



International Standard

ISO 4823

Dentistry — Elastomeric impression and bite registration materials

*Médecine bucco-dentaire — Produits pour empreintes et
matériaux pour enregistrement des rapports intermaxillaires à
base d'élastomères*

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Foreword

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This sixth edition cancels and replaces the fifth edition (ISO 4823:2021), which has been technically revised.

The main changes are as follows:

- packaging and instructions for use requirements have been updated;
- editorial corrections have been made.

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