



## Slovak Institute of Metrology

Notified body No. 1781

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**Number of Application:**

(to be filled by the Slovak Institute of Metrology):

### APPLICATION FOR CONFORMITY ASSESSMENT

In accordance with the Directive 2014/32/EU of the European Parliament and the Council of 26 February 2014 on the harmonisation of the laws of the Member States relating to the making available on the market of measuring instruments (recast)

#### 1. Applicant (manufacturer or authorised representative)

<b>Company name:</b>	
<b>Address:</b>	<b>ID:</b>
<b>Representative of the company (Name and Surname):</b>	<b>Phone:</b> <b>Email:</b> <b>I agree to receive information:</b> <input type="checkbox"/> yes <input type="checkbox"/> no
<b>Representative authorised for the meeting:</b>	<b>Phone:</b>
	<b>Fax:</b>
	<b>Email:</b>

#### 2. Manufacturer

<b>Company name:</b>	<b>Phone:</b>
	<b>Fax:</b>
<b>Address:</b>	<b>Email:</b>

#### 3. Procedure/Modules

- Module H - QUALITY ASSURANCE OF FINAL INSTRUMENT INSPECTION AND TESTING**
- Approval of the quality system
- Amendment/extension of the quality system

#### 4. Declaration of the manufacturer/applicant:

Signing this application I confirm that the information provided in this application is correct and that I have not applied to any other notified body (authorised body) for conformity assessment. I agree with the requirements for conformity assessment according to the Directive 2014/32/EU. I undertake to comply with all the requirements for conformity and to supply all the necessary information.

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<i>Number of Application:</i>	
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Date

Name and Signature

**5. For the following sites:**

<b>Full address</b>	

**6. Information for the instruments category envisaged:**

Measuring instruments categories		
Name/designation of the instrument	Type designation	Number certificate EU Type examination

**6. Description of quality assurance \*/**

*\*/ In case of lack of place, please add in the enclosure*

**Quality management system of manufacturer is certified in accordance with the EN ISO 9001:2015**

yes       no

**Testing laboratory of manufacturer is accredited in accordance with the EN ISO/IEC 17025:2005**

yes       no

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**7. Information concerning all outsourced processes used by the organisation that will affect conformity to requirements**

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**8. Information concerning the use of consultancy to the management system**

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**9. List of enclosed documentation \*/**

No.	Title	Marking
1	Quality manual	
2	Copies of already existing quality systems certificates	
3	Copies of certificate EU- type examination	
4	Quality objectives and the organisational structure, responsibilities and powers of the management with regard to product quality;	
5	Corresponding manufacturing, quality control and quality assurance techniques, processes and systematic actions that will be used;	
6	Examinations and tests that will be carried out before, during, and after manufacture, and the frequency with which they will be carried out;	
7	Quality records, such as inspection reports and test data, calibration data, qualification reports on the personnel concerned;	
8	Means of monitoring the achievement of the required product quality and the effective operation of the quality system.	

\*/ Further documents may be request within the scope of the recognition (assessment) procedure

Date:

Name and Signature (Applicant):