
**Clinical laboratory testing and *in vitro*
diagnostic test systems — *In vitro*
diagnostic medical devices for
professional use — Summary of
regulatory requirements for information
supplied by the manufacturer**

Essais cliniques de laboratoire et systèmes d'essai de diagnostic in vitro — Dispositifs de diagnostic médical in vitro à usage professionnel — Résumé des exigences de régulation pour les informations fournies par le fabricant



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