

TEVA PHARMACEUTICAL INDUSTRIES LTD  
Form 6-K  
August 02, 2011

**FORM 6-K**

SECURITIES AND EXCHANGE COMMISSION

Washington, D.C. 20549

**Report of Foreign Private Issuer**

**Pursuant to Rule 13a-16 or 15d-16  
under the Securities Exchange Act of 1934**

For the month of August 2011

Commission File Number 0-16174

**Teva Pharmaceutical Industries Limited**

(Translation of registrant's name into English)

**5 Basel Street, P.O. Box 3190**

**Petach Tikva 49131 Israel**

(Address of principal executive offices)

Indicate by check mark whether the registrant files or will file annual reports under cover of Form 20-F or Form 40-F:

Form 20-F   X  

Form 40-F           

Indicate by check mark if the registrant is submitting the Form 6-K in paper as permitted by Regulation S-T Rule 101(b)(1):                   

Indicate by check mark if the registrant is submitting the Form 6-K in paper as permitted by Regulation S-T Rule 101(b)(7):

**Teva Receives European Marketing Authorization for  
ZOELY(TM) Oral Contraceptive**

*-- New OC Offers Innovative Combination which Includes  
a Natural Estrogen in a Convenient Dosing Regimen --*

*-- Teva Holds Marketing Rights in Four Key European Countries --*

*-- Launch Expected Before End 2011 --*

**Jerusalem, Israel, August 2, 2011** - Teva Pharmaceutical Industries Ltd (NASDAQ: TEVA) announced today that its new oral contraceptive, ZOELY<sup>(TM)</sup> (NOMAC/E2 -nomegestrol acetate 2.5 mg/17 -estradiol 1.5 mg), has received a Marketing Authorization in the European Union (EU) for the prevention of pregnancy. This clears the way to allow the product to be launched in Europe by the end of 2011.

Unlike currently available contraceptive pills, ZOELY<sup>(TM)</sup> is a combined oral contraceptive (COC) tablet containing a unique monophasic combination of 2 steroid hormones; 17-beta estradiol, an estrogen that is structurally identical to the major estrogen produced by the ovaries of healthy non-pregnant women, and nomegestrol acetate, a highly selective progesterone-derived progestin. This innovative combination will be made available in a convenient 24 day active and 4 day placebo dosing regimen.

"The EU Marketing Authorization for ZOELY<sup>(TM)</sup> marks an important step in creating a strong women`s healthcare franchise for Teva in Europe," said **Dr Gerard van Odijk, President and CEO of Teva Europe**. "The acquisition of Théramex in January this year gives us a platform to leverage our capability in women`s healthcare, and the launch of ZOELY<sup>(TM)</sup> is part of our developing business in this area."

Teva holds exclusive marketing rights for ZOELY<sup>(TM)</sup> in France, Italy, Belgium and Spain and expects the first launch by the end of the year.

The approval of ZOELY<sup>(TM)</sup> applies to the 27 EU Member States plus Norway, Iceland and Liechtenstein.

## **About Teva**

Teva Pharmaceutical Industries Ltd. (NASDAQ:TEVA) is a leading global pharmaceutical company, committed to increasing access to high-quality healthcare by developing, producing and marketing affordable generic drugs as well as innovative and specialty pharmaceuticals and active pharmaceutical ingredients. Headquartered in Israel, Teva is the world's largest generic drug maker, with a global product portfolio of more than 1,300 molecules and a direct presence in 60 countries. Teva's branded businesses focus on neurological, respiratory and women's health therapeutic areas as well as biologics. Teva currently employs approximately 42,000 people around the world and reached \$16.1 billion in net sales in 2010.

## **Teva's Safe Harbor Statement under the U. S. Private Securities Litigation Reform Act of 1995:**

This release contains forward-looking statements, which express the current beliefs and expectations of management. Such statements are based on management's current beliefs and expectations and involve a number of known and unknown risks and uncertainties that could cause our future results, performance or achievements to differ significantly from the results, performance or achievements expressed or implied by such forward-looking statements. Important factors that could cause or contribute to such differences include risks relating to: our ability to successfully develop and commercialize additional pharmaceutical products, the introduction of competing generic equivalents, the extent to which we may obtain U.S. market exclusivity for certain of our new generic products and regulatory changes that may prevent us from utilizing exclusivity periods, potential liability for sales of generic products prior to a final resolution of outstanding patent litigation, including that relating to the generic version of Protonix<sup>&reg</sup>, the extent to which any manufacturing or quality control problems damage our reputation for high quality production, the effects of competition on sales of our innovative products, especially Copaxone<sup>&reg</sup> (including potential generic and oral competition for Copaxone<sup>&reg</sup>), the impact of continuing consolidation of our distributors and customers, our ability to identify, consummate and successfully integrate acquisitions (including the acquisition of Cephalon), interruptions in our supply chain or problems with our information technology systems that adversely affect our complex manufacturing processes, intense competition in our specialty pharmaceutical businesses, any failures to comply with the complex Medicare and Medicaid reporting and payment obligations, our exposure to currency fluctuations and restrictions as well as credit risks, the effects of reforms in healthcare regulation, adverse effects of political or economical instability, major hostilities or acts of terrorism on our significant worldwide operations, increased government scrutiny in both the U.S. and Europe of our agreements with brand companies, dependence on the effectiveness of our patents and other protections for innovative products, our ability to achieve expected results through our innovative R&D efforts, the difficulty of predicting U.S. Food and Drug Administration, European Medicines Agency and other regulatory authority approvals, uncertainties surrounding the legislative and regulatory pathway for the registration and approval of biotechnology-based products, potentially significant impairments of intangible assets and goodwill, potential increases in tax liabilities resulting from challenges to our intercompany

arrangements, our potential exposure to product liability claims to the extent not covered by insurance, the termination or expiration of governmental programs or tax benefits, current economic conditions, any failure to retain key personnel or to attract additional executive and managerial talent, environmental risks and other factors that are discussed in this report and in our other filings with the U.S. Securities and Exchange Commission.

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Teva Pharmaceutical Industries Ltd. Web Site: [www.tevapharm.com](http://www.tevapharm.com)

**SIGNATURES**

Pursuant to the requirements of the Securities Exchange Act of 1934, the registrant has duly caused this report to be signed on its behalf by the undersigned, thereunto duly authorized.

TEVA PHARMACEUTICAL INDUSTRIES LIMITED

(Registrant)

By: /s/ Eyal Desheh

Name: Eyal Desheh  
Title: Chief Financial Officer

Date: August 2, 2011

