

**K120284 DIGITEX SUTURE DELIVERY SYSTEM,
DIGITEXPOLYPROPYLENE SUTURE CARTRIDGE SIZE 0, WITH
NEEDLE AND WITHOUT NEEDLE, DIGITEX**May 23, 2012
113 days to decisionK120284 · Product code: **NEW** · General & Plastic Surgery
Source: <https://www.510kdatabase.net/k120284/>**SUBMISSION DETAILS**

Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Suture, Surgical, Absorbable, Polydioxanone (NEW)
Date received	Jan 31, 2012
Decision date	May 23, 2012
Days to decision	113 days
Third-party review	No
Summary / Statement	Summary

APPLICANT

Company	Coloplast A/S
Location	Mchenry, IL, US
Contact	JANELL COLLEY
Website	http://www.coloplast.com/
510(k) history	71 submissions · 68 cleared · 1983-2023

Coloplast A/S is a Danish multinational medical device manufacturer based in McHenry, US. The company develops and markets devices for ostomy, urology, continence, and wound care. Coloplast has received FDA 510(k) clearances from total submissions since its first clearance in 1983. The company's regulatory portfolio is dominated by Gastroenterology & Urology devices, including catheter systems, guidewires, and access sheaths. The latest clearance on record dates to 2023, reflecting the company's historical engagement with FDA regulatory pathways. Notable cleared devices i...