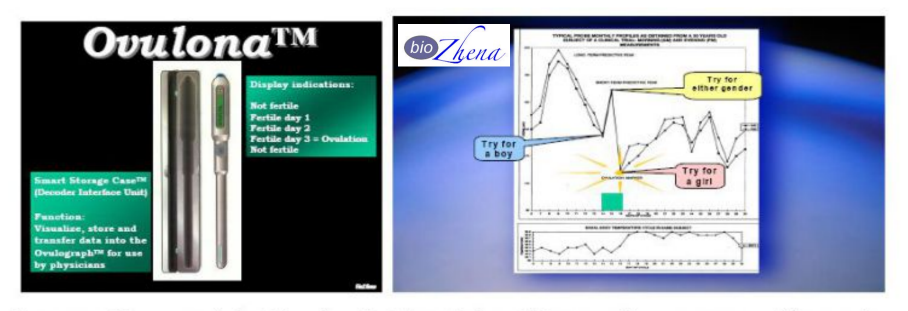


Summary of the first application



The personal home-use device determines the 3 days during which conception can occur , and it generates a new type of profile for use by the medical profession

Citing from FDA 510K Letter Number K973860: ... The Monitor serves as an independent information aid to the woman by helping her to define the fertile window... whereby she may choose the **proper timing** for vaginal intercourse. ... The Monitor may serve to provide the user **and her physician** with data to better time artificial insemination or other interventional techniques.