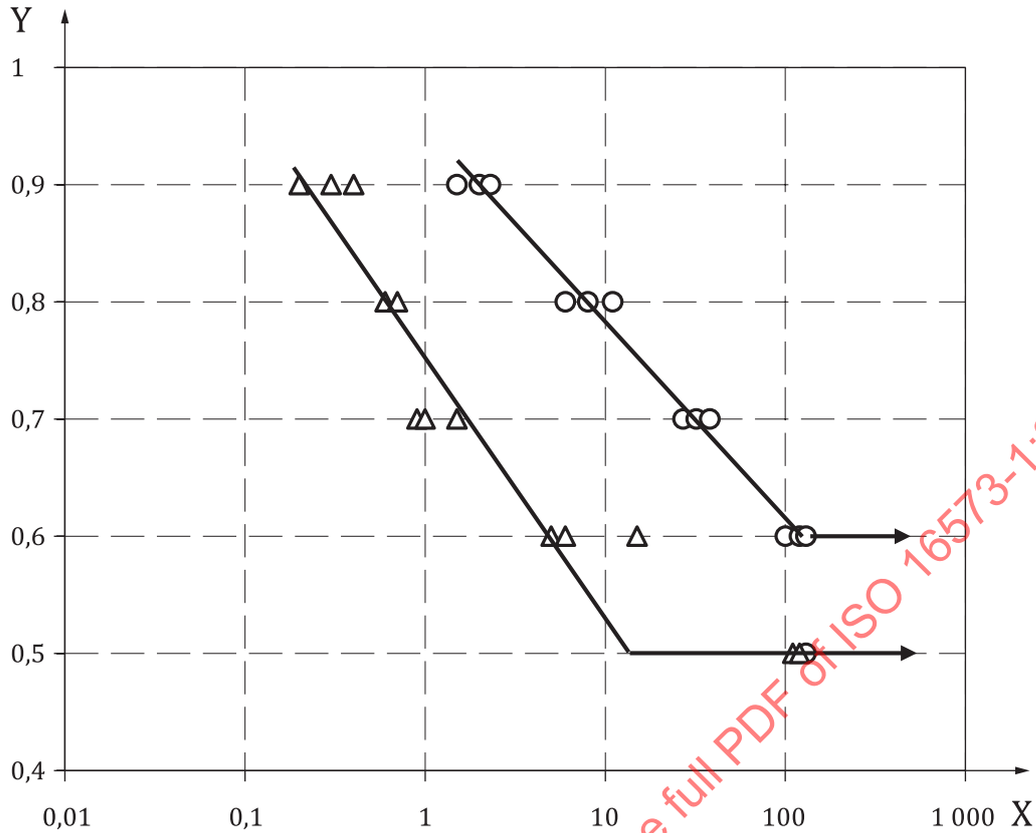
Key

X time to failure, h

Y diffusible hydrogen content, % (mass fraction)

FP16T and TM16T denote full pearlite microstructure and tempered martensite microstructure, respectively, having the same tensile strength of 1 600 MPa^[6]

Figure 3 — Example of the results of the constant loading test (time to failure vs diffusible hydrogen content)



Key

X time to failure, h

Y stress ratio, σ_{ap}/σ_{nb}

Stress ratio (σ_{ap}/σ_{nb}) represents applied stress/notch tensile stress

Figure 4 — Example of the results of the constant loading test

9 Post-test specimen treatment

For fractured specimens or specimens not fractured after 100 h or more have elapsed, the following process shall be applied.

- a) To prevent hydrogen desorption, keep fractured specimens or specimens not fractured after 100 h or more have elapsed in liquid nitrogen, or freezer or cooler box containing dry ice below $-50\text{ }^{\circ}\text{C}$.
- b) Samples for the hydrogen content analysis are prepared as follows (see [Figure 5](#)).

In case of broken specimen, cut out at a predetermined length from the flat part closest to the broken surface.

In case of unbroken specimen, cut out at a predetermined length from the centre of the specimen.