INTERNATIONAL STANDARD

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Corrosion of metals and alloys — Accelerated cyclic corrosion tests with exposure to synthetic ocean water salt-deposition process — "Dry" and "wet" conditions at constant absolute humidity

Corrosion des métaux et alliages — Essais de corrosion cyclique accélérée avec exposition à l'eau de mer synthétique par procédé de dépôt de sel — Conditions "sèches" et conditions "humides" à taux d'humidité absolue constant



Reference number ISO 16539:2013(E)

ISO 16539:2013(E)



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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.

International Standards are drafted in accordance with the rules given in the ISO/IEC Directives, Part 2.

The main task of technical committees is to prepare International Standards. 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 casting a vote.

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.

ISO 16539 was prepared by Technical Committee ISO/TC 156, Corrosion of metals and alloys.

Introduction

Corrosion of metallic materials, with or without corrosion protection, is influenced by many environmental factors, the importance of which can vary with the type of metallic materials and with the type of environment. It is impossible, therefore, to design accelerated laboratory corrosion tests in such a way that all environmental factors influencing resistance to corrosion are taken into account. Laboratory tests are, therefore, designed to simulate the effects of most important factors, which enhance the corrosion of metallic materials.

The accelerated corrosion tests described in this International Standard are designed to simulate and enhance the environmental influence on a metallic material of exposure to outdoor climates, where exposure to salt-contaminated conditions occurs and can promote corrosion. It has been prepared by reference to technical papers and reports (see the Bibliography).

The particular advantages of two tests described in this International Standard over conventional accelerated tests, such as the neutral salt spray test (NSS) as specified in ISO 9227 and the wet (salt fog)/dry/humidity test as specified in ISO 14993, lies in their better ability to reproduce the corrosion that occurs in atmospheric environments which contain much sea salt.

Accelerated corrosion tests to simulate atmospheric corrosion in such environments are intended/expected to include the following requirements.

- a) Constant absolute humidity: It is generally observed that temperature and relative humidity change under a constant absolute humidity in outdoor environments. The water absorption of deposited salts is an important factor affecting atmospheric corrosion behaviour. The same relationship, constant absolute humidity, as an actual environment is intended/expected to exist for temperature and relative humidity during dry/wet cycles.
- b) Control of the amount of salt deposition: The amount of the salt deposition on test specimens is intended/expected to be changed according to the corrosivity of the atmosphere in which the tested metallic material can be used. A salt-containing test solution is intended/expected to be diluted or the spraying time is intended/expected to be adjusted to provide the same yearly average amount of the salt deposition as an actual environment.

Therefore, the tests described in this International Standard involve the salt deposition and cyclic dry/wet conditions under a constant absolute humidity.

The results obtained do not permit far-reaching conclusions on the corrosion resistance of the tested metallic material under the whole range of environmental conditions in which it can be used. Nevertheless, the tests provide valuable information on the relative performance of materials exposed to salt-contaminated environments similar to those used in the tests.

The International Organization for Standardization (ISO) draws attention to the fact that it is claimed that compliance with this International Standard can involve the use of a patent concerning the tests given in Clause 8, Table 3, and Figure 2.

ISO takes no position concerning the evidence, validity, and scope of this patent right.

The holder of this patent right has ensured the ISO that he/she is willing to negotiate licences under reasonable and non-discriminatory terms and conditions with applicants throughout the world. In this respect, the statement of the holder of this patent right is registered with ISO. Information can be obtained from:

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Corrosion of metals and alloys — Accelerated cyclic corrosion tests with exposure to synthetic ocean water salt-deposition process — "Dry" and "wet" conditions at constant absolute humidity

1 Scope

This International Standard specifies two accelerated corrosion test procedures, Methods A and B, for the evaluation of corrosion behaviour of surface-treated metals and their alloys with and without paint on them in atmospheric environments. It also specifies the apparatus used. The two tests involve salt deposition and dry/wet conditions at a constant absolute humidity.

Method A applies to:

metals and their alloys (including corrosion-resistance alloys)

Method B applies to:

metals and their alloys

metals and their alloys with coatings [including metallic coatings (anodic or cathodic), organic coatings, and conversion coatings]

2 Normative references

The following documents, in whole or in part, are normatively referenced in this document and are indispensable for its application. For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.

ISO 4628-1, Paints and varnishes — Evaluation of degradation of coatings — Designation of quantity and size of defects, and of intensity of uniform changes in appearance — Part 1: General introduction and designation system

ISO 4628-2, Paints and varnishes — Evaluation of degradation of coatings — Designation of quantity and size of defects, and of intensity of uniform changes in appearance — Part 2: Assessment of degree of blistering

ISO 4628-3, Paints and varnishes — Evaluation of degradation of coatings — Designation of quantity and size of defects, and of intensity of uniform changes in appearance — Part 3: Assessment of degree of rusting

ISO 4628-4, Paints and varnishes — Evaluation of degradation of coatings — Designation of quantity and size of defects, and of intensity of uniform changes in appearance — Part 4: Assessment of degree of cracking

ISO 4628-5, Paints and varnishes — Evaluation of degradation of coatings — Designation of quantity and size of defects, and of intensity of uniform changes in appearance — Part 5: Assessment of degree of flaking

ISO 8407, Corrosion of metals and alloys — Removal of corrosion products from corrosion test specimens

ISO 8993, Anodizing of aluminium and its alloys — Rating system for the evaluation of pitting corrosion — Chart method

ISO 9227, Corrosion tests in artificial atmospheres — Salt spray tests

ISO 10289, Methods for corrosion testing of metallic and other inorganic coatings on metallic substrates — Rating of test specimens and manufactured articles subjected to corrosion tests

ISO 11130, Corrosion of metals and alloys — Alternate immersion test in salt solution

ISO 14993, Corrosion of metals and alloys — Accelerated testing involving cyclic exposure to salt mist, "dry" and "wet" conditions

ISO 17872, Paints and varnishes — Guidelines for the introduction of scribe marks through coatings on metallic panels for corrosion testing

Test solution 3

Preparation of the mixed salt solution

The mass of reagents shown in Table 1 shall be dissolved in distilled water or deionized water, with a conductivity of not higher than 20 μS /cm at 25 °C ± 2 °C, to produce a mixed salt solution with a concentration of 36,0 g/l \pm 3,6 g/l.

The composition of the mixed salt solution is the same as typical synthetic ocean water shown in ISO 11130:2010, A.3 (test solution for simulating the corrosive effects of ocean water).

Table 1 — Reagents and their concentrations of a mixed salt solution

Reagents	Concentration (g/l)
NaCl	24,53
MgCl ₂	5,20
Na ₂ SO ₄	4,09
CaCl ₂	1,16
KCl	0,695
NaHCO ₃	0,201
KBr	0,101
H ₃ BO ₃	0,027
SrCl ₂	0,025
NaF	0,003

WARNING - Handling of SrCl2 and NaF can be hazardous and shall be restricted to use by skilled chemists or conducted under their control.

pH adjustment

To adjust the pH of the solution, 5.0 g ± 0.5 g of NaOH shall be dissolved in water and diluted to total volume of 1 L to make 0,125 mol/l NaOH solution. It shall be added to the mixed salt solution to adjust the pH to 8.2 ± 0.1 at $25 \,^{\circ}\text{C} \pm 2 \,^{\circ}\text{C}$.

3.3 Preparation of the test solution

The test solution shall be used as the mixed salt solution or shall be diluted by 1:10 and 1:100 to the mixed salt solution with a concentration of 36,0 g/l \pm 3,6 g/l, 3,60 g/l \pm 0,36 g/l, and 0,360 g/l \pm 0,036 g/l. When not specified, the concentration shall be agreed by the interested parties.

4 Apparatus

4.1 Component protection

All components in contact with the test solution shall be made of, or lined with, materials resistant to corrosion by the test solution and which do not influence the corrosivity of the atomized test solution. The apparatus shall include the components described in the following.

4.2 Exposure cabinet

It is essential that temperature and humidity in exposure cabinet are controlled constantly.

4.3 Salt deposition device

The atomizer shall be made of corrosion resistant materials, e.g. glass, plastic, or titanium. The atomized test solution shall be deposited homogeneously on specimens. The amount of deposited test solution shall be controlled. The diameter of the atomized test solution should be equal to or less than $100 \, \mu m$.

The amount of deposited test solution can be controlled in the following manners:

- a) by controlling the test period with a continuous atomization;
- b) by controlling the amount of test solution, the atomized pressure, and the moving speed of the atomizer.

The compressed air supplied to the atomizer shall be passed through a filter to remove all trace of oil or solid matter.

4.4 Temperature and humidity control device

The device shall be capable of detecting and controlling the temperature and humidity around specimens. In the transition period of temperature and humidity, the device shall be capable of controlling the dry bulb temperature linear to target value and test period. The device shall be capable of controlling the humidity by keeping the absolute humidity constant to dry bulb temperature at least in every minute.

4.5 Rinse treatment of specimens

After the temperature and humidity cycle, the rinse treatment is conducted before re-depositing the test solution. Firstly, the test specimens are washed with drinking water, and then with deionized water. Water drops are then removed carefully with clean air in order not to take off corrosion products from the surface of the test specimens.

4.6 Types of apparatus

The following are three types of apparatus that satisfy the requirements specified in 4.1 to 4.5.

a) Two-cabinet type (automatic procedures)

Specimens move between salt deposition cabinet and exposure cabinet. The atomizer moves front to back and side to side, making it possible to change the amount of salt deposition on each specimen. Then, the specimens are moved to the exposure cabinet and wet/dry cycles and rinse treatment are conducted automatically (see $\underline{\text{Annex A}}$).

b) One-cabinet type (automatic procedures)

Specimens are settled in one cabinet. The atomizer is set, and wet/dry cycles and rinse treatment are conducted automatically (see Annex B).

c) One-cabinet type (manual procedures)

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After salt deposition (4.3) by hand, the test specimens shall be settled in the cabinet for wet/dry cycles. Then, the test specimens shall be taken out from the cabinet and rinse treatment conducted. After the salt deposition, the test specimens shall be settled up in the cabinet as quickly as possible (see Annex C).

5 **Test specimens**

- The number and type of test specimens shall be selected according to the specification for the material or product being tested. When not so specified, these details shall be mutually agreed between the interested parties.
- The test specimens shall be carefully cleaned prior to testing so as to remove those traces (dirt, oil, or other foreign matter), which could influence the result. The cleaning method employed shall depend on the nature of the materials and the contaminants, but shall not include the use of any abrasives or solvents which can attack the surface of the specimens.
- For the metals or alloy without organic coatings and inorganic coating materials, the specimens shall be thoroughly cleaned with an appropriate organic solvent using a clean soft brush or an ultrasonic cleaning device. The cleaning shall be carried out in a vessel full of solvent. After cleaning, the specimens shall be rinsed with fresh solvent, then dried.
- Unless otherwise specified, specimens intentionally coated with protective organic films should not be cleaned prior to the test. If cleaning is necessary, specimens shall be wiped with gauze impregnated with ethanol, taking care not to damage the surface of the specimens.

Care shall be taken that specimens are not contaminated after cleaning by careless handling.

If the test specimens are cut from a larger coated article, cutting shall be carried out in such a way that the coating is not damaged in the area adjacent to the cut. Unless otherwise specified, the cut edges shall be adequately protected by coating them with a suitable material, stable under the conditions of the test, such as paint, wax, or adhesive tape.

Salt deposition measurement method

The test solution shall be applied to make salt deposition on the surface of the specimen. The average amount of salt deposition shall be measured by the mass change of the specimen before and after saltdeposition process to the nearest 1 mg. These measurements shall be carried out as quickly as possible in order to prevent the specimen from drying.

Arrangement of the test specimens

After the salt deposition on the test specimens, the test specimens shall be placed in the cabinet. The test specimens shall be arranged so that they do not come into contact with the cabinet.

The specimen should be placed horizontally.

The specimens may be placed at different levels within the cabinet, as long as the droplet of the moisture does not drip from specimens or their supports at one level onto specimens placed below them.

Operating conditions and procedure

Operating conditions shall be in accordance with those given in Table 2 for Method A, and Table 3 for Method B. Operation procedure shall be in accordance with those given in Figure 1 for Method A, and Figure 2 for Method B.

A test solution that has used salt deposition shall not be re-used.

During the test, pressure in the cabinet shall maintain the atmospheric pressure.

Table 2 — Test conditions in Method A

Procedure	Con	ditions
a) Salt deposition 1) Temperature 2) Test solution 3) Frequency 4) Salt deposition	1) Room temperature 2) Described in Clause 3 3) After every cycle 4) The amount of deposited test solution on a specimen shall be 250 g/m ² ± 50 g/m ² . The concentration of the solution should be equal to or greater than 1/1 000 of synthetic ocean water.	
b) "Dry" condition	Temperature	Relative humidity
1) A	(49 ± 1) °C	(32 ± 5) %
2) B	(54 ± 1) °C	(25 ± 5) %
3) C	(55 ± 1) °C	(24 ± 5) %
4) D	(54 ± 1) °C	(25 ± 5) %
5) E	(49 ± 1) °C	(32 ± 5) %
c) "Wet" condition	(30 ± 1) °C	(90 ± 5) %
d) Rinse treatment	Clean drinking water Not exceeding 40 °C	
e) Length and composition of a single exposure cycle (A single exposure cycle is 24 h.)	"Dry" 10 h 48 min	
f) Time to reach the specified condition within a single exposure cycle	"Wet" to "Dry (A)" "Dry (A)" to "Dry (I "Dry (B)" to "Dry (I "Dry (C)" to "Dry (I "Dry (D)" to "Dry (I "Dry (E)" to "Wet"	C)" 1 h 22 min O)" 1 h 22 min

NOTE 1 Temperature and humidity shall be changed linearly at specified time intervals [see f) in this table]. Consecutive dry conditions A to E give the same absolute humidity, dew point 30° C.

NOTE 2 The tolerances (±) given for temperature and relative humidity are the allowable fluctuations of the parameter concerned about the given value under equilibrium conditions. This does not mean that the value can vary by plus/minus the amount indicated from the given value.

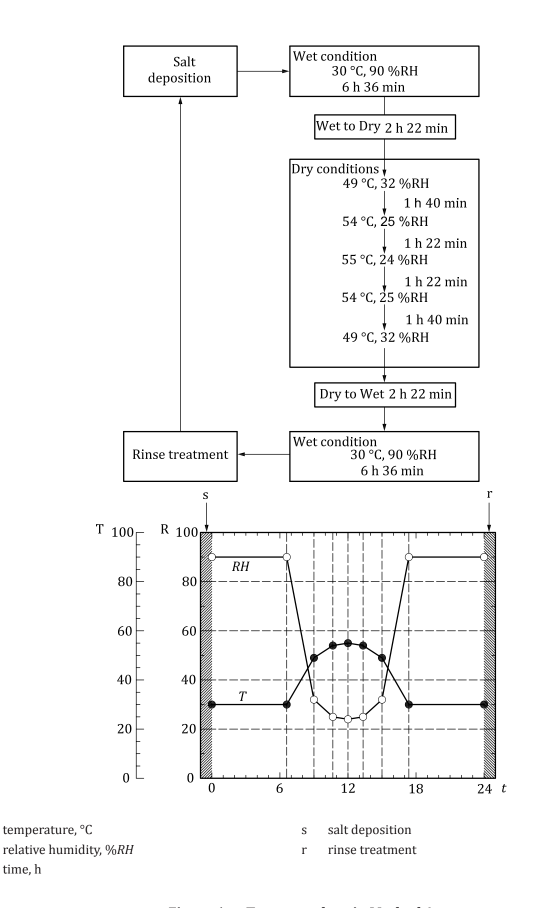


Figure 1 — Test procedure in Method A

Key

R

time, h

Table 3 — Test conditions in Method B

Procedure	Conditions	
a) Salt deposition 1) Temperature 2) Test solution 3) Frequency 4) Salt deposition	1) Room temperature 2) Described in Clause 3 3) After 8 and 11 cycles 4) The amount of deposited test solution on a specimen shall be 28,0 g/m ² ± 2,8 g/m ² . The concentration of the solution should be equal to or greater than 1/1 000 of synthetic ocean water.	
b) "Dry" condition	Temperature	Relative humidity
	(60 ± 1) °C	(35 ± 5) %
c) "Wet" condition	(40 ± 1) °C	(95 ± 5) %
d) Rinse treatment	Clean drinking water Not exceeding 40 °C	
e) Length and composition of a single exposure cycle (A single exposure cycle is 8 h)	"Dry" 3 h "Wet" 3 h Time to reach the specified conditions "Dry" to "Wet" 1 h "Wet" to "Dry" 1 h	
f) Test procedures	"Wet" to "Dry" 1 h 1. Salt deposition. 2. Dry/wet exposure cycle: 8 cycles. After dry/wet cycle, the specimens shall be removed from the cabinet immediately after the temperature reaches 60 °C. 3. Rinse treatment. 4. Salt deposition. Rinse treatment and salt deposition shall be accomplished during dry conditions. 5. Dry/wet exposure cycle: 11 cycles. After dry/wet cycle, the specimens shall be removed from the cabinet immediately after the temperature reaches	
	60 °C. 6. Rinse treatment.	

NOTE 1 $\,\,$ Temperature and humidity shall be changed at the same absolute humidity, dew point 39 °C.

NOTE 2 The tolerances (±) given for temperature and relative humidity are the allowable fluctuations of the parameter concerned about the given value under equilibrium conditions. This does not mean that the value can vary by plus/minus the amount indicated from the given value.

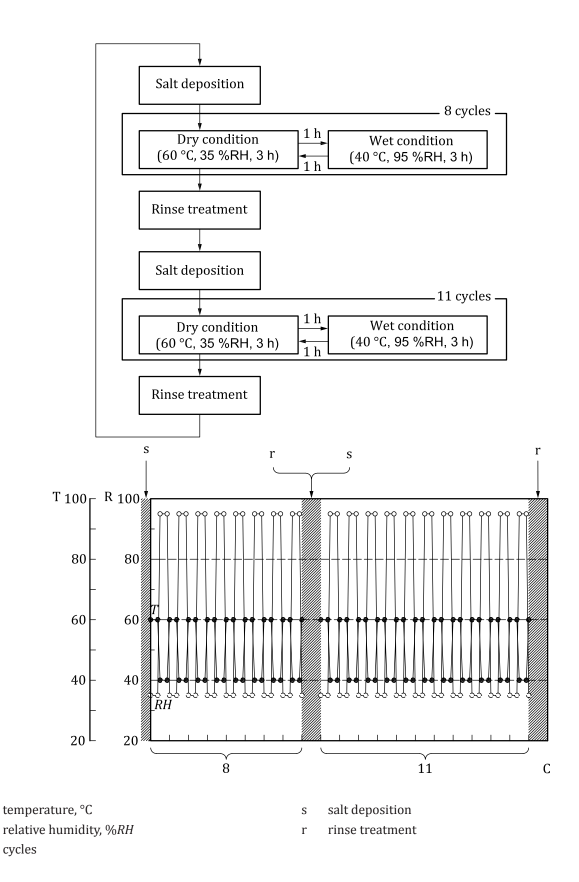


Figure 2 — Test procedure in Method B

Rinse treatment shall be conducted immediately after the temperature reaches 60 °C.

Key T

R C

9 Treatment of specimens after test

At the end of the test period, the test specimens shall be removed from the cabinet and allowed to dry for 0,5 h to 1 h before rinsing in order to reduce the risk of removing corrosion products. Before they are examined, the residues of sprayed solution shall be carefully removed from their surfaces. This may be accomplished by dipping the test specimens gently in clean running water at a temperature not exceeding 40 °C, and then drying them immediately in a stream of air at an overpressure not exceeding 200 kPa.

10 Continuity of tests

Tests shall preferably be continued without interruption through the test duration. When it is necessary to interrupt the operation, the period of interruption shall be minimized.

Where it is necessary to interrupt tests, the test specimens shall be treated as follows:

The test specimens shall be taken out from the test cabinet and treated in the manner described in Clause 9, and thereafter stored in a desiccator until testing is resumed.

11 Duration of tests

The period of test shall be as designated by the specification covering the material or product being tested. When not specified, the period shall be agreed by the interested parties.

Recommended periods of test are:

- d) for Method A: 3 cycles (72 h), 7 cycles (168 h), 21 cycles (504 h), 42 cycles (1008 h);
- e) for Method B: 7 days, 14 days, 21 days, 28 days, 42 days, 56 days, 70 days.

12 Evaluation of results

Many different criteria for the evaluation of the test results may be applied to meet particular requirements, for example:

- f) time elapsing before appearance of the first signs of corrosion;
- g) appearance after the test;
- h) number and distribution of corrosion defects (i.e. pits, cracks, rusting, or creep from scratches in the case of paints; these may be assessed by methods described in ISO 8993 or ISO 10289 and for paints in ISO 4628 (all parts) and ISO 17872;
- i) appearance after removing superficial corrosion products (see ISO 8407);
- j) change in mass;
- k) alteration revealed by micrographic examination;
- l) changes in mechanical or electrical properties.

13 Test report

The test report shall provide the following information:

- m) reference to this International Standard (i.e. ISO 16539);
- n) description of the test equipment;
- o) description of the material tested;

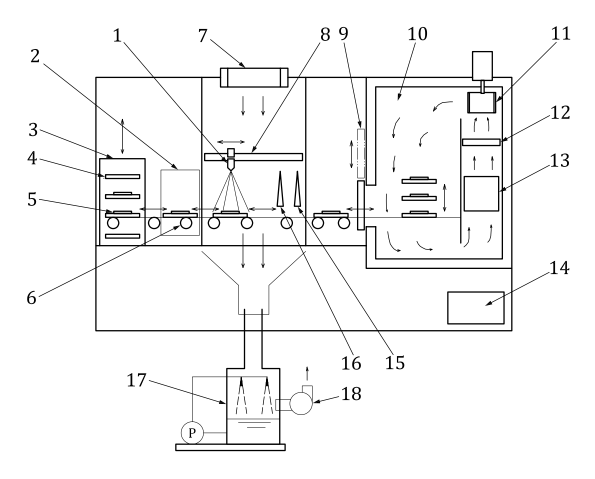
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- dimensions and shape of the test specimens and the nature and area of the surface tested;
- details of the preparation of the test specimens, including any cleaning treatment applied before testing and any protection given to edges;
- concentration of test solution, amount of salt deposition, and salt deposition procedure; r)
- temperature and relative humidity in each of the salt-deposition process and "dry" and "wet" s) conditions during test;
- frequency and duration of any interruptions; t)
- duration of the test;
- method used to clean the specimens after testing and, where appropriate, an indication of the mass loss resulting from the cleaning operation and the method used for correcting for this mass loss;
- tests results, such as mass and thickness losses for uncoated specimens and blister width and peeling width for paint specimens;
- if necessary, photographic records and/or descriptions of the appearance of tested specimens. x)

Annex A

(informative)

Combined cyclic test instrument with salt deposition unit (two cabinets)



T7
KOV
IXCV

- 1 atomizer
- 2 dry chamber
- 3 specimen storage
- 4 specimen tray
- 5 specimen
- 6 roller for specimen movement / specimen drive unit
- 7 filte
- 8 nozzle drive unit
- 9 hosting door

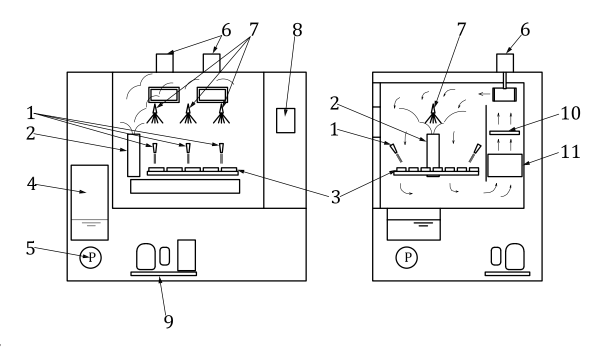
- 10 exposure cabinet
- 11 blower motor
- 12 air heater
- 13 cooling unit
- 14 refrigerator
- 15 rinse nozzle
- 16 blower (to remove water droplet)
- 17 exhaust treatment unit
- 18 exhaust blower

Figure A.1 — Combined cyclic test instrument with salt deposition unit (two-cabinet type)

Annex B

(informative)

Combined cyclic test instrument with salt deposition unit (one cabinet)



Key

- blower (to remove water droplet)
- 2 atomizer
- 3 specimen
- 4 solution supply tank
- 5 solution supply pump
- 6 blower motor

- rinse nozzle
- 8 control panel
- 9 refrigerator
- 10 air heater
- cooling unit

Figure B.1 — Combined cyclic test instrument with salt deposition unit (one-cabinet type)

Annex C (informative)

Salt deposition method by manual spraying

C.1 Apparatus

The device for spraying the test solution should comprise a supply of clean air, a supply of controlled pressure, and a reservoir containing the solution to be sprayed.

The atomizing pressure should be at an overpressure of 0,12 MPa to 0,3 MPa and preferably be controlled constantly at 0,15 MPa.

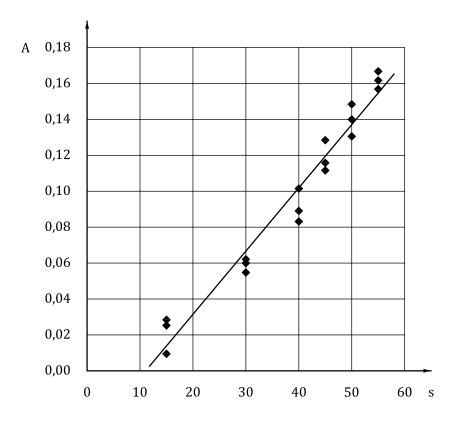
C.2 Salt deposition method

The dried specimens should be settled up vertically in front of the salt spraying device.

The distance from the spraying nozzle to the specimen should be 300 mm.

The test solution should be sprayed during the desired time to make salt deposition on the surface of the specimen. An example of the relationship between the amount of salt deposition and the spraying time is shown in Figure C.1.

The average amount of salt deposition should be measured by the mass change of the specimen before and after spraying to the nearest 1 mg. These measurements should be carried out as quickly as possible in order to prevent the specimen from drying.



Key

amount of salt deposition, (g) Α

S spraying time, (s)

The specimen used is a zinc-coated steel sheet covered chromate conversion coating, which has a NOTE dimension of 70 mm x 70 mm x 0,8 mm. The concentration of the salt solution is 3,60 g/l \pm 0,36 g/l. Atomizing pressure is 0,15 MPa. The distance from the spraying nozzle to the specimen is about 300 mm.

Figure C.1 — Relationship between the amount of salt deposition and the spraying time

Annex D

(informative)

Recommended periods of testing

Correlations between corrosion of specimens exposed in a marine environment (Okinawa in Japan), which is classified C4 according to corrosivity categories in ISO 9223, and the accelerated corrosion test were investigated.

The representative data are shown as below.

<u>Table D.1</u> shows correlations between the rusted area for stainless steels exposed in the marine environment (Okinawa in Japan) and those obtained by Method A. In general, a 13-cycle test of Method A is recommended for the ranking of stainless steels.

<u>Table D.2</u> shows mass loss of uncoated cold rolled carbon steel sheets and coated steel sheets exposed in the marine environment (Okinawa in Japan) and those obtained by Method B.

In general, a 28-day test or 56-day test of Method B is recommended for the ranking of any bare metal (alloy) or a metal protected with a thin conversion coating or a metallic, inorganic or organic coating. A 112-day test of Method B should be sufficient to rank high-quality coating systems.

Table D.1 — Comparison of percentage of area rusted due to corrosion in different tests

	Percentage of area rusted after test (%)		
Material tested code (ISO number)	Test Method A salt deposition (Cl ⁻ : 1 000 mg/m ²) 13 cycles	Exposure period 18 months (Okinawa in Japan), according to corrosivity class C4 in ISO 9223	
S304 (4301-304-00-I)	5 to 9	9 to 20 (unsheltered)	
S410L (4030-410-90-X)	2 to 5	20 to 40 (unsheltered) 40 to 100 (sheltered)	
S436L (4513-436-00-J)	2 to 5	2 to 5 (unsheltered) 5 to 9 (sheltered)	
S444 (4521-444-00-I)	0,2 to 0,4	0,2 to 0,4 (unsheltered) 0,9 to 2 (sheltered)	
S445J1 (4128-445-92-J)	0,9 to 2	0,2 to 0,4 (unsheltered) 0,2 to 0,4 (sheltered)	
S447J1 (4135-447-92-C)	0,2 to 0,4	0,09 to 0,2 (unsheltered)	
S316 (4401-316-00-I)	2 to 5	0,9 to 2 (unsheltered) 0,9 to 2 (sheltered)	
S312L (4547-312-54-I)	0,4 to 0,9	0,09 to 0,2 (unsheltered) 0,09 to 0,2 (sheltered)	

Table D.2 — Comparison of mass losses of cold rolled carbon steel sheet and coated steel sheets due to corrosion in different tests

	Mass loss after test (g/m²)			
Material tested	Test Method B salt deposition (100 mg/m²)	Exposure period 21 months (Okinawa in Japan)	According to corrosivity class C4 in ISO 9223 first year data	
CRS	130 to 230 (28 days of exposure) 300 to 500 (56 days of exposure)	520 to 800(unsheltered) 900 to 1400 (sheltered)	400 to 650	
GI	10 to 18 (28 days of exposure) 18 to 28 (56 days of exposure)	20 to 32 (unsheltered) 60 to 100 (sheltered)		
Zn-5 % Al	10 to 16 (28 days of exposure) 16 to 26 (56 days of exposure)	18 to 28 (unsheltered) 43 to 65 (sheltered)	15 to 30 (Pure zinc)	
55 % Al-Zn	2 to 4 (28 days of exposure) 3 to 5 (56 days of exposure)	7 to 12 (unsheltered) 9 to 15 (sheltered)		

CRS: Cold rolled carbon steel sheet

GI: Hot dipped galvanized steel sheet (coating weight of both sides 270 g/m²)

Zn-5 % Al: Zn-5 % Al alloyed coated steel sheet (coating weight of both sides 180 g/m²)

55 % Al-Zn: 55 % Al-Zn coated steel sheet (coating weight of both sides 150 g/m²)

Annex E

(informative)

Methods for evaluation of corrosivity of test

E.1 Reference specimens

To verify the apparatus, four reference specimens of the following material shall be used:

CR 4 grade steel according to ISO 3574, with a surface free from pores, slight marks, and scratches, and with a matt finish (arithmetical mean deviation of the profile $Ra = 0.8 \mu m \pm 0.3 \mu m$).

The reference specimens should have a dimension of 70 mm × 70 mm.

The reference specimens shall be carefully cleaned immediately before testing. Cleaning shall eliminate all those traces (dirt, oil, or other foreign matter) which could influence the test results.

After drying, the mass of the reference specimens shall be determined to the nearest 1 mg.

Protect back side and edge face of the reference specimens with a removal coating, for example, an adhesive plastic film.

E.2 Arrangement of the reference specimens

Four reference specimens shall be positioned in four quadrants in the zone of the cabinet where the test specimens are placed. The specimen shall be flat and placed horizontally.

The supports of the reference test specimens shall be made of or coated with plastics, glass, or other inert insulating materials, which would not influence corrosion of the specimen being tested.

E.3 Determination of mass loss

At the end of the test, the reference specimens shall be immediately taken out of the test chamber. Then the corrosion products shall be removed by repetitive cleaning in accordance with ISO 8407.

The following chemical cleaning procedures shall be used.

For steel, a 20 % (volume fraction) solution of diammonium citrate [(NH₄)₂HC₆H₅O₇], of analytical reagent grade, in water, for 10 min at 23 °C. After each cleaning cycle, the specimens shall be rinsed under running water at ambient temperature with light brushing followed by drying.

The mass of the reference specimens shall be determined to the nearest 1 mg.

From the plot of mass versus number of cleaning cycles, determine the true mass of the specimen after removal of the corrosion products as described in ISO 8407. Subtract this number from the initial mass of the reference specimen prior to testing and divide the resulting number by the area of the exposed surface of the reference specimen to calculate the metal mass loss per square metre of the reference specimen.

E.4 Satisfactory performance of cabinet

The operation of the test apparatus is satisfactory if the mass loss of each reference specimen is within the range in <u>Tables E.1</u> and <u>E.2</u>.

Table E.1 — Allowed range of mass loss of the reference specimens (Test Method A)

True	Salt deposition	Mass loss (g/m²)	
Туре	(mg/m ²)	5 days	10 days
M-41 J A	100	12 ± 2	30 ± 8
Method A	1 000	40 ± 10	90 ± 30

Table E.2 — Allowed range of mass loss of the reference specimens (Test Method B)

Type	Salt deposition	Mass loss (g/m²)	
Type	(mg/m ²)	28 days	56 days
	10	80 ± 20	140 ± 30
Method B	100	180 ± 50	400 ± 100
	1 000	800 ± 200	2 000 ± 500

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- NOTE References [1], [2], [6], [9], [10], and [11] are for Method A; references [3], [4], [7], and [8] are for Method B; and reference [5] is for Method A and Method B.

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