INTRODUCING CONTRACEPTIVE METHODS IN LOW-RESOURCE SETTINGS: NEW OPPORTUNITIES IN SENEGAL

PROJECT DESCRIPTION

In October 2013, USAID awarded the Delivering Contraceptive Vaginal Rings project to the Population Council to foster the introduction of two contraceptive methods. The objectives of this project are to:

1) improve availability and affordability of a three-month Progesterone Vaginal Ring (PVR) for postpartum breastfeeding women, and
2) facilitate regulatory approval and introduction of a one-year contraceptive vaginal ring containing Nestorone® and ethinyl estradiol (NES®/EE)—a long-acting method for non-breastfeeding women that is in the late stages of development.

Three-Month Progesterone Vaginal Ring

The three-month PVR, a user-initiated contraceptive, can be used safely by postpartum breastfeeding women to help space pregnancies. The ring contains natural progesterone and does not affect a woman’s ability to produce breast milk. Each ring can be used continuously for three months, and four rings can be used successively the first year after childbirth. Fertility generally returns shortly after discontinuation. The PVR is approved in eight Latin American countries, and efforts are underway to introduce it in more countries where contraceptive choices during breastfeeding are limited and where prolonged lactation is prevalent.

One-Year NES/EE Contraceptive Vaginal Ring

The one-year NES/EE CVR is a long-acting contraceptive in the late stages of development. It contains Nestorone® (NES), a new progestin that has been shown to be effective in suppressing ovulation, and ethinyl estradiol (EE), an approved, marketed hormonal compound. Once inserted by the woman, the NES/EE CVR is designed to be left in place for three weeks, removed for one week to allow for a monthly menstrual cycle, and then reinserted by the woman. The same ring can be used for up to 13 cycles (one year), with the goal of minimizing product cost and time spent visiting a provider or pharmacist. Phase 3 clinical trials have been completed, as have additional clinical and product specification studies required by the FDA. The Population Council is preparing a New Drug Application to submit to the FDA for regulatory review.

MILESTONES ACHIEVED

Continued progress was made in preparing the New Drug Application for the investigational NES®/EE CVR for submission to the US Food & Drug Administration (FDA).

Findings from Acceptability and Willingness to Pay studies were widely disseminated at national and global levels.

The PVR has been added to the World Health Organization’s Expression of Interest (EOI) following inclusion in WHO’s MEC and EML.

Project partner WomanCare Global (WCG) began compilation of the PVR dossier in preparation for registration with Nigeria’s regulatory body, the National Agency for Food and Drug Administration and Control (NAFDAC).

WCG also identified a Market Authorization Holder (MAH) for the PVR to ensure compliance with relevant regulatory requirements, post-registration.

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YEAR 4 ACTIVITIES
(OCTOBER 2016 – SEPTEMBER 2017)

2.1 Implementation research on product introduction
No activities proposed.

2.2 Register the PVR in Kenya, Malawi, Nigeria, Senegal, and Zambia
Project partner WomanCare Global (WCG) will:

- Complete registration with the Nigerian regulatory body and continue on the pathway toward country registration in Kenya and Senegal.
- Work closely with the selected MAH, which will be responsible for pharmacovigilance of the PVR post-registration.
- Conduct a regulatory audit of the PVR manufacturing plant, a requirement of regulators and national ministries of health.

2.3 Technical assistance for WHO Pre-Qualification of the PVR
- Project staff continue to provide technical assistance to PVR manufacturer Grünenthal in many thematic areas including a PK study, stability testing, and validation of analytical methods. Grünenthal will seek WHO-PQ to facilitate PVR procurement by a wider number of agencies including UNFPA, USAID, ministries of health, and social marketing organizations.
- Results of a clinical trial conducted by the Population Council in India in collaboration with the Indian Council of Medical Research comparing the safety and efficacy of the PVR versus the IUD will be finalized for inclusion in the WHO-PQ application.
- Project staff will continue analysis of used PVRs and compare with data collected from acceptability studies in each country. The secondary analysis will reveal the extent to which women adhere to PVR use; results will inform refinement of user instructions, including those related to ring disposal.

2.4 Design introductory strategies for contraceptive vaginal rings

- Foster partnerships with a wide range of stakeholders, implementing partners, USAID-funded projects, private sector social marketing organizations, professional associations, and reproductive supply networks to support subsequent introduction of vaginal rings.
- Advocate for inclusion of vaginal rings in national EMLs, and identify policy and program opportunities for integrating new contraceptives into existing implementation plans, providing a unified platform for introduction planning.
- Convene meetings to develop strategies to introduce vaginal rings, with an emphasis on identifying and training provider cadres from multiple distribution channels within the private and public sectors.

KEY PROJECT CONTACTS

The Population Council will lead the proposed project activities, along with a consortium that includes WomanCare Global, Grünenthal, and QPharma. For more information about the products or the Delivering Contraceptive Vaginal Rings project, contact Dr. Nafi Diop, Dr. John Townsend, or the USAID/Washington project management staff.

Population Council staff leading this effort in Senegal:
- Dr. Nafi Diop (ndiop@popcouncil.org)
- Mr. Babacar Mané

US-based Population Council staff directing the project:
- Dr. John Townsend, Principal Director (jtownsend@popcouncil.org)
- Dr. Saumya RamaRao
- Dr. Ruth Merkatz

Office of Population and Reproductive Health, USAID/Washington project management staff:
- Dr. Judy Manning, Agreement Officer’s Representative (jmanning@usaid.gov)
- Ms. Tabitha Sripipatana, Technical Advisor (tsripipatana@usaid.gov)
- Mr. Clifton Kenon, Technical Advisor (ckenon@usaid.gov)