

K030360 RANDOX EVIDENCEDec 23, 2003
323 days to decisionK030360 · Product code: **DIO** · Chemistry
Source: <https://www.510kdatabase.net/k030360/>**SUBMISSION DETAILS**

Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Enzyme Immunoassay, Cocaine And Cocaine Metabolites (DIO)
Date received	Feb 3, 2003
Decision date	Dec 23, 2003
Days to decision	323 days
Third-party review	No
Summary / Statement	Statement

APPLICANT

Company	Radox Laboratories, Ltd.
Location	Antrim, N. Ireland, IR
Contact	PAULINE ARMSTRONG
Website	http://www.radox.com/
510(k) history	116 submissions · 115 cleared · 1992-2025

Radox Laboratories, Ltd. is a global diagnostic company specializing in chemistry devices and laboratory solutions. The company operates with a manufacturing facility in Antrim, Northern Ireland, and has over 40 years of expertise in diagnostic innovation. Radox has received FDA 510(k) clearances from total submissions since its first clearance in 1992. The company's regulatory portfolio is dominated by chemistry devices, including clinical chemistry analyzers, reagents, controls, and electrodes. The latest clearance on record dates to 2023. Recent FDA 510(k) cleared de...

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