

K101203 TINA-QUANT ALBUMIN GEN 2Sep 10, 2010
134 days to decisionK101203 · Product code: **DCF** · Chemistry
Source: <https://www.510kdatabase.net/k101203/>**SUBMISSION DETAILS**

| | |
|-----------------------|--|
| Decision | Substantially Equivalent (Cleared) |
| Submission type | Traditional |
| Device classification | Albumin, Antigen, Antiserum, Control (DCF) |
| Date received | Apr 29, 2010 |
| Decision date | Sep 10, 2010 |
| Days to decision | 134 days |
| Third-party review | No |
| Summary / Statement | Summary |

APPLICANT

| | |
|----------------|---|
| Company | Roche Diagnostics |
| Location | Indianapolis, IN, US |
| Contact | KATHIE J GOODWIN |
| Website | https://diagnostics.roche.com |
| 510(k) history | 182 submissions · 180 cleared · 2005-2026 |

Roche Diagnostics is a Swiss multinational healthcare company specializing in diagnostic devices and solutions. The company operates its U.S. diagnostics division from Indianapolis. Roche Diagnostics maintains a strong FDA 510(k) regulatory record with FDA 510(k) cleared devices from total submissions since 2005. The company's portfolio spans chemistry devices, immunology assays, microbiology testing, and hematology systems. The latest clearance in 2026 reflects continued innovation and regulatory engagement. Recent cleared devices include glucose monitoring systems, elec...
