

K922575 ALWAYS CONTOURS UNSCENTED MENSTRUAL PADSFeb 2, 1993
246 days to decisionK922575 · Product code: **HHD** · Obstetrics & Gynecology
Source: <https://www.510kdatabase.net/k922575/>**SUBMISSION DETAILS**

Decision	Substantially Equivalent (Cleared)
Submission type	Traditional
Device classification	Pad, Menstrual, Unscented (HHD)
Date received	Jun 1, 1992
Decision date	Feb 2, 1993
Days to decision	246 days
Third-party review	No
Summary / Statement	Summary

APPLICANT

Company	Procter & Gamble Co.
Location	Cincinnati, OH, US
Contact	JAMES T O'NEILL
Website	http://www.pg.com/
510(k) history	23 submissions · 23 cleared · 1988-2014

Procter & Gamble Co. is a consumer health and personal care company headquartered in Cincinnati, US. The company develops and markets a broad range of health and wellness products globally. Procter & Gamble has received FDA 510(k) clearances from total submissions, with no denied submissions on record. The company's regulatory focus has centered on Obstetrics & Gynecology devices, which represent the dominant category of its submissions. FDA 510(k) clearances span from 1988 to 2014, establishing a historical regulatory record in feminine care and oral health device categories.

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Device record: <https://www.510kdatabase.net/k922575/> Data sourced from FDA 510(k) public records (accessdata.fda.gov).

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