

K792631 STERILIZATION PROCESS/DBL. GRIP CORD CLPJan 28, 1980
40 days to decisionK792631 · Product code: **KNA** · Obstetrics & GynecologySource: <https://www.510kdatabase.net/k792631/>**SUBMISSION DETAILS**

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|-----------------------|---|
| Decision | Substantially Equivalent (Cleared) |
| Submission type | Traditional |
| Device classification | Instrument, Manual, Specialized Obstetric-gynecologic (KNA) |
| Date received | Dec 19, 1979 |
| Decision date | Jan 28, 1980 |
| Days to decision | 40 days |
| Third-party review | No |

APPLICANT

| | |
|----------------|---|
| Company | Procter & Gamble Mfg. Co. |
| Location | Mchenry, IL, US |
| Website | https://www.pg.com |
| 510(k) history | 72 submissions · 72 cleared · 1976-1988 |

Procter & Gamble Mfg. Co. is a consumer health and personal care manufacturer headquartered in McHenry, US. The company has a long history of developing products across multiple healthcare categories. The company received FDA 510(k) clearances from total submissions, with no denied submissions on record. Regulatory activity spans from 1976 to 1988, establishing a historical record primarily in Obstetrics & Gynecology and Gastroenterology & Urology device categories. The company is inactive in the FDA 510(k) clearance database, with no submissions recorded in the past seven...