Factory Inspection Report

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This document contains:

- two cover pages
- a report form of 14 pages
- Inspector’s Evaluation – Findings
- Inspector’s Evaluation - Informative
- TEST DATA SHEET - Product Verification Test
- TEST DATA SHEET  Routine Tests
- IDENTIFICATION OF SELECTED SAMPLE

PD CIG 023 reports shall not contain any unauthorised modifications which change the original meaning or the requirements. Any additions created to any document in the series shall be shown in an Appendix.
FACTORY INSPECTION REPORT

Inspection carried out by (Name of Inspection Body):
Reference number of the Body carrying out the inspection:
– For page control, please write this number in the header of each page (including the attachments)

GENERAL GUIDANCE
– The questions of this factory inspection report are based on the requirements given in Permanent Document CIG 021.
– Guidance for the inspector is given in Permanent Document CIG 024.
– Both documents, PD CIG 021 and PD CIG 024 shall be taken into account during inspection.
– Instructions to the Inspector are shown in italics
– The report shall be completed even if there is no production at the time of the visit.
– For all ‘NO’ answers details shall be provided on the INSPECTORS EVALUATION-Findings page
– For all ‘N/A’ answers rationale shall be provided as to why the item is not applicable
– Details should be given on INSPECTOR’S EVALUATION-Informative page.

1.0 GENERAL INFORMATION

1.1 Manufacturer’s registered name and factory location

Manufacturer’s registered name:
Street address of the factory and Number:
Postal code:
City:
County:
Country:
GPS-coordinates: (optional)

1.2 Manufacturer’s representative name and contact data

Manufacturer’s representative name:
Position:
Telephone:
Fax:
E-Mail:
Reference number of the body carrying out the inspection:

1.3 Record below the names and position held of the main people involved in the inspection

- [ ] same as mentioned under 1.2

If not the same as mentioned under 1.2 please give details

<table>
<thead>
<tr>
<th>Name</th>
<th>Position</th>
<th>Telephone</th>
<th>Fax</th>
<th>E-Mail</th>
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1.4 [ ] Pre-Licence    [ ] Routine    [ ] ENEC

[ ] HAR    [ ] EMC    Others:

1.5 Pre-Licence only: Is the information given in the Questionnaire CIG 022 Section B accurate and complete?

If ‘no’, amend the Questionnaire as appropriate and attach a copy to this report.

<table>
<thead>
<tr>
<th>YES</th>
<th>N/A</th>
<th>NO</th>
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<tbody>
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</table>

1.6 Inspection Details:

<table>
<thead>
<tr>
<th>Certification Body requesting inspection</th>
<th>Inspection X of Y</th>
<th>File Reference No.</th>
<th>Product Category</th>
<th>Type of Product</th>
</tr>
</thead>
<tbody>
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</tbody>
</table>

1.7 Name of Inspector Date of inspection:

(YYYY – MM – DD )
### Verification of purchased components and materials which have a safety implication on the certified product (Incoming Inspection)

#### 2.1 Are materials, components and sub-assemblies verified by the manufacturer as complying with appropriate specification?

<table>
<thead>
<tr>
<th></th>
<th>YES</th>
<th>N/A</th>
<th>NO</th>
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</thead>
<tbody>
<tr>
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</table>

#### 2.2 Does this verification also include the verification of the Certification Marks?

<table>
<thead>
<tr>
<th></th>
<th>YES</th>
<th>N/A</th>
<th>NO</th>
</tr>
</thead>
<tbody>
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</tbody>
</table>

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**Description of the procedure (one or more boxes may be ticked)**

- Rely on suppliers’ out-going inspection / Suppliers’ quality plan
- Audit conducted at the suppliers’ premises
- Supplier control based on manufacturers’ check list
- Conduct own incoming inspection
- Identification check
  - Checked for correct type
  - Comparison to a reference
  - Rating
  - Certification mark
- Certificate of conformity
- Others
- Details given on INSPECTOR’S EVALUATION-Informative page

**Description of the procedure or ref. of documented procedure & revision or issue date:**

- Details are given on INSPECTOR’S EVALUATION-Informative page.

#### 2.3 If the manufacturer relies on Certificates of Conformity, do they clearly identify the product, quantity of items covered, the specification to which the products conform, the production date and are they properly issued?

<table>
<thead>
<tr>
<th></th>
<th>YES</th>
<th>N/A</th>
<th>NO</th>
</tr>
</thead>
<tbody>
<tr>
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</tbody>
</table>

#### 2.4 Is there a procedure covering the way to handle non-conforming components and materials?

<table>
<thead>
<tr>
<th></th>
<th>YES</th>
<th>N/A</th>
<th>NO</th>
</tr>
</thead>
<tbody>
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</tbody>
</table>

**Description of the procedure or ref. of documented procedure & revision or issue date:**

- Details are given on INSPECTOR’S EVALUATION-Informative page.

#### 2.5 Is the procedure and the way in which it is applied satisfactory?

- (e.g.: components and materials clearly identified and/or segregated to prevent unauthorised use?)

<table>
<thead>
<tr>
<th></th>
<th>YES</th>
<th>N/A</th>
<th>NO</th>
</tr>
</thead>
<tbody>
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</tbody>
</table>

#### 2.6 Are records of the incoming inspection maintained and satisfactory?

<table>
<thead>
<tr>
<th></th>
<th>YES</th>
<th>N/A</th>
<th>NO</th>
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</tbody>
</table>
## Production Control, Inspection and Routine Tests

### 3.1 Are the Quality Assurance and manufacturing Personnel adequately briefed on their duties?

- **YES**
- **N/A**
- **NO**

### 3.2 Do they have readily available up-to-date documents, manufacturing and test instructions, photographs, drawings or samples on all those parts which have an impact on the safety of the finished products?

- **YES**
- **N/A**
- **NO**

### 3.3 Is there evidence that the production process ensures that the final product is identical to the reference version as described in clause 15.1?

- **YES**
- **N/A**
- **NO**

### 3.4 Is there a procedure to ensure that all products will be tested or inspected according to the manufacturer’s requirements?

- **YES**
- **N/A**
- **NO**

**Description of the procedure or ref. of documented procedure & revision or issue date:**

Details are given on INSPECTOR’S EVALUATION-Informative page.

### 3.5 Is the production process controlled at appropriate stages?

- **YES**
- **N/A**
- **NO**

### 3.6 Are products inspected at appropriate stages of manufacture (Production Line Inspection)?

- **YES**
- **N/A**
- **NO**

**Give details of all tests and inspections performed by the manufacturer and enter in the routine test table on the TEST DATA SHEET**

### 3.7 Do the Routine Tests entered on the TEST DATA SHEET sufficiently cover all the Certification Bodies’ requirements?

- **YES**
- **N/A**
- **NO**

### 3.8 Is there a procedure covering the way to handle non-conforming products?

- **YES**
- **N/A**
- **NO**

**Description of the procedure or ref. of documented procedure & revision or issue date:**

Details are given on INSPECTOR’S EVALUATION-Informative page.
### Procedure of handling non-conforming products (one or more boxed may be ticked)

- Automated segregation process
- Manual segregation process
- Non-conforming products are destroyed
- Non-conforming products are repaired
- Others (please give details)
- Details given on INSPECTOR’S EVALUATION-Informative page

| 3.9 | Is the procedure and the way in which it is applied satisfactory? (e.g. non-conforming products clearly identified or segregated to prevent unauthorised use?) | YES | N/A | NO |

| 3.10 | Are repaired and reworked (corrected) items **again** subjected to appropriate tests/inspections in accordance with procedures? | YES | N/A | NO |

**Description of the procedure or ref. of documented procedure & revision or issue date:**
- Details are given on INSPECTOR’S EVALUATION-Informative page.

| 3.11 | Are test records of the routine tests maintained and satisfactory? | YES | N/A | NO |

| 3.12 | Are records kept at least for the period between two inspection visits? | YES | N/A | NO |

### Functional Check on Test and Measuring Equipment used for Safety Tests (Dummy Test)

| 4.1 | Is there a procedure describing how the functional checks shall be conducted? | YES | N/A | NO |
- Automated process
- Manual process

**Description of the procedure or ref. of documented procedure & revision or issue date:**
- Details are given on INSPECTOR’S EVALUATION-Informative page.

| 4.2 | Is there evidence that the functional check of the equipment is conducted properly, even if certified products were not in production? | YES | N/A | NO |

| 4.3 | Is a functional check conducted with intervals which will allow previous production to be retested if incorrect functioning is detected before it leaves the factory? | YES | N/A | NO |
## 4.4 Is the proper function of the test equipment verified with a simulated failure (dummy) or by other equivalent means?
- **YES**
- **N/A**
- **NO**

- Simulated failure (dummy)
- Test procedure according to the equipment manual
- Internal self test; test program included in equipment certification
- Internal self test; verified by the inspector

## 4.5 Is there evidence that the simulated failure (dummy) (if used) represents the tripping limits?
- **YES**
- **N/A**
- **NO**

## 4.6 Is there a procedure requiring appropriate actions to be taken by the operator if a functional check is found to be unsatisfactory?
- **YES**
- **N/A**
- **NO**

**Description of the procedure or ref. of documented procedure & revision or issue date:**
- Details are given on INSPECTOR’S EVALUATION-Informative page.

## 4.7 Is this procedure appropriate to ensure that improperly checked products are re-tested?
- **YES**
- **N/A**
- **NO**

## 4.8 Are subsequent corrective actions taken recorded in all cases?
- **YES**
- **N/A**
- **NO**

## 4.9 Are the test records of results of functioning checks of test and measuring equipment maintained and satisfactory?
- **YES**
- **N/A**
- **NO**

## 4.10 Are records kept at least for the period between two inspection visits?
- **YES**
- **N/A**
- **NO**

## 5 Products seen in Production during visit

*Identify type number and any certification mark that appeared on products seen in production at the time of the visit. If no certified products were seen, indicate what kinds of products were manufactured at the time of visit.*

*The manufacturing process should nevertheless be examined.*

*At least one kind of product per product category and electrical insulation class shall be listed.*

- No production
- Production seen

*Complete TEST DATA SHEET for each kind of product per product category and electrical insulation class even if there is no production.*
## Calibration of Safety Test and Measuring Equipment

### 6.1 Is test and measuring equipment used calibrated or verified?

<table>
<thead>
<tr>
<th></th>
<th>YES</th>
<th>N/A</th>
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(Several boxes may be ticked)
- [ ] **Verification** done by the manufacturer by means of calibrated reference equipment
- [ ] **Calibration** done by:
  - [ ] Laboratory accredited according to ISO/IEC 17025
  - [ ] Test equipment manufacturer/supplier
  - [ ] National metrology institute
  - [ ] Other (please provide details):

### 6.2 Is reference equipment (used for verification) calibrated?

<table>
<thead>
<tr>
<th></th>
<th>YES</th>
<th>N/A</th>
<th>NO</th>
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</tbody>
</table>

(Several boxes may be ticked)
Calibration of reference equipment done by:
- [ ] Laboratory accredited according to ISO/IEC 17025
- [ ] Test equipment manufacturer/supplier
- [ ] National metrology institute
- [ ] Other (please provide details):

### 6.3 Is the equipment provided with a label or similar indicating the next calibration/verification due date?

<table>
<thead>
<tr>
<th></th>
<th>YES</th>
<th>N/A</th>
<th>NO</th>
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### 6.4 Do the calibration/verification records indicate that calibration is traceable to national/international standards of measurement?

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<thead>
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<th>YES</th>
<th>N/A</th>
<th>NO</th>
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### 6.5 Are the records for calibration/verification of test and measuring equipment maintained and satisfactory?

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<tr>
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<th>YES</th>
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### 6.6 Are records kept at least for the period between two inspection visits?

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<tr>
<th></th>
<th>YES</th>
<th>N/A</th>
<th>NO</th>
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</table>
## 7 Handling and Storage

7.1 Are the components and materials to be used for production stored and handled in such a way as to ensure that they will continue to comply with the applicable standards?  
- **YES**  
- **N/A**  
- **NO**

7.2 Are the finished products stored and handled in such a way as to ensure that they will continue to comply with the applicable standards?  
- **YES**  
- **N/A**  
- **NO**

## 8 Product Verification Tests / Periodic Tests (PVT)

8.1 Are required PVT conducted? (one or more boxes may be ticked)  
- **NO** PVT required, all questions of this section shall be marked with ‘N/A’  
- PVT conducted at the factory location  
- PVT conducted at a external laboratory owned by the manufacturer  
- PVT conducted at a external laboratory owned by the license holder  
- PVT conducted by independent external laboratory  
- PVT conducted by certification body’s laboratory  
- Others (please provide details):  
  - Details are given on INSPECTOR’S EVALUATION-Informative page.  
If conducted at a location other than the manufacturers premises, then specify where performed:  
- Details are given on INSPECTOR’S EVALUATION-Informative page.

**Note:** Product Verification Tests shall be conducted under the responsibility of the manufacturer and may be named also as Periodic Tests or Sample Tests depending on the certification scheme.

Describe which tests(required by the Certification Body/certification scheme) are conducted and at what sampling rate on TEST DATA SHEET – **Product Verification Tests**

**Note:** Details of any additional product verification tests should be entered by the Inspector on the INSPECTOR’S EVALUATION instead of the TEST DATA SHEET.

8.2 Are the tests conducted in accordance with procedures?  
- **YES**  
- **N/A**  
- **NO**

**Description of the procedure or ref. of documented procedure & revision or issue date:**  
- Details are given on INSPECTOR’S EVALUATION-Informative page.

8.3 Is appropriate equipment that is required for conducting tests available?  
- **YES**  
- **N/A**  
- **NO**

8.4 Are the tests described in TEST DATA SHEET – **Product Verification Tests** in compliance with the requirements of the Certification Schemes and/or the requesting Certification Body?  
- **YES**  
- **N/A**  
- **NO**
Reference number of the body carrying out the inspection:

8.5 Is there a procedure requiring actions to be taken if PVT are found to be unsatisfactory? 

<table>
<thead>
<tr>
<th>YES</th>
<th>N/A</th>
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</thead>
</table>

Description of the procedure or ref. of documented procedure & revision or issue date:

Details are given on INSPECTOR’S EVALUATION- Informative page.

8.6 Are the records of product verification tests maintained and satisfactory? 

<table>
<thead>
<tr>
<th>YES</th>
<th>N/A</th>
<th>NO</th>
</tr>
</thead>
</table>

8.7 Are records kept at least for the period between two inspection visits? 

<table>
<thead>
<tr>
<th>YES</th>
<th>N/A</th>
<th>NO</th>
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</thead>
</table>

9 Void

10 Corrective actions in response to inspector’s evaluation

If there were any unsatisfactory findings entered in the previous inspection report, have these been corrected?

<table>
<thead>
<tr>
<th>YES</th>
<th>N/A</th>
<th>NO</th>
</tr>
</thead>
</table>

Provide details of each unsatisfactory finding and how each has been resolved

11 Quality Management System

If the manufacturer has a Quality Management System certified or assessed by an accredited Body, provide details of QMS standard, scope, name of certification body and certificate expiry date.

or provide copy of the certificate.

- Quality Management System NOT certified
- Quality Management System certified by an accredited Body
- Quality Management System certified by a non accredited Body
- Copy of the certificate provided as appendix to this report

Details of QMS standard:

Does the scope covers the production of the certified product: ☐ YES ☐ NO

Name of certification body:

Certificate no.:

Certificate issued date:

Certificate expiry date:
## 12 Manufacturer’s self assessment of the manufacturing- and control process of certified products (Former: Audits of the Quality System)

| 12.1 Does the manufacturer regularly check that all procedures as required by the Certification Body(ies) and the harmonised inspection scheme (PD CIG 021) are followed? | YES | N/A | NO |

| 12.2 Are records regarding results and actions taken available? Note: The use of PD CIG 023 to document the results of the self assessment is acceptable | YES | N/A | NO |

| 12.3 Are the personnel carrying out above required checks appropriately trained and independent of the process being assessed? | YES | N/A | NO |

## 13 Void

## 14 Customer Complaints

The Manufacturer shall record any technical complaint regarding the certified product. The questions in this section shall be answered even if no customer complaints have been received. In this case the questions should be applied to the process.

| 14.1 Is there a procedure regarding how to handle customer complaints? | YES | N/A | NO |

| 14.2 Are the received complaints reviewed on a regular basis regarding whether they are related to single errors or system errors? Actual case checked | YES | N/A | NO |

| 14.3 Are corrective actions and decisions regarding customer complaints recorded? Actual case checked | YES | N/A | NO |

| 14.4 Is the originator of the complaint informed about the handling and the result of the complaint? Actual case checked | YES | N/A | NO |
14.5 Are the records of customer complaints maintained and satisfactory? | YES | N/A | NO
|   |   |   |

14.6 Are records kept at least for the period between two inspection visits? | YES | N/A | NO
|   |   |   |

### 15 Changes to Certified Products

15.1 Is reference about the certified version available?  
(One or more boxes may be ticked)  
- Set of drawings  
- Parts list  
- Product description  
- Reference sample  
- Photo-documentation  
- Other specification (Please provide details):  
  Details are given on INSPECTOR'S EVALUATION-Informative page.

15.2 Is this reference under control of the licence holder? | YES | N/A | NO
|   |   |   |

15.3 Is there a procedure ensuring that no changes to the construction of certified products will be implemented prior to acceptance by the License Holder? | YES | N/A | NO
|   |   |   |

*Description of the procedure or ref. of documented procedure & revision or issue date:*  
- Details are given on INSPECTOR'S EVALUATION-Informative page.

15.4 If the manufacturer is also the licence holder:  
Is there a procedure ensuring that constructional changes of the certified product will be made only after approval by the Certification Body? | YES | N/A | NO
|   |   |   |

*Description of the procedure or ref. of documented procedure & revision or issue date:*  
- Details are given on INSPECTOR'S EVALUATION-Informative page.

15.5 Are any changes made to the certified version since the last inspection? | YES | N/A | NO
|   |   |   |

- no changes  
- changes authorised by the license holder

### 16 Selection and Shipping of Re-Examination Sample(s)

*Regarding samples requested by the Certification Body(ies) please refer to the table IDENTIFICATION OF SELECTED SAMPLES and enter details as appropriate*
16.1 Please give reasons why no samples were selected during the inspection:
(one or more boxes may be ticked)
☐ None required by the certification body:
☐ No production, no stock:
☐ Build to clients’ order
☐ No access to warehouse
☐ Warehouse not at manufacturer’s location
☐ Manufacturer has been instructed to send re-examination samples:
☐ Others (Please provide details):
☐ Details are given on INSPECTOR’S EVALUATION-Informative page

16.2 If the selected sample(s) do not bear the Certification Mark then provide the reason for selection in the table IDENTIFICATION OF SELECTED SAMPLES
(one or more boxes may be ticked)
☐ Type reference is mentioned on the certification bodies certification list
☐ Mark is applied on the package, catalogue or by other means
☐ Special sample selection order
☐ Others (Please provide details):
☐ Details are given on INSPECTOR’S EVALUATION-Informative page.
17 Inspector’s Evaluation

17.1 List your findings on the INSPECTORS EVALUATION – Findings page(s) by referencing the applicable clauses in this report (including comments, recommendations, etc.) and explain them to the manufacturer.

If possible indicate also the corrective actions the manufacturer intends to take.

17.2 Give your recommendations by ticking the appropriate box

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<thead>
<tr>
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</thead>
<tbody>
<tr>
<td>1</td>
<td>No unsatisfactory findings.</td>
<td>Grant or continue certification.</td>
</tr>
<tr>
<td>2</td>
<td>Minor unsatisfactory finding(s).</td>
<td>Manufacturer’s corrective action(s) will be checked at next visit. Grant or continue certification.</td>
</tr>
<tr>
<td>3</td>
<td>Major unsatisfactory finding(s). Safety not directly affected.</td>
<td>Manufacturer shall confirm corrective action(s). Grant or continue certification. Special or early routine inspection recommended for checking corrective action(s).</td>
</tr>
<tr>
<td>4</td>
<td>Critical unsatisfactory finding(s), Safety directly affected.</td>
<td>Certification refused/suspended and repeated factory inspection recommended after the manufacturer has confirmed implementation of corrective action(s).</td>
</tr>
</tbody>
</table>

17.3 Attachments:

For page control, please write the reference number in the header of each attachment page.

- PD CIG 023 - Signature page
- ENEC Appendix to PD CIG 023
- Copy of Quality Management Certificate
- Others

Total no. of pages of this report including all attachment pages:

A copy of this report shall be provided to the undersigned contact person who should be aware of the contents and sign for its receipt.

- Printed copy provided
- Electronic copy provided

Inspection duration: hours.

The responsibility for ensuring that a product is manufactured in accordance with the standard to which it was originally approved rests with the licence holder.

Date: Date:

Inspector’s name (printed letters): Contact person’s name (printed letters):
Reference number of the body carrying out the inspection:

<table>
<thead>
<tr>
<th>Signature:</th>
<th>Signature:</th>
</tr>
</thead>
</table>

☐ For signature see attached signature page
Reference number of the body carrying out the inspection:

Inspector's Evaluation

<table>
<thead>
<tr>
<th>Findings</th>
<th>Inspector's points requiring corrective action from the manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Use separate Supplementary Page for different Certification Bodies if necessary</td>
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</tbody>
</table>
Reference number of the body carrying out the inspection:
Inspector's Evaluation

Informative
Use separate Supplementary Page for different Certification Bodies if necessary
Reference number of the body carrying out the inspection:
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<thead>
<tr>
<th>CB</th>
<th>Product, Sampling rate, Standards Clause or Test-parameters, Results</th>
</tr>
</thead>
<tbody>
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### TEST DATA SHEET - Routine Tests

<table>
<thead>
<tr>
<th>TESTS</th>
<th>% check</th>
<th>Test value applied</th>
<th>Time</th>
<th>Factory limits applied</th>
<th>Failure indicated by</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>a Earth continuity</td>
<td></td>
<td>V A</td>
<td>s</td>
<td>Ohm (max.)</td>
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<td></td>
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<tr>
<td>b Insulation resistance</td>
<td></td>
<td>V d.c.</td>
<td>s</td>
<td>MOhm (min.)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>c Leakage current</td>
<td></td>
<td>V</td>
<td></td>
<td>mA (max.)</td>
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</table>

#### Dielectric strength

- Basic insulation
  - V
  - s
  - mA (max.)

- Supplementary insulation
  - V
  - s
  - mA (max.)

- Reinforced insulation
  - V
  - s
  - mA (max.)

### Remarks

- W
- R
Reference number of the body carrying out the inspection:

- Indicate method used (hot/cold, at mains voltage, low voltage resistance check, etc.).
- Are all controls and components checked during the test?
- Test witnessed by the inspector, R = according to records
**IDENTIFICATION OF SELECTED SAMPLES**

<table>
<thead>
<tr>
<th>Selected for</th>
<th>Label No.</th>
<th>Quantity</th>
<th>Product/Type/Technical data</th>
<th>Licence No.</th>
<th>Production period</th>
<th>Code letters</th>
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Code letters: P = Sample from Production or S = Stock; F = Forwarded by the Manufacturer; T = Transported to the Certification Body by the Inspector; A = Shipped by the Inspection Agency
Reference number of the body carrying out the inspection: