



Appointment to act as a Conformity Assessment Body

Legislation

***The Non-automatic Weighing Instruments Regulations 2016 (SI 2016/1152),
as amended.***

Conformity assessment body details

Body Name	EETSA (East of England Trading Standards Authority)
Appointment Type	Approved Body
Approved Body Number	0922

Appointment details

Appointment	Having considered the recommendation made to the Department for Business and Trade, the Secretary of State has appointed the above Conformity Assessment Body to assess the conformity of the product categories and for the modules identified below.
Northern Ireland	In addition, the body is appointed to act for the purposes of conformity assessment of products for Northern Ireland, the scope and extent of appointment mirroring the categories and conformity assessment activities indicates below. This will require the Company to perform conformity assessment in line with EU requirements for goods assessed for the Northern Ireland market.

Assessment details

Assessment Route	Local Authority Protocol
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Revision details

Version	Date	Details
1	27 February 2025	Initial issue

The current validity of this document should be confirmed through the CAB's current listing on the UK Market Conformity Assessment Database.

Product Categories and Modules of Appointment

<i>Product categories</i>	<i>Assessment Procedure as defined in Schedule 7 of the Regulations</i>
<p>Non-automatic weighing instruments for the purpose of the:</p> <ul style="list-style-type: none"> (a) determination of mass for commercial transactions; (b) determination of mass for the calculation of a toll, tariff, tax, bonus, penalty, remuneration, indemnity or similar type of payment; (c) determination of mass for the application of laws or regulations or for an expert opinion given in court proceedings; (d) determination of mass in the practice of medicine for weighing patients for the purposes of monitoring, diagnosis and medical treatment; (e) determination of mass for making up medicines on prescription in a pharmacy and determination of mass in analyses carried out in medical and pharmaceutical laboratories; (f) determination of price on the basis of mass for the purposes of direct sales to the public and the making-up of pre-packages; 	<p style="text-align: center;">-</p> <p>Module F (Paragraph 4) Conformity to type based on product verification</p>

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