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- "Addition" means that the relevant text of the Particular Standard is a new element (e.g. subclause, 3rd element, note, table, figure) additional to the text of the General Standard.
- "Amendment" means that a clause, subclause, table, figure, etc. of the General Standard is partially modified by deletion and/or addition as indicated by the text of this Particular Standard.

To avoid confusion with any amendments to the General Standard, a particular numbering has been employed for elements added by this International Standard: clauses, subclauses, tables and figures are numbered starting from 101; additional tables are lettered AA, BB, etc. and additional annexes are lettered AA, BB, etc.

In this International Standard, the following print types are used:

- requirements, compliance with which can be tested, and definitions: roman type;
- notes and examples: smaller roman type;
- description of type of document, source, and test specifications: *italic type*;
- terms defined in Clause 2 of the General Standard IEC 60601-1:1983 or in this Particular Standard: **bold**.

Throughout this Particular Standard, text for which a rationale is provided in Annex AA, is indicated by an asterisk (*).