
**Molecular in vitro diagnostic
examinations — Specifications for pre-
examinations processes for formalin-
fixed and paraffin-embedded (FFPE)
tissue —**

**Part 2:
Isolated proteins**

*Analyses de diagnostic moléculaire in vitro — Spécifications relatives
aux processus préanalytiques pour les tissus fixés au formol et inclus
en paraffine (FFPE) —*

Partie 2: Protéines extraites

ISO 20166-2:2018

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Foreword

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