

**K951359 RUBELLA IGG ASSAY FOR THE TECHNICON IMMUNO
1 SYSTEM IN-VITRO DIAGNOSTIC DEVICE**Apr 15, 1996
385 days to decisionK951359 · Product code: LFX · Microbiology
Source: <https://www.510kdatabase.net/k951359/>**SUBMISSION DETAILS**

| | |
|-----------------------|----------------------------------------------------|
| Decision | Substantially Equivalent (Cleared) |
| Submission type | Traditional |
| Device classification | Enzyme Linked Immunoabsorbent Assay, Rubella (LFX) |
| Date received | Mar 27, 1995 |
| Decision date | Apr 15, 1996 |
| Days to decision | 385 days |
| Third-party review | No |
| Summary / Statement | Summary |

APPLICANT

| | |
|----------------|-----------------------------------------|
| Company | Bayer Corp. |
| Location | Elkhart, IN, US |
| Contact | GABRIEL J MURACA, JR. |
| 510(k) history | 96 submissions · 96 cleared · 1989-2003 |

Bayer Corp. is the American subsidiary of Bayer AG, headquartered in Whippany, New Jersey. The company operates 40 fully consolidated subsidiaries across 19 states. Bayer Corp. received FDA 510(k) clearances from total submissions, with no denied submissions on record. The company's regulatory activity spans from 1989 to 2003, with a primary focus on chemistry devices and immunology assays. Notable cleared devices include the ASCENSIA BREEZE BLOOD GLUCOSE METER, CLINITEST PREGNANCY TEST, and the ADVIA CENTAUR immunoassay system. This represents a historical regulatory rec...