STATISTICAL REVIEW AND EVALUATION

CLINICAL STUDIES

NDA/Serial Number: 22-501 / Resubmission

Drug Name: LO LOESTRIN FE (Norethindrone acetate and Ethinyl Estradiol tablets, and Ferrous Fumarate tablets)

Indication(s): Prevention of Pregnancy

Applicant: Warner Chilcott Company, Inc.

Date(s): Submission Date: 4/21/2010
PDUFA Due Date: 10/21/2010

Review Priority: Priority

Biometrics Division: Division of Biometrics III

Statistical Reviewer: Kate Dwyer, Ph.D.

Concurring Reviewers: Mahboob Sobhan, Ph.D.

Medical Division: Division of Reproductive and Urologic Drug Products

Clinical Team: Ronald Orleans, M.D., Medical Reviewer
Lisa Soule, M.D., Team Leader

Project Manager: Karl Stiller

Keywords: Labeling review
**BACKGROUND**

The Applicant has resubmitted this supplement in response to the Complete Response Letter issued by the Agency on January 26, 2010 regarding several manufacturing deficiencies. This is a labeling review for LO LOESTRIN FE, an oral contraceptive, and no additional clinical trial data was included in this submission. The proposed indication is for the prevention of pregnancy in women who elect to use oral contraceptive.

**CONCLUSION**

The efficacy (using Pearl Index) result in the label was evaluated and verified by this reviewer in the original statistical review of this NDA. Since no additional efficacy data was included in this resubmission, this reviewer agrees with the final version of the label.
This is a representation of an electronic record that was signed electronically and this page is the manifestation of the electronic signature.

/s/

KATE L DWYER
10/18/2010
re-check in as general review.

MAHBOOB SOBHAN
10/18/2010