



Institut luxembourgeois de la normalisation
de l'accréditation, de la sécurité et qualité
des produits et services

ILNAS-EN ISO 20166-3:2019

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

Molekularanalytische in-vitro-
diagnostische Verfahren - Spezifikationen
für präanalytische Prozesse für
formalinfixierte und paraffineingebettete

Analyses de diagnostic moléculaire in
vitro - Spécifications relatives aux
processus préanalytiques pour les tissus
fixés au formol et inclus en paraffine

01/2019



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**Molecular in vitro diagnostic examinations - Specifications
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paraffin-embedded (FFPE) tissue - Part 3: Isolated DNA
(ISO 20166-3:2018)**

Analyses de diagnostic moléculaire in vitro -
Spécifications relatives aux processus préanalytiques
pour les tissus fixés au formol et inclus en paraffine
(FFPE) - Partie 3: ADN extrait (ISO 20166-3:2018)

Molekularanalytische in-vitro-diagnostische Verfahren
- Spezifikationen für präanalytische Prozesse für
formalinfixierte und paraffineingebettete (FFPE)-
Gewebebeobachten - Teil 3: Isolierte DNS (ISO 20166-
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European foreword

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

**Part 3:
Isolated DNA**

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

Partie 3: ADN extrait



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