

K810772 MEDIUM 199 W/EARLE’S SALTS & GLUTAMINE

May 13, 1981
51 days to decision

K810772 · Product code: **KIT** · Chemistry
Source: <https://www.510kdatabase.net/k810772/>

SUBMISSION DETAILS

Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Media And Components, Synthetic Cell And Tissue Culture (KIT)
Date received	Mar 23, 1981
Decision date	May 13, 1981
Days to decision	51 days
Third-party review	No

APPLICANT

Company	Dutchland Laboratories, Inc.
Location	Mchenry, IL, US
510(k) history	75 submissions · 75 cleared · 1981-1982

510k Database - www.510kdatabase.net
 Device record: <https://www.510kdatabase.net/k810772/> Data sourced from FDA 510(k) public records (accessdata.fda.gov).
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