

K090704 MULTILINK IMPLANTJun 17, 2009
92 days to decisionK090704 · Product code: **EBF** · DentalSource: <https://www.510kdatabase.net/k090704/>**SUBMISSION DETAILS**

Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Material, Tooth Shade, Resin (EBF)
Date received	Mar 17, 2009
Decision date	Jun 17, 2009
Days to decision	92 days
Third-party review	No
Summary / Statement	Statement

APPLICANT

Company	Ivoclar Vivadent, Inc.
Location	Amherst, NY, US
Contact	DONNA M HARTNETT
Website	https://www.ivoclar.com
510(k) history	65 submissions · 65 cleared · 2001-2026

Ivoclar Vivadent, Inc. is a dental solutions provider based in Amherst, US. The company develops modern products for dental practitioners and laboratory technicians. The company has received FDA 510(k) clearances from total submissions since 2001. Dental devices represent 97% of its regulatory portfolio. The latest clearance was issued in 2026, confirming active market engagement. Recent cleared devices include restorative materials, adhesives, CAD-on restorations, bulk-fill composites, and curing lights. The product portfolio spans chairside and laboratory workflows for ...