

ENDO PHARMACEUTICALS HOLDINGS INC

Form 10-Q

May 04, 2010

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UNITED STATES
SECURITIES AND EXCHANGE COMMISSION

Washington, DC 20549

FORM 10-Q

(Mark One)

QUARTERLY REPORT PURSUANT TO SECTION 13 OR 15(d) OF THE SECURITIES EXCHANGE ACT OF 1934

FOR THE QUARTERLY PERIOD ENDED MARCH 31, 2010.

OR

TRANSITION REPORT PURSUANT TO SECTION 13 OR 15(d) OF THE SECURITIES EXCHANGE ACT OF 1934

FOR THE TRANSITION PERIOD FROM TO .

Commission file number: 001-15989

ENDO PHARMACEUTICALS HOLDINGS INC.

(Exact Name of Registrant as Specified in Its Charter)

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Delaware
(State or other jurisdiction of
incorporation or organization)

13-4022871
(I.R.S. Employer
Identification Number)

100 Endo Boulevard Chadds Ford, Pennsylvania
(Address of Principal Executive Offices)

19317
(Zip Code)

(610) 558-9800

(Registrant's Telephone Number, Including Area Code)

Not applicable

(Former name, former address and former fiscal year, if changed since last report)

Indicate by check whether the registrant: (1) has filed all reports required to be filed by Sections 13 or 15(d) of the Securities Exchange Act of 1934 during the preceding 12 months (or for such shorter period that the registrant was required to file such reports), and (2) has been subject to such filing requirements for the past 90 days. Yes No

Indicate by check whether the registrant has submitted electronically and posted on its corporate Web site, if any, every Interactive Data File required to be submitted and posted pursuant to Rule 405 of Regulation S-T (§232.405 of this chapter) during the preceding 12 months (or for such shorter period that the registrant was required to submit and post such files). Yes No

Indicate by check mark whether the registrant is a large accelerated filer, an accelerated filer, or a smaller reporting company. See definitions of large accelerated filer, accelerated filer and smaller reporting company in Rule 12b-2 of the Exchange Act. (Check one):

Large accelerated filer

Accelerated filer

Non-accelerated filer (Do not check if a smaller reporting company)

Smaller reporting company

Indicate by check mark whether the registrant is a shell company (as defined in Rule 12b-2 of the Act). YES NO

Indicate the number of shares outstanding of each of the issuer's classes of common stock, as of the latest practical date.

Common Stock, \$0.01 par value

Shares outstanding as of April 23, 2010: 116,266,242

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FORWARD LOOKING STATEMENTS

Statements contained or incorporated by reference in this Quarterly Report on Form 10-Q contain information that includes or is based on forward-looking statements within the meaning of Section 27A of the Securities Act and Section 21E of the Exchange Act. These statements, including estimates of future revenues, future expenses, future net income and future earnings per share, contained in the section titled Management's Discussion and Analysis of Financial Condition and Results of Operations, in our Annual Report on Form 10-K for the year ended December 31, 2009, filed with the Securities and Exchange Commission on February 26, 2010, are subject to risks and uncertainties. Forward-looking statements include the information concerning our possible or assumed results of operations. Also, statements including words such as believes, expects, anticipates, intends, estimates, plan, will, may or similar expressions are forward-looking statements. We make these forward-looking statements on our current expectations and projections about the growth of our business, our financial performance and the development of our industry. Because these statements reflect our current views concerning future events, these forward-looking statements involve risks and uncertainties. Investors should note that many factors, as more fully described under the caption Risk Factors in Item 1A of our Annual Report on Form 10-K for the year ended December 31, 2009 and as otherwise enumerated herein or therein, could affect our future financial results and could cause our actual results to differ materially from those expressed in forward-looking statements contained in our Annual Report on Form 10-K. Important factors that could cause our actual results to differ materially from the expectations reflected in the forward-looking statements in our Annual Report on Form 10-K include those factors described herein under the caption Risk Factors and in documents incorporated by reference, including, among others:

our ability to successfully develop, commercialize and market new products;

timing and results of pre-clinical or clinical trials on new products;

our ability to obtain regulatory approval of any of our pipeline products;

competition for the business of our branded and generic products, and in connection with our acquisition of rights to intellectual property assets;

market acceptance of our future products;

government regulation of the pharmaceutical industry;

our dependence on a small number of products;

our dependence on outside manufacturers for the manufacture of most of our products;

our dependence on third parties to supply raw materials and to provide services for certain core aspects of our business;

new regulatory action or lawsuits relating to our use of narcotics in most of our core products;

our exposure to product liability claims and product recalls and the possibility that we may not be able to adequately insure ourselves;

our ability to protect our proprietary technology;

the successful efforts of manufacturers of branded pharmaceuticals to use litigation and legislative and regulatory efforts to limit the use of generics and certain other products;

our ability to successfully implement our acquisition and in-licensing strategy;

regulatory or other limits on the availability of controlled substances that constitute the active ingredients of some of our products and products in development;

the availability of third-party reimbursement for our products;

the outcome of any pending or future litigation or claims by third parties or the government, and the performance of indemnitors with respect to claims for which we have the right to be indemnified;

our dependence on sales to a limited number of large pharmacy chains and wholesale drug distributors for a large portion of our total revenues;

significant litigation expenses to defend or assert patent infringement claims;

any interruption or failure by our suppliers, distributors and collaboration partners to meet their obligations pursuant to various agreements with us;

a determination by a regulatory agency that we are engaging or have engaged in inappropriate sales or marketing activities, including promoting the off-label use of our products;

existing suppliers become unavailable or lose their regulatory status as an approved source, causing an inability to obtain required components, raw materials or products on a timely basis or at commercially reasonable prices;

the loss of branded product exclusivity periods and related intellectual property;

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our exposure to securities that are subject to market risk including auction-rate securities that are currently illiquid due to an inactive auction-rate market;

our ability to successfully execute our strategy;

disruption of our operations if our information systems fail or if we are unsuccessful in implementing necessary upgrades or new software; and

our ability to maintain or expand our business if we are unable to retain or attract key personnel and continue to attract additional professional staff.

We do not undertake any obligation to update our forward-looking statements after the date of this Report for any reason, even if new information becomes available or other events occur in the future. You are advised, however, to consult any further disclosures we make on related subjects in our 10-Q, 10-K, and 8-K reports to the Securities and Exchange Commission (SEC). Also note that we provide the preceding cautionary discussion of risks, uncertainties and possibly inaccurate assumptions relevant to our business. These are factors that, individually or in the aggregate, we think could cause our actual results to differ materially from expected and historical results. We note these factors for investors as permitted by the Private Securities Litigation Reform Act of 1995. You should understand that it is not possible to predict or identify all such factors. Consequently, you should not consider the preceding to be a complete discussion of all potential risks or uncertainties.

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Additional paid-in capital	822,614	817,467
Retained earnings	1,165,646	1,105,291
Accumulated other comprehensive loss	(1,413)	(1,881)
Treasury stock, 18,920,003 and 17,716,303 shares at March 31, 2010 and December 31, 2009, respectively	(453,824)	(424,816)
Total stockholders' equity	1,534,377	1,497,411
TOTAL LIABILITIES AND STOCKHOLDERS' EQUITY	\$ 2,539,651	\$ 2,488,803

See Notes to Condensed Consolidated Financial Statements.

Table of Contents**ENDO PHARMACEUTICALS HOLDINGS INC.****CONDENSED CONSOLIDATED STATEMENTS OF OPERATIONS (UNAUDITED)****(In thousands, except per share data)**

	Three Months Ended March 31,	
	2010	2009
REVENUES:		
Net sales	\$ 360,349	\$ 335,300
Royalty and other revenue	\$ 4,063	
TOTAL REVENUES	\$ 364,412	\$ 335,300
COSTS AND EXPENSES:		
Cost of revenues	94,073	83,009
Selling, general and administrative	133,335	120,006
Research and development	29,168	28,414
Acquisition-related items	1,529	26,405
OPERATING INCOME	106,307	77,466
INTEREST EXPENSE, NET	9,804	7,593
OTHER (INCOME) EXPENSE, NET	(219)	1,105
INCOME BEFORE INCOME TAX	96,722	68,768
INCOME TAX	36,367	29,731
NET INCOME	\$ 60,355	\$ 39,037
NET INCOME PER SHARE:		
Basic	\$ 0.51	\$ 0.33
Diluted	\$ 0.51	\$ 0.33
WEIGHTED AVERAGE SHARES:		
Basic	117,347	116,822
Diluted	118,031	117,209

See Notes to Condensed Consolidated Financial Statements.

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SUPPLEMENTAL INFORMATION:

Interest paid	\$ 2,843	\$ 6
Income taxes paid	\$ 13,572	\$ 1,792

SCHEDULE OF NON-CASH INVESTING AND FINANCING ACTIVITIES

Purchases of property and equipment financed by capital leases	\$ 162	\$ 40
Accrual for purchases of property and equipment	\$ 2,291	\$ 493

In connection with the purchase of all of the capital stock of Indevus Pharmaceuticals, Inc., liabilities were assumed as follows:

Fair value of assets acquired	\$	\$ 1,013,724
Cash paid for the capital stock		(367,221)
Contingent consideration		(174,350)

Liabilities assumed	\$	\$ 472,153
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See Notes to Condensed Consolidated Financial Statements.

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agreement. This pronouncement did not have a material impact on the Company's consolidated financial statements.

The Company adopted new authoritative guidance on the fair value option for financial assets and financial liabilities which became effective for fiscal years beginning after November 15, 2007. The Standard's objective is to reduce both complexity in accounting for financial instruments and the volatility in earnings caused by measuring related assets and liabilities differently.

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identified events or changes in circumstances that would have a significant adverse effect on the carrying value of our one \$20.0 million cost method investment.

As of March 31, 2010, the Company held certain assets and liabilities that are required to be measured at fair value on a recurring basis, including money market funds, available-for-sale securities and trading securities, auction-rate securities rights, and acquisition-related contingent consideration. Fair value guidance establishes a three-tier fair value hierarchy, which prioritizes the inputs used in measuring fair value. These tiers include:

Level 1 Quoted prices in active markets for identical assets or liabilities.

Level 2 Inputs other than Level 1 that are observable, either directly or indirectly, such as quoted prices for similar assets or liabilities; quoted prices in markets that are not active; or other inputs that are observable or can be corroborated by observable market data for substantially the full term of the assets or liabilities.

Level 3 Unobservable inputs that are supported by little or no market activity and that are significant to the fair value of the assets or liabilities.

market makers, thus reducing the potential usefulness of those observations. In addition, the current lack of liquidity prevents the Company from comparing our securities directly to securities with quoted market prices.

Overview of Auction-Rate Securities Rights

In October 2008, UBS AG (UBS) made an offer (the UBS Offer) to the Company and other clients of UBS Securities LLC and UBS Financial Services Inc. (collectively, the UBS Entities), pursuant to which the Company received auction-rate securities rights (the Rights) to sell to UBS all auction-rate securities held by the Company as of February 13, 2008 in a UBS account (the Eligible Auction-Rate Securities). The Rights permit the Company to require UBS to purchase the Eligible Auction-Rate Securities for a price equal to par value plus any accrued but unpaid dividends or interest beginning on June 30, 2010 and ending on July 2, 2012. As of March 31, 2010, we had Eligible Auction-Rate Securities with a par value of \$197.9 million, representing 91% of our total auction-rate securities portfolio at par. The remaining nine percent (9%), or \$18.8 million at par, of our auction-rate securities portfolio are not held in a UBS account and therefore are not subject to the UBS Offer.

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Auction-Rate Securities. However, in management's view, the auction-rate securities rights act as an economic hedge against further fair value changes in the Eligible Auction-Rate Securities. At March 31, 2010, the fair value of our auction-rate securities rights were \$13.7 million. The decrease in fair value from December 31, 2009 to March 31, 2010 of \$1.9 million was recognized as a charge to earnings and included in Other (income) expense, net in the Condensed Consolidated Statements of Operations. Future changes in fair value will also be recognized in earnings.

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issuers, our intent and ability to retain our investment in the issuers for a period of time sufficient to allow for any anticipated recovery in market value and based on the extent to which fair value is less than par. Accordingly, we recorded a \$0.01 million gain and a \$0.6 million gain in shareholders' equity in accumulated other comprehensive loss as of March 31, 2010 and December 31, 2009, respectively. Securities not subject to the UBS Offer are analyzed each reporting period for other-than-temporary impairment factors. Any future fluctuation in fair value related to these instruments that the Company judges to be temporary, including any recoveries of previous write-downs, would be recorded to other comprehensive income. If the Company determines that any future valuation adjustment was other-than-temporary, it would record a charge to earnings as appropriate. However, there can be no assurance that our current belief that the securities not subject to the UBS Offer will recover their value will not change.

	Fair Value Measurements Using Significant Unobservable Inputs (Level 3) Acquisition-related Contingent Consideration
Liabilities:	
Balance at January 1, 2010	\$ (58,470)
Amounts acquired or issued	
Transfers in and/or (out) of Level 3	
Changes in fair value recorded in earnings	(890)
Balance at March 31, 2010	\$ (59,360)

post-injection anaphylactic reaction and pulmonary oil microembolism. The letter also specified that our proposed Risk Evaluation and Mitigation Strategy with respect to the product is not sufficient. We believe that significant regulatory uncertainty currently exists with respect to the timing, label and regulatory path forward for Aveed™, and accordingly determined that a review for asset impairment was appropriate. Although the Company is continuing to evaluate the FDA's findings to better understand the agency's concerns, we were required to estimate the fair value of our Aveed™ indefinite-lived intangible asset as of the date we received the Complete Response letter. To estimate fair value we assessed the possible changes to the product's indication and targeted population of eligible recipients, the future probability of regulatory approval, relative timing of commercialization, and estimates of the amount and timing of future cash flows. In January 2010, the Company was notified that the U.S. patent office had issued a Notice of Allowance on a patent covering the Aveed™ formulation. Therefore, management considered the likely benefit of patent exclusivity when estimating these future cash flows. To calculate the fair value of the Aveed™ intangible asset, the Company used an income approach using a discounted cash flow model considering management's current evaluation of the above mentioned factors. The Company utilized probability-weighted cash flow models using a present value discount factor of 15% which we believe to be commensurate with the overall risk associated with this particular product. The cash-flow models included our best estimates of future FDA approval associated with each potential indication and population of eligible recipients. The Company presently believes that the level and timing of cash flows assumed, discount rate, and probabilities of success appropriately reflect market participant assumptions.

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The fair value of the Aveed intangible asset was determined to be \$35 million. Accordingly, the Company recorded a pre-tax non-cash impairment charge of \$65 million for the year ended December 31, 2009, representing the difference between the carrying value of the intangible asset and its estimated fair value. The impairment charge was recognized in earnings and included the Impairment of other intangible assets line item in the Consolidated Statements of Operations during the three months ended December 31, 2009. During the three months ended March 31, 2010, there have been no events, changes in circumstances or indicators of impairment that would have triggered an additional impairment review of the Aveed intangible asset. The Company expects to have an open dialogue in a meeting with the FDA in the second quarter of 2010, the results of which may impact those assumptions.

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Changes in any of our assumptions may result in a further reduction to the estimated fair value of the Aveed™ intangible asset resulting in additional and potentially full future impairment charges. Such additional impairment charges could materially impact our results of operations in future periods.

NOTE 4. INVENTORIES

Inventories are comprised of the following at March 31, 2010 and December 31, 2009, respectively (in thousands):

	March 31, 2010	December 31, 2009
Raw materials	\$ 10,790	\$ 8,510
Work-in-process	23,509	25,799
Finished goods	52,751	50,584
Total	\$ 87,050	\$ 84,893

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method. The relief from royalty method is based on a hypothetical royalty stream that would be received if the Company were to out-license the technology. The Hydron[®] Polymer Technology is currently used in the following products: Vantas[®], Supprelin[®] LA and octreotide. Thus, we derived the hypothetical royalty income from the projected revenues of those drugs. The fair value of the Hydron[®] Polymer Technology also includes an existing royalty payable by the Company to certain third party partners based on the net sales derived from drugs that use the Hydron[®] Polymer Technology. Discount rates applied to the estimated cash flows for all intangible assets acquired ranged from 13% to 20%, depending on the current stage of development, the overall risk associated with the particular project or product and other market factors. We believe the discount rates used are consistent with those that a market participant would use.

The \$121.5 million of goodwill was assigned to our pharmaceutical products segment, which was our only reportable segment as of December 31, 2009. The goodwill recognized is attributable primarily to the potential additional applications for the Hydron[®] Polymer Technology, expected corporate synergies, the assembled workforce of Indevus and other factors. None of the goodwill is expected to be deductible for income tax purposes.

The deferred tax assets of \$167.7 million are related primarily to federal net operating loss and credit carryforwards of Indevus and its subsidiaries. The deferred tax liabilities of \$210.6 million are related primarily to the difference between the book basis and tax basis of identifiable intangible assets.

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Novartis has the exclusive right, at its sole discretion, to effect a switch of the Licensed Product from a prescription product to an over-the-counter (OTC) product in the United States (an OTC Switch) by filing an amendment or supplement to the Licensed Product New Drug Application or taking any other action necessary or advisable in connection therewith to effect the OTC Switch, and thereafter to commercialize such OTC product. Notwithstanding the foregoing, Novartis shall not launch an OTC equivalent product prior to a time specified in the Voltaren® Gel Agreement, and Novartis shall not take any action that results in the loss of the prescription product status for the Licensed Product prior to such time. Novartis will notify Endo if it submits a filing to the FDA in respect of an OTC equivalent product. In the event that Novartis gains approval of an OTC equivalent product that results in the Licensed Product being declassified as a prescription product, then Novartis will make certain royalty payments to Endo on net sales

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and commercialize Sanctura XR[®] in that country. The term of the Madaus Agreement for Sanctura XR[®] extends until the expiration, on a country-by-country basis, of all royalty obligations owed to the Company from Madaus which ceases upon the last to expire applicable patent in the Madaus Territory. Either party may terminate the amended Madaus Agreements in the event of a material breach by the other party.

Supernus

In March 2003, Indevus entered into a Development and License Agreement (the Supernus Agreement) with Supernus Pharmaceuticals, Inc. (Supernus) pursuant to which Supernus agreed to develop Sanctura XR[®] and granted exclusive, worldwide rights under certain Supernus patents and know-how to Indevus. The Supernus agreement includes potential future development and commercialization milestone payments from the Company to Supernus, including royalties based on sales of Sanctura XR[®], and potential future development and commercialization milestone payments for up to an aggregate of \$2.4 million upon the launch of Sanctura XR[®] in certain geographic areas. In addition, the Supernus agreement includes potential future development and

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\$26.4 million of acquisition related costs, which were attributable to transaction fees, professional service fees, employee retention and separation arrangements and other costs related to the acquisition. This compares to \$1.5 million in 2010 primarily reflecting changes in the fair value of the acquisition-related contingent consideration.

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manufacturers which includes an anticipated impact from foreign exchange. Selling, general and administrative expenses, as a percentage of revenues, are expected to decline in 2010, relative to 2009, reflecting new approaches to customer segmentation and marketing as well as annualized effects of the prior year's cost reduction efforts. We will continue to provide promotional support behind our key on-market products, including those acquired as part of our acquisition of Indevus. R&D expenses are expected to increase as we invest in clinical development programs in support of our third party collaboration agreements as well as the further advancement of the development products being acquired from Indevus. Of course, there can be no assurance that the Company will achieve these results.

LIQUIDITY AND CAPITAL RESOURCES

Our principal source of liquidity is cash generated from operations. Our principal liquidity requirements are for working capital for operations, acquisitions, licenses, milestone payments, capital expenditures and debt service payments. The Company continues to maintain a sufficient level of working capital, which was approximately \$895.9 million at March 31, 2010, increasing from \$808.4 million at December 31, 2009. Cash, cash equivalents and current marketable securities were approximately \$835.0 million at March 31, 2010 compared to \$733.7 million at December 31, 2009.

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On November 10, 2008, the Company accepted the UBS Offer. As a result, the Company granted to the UBS Entities, the sole discretion and right to sell or otherwise dispose of, and/or enter orders in the auction process with respect to the Eligible Auction-Rate Securities on the Company's behalf until the Expiration Date, without prior notification, so long as the Company receives a payment of par value plus any accrued but unpaid dividends or interest upon any sale or disposition.

In addition, as part of the UBS Offer, Endo is eligible for no net cost loans, should we desire to borrow money prior to the commencement of the exercise period for the Rights. Under the terms of the UBS Offer, Endo may be eligible for no net cost loans for an amount up to 75% of the market value of the Eligible Auction-Rate Securities at the time of the loan. The loans would become fully payable as soon as UBS receives the proceeds from a purchase of the Eligible Auction-Rate Securities. Our Rights pursuant to the UBS Offer, including the no net cost loans are not secured by UBS. As a result, in the event UBS becomes insolvent, secured creditors of UBS may be able to attach their secured interests to our no net cost loans.

	Three Months Ended	
	March 31,	
	2010	2009
Cash Flow Data-Operating Activities:		
Net income	\$ 60,355	\$ 39,037
Depreciation and amortization	21,521	14,915
Stock-based compensation	3,791	1,937
Change in fair value of acquisition-related contingent consideration	890	

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make royalty payments based on a percentage of future sales of the products in the event regulatory approval for marketing is obtained. From a business perspective, we view these payments favorably as they signify that the products are moving successfully through the development phase toward commercialization. For a complete description of our contingent payments involving our license and collaboration agreements, see Note 6, and Note 10 of the Condensed Consolidated Financial Statements included in Part I, Item 1 of this Quarterly Report on Form 10-Q.

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Acquisitions. As part of our business strategy, we plan to consider and, as appropriate, make acquisitions of other businesses, products, product rights or technologies. Our cash reserves and other liquid assets may be inadequate to consummate such acquisitions and it may be necessary for us to issue stock or raise substantial additional funds in the future to complete future transactions. In addition, as a result of our acquisition efforts, we are likely to experience significant charges to earnings for merger and related expenses (whether or not our efforts are successful) that may include transaction costs, closure costs or costs of restructuring activities.

Indevus Acquisition. On February 23, 2009 (the Acquisition Date), the Company completed its initial tender offer (the Offer) for all outstanding shares of common stock of Indevus. Through purchases in subsequent offering periods, the exercise of a top-up option and a subsequent merger (the Merger), the Company completed its acquisition of Indevus on March 23, 2009, at which time Indevus became a wholly-owned subsidiary of the Company.

The Indevus Shares were purchased at a price of \$4.50 per Indevus Share, net to the seller in cash, plus contractual rights to receive up to an additional \$3.00 per Indevus Share in contingent cash consideration payments, pursuant to the terms of the Agreement and Plan of Merger, dated as of January 5, 2009. Accordingly, the Company paid approximately \$368 million in aggregate initial cash consideration for the Indevus Shares and entered into the Aveed™ Contingent Cash Consideration Agreement and the Octreotide Contingent Cash Consideration Agreement (each as defined in the Merger Agreement), providing for the payment of up to an additional \$3.00 per Indevus Share in contingent cash consideration payments, in accordance with the terms of the Offer.

The total cost to acquire all outstanding Indevus Shares pursuant to the Offer and the Merger could be up to an additional approximately \$267 million, if Endo is obligated to pay the maximum amounts under the Aveed™ Contingent Cash Consideration Agreement and the Octreotide Contingent Cash Consideration Agreement.

Indevus was a specialty pharmaceutical company engaged in the acquisition, development, and commercialization of products to treat conditions in urology, endocrinology and oncology. Following the completion of the Merger, Indevus was renamed Endo Pharmaceuticals Solutions Inc.

Approved products include the following:

Sanctura® (trospium chloride) was launched in August 2004. Sanctura® is indicated for the treatment of overactive bladder (OAB) with symptoms of urge urinary incontinence, urgency and urinary frequency. Sanctura® is currently promoted in the U.S. by Allergan Inc.

Sanctura XR® (trospium chloride extended release capsules) is a 60 mg, once-daily formulation of Sanctura®, the only approved quaternary amine compound clinically proven to effectively treat OAB symptoms in as early as one week, with a low incidence of side effects. Sanctura XR® is currently promoted in the U.S. by Allergan Inc. and by Madaus AG in Europe.

Supprelin® LA (histrelin acetate) was launched in June 2007. Supprelin® LA is a 12-month hydrogel implant for treating central precocious puberty (CPP) or the early onset of puberty in children. Supprelin® LA utilizes our patented Hydron® Polymer Technology, designed to provide the continuous 12-month administration of a controlled dose of histrelin, a GnRH agonist.

Vantas® (histrelin) was launched in the U.S. in November 2004. Vantas® is a soft and flexible 12-month hydrogel implant currently marketed in the U.S. that provides histrelin, a luteinizing hormone releasing hormone (LHRH) agonist, for the palliative treatment of advanced prostate cancer. The product utilizes our patented Hydron® Polymer Technology that allows for a controlled delivery of medicine over a 12-month period. In November 2005, Vantas® was approved in Denmark, and in March 2006, received approval for marketing in Canada from Health Canada. Regulatory approval was granted in May 2007 in Germany, Ireland, Italy, Spain and the United Kingdom. As of August 2007, Vantas® was approved in Thailand, Singapore, and Malaysia and approval is pending in Taiwan, Korea, Hong Kong and China. Additionally, Vantas® received approval in Argentina in January 2007 and is currently being marketed in that country.

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Delatestryl[®] (testosterone enanthate) is a marketed injectable testosterone preparation for the treatment of male hypogonadism. Delatestryl[®] provides testosterone enanthate, a derivative of the primary endogenous androgen testosterone, for intramuscular injection.

Hydron[®] Implant is a subcutaneous, retrievable, non-biodegradable, hydrogel reservoir drug delivery device. The Hydron[®] Implant is designed to provide sustained release of a broad spectrum of drugs continuously, at constant, predetermined rates. The Hydron[®] Implant is the only soft, flexible, reservoir-based drug delivery system available for parenteral administration. The hydrogel polymer compositions possess flexible, tissue-like characteristics providing excellent biocompatibility and patient comfort. This technology serves as the basis for two of our currently marketed products including Vantas[®] and Supprelin[®] LA.

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Valstar® (valrubicin) is a sterile solution of valrubicin for intravesical instillation and is the only product approved by the FDA for therapy of bacillus Calmette-Guerin (BCG)-refractory carcinoma *in situ* (CIS) of the bladder. Valstar®, originally approved by the FDA in 1998, was withdrawn from the market due to a manufacturing problem involving impurity issues in the original formulation and was placed on the FDA Drug Shortages List. In April 2007, the Company submitted a supplemental New Drug Application (sNDA) to the FDA seeking approval to reintroduce Valstar® and in February 2009 obtained FDA approval of its sNDA for Valstar®. In September 2009, we launched Valstar® for the treatment of patients with BCG-refractory CIS of the bladder. We continue to work closely with the manufacturer to build quantities of the product to support our newly launched product.

As of March 31, 2010, primary development products included the following from the Indevus acquisition:

Aveed™ (testosterone undecanoate) is expected to be the first long-acting injectable testosterone preparation available in the U.S. for the treatment of male hypogonadism in the growing market for testosterone replacement therapies. Aveed™ had historically been referred to as Nebido®. On May 6, 2009, we received notice from the FDA that Nebido® was unacceptable as a proprietary name for testosterone undecanoate. In August 2009, we received approval from FDA to use the name Aveed™. The Company acquired U.S. rights to Aveed™ from Schering AG, Germany, in July 2005. In June 2008, we received an approvable letter from the FDA indicating that the NDA may be approved if the Company is able to adequately respond to certain clinical deficiencies related to the product. In September 2008, agreement was reached with the FDA with regard to the additional data and risk management strategy. In March 2009, the FDA accepted for review the complete response submission to the new drug application for Aveed™ intramuscular injection. On December 2, 2009, we received a complete response letter from the FDA regarding Aveed™ in response to our March 2009 complete response submission. In the complete response letter, the FDA has requested information from Endo to address the agency's concerns regarding very rare but serious adverse events, including post-injection anaphylactic reaction and pulmonary oil microembolism. The letter also specified that the proposed Risk Evaluation and Mitigation Strategy (REMS) is not sufficient. The Company is continuing to evaluate how best to address the concerns of the FDA and intends to have future dialogue with the agency regarding a possible regulatory pathway. We are expected to meet with the FDA in the second quarter of 2010 to discuss the future path of our development efforts. The outcome of this meeting and future communications with the FDA could have a material impact on (1) management's assessment of the overall probability of approval, (2) the timing of such approval, (3) the targeted indication or patient population and (4) the likelihood of additional clinical trials. In April of 2010, we received an Issue Notification for the Aveed patent setting forth a projected issue date of May 18, 2010. If the patent issues on that date, its projected expiration date would be March 14, 2027.

Octreotide implant, currently in Phase III clinical trials for the treatment of acromegaly, utilizes our patented Hydron® Polymer Technology to deliver six months of octreotide, a long-acting octapeptide that mimics the natural hormone somatostatin to block production of growth hormone (GH). The octreotide implant is also currently in Phase II trials for the treatment of carcinoid syndrome.

The table below provides estimates as to the timing associated with completion of development for the remaining primary development products.

Product	Indication	Development Phase	Anticipated Year of Completion
Aveed™	Hypogonadism (Testosterone Deficiency)	NDA filed	2011 - 2013
Octreotide implant	Acromegaly	Phase III	2012
Octreotide implant	Carcinoid Syndrome	Phase II	2013

The anticipated year of completion shown in the above table represents our current best estimate as to the year in which the Company anticipates product approval from the FDA. This estimate assumes successful and timely completion of all clinical trials in preparation of an NDA filing. However, these anticipated completion dates are subject to significant change, particularly for those products not yet in Phase III clinical development due to uncertainty of the number, size, and duration of the trials which may be required to complete development. Once an NDA is filed with the FDA, there can be no assurance that the FDA will approve the NDA to permit the Company to market and sell the relevant product.

Management believes the Company's acquisition of Indevus is particularly significant because it reflects our commitment to expand our business beyond pain management into complementary medical areas where we believe we can be innovative and competitive. The combined company markets products through four field sales forces and has the capability to develop innovative new therapies using a novel drug delivery

technology.

The operating results of Indevus from February 23, 2009 are included in the accompanying condensed consolidated statements of operations. The consolidated balance sheet as of December 31, 2009 reflects the acquisition of Indevus, effective February 23, 2009, the date the Company obtained control of Indevus.

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The acquisition date fair value of the total consideration transferred was \$540.9 million, which consisted of the following (in thousands):

	Fair Value of Consideration Transferred
Cash	\$ 368,034
Contingent consideration	172,860
Total	\$ 540,894

The following table summarizes the fair values of the assets acquired and liabilities assumed at the Acquisition Date (in thousands):

	February 23, 2009
Cash and cash equivalents	\$ 117,675
Accounts receivable	14,591
Inventories	17,157
Prepaid and other current assets	8,322
Property, plant and equipment	8,856
Other intangible assets	532,900
Deferred tax assets	167,749
Other non-current assets	1,331
Total identifiable assets	\$ 868,581
Accounts payable	\$ (5,116)
Accrued expenses	(26,725)
Convertible notes	(72,512)
Non-recourse notes	(115,235)
Deferred tax liabilities	(210,647)
Other non-current liabilities	(18,907)
Total liabilities assumed	(449,142)
Net identifiable assets acquired	\$ 419,439
Goodwill	\$ 121,455
Net assets acquired	\$ 540,894

The above estimated fair values of assets acquired and liabilities assumed are based on the information that was available as of the Acquisition Date to estimate the fair value of assets acquired and liabilities assumed.

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The valuation of the intangible assets acquired and related amortization periods are as follows:

	Valuation (in millions)	Amortization Period (in years)
In Process Research & Development:		
Valstar ^{®(1)}	\$ 88.0	n/a
Aveed [™]	100.0	n/a
Octreotide	31.0	n/a
Pagoclone	21.0	n/a
Pro2000(2)	4.0	n/a
Other	11.9	n/a
Total	\$ 255.9	n/a
License Rights:		
Hydron [®] Polymer	\$ 22.0	10
Vantas [®]	36.0	10
Sanctura [®] Franchise	94.0	12
Supprelin [®] LA	124.0	10
Other	1.0	4
Total	\$ 277.0	11
Total other intangible assets	\$ 532.9	

- (1) The FDA approved the sNDA for Valstar[®] subsequent to the Acquisition Date. Therefore, Valstar[®] was initially classified as in-process research and development and subsequently transferred to License Rights upon obtaining FDA approval and is being amortized over a 15 year useful life.
- (2) In December 2009, the Company's Phase III clinical trials for Pro2000 provided conclusive results that the drug was not effective. The Company concluded there was no further value or alternative future uses associated with this indefinite-lived asset. Accordingly, we recorded a \$4.0 million impairment charge to write-off the Pro2000 intangible asset in its entirety.

The fair value of the in-process research and development assets and License Rights assets, with the exception of the Hydron[®] Polymer Technology, were estimated using an income approach. Under this method, an intangible asset's fair value is equal to the present value of the incremental after-tax cash flows (excess earnings) attributable solely to the intangible asset over its remaining useful life. To calculate fair value, the Company used probability-weighted cash flows discounted at rates considered appropriate given the inherent risks associated with each type of asset. The Company believes that the level and timing of cash flows appropriately reflect market participant assumptions. Cash flows were generally assumed to extend either through or beyond the patent life of each product, depending on the circumstances particular to each product. The fair value of the Hydron[®] Polymer Technology was estimated using an income approach, specifically known as the relief from royalty method. The relief from royalty method is based on a hypothetical royalty stream that would be received if the Company were to out-license the technology. The Hydron[®] Polymer Technology is currently used in the following products: Vantas[®], Supprelin[®] LA and octreotide. Thus, we derived the hypothetical royalty income from the projected revenues of those drugs. The fair value of the Hydron[®] Polymer Technology also includes an existing royalty payable by the Company to the certain third party partners based on the net sales derived from drugs that use the Hydron[®] Polymer Technology. Discount rates applied to the estimated cash flows for all intangible assets acquired ranged from 13% to 20%, depending on the current stage of development, the overall risk associated with the particular project or product and other market factors. We believe the discount rates used are consistent with those that a market participant would use.

The \$121.5 million of goodwill was assigned to our pharmaceutical products segment, which is our only reportable segment as of December 31, 2009. The goodwill recognized is attributable primarily to the potential additional applications for the Hydron[®] Polymer Technology, expected corporate synergies, the assembled workforce of Indevus and other factors. None of the goodwill is expected to be deductible for income tax purposes.

Acquisition-Related Contingent Consideration

As of March 31, 2010 and December 31, 2009, the fair value of the contingent consideration is \$59.4 million and \$58.5 million, respectively.

In the event that the Company receives an approval letter from the FDA with respect to the Aveed™ NDA on or before the third anniversary of the time at which we purchased the Indevus Shares in the Offer, then the Company will, subject to the terms described below, (i) pay an additional \$2.00 per Indevus Share to the former stockholders of Indevus, if such approval letter grants the right to market and sell Aveed™ immediately and provides labeling for Aveed™ that does not contain a boxed warning (Aveed™ With Label) or alternatively, (ii) pay an additional \$1.00 per Indevus Share, if such approval letter grants the right to market and sell

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Aveed™ immediately and provides labeling for Aveed™ that contains a boxed warning (Aveed™ Without Label). In the event that either an Aveed™ With Label approval or an Aveed™ Without Label approval has not been obtained prior to the third anniversary of the closing of the Offer, then the Company will not pay, and the former Indevus stockholders will not receive, any payments under the Aveed™ Contingent Cash Consideration Agreement.

Further, in the event that the Aveed™ Without Label approval is received and subsequently, Endo and its subsidiaries publicly report audited financial statements which reflect cumulative net sales of Aveed™ of at least \$125.0 million for four consecutive calendar quarters on or prior to the fifth anniversary of the date of the first commercial sale of Aveed™ (Aveed™ Net Sales Event), then the Company will, subject to the terms described below, pay an additional \$1.00 per Indevus Share to the former stockholders of Indevus. In the event that the Aveed™ Net Sales Event does not occur prior to the fifth anniversary of the date of the first commercial sale of Aveed™ then the Company will not pay, and former Indevus stockholders will not receive, any additional amounts under the Aveed™ Contingent Cash Consideration Agreement.

The range of the undiscounted amounts the Company could pay under the Aveed™ Contingent Cash Consideration Agreement is between \$0 and approximately \$175 million. The fair value of the contractual obligation to pay the Aveed™ contingent consideration recognized on the Acquisition Date was \$133.1 million. We determined the fair value of the obligation to pay the Aveed™ contingent consideration based on a probability-weighted income approach. This fair value measurement is based on significant inputs not observable in the market and thus represents a Level 3 measurement within the fair value hierarchy. Under the Aveed™ Contingent Cash Consideration Agreement, there are three scenarios that could potentially lead to amounts being paid to the former stockholders of Indevus. These scenarios are (1) obtaining an Aveed™ With Label approval, (2) obtaining an Aveed™ Without Label approval and (3) achieving the \$125.0 million sales milestone on or prior to the fifth anniversary of the date of the first commercial sale of Aveed™ should the Aveed™ Without Label approval be obtained. The fourth scenario is Aveed™ not receiving approval within three years of the closing of the Offer, which would result in no payment to the former stockholders of Indevus. Each scenario was assigned a probability based on the current regulatory status of Aveed™. The resultant probability-weighted cash flows were then discounted using a discount rate of U.S. Prime plus 300 basis points, which the Company believes is appropriate and is representative of a market participant assumption.

Similarly, in the event that an approval letter from the FDA is received with respect to an octreotide NDA (such approval letter, the Octreotide Approval) on or before the fourth anniversary of the closing of the Offer, then the Company will, subject to the terms described below, pay an additional \$1.00 per Indevus Share to the former stockholders of Indevus (such payment, the Octreotide Contingent Cash Consideration Payment). In the event that an Octreotide Approval has not been obtained prior to the fourth anniversary of the closing of the Offer, then the Company will not pay, and the former Indevus stockholders shall not receive, the Octreotide Contingent Cash Consideration Payment.

The range of the undiscounted amounts the Company could pay under the Octreotide Contingent Cash Consideration Agreement is between \$0 and approximately \$91 million. The fair value of the octreotide contractual obligation to pay the contingent consideration recognized on the Acquisition Date was \$39.8 million. We determined the fair value of the contractual obligation to pay the Octreotide Contingent Consideration Payment based on a probability-weighted income approach. This fair value measurement is based on significant inputs not observable in the market and thus represents a Level 3 measurement within the fair value hierarchy. Under the Octreotide Contingent Cash Consideration Agreement, the two scenarios that require consideration are (1) Octreotide Approval on or before the fourth anniversary of the closing of the Offer or (2) no Octreotide Approval on or before the fourth anniversary of the closing of the Offer. Each scenario was assigned a probability based on the current development stage of octreotide. The resultant probability-weighted cash flows were then discounted using a discount rate of U.S. Prime plus 300 basis points, which the Company believes is appropriate and is representative of a market participant assumption.

In addition to the potential contingent payments under the Aveed™ Contingent Cash Consideration Agreement and the Octreotide Contingent Cash Consideration Agreement, the Company has assumed a pre-existing contingent consideration obligation relating to Indevus's acquisition of Valera Pharmaceuticals, Inc. (the Valera Contingent Consideration), which was consummated on April 18, 2007. The Valera Contingent Consideration entitles former Valera shareholders to receive additional Indevus Shares based on an agreed upon conversion factor if FDA approval of the octreotide implant for the treatment for acromegaly is achieved on or before April 18, 2012. Upon Endo's acquisition of Indevus, each Valera shareholder's right to receive additional Indevus Shares was converted into the right to receive \$4.50 per Indevus Share that such former Valera shareholder would have received plus contractual rights to receive up to an additional \$3.00 per Indevus Share that such former Valera shareholder would have received in contingent cash consideration payments under the Aveed™ Contingent Cash Consideration Agreement and the Octreotide Contingent Cash Consideration Agreement. These amounts would only be payable to former Valera shareholders if there were Octreotide Approval. The range of the undiscounted amounts the Company could pay with respect to the Valera Contingent Consideration is between \$0 and approximately \$33 million.

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The Company is accounting for the Valera Contingent Consideration in the same manner as if it had entered into that arrangement with respect to its acquisition of Indevus. Accordingly, the fair value of the Valera Contingent Consideration recognized on the Acquisition Date was \$13.7 million. Fair value was estimated based on a probability-weighted discounted cash flow model, or income approach. This fair value measurement is based on significant inputs not observable in the market and thus represents a Level

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3 measurement within the fair value hierarchy. The fair value of the Valera Contingent Consideration is estimated using the same assumptions used for the Aved™ Contingent Cash Consideration Agreement and Octreotide Contingent Cash Consideration Agreement, except that the probabilities associated with the Valera Contingent Consideration take into account the probability of obtaining the Octreotide Approval on or before the fourth anniversary of the closing of the Offer. This is due to the fact that the Valera Contingent Consideration will not be paid unless Octreotide for the treatment of acromegaly is approved prior to April 18, 2012.

As of March 31, 2010, the fair value of the acquisition-related contingent consideration increased by approximately 0.9 million from December 31, 2009 primarily reflecting changes of our present value assumptions associated with our valuation model. There have been no changes to management's December 31, 2009 assessment of the probabilities or anticipated timelines that we will be obligated to make contingent consideration payments under the Aved™ Contingent Cash Consideration Agreement within the specified contractual timeframe, as well as the anticipated timeline for the NDA filing and FDA approval of octreotide. The increase in the liability was recorded as a loss and is included in the Acquisition-related items line item in the accompanying Condensed Consolidated Statements of Operations. Changes in any of our assumptions may result in a further volatility to the estimated fair value of the acquisition-related contingent consideration. Such additional changes to fair value could materially impact our results of operations in future periods. The Company expects to have an open dialogue in a meeting with the FDA in the second quarter of 2010, the results of which could materially impact the fair value of the Aved contingent consideration liability.

As of March 31, 2010, there were no changes to the range of the undiscounted amounts the Company may be required to pay under the Aved™ Contingent Cash Consideration Agreement and the Octreotide Contingent Consideration Agreement or related to the Valera Contingent Consideration.

Convertible Notes due 2009. As discussed in Note 12 to the Condensed Consolidated Financial Statements in Part I, Item 1 of this Report, as a result of our acquisition of Indevus Pharmaceuticals, Inc., the Company assumed Indevus's 6.25% Convertible Senior Notes due July 2009 (the Notes). Pursuant to the Indenture governing the Notes, within 30 days of the effective date of the Merger, holders of the Notes had the right to tender their Notes for the principal amount of the Notes plus any accrued and unpaid interest. During this 30-day period, approximately \$3.6 million in aggregate principal amount of Notes were tendered and the Company paid this amount in April 2009.

The Notes matured on July 15, 2009. Accordingly, in July 2009, the Company paid \$68.3 million in outstanding principal to satisfy the Notes in their entirety.

Convertible Senior Subordinated Notes due 2015. As discussed in Note 12 to the Condensed Consolidated Financial Statements in Part I, Item 1 of this Report, in April 2008, we issued \$379.5 million in aggregate principal amount of 1.75% Convertible Senior Subordinated Notes due April 15, 2015 (the Convertible Notes) in a private offering for resale to qualified institutional buyers pursuant to Rule 144A under the Securities Act of 1933, as amended.

We received proceeds of approximately \$370.7 million from the issuance, net of the initial purchaser's discount and certain other costs of the offering. Interest is payable semi-annually in arrears on each April 15 and October 15 with the first interest payment being made on October 15, 2008. The Convertible Notes will mature on April 15, 2015, unless earlier converted or repurchased by us.

Holders of the Convertible Notes may convert their notes based on a conversion rate of 34.2466 shares of our common stock per \$1,000 principal amount of notes (the equivalent of \$29.20 per share), subject to adjustment upon certain events, only under the following circumstances as described in the Indenture for the Convertible Notes (the Indenture): (1) during specified periods, if the price of our common stock reaches specified thresholds; (2) if the trading price of the Convertible Notes is below a specified threshold; (3) at any time after October 15, 2014; or (4) upon the occurrence of certain corporate transactions. We will be permitted to deliver cash, shares of Endo common stock or a combination of cash and shares, at our election, to satisfy any future conversions of the notes. It is our current intention to settle the principal amount of any conversion consideration in cash.

Non-recourse Notes. As discussed in Note 12 to the Condensed Consolidated Financial Statements in Part I, Item 1 of this Report, on August 26, 2008, Indevus closed a private placement to institutional investors of \$105.0 million in aggregate principal amount of 16% non-convertible, non-recourse, secured promissory notes due 2024 (Non-recourse Notes). The Non-recourse Notes were issued by Ledgemont Royalty Sub LLC (Royalty Sub), which was a wholly-owned subsidiary of Indevus at the time of the note issuance and subsequently became a wholly-owned subsidiary of the Company upon our acquisition of Indevus. As of the Acquisition Date, the Company recorded these notes at their fair value of approximately \$115.2 million. The Company was amortizing these notes to their face value of \$105.0 million at maturity in 2024.

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In connection with the issuance of the Non-recourse Notes, Indevus and Royalty Sub entered into a Purchase and Sale Agreement pursuant to which Indevus sold to Royalty Sub its rights to receive royalty payments from Allergan arising under the Allergan Agreement (as described in Note 6 of the Condensed Consolidated Financial Statements in Part I, Item 1 of this Report) for sales in the U.S. of Sanctura® and Sanctura XR®. To secure repayment of the Non-recourse Notes, Royalty Sub granted a continuing security interest to the trustee for the benefit of the noteholders in, among other things, the royalty payments made by Allergan under

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the Allergan Agreement discussed above, all of its rights under the Purchase and Sale Agreement and any accounts established in accordance with the Indenture (and all amounts from time to time credited to such accounts). The Non-recourse Notes have not been guaranteed by Indevus or the Company. Principal on the Non-recourse Notes is required to be paid in full by the final legal maturity date of November 5, 2024, unless repaid or redeemed earlier. In the event the Non-recourse Notes are repaid or redeemed prior to November 5, 2024, the noteholders will be entitled to a redemption premium (as described below). The interest rate applicable to the Non-recourse Notes is 16% per year and is payable quarterly in arrears and commenced on November 5, 2008.

Principal and interest on the Non-recourse Notes will be paid from the royalties from Allergan. Payments may also be made from the interest reserve account (described below) and certain other accounts established in accordance with the Indenture. In connection with the issuance of the Non-recourse Notes, a \$10.0 million interest reserve account was established to fund potential interest payment shortfalls. As of March 31, 2010, there was no remaining restricted cash on the Company's consolidated balance sheet. Royalty Sub will receive directly all royalties payable to the Company until the Non-recourse Notes have been repaid in full.

In August 2009, the Company commenced a cash tender offer for any and all outstanding Non-recourse notes. The purpose of the tender offer was to acquire any and all Notes to reduce our consolidated interest expense. The tender offer included an early tender deadline, whereby holders of the Non-recourse notes could early tender and receive the total early consideration of \$1,000 per \$1,000 principal amount of the Non-recourse notes. Holders who tendered their Non-recourse notes after such time and at or prior to the expiration of the tender offer period were eligible to receive the tender offer consideration of \$950 per \$1,000 principal amount of Non-recourse notes, which was the total early consideration less the early tender payment. The tender offer expired on September 24, 2009, at 5:00 p.m., New York City time (the Expiration Time). As of the Expiration Time, \$48 million Non-recourse notes had been validly tendered and not withdrawn. The Company accepted for payment and purchased Non-recourse notes at a purchase price of \$1,000 per \$1,000 principal amount, for a total amount of approximately \$48 million (excluding accrued and unpaid interest up to, but not including, the payment date for the Notes, fees and other expenses in connection with the tender offer). The aggregate principal amount of Non-recourse notes purchased represents approximately 46% of the \$105 million aggregate principal amount of Non-recourse notes that were outstanding prior to the Expiration Time. Accordingly, the Company recorded a \$4 million gain on the extinguishment of debt, net of transaction costs. The gain was calculated as the difference between the aggregate amount paid to purchase the Non-recourse notes and their carrying amount.

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If the royalty payments from Allergan and amounts in the interest reserve account are insufficient to pay all of the interest and principal, if any, due on a payment date, the shortfall will accrue interest at the interest rate applicable to the Non-recourse Notes (16%) compounded quarterly. If any interest payment shortfall is not paid in full by the succeeding payment date, an Event of Default under the Indenture will occur, unless the Company contributes cash to a capital account of Royalty Sub in an amount sufficient to satisfy any such shortfall. Pursuant to the Indenture, the Company has the right, but not the obligation, to contribute cash in an amount equal to the shortfall to the capital account for distribution by the trustee to the noteholders. The Company has the right to satisfy such an interest payment shortfall no more than six times over the life of the Non-recourse Notes and no more than three consecutive times. In the event that the Company is no longer permitted to fund the capital account to satisfy an interest payment shortfall, and the Company does not redeem the Non-recourse Notes (as described below), an Event of Default will occur and the noteholders may accelerate the obligations of Royalty Sub under the Non-recourse Notes and exercise their remedies thereunder, including assuming all rights to future royalty payments from Allergan. Based on current expectations, it is reasonably possible that we may exceed the maximum number of times we can fund the capital account to satisfy an interest payment shortfall as early as November 2010.

The Non-recourse Notes will be subject to redemption at the option of Royalty Sub. If the applicable redemption of the Non-recourse Notes occurs on or prior to August 5, 2010, the redemption price will be equal to the greater of (x) the outstanding principal balance of the Non-recourse Notes being redeemed or (y) the present value, discounted at the rate on U.S. Treasury obligations with a comparable maturity to the remaining weighted average life of the Non-recourse Notes plus 1.00%, of the principal payment amounts and interest at the rate applicable to the Non-recourse Notes on the outstanding principal balance of the Non-recourse Notes. If the applicable redemption of the Non-recourse Notes occurs after August 5, 2010, the redemption price will be equal to the percentage of the outstanding principal balance of the Non-recourse Notes being redeemed specified below for the period in which the redemption occurs:

Payment Dates (between indicated dates)	Redemption Percentage
From November 5, 2010 to and including August 5, 2011	108%
From November 5, 2011 to and including August 5, 2012	104%
From November 5, 2012 and thereafter	100%

Legal Proceedings. We are subject to various patent, product liability, government investigations and other legal proceedings in the ordinary course of business. Contingent accruals are recorded when we determine that a loss related to a litigation matter is both probable and reasonably estimable. Due to the fact that legal proceedings and other contingencies are inherently unpredictable, our assessments involve significant judgments regarding future events. For a complete description of legal proceedings, see Note 10 of the Condensed Consolidated Financial Statements included in Part I, Item 1 of this Quarterly Report on Form 10-Q.

Fluctuations. Our quarterly results have fluctuated in the past, and may continue to fluctuate. These fluctuations may be to the timing of new product launches, purchasing patterns of our customers, market acceptance of our products, the impact of competitive products and pricing, impairment of intangible assets, separation benefits, business combination transaction costs, upfront, milestone and certain other payments made or accrued pursuant to licensing agreements and changes in the fair value of financial instruments and contingent assets and liabilities recorded as part of a business combination. Further, a substantial portion of our net sales are through three wholesale drug distributors who in turn supply our products to pharmacies, hospitals and physicians. Accordingly, we are potentially subject to a concentration of credit risk with respect to our trade receivables.

Growth Opportunities. We continue to evaluate growth opportunities including strategic investments, licensing arrangements, acquisitions of businesses, product rights or technologies, and strategic alliances and promotional arrangements which could require significant capital resources. We intend to continue to focus our business development activities on further diversifying our revenue base through product licensing and company acquisitions, as well as other opportunities to enhance stockholder value. Through execution of our business strategy we intend to focus on developing new products through both an internal and a virtual research and development organization with greater scientific and clinical capabilities; expanding the Company's product line by acquiring new products and technologies in existing therapeutic and complementary areas; increasing revenues and earnings through sales and marketing programs for our innovative product offerings and effectively using the Company's resources; and providing additional resources to support our generics business.

Non-U.S. Operations. We currently have no operations outside of the United States. As a result, fluctuations in foreign currency exchange rates do not have a material effect on our financial statements.

Inflation. We do not believe that inflation had a material adverse effect on our financial statements for the periods presented.

Off-Balance Sheet Arrangements. We have no off-balance sheet arrangements as defined in Item 303(a) (4) of Regulation S-K.

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CRITICAL ACCOUNTING ESTIMATES

For a complete discussion of the Company's critical accounting estimates, see "Critical Accounting Estimates" in Item 7 of our Annual Report on Form 10-K for the year ended December 31, 2009, filed with the Securities and Exchange Commission on February 26, 2010.

Item 3. Quantitative and Qualitative Disclosures about Market Risk.

For quantitative and qualitative disclosures about market risk, see Item 7A, "Quantitative and Qualitative Disclosures about Market Risk," of our annual report on Form 10-K for the year ended December 31, 2009, filed with the Securities and Exchange Commission on February 26, 2010. Our exposures to market risk have not changed materially since December 31, 2009.

Item 4. Controls and Procedures.

Evaluation of Disclosure Controls and Procedures

The Company's management, with the participation of the Company's Chief Executive Officer and Principal Financial Officer, has evaluated the effectiveness of the Company's disclosure controls and procedures, as defined in Rules 13a-15(e) and 15d-15(e) of the Securities Exchange Act of 1934, as of March 31, 2010. Based on that evaluation, the Company's Chief Executive Officer and Principal Financial Officer concluded that the Company's disclosure controls and procedures were effective as of March 31, 2010.

Changes in Internal Control over Financial Reporting

There were no changes in the Company's internal control over financial reporting during the first quarter of 2010 that have materially affected, or are reasonably likely to materially affect, the Company's internal control over financial reporting.

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PART II

OTHER INFORMATION

Item 1. Legal Proceedings.

The disclosures under Note 10. Commitments and Contingencies-Legal Proceedings included in Part I Item I of this Report is incorporated in this Part II, Item 1 by reference.

Item 1A. Risk Factors

The risk factor listed below is included for the purpose of supplementing the risk factors disclosed in the section entitled "Risk Factors" in Part I, Item 1A of our Annual Report on Form 10-K for the year ended December 31, 2009 filed with the SEC on February 26, 2010. Other than this new risk factor, there have been no material changes to the risk factors disclosed in the Company's Annual Report on Form 10-K for the year ended December 31, 2009.

While healthcare reform may increase the number of patients who have insurance coverage for our products, its cost containment measures may adversely affect reimbursement for our products.

On March 23, 2010, President Obama signed into law H.R. 3590, the Patient Protection and Affordable Care Act (PPACA). On March 30, 2010, the President signed H.R. 4872, the Healthcare and Education Reconciliation Act of 2010 (Reconciliation Act), which included a package of fixes to the PPACA as well as additional elements to reform healthcare in the United States.

The passage of the PPACA and the Reconciliation Act is expected to result in a transformation of the delivery and payment for healthcare services in the U.S. The combination of these measures will expand health insurance coverage to an estimated 32 million Americans. In addition, there are significant health insurance reforms that will improve patients' ability to obtain and maintain health insurance. Such measures include the elimination of lifetime caps, no rescission of policies, and no denial of coverage due to preexisting conditions. The expansion of healthcare insurance and these additional market reforms should result in greater access to our products.

However, a number of provisions contained in the healthcare reform package may adversely affect reimbursement for our products. In 2010, the new law will increase the minimum basic Medicaid rebate for brand name prescription drugs to 23.1%, increase the minimum basic Medicaid rebate for generic drugs to 13%, require pharmaceutical manufacturers to pay states rebates on prescription drugs dispensed to Medicaid managed care enrollees, increase the additional Medicaid rebates for "new formulations", and expand the entities eligible for 340B pricing and the revision of the AMP definition to remove physician class of trade. Further, the best price requirements with respect to Medicaid rebates have traditionally been a significant consideration with respect to the level of rebates in our Medicare and commercial contracting which could adversely impact our future results of operations.

Item 2. Unregistered Sale of Equity Securities and Use of Proceeds.

None.

Item 3. Defaults Upon Senior Securities.

None.

Item 4. Removed and Reserved.

Item 5. Other Information.

None.

Item 6. Exhibits.

The information called for by this item is incorporated by reference to the Exhibit Index of this Report.

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SIGNATURES

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

ENDO PHARMACEUTICALS HOLDINGS INC.

(Registrant)

Name: */s/ DAVID P. HOLVECK*
David P. Holveck
Title: **President and Chief Executive Officer**

(Principal Executive Officer)

Name: */s/ ALAN G. LEVIN*
Alan G. Levin
Title: **Executive Vice President, Chief Financial Officer**

(Principal Financial Officer)

Name: */s/ EDWARD J. SWEENEY*
Edward J. Sweeney
Title: **Vice President, Controller and Principal Accounting Officer**

(Principal Accounting Officer)

Date: May 3, 2010

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No.	Title
3.1	Amended and Restated Certificate of Incorporation of Endo Pharmaceuticals Holdings Inc. (Endo) (incorporated herein by reference to Exhibit 10.32 of the Form 10-Q for the Quarter ended June 30, 2008 filed with the Commission on August 1, 2008)
3.2	Amended and Restated By-laws of Endo (incorporated herein by reference to Exhibit 3.2 of the Form 10-K for the year ended December 31, 2009 filed with the Commission on February 26, 2010)
4.1	Amended and Restated Executive Stockholders Agreement, dated as of July 7, 2003, by and among Endo, Endo Pharma LLC (Endo LLC), Kelso Investment Associates V, L.P. (KIA V), Kelso Equity Partners V, L.P. (KEP V) and the Management Stockholders (as defined therein) (incorporated herein by reference to Exhibit 4.1 of the Form 10-Q for the Quarter ended June 30, 2003 filed with the Commission on August 14, 2003)
4.1.2	Amendment to Amended and Restated Executive Stockholders Agreement, dated as of June 28, 2004, by and among Endo, Endo LLC, KIA V, KEP V and the Management Stockholders (as defined therein) (incorporated herein by reference to Exhibit 4.1 of the Form 10-Q for the Quarter ended September 30, 2004 filed with the Commission on November 5, 2004) the Commission on July 1, 2003)
4.1.3	Amendment 2 to the Amended and Restated Stockholders Agreement, dated September 20, 2005, by and among the Company, Endo LLC, Kelso and certain Amending Stockholders (as defined therein) (incorporated herein by reference to Exhibit 4.1.3 of the Current Report on Form 8-K filed with the Commission on September 22, 2005)
4.2	Amended and Restated Employee Stockholders Agreement, dated as of June 5, 2003, by and among Endo, Endo LLC, KIA V, KEP V and the Employee Stockholders (as defined therein) (incorporated herein by reference to Exhibit 10.2 of Amendment No. 2 to the Form S-3 Registration Statement (Registration No. 333-105338) filed with the Commission on July 1, 2003)
4.2.2	Amendment to Amended and Restated Employee Stockholders Agreement, dated as of June 28, 2004, by and among Endo, Endo LLC, KIA V, KEPV and the Management Stockholders (as defined therein) (incorporated herein by reference to Exhibit 4.1 of the Form 10-Q for the Quarter ended September 30, 2004 filed with the Commission on November 5, 2004)
4.2.3	Amendment 2 to the Amended and Restated Employee Stockholders Agreement, dated September 20, 2005, by and among the Company, Endo LLC, Kelso and certain Amending Stockholders (as defined therein) (incorporated herein by reference to Exhibit 4.2.3 of the Current Report on Form 8-K filed with the Commission on September 22, 2005)
4.3	Employee Stockholders Consent and Release, effective September 20, 2005, by and among the Company, Endo LLC, Kelso and certain Employee Stockholders (as defined therein) signatory thereto (incorporated herein by reference to Exhibit 4.3 of the Current Report on Form 8-K filed with the Commission on September 22, 2005)
4.4	Registration Rights Agreement, dated as of July 17, 2000, by and between Endo and Endo LLC (incorporated herein by reference to Exhibit 4.4 of the Form 10-Q for the Quarter ended June 30, 2000 filed with the Commission on August 15, 2000)
4.5	Amendment to Registration Rights Agreement, dated as of June 30, 2003, by and between Endo and Endo LLC (incorporated herein by reference to Exhibit 10.1 of Amendment No. 2 to the Form S-3 Registration Statement (Registration No. 333-105338) filed with the Commission on July 1, 2003)
4.8	Indenture dated as of August 6, 2007 between Indevus and The Bank of New York Trust Company, N.A, as trustee (incorporated herein by reference to Exhibit 4.1 of the Indevus Current Report on Form 8-K filed with the Commission on August 7, 2007)
4.8.1	Supplemental Indenture, dated as of March 23, 2009, by and between Indevus and the The Bank of New York Mellon Trust Company, N.A. (formerly known as The Bank of New York Trust Company, N.A.) (incorporated herein by reference to Exhibit 10.1 to the Indevus Current Report on Form 8-K, dated March 23, 2009)
10.1	Shelf Registration Agreement, dated September 21, 2005, by and between Endo, Endo LLC and certain Management Stockholders (as defined therein) (incorporated herein by reference to Exhibit 10.1 of the Current Report on Form 8-K filed with the Commission on September 22, 2005)

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No.	Title
10.2	Shelf Registration Agreement, dated April 30, 2004, between Endo Pharmaceuticals Holdings Inc. and Endo Pharma LLC (incorporated herein by reference to Exhibit 10.2 of Amendment No. 1 to the Form S-3 Registration Statement (Registration No. 333-115032) filed with the Commission on June 10, 2004)
10.3	Amendment to Shelf Registration Agreement, dated June 10, 2004 between Endo Pharmaceuticals Holdings Inc. and Endo Pharma LLC (incorporated herein by reference to Exhibit 10.3 of Amendment No. 1 to the Form S-3 Registration Statement (Registration No. 333-115032) filed with the Commission on June 10, 2004)
10.4	Agreement dated April 29, 2008 between Endo Pharmaceuticals Holdings Inc. and D. E. Shaw Valence Portfolios, L.L.C. (on behalf of itself and its affiliates that are members of the 13D Group with respect to the Endo common stock) (incorporated herein by reference to Exhibit 99.1 of the Current Report on Form 8-K/A dated May 1, 2008)
10.5	[Intentionally Omitted.]
10.6	Amended and Restated Tax Sharing Agreement, dated as of April 30, 2004 by and among Endo, Endo Inc. and Endo LLC (incorporated herein by reference to Exhibit 10.6 of the Form 10-Q for the Quarter ended March 31, 2004 filed with the Commission on May 10, 2004)
10.7	Convertible Bond Hedge Transaction Confirmation entered into by and between the Company and Deutsche Bank AG, London Branch, dated April 9, 2008 (incorporated herein by reference to Exhibit 10.7 of the Form 10-Q for the Quarter ended March 31, 2008 filed with the Commission on May 2, 2008)
10.8	Issuer Warrant Transaction Confirmation entered into by and between the Company and Deutsche Bank AG, London Branch, dated April 9, 2008 (incorporated herein by reference to Exhibit 10.8 of the Form 10-Q for the Quarter ended March 31, 2008 filed with the Commission on May 2, 2008)
10.9	Issuer Share Repurchase Transaction Confirmation entered into by and between the Company and Deutsche Bank AG, London Branch, dated April 9, 2008 (incorporated herein by reference to Exhibit 10.9 of the Form 10-Q for the Quarter ended March 31, 2008 filed with the Commission on May 2, 2008)
10.10	Sole and Exclusive License Agreement, dated as of November 23, 1998, by and between Endo Pharmaceuticals Inc. (Endo Pharmaceuticals) and Hind HealthCare, Inc. (incorporated herein by reference to Exhibit 10.10 of the Registration Statement filed with the Commission on June 9, 2000)
10.11	Endo Pharmaceuticals Holdings Inc. Executive Deferred Compensation Plan (incorporated herein by reference to Exhibit 10.1 of the Current Report on Form 8-K dated December 19, 2007)
10.12	Endo Pharmaceuticals Holdings Inc. 401(k) Restoration Plan (incorporated herein by reference to Exhibit 10.2 of the Current Report on Form 8-K dated December 19, 2007)
10.13	[Intentionally Omitted.]
10.14	Supply and Manufacturing Agreement, dated as of November 23, 1998, by and between Endo Pharmaceuticals and Teikoku Seiyaku Co., Ltd (incorporated herein by reference to Exhibit 10.14 of the Registration Statement filed with the Commission on June 9, 2000)
10.14.1	First Amendment, dated April 24, 2007, to the Supply and Manufacturing Agreement, dated as of November 23, 1998, by and between Endo Pharmaceuticals Inc. and Teikoku Seiyaku Co., Ltd. / Teikoku Pharma USA, Inc. (incorporated herein by reference to Exhibit 10.14.1 of the Current Report on Form 8-K dated April 30, 2007)
10.14.2	Second Amendment, dated January 6, 2010, to the Supply and Manufacturing Agreement, dated as of November 23, 1998, by and between Endo Pharmaceuticals Inc. and Teikoku Seiyaku Co., Ltd. / Teikoku Pharma USA, Inc. (incorporated herein by reference to Exhibit 10.14.2 of the Current Report on Form 8-K dated January 11, 2010)
10.15	Supply Agreement, dated as of July 1, 1998, by and between Endo Pharmaceuticals and Mallinckrodt Inc. (Mallinckrodt) (incorporated herein by reference to Exhibit 10.15 of the Registration Statement filed with the Commission on June 9, 2000)
10.16	Supply Agreement for Bulk Narcotics Raw Materials, dated as of July 1, 1998, by and between Endo Pharmaceuticals and Mallinckrodt (incorporated herein by reference to Exhibit 10.16 of the Registration Statement filed with the Commission on June 9, 2000)
10.16.1	

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First Amendment, effective July 1, 2000, to the Supply Agreement for Bulk Narcotics Raw Materials, dated as of July 1, 1998, by and between Endo Pharmaceuticals and Mallinckrodt (incorporated herein by reference to Exhibit 10.16.1 of the Current Report on Form 8-K dated April 14, 2006)

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No.	Title
10.16.2	Second Amendment, dated April 10, 2006, to the Supply Agreement for Bulk Narcotics Raw Materials, dated as of July 1, 1998, by and between Endo Pharmaceuticals and Mallinckrodt (incorporated herein by reference to Exhibit 10.16.2 of the Current Report on Form 8-K dated April 14, 2006)
10.17	[Intentionally Omitted.]
10.18	Amended and Restated Strategic Alliance Agreement, dated as of April 2, 2002, by and between Endo Pharmaceuticals and Penwest Pharmaceuticals Co. (incorporated herein by reference to Exhibit 10.18 of the Quarterly Report on Form 10-Q for the Quarter Ended March 31, 2002 filed with the Commission on May 14, 2002)
10.18.1	Amendment, dated January 7, 2007, to the Amended and Restated Strategic Alliance Agreement, dated as of April 2, 2002, by and between Endo Pharmaceuticals Inc. and Penwest Pharmaceuticals Co. (incorporated herein by reference to Exhibit 10.18.1 of the Current report on Form 8-K dated January 11, 2007)
10.18.2	Third Amendment to the Amended and Restated Strategic Alliance Agreement by and between Penwest Pharmaceuticals Co. and Endo Pharmaceuticals Inc., dated as of March 31, 2009 (incorporated herein by reference to Exhibit 10.18.2 of the Current report on Form 8-K dated April 6, 2009)
10.18.3	Fourth Amendment to the Amended and Restated Strategic Alliance Agreement by and between Penwest Pharmaceuticals Co. and Endo Pharmaceuticals Inc., dated as of April 8, 2010.
10.19	Agreement, dated as of February 1, 2000, by and between Endo Pharmaceuticals and UPS Supply Chain Solutions, Inc. (f/d/b/a Livingston Healthcare Services Inc.) (incorporated herein by reference to Exhibit 10.19 of the Registration Statement filed with the Commission on June 9, 2000)
10.20	Medical Affairs Support Services Agreement, dated as of June 1, 1999, by and between Endo Pharmaceuticals and Kunitz and Associates, Inc. (incorporated herein by reference to Exhibit 10.20 of the Registration Statement filed with the Commission on June 9, 2000)
10.21	Endo Pharmaceuticals Holdings Inc. 2000 Stock Incentive Plan (incorporated herein by reference to Exhibit 10.21 of the Quarterly Report on Form 10-Q for the Quarter Ended September 30, 2000 filed with the Commission on November 13, 2000)
10.22	Endo LLC Amended and Restated 1997 Employee Stock Option Plan (incorporated herein by reference to Exhibit 10.22 of the Quarterly Report on Form 10-Q for the Quarter Ended September 30, 2000 filed with the Commission on November 13, 2000)
10.23	Endo LLC Amended and Restated 1997 Executive Stock Option Plan (incorporated herein by reference to Exhibit 10.23 of the Quarterly Report on Form 10-Q for the Quarter Ended September 30, 2000 filed with the Commission on November 13, 2000)
10.24	Endo LLC 2000 Amended and Restated Supplemental Employee Stock Option Plan (incorporated herein by reference to Exhibit 10.24 of the Quarterly Report on Form 10-Q for the Quarter Ended September 30, 2000 filed with the Commission on November 13, 2000)
10.25	Endo LLC 2000 Amended and Restated Supplemental Executive Stock Option Plan (incorporated herein by reference to Exhibit 10.25 of the Quarterly Report on Form 10-Q for the Quarter Ended September 30, 2000 filed with the Commission on November 13, 2000)
10.26	Separation Agreement, dated as of September 8, 2008, between the Endo Pharmaceuticals Holdings Inc. and Charles A. Rowland, Jr. (incorporated herein by reference to Exhibit 10.1 of the Current Report on Form 8-K dated September 8, 2008)
10.27	Executive Employment Agreement between Endo Pharmaceuticals Holdings Inc. and Ivan Gergel, M.D., dated as of April 29, 2008 (incorporated herein by reference to Exhibit 10.1 of the Current Report on Form 8-K dated March 25, 2009)
10.28	Amended and Restated Employment Agreement, dated as of December 19, 2007, by and between the Company and Nancy J. Wysenski (incorporated herein by reference to Exhibit 10.29 of the Form 10-K for the year ended December 31, 2007 filed with the Commission on February 26, 2008)
10.29	Auction-Rate Securities Rights Agreement, dated November 10, 2008, by and between Endo Pharmaceuticals and UBS AG (incorporated herein by reference to Exhibit 10.29 to the Form 10-K for the year ended December 31, 2008 filed with the Commission on March 2, 2009)
10.30	Employment Agreement, dated as of April 1, 2008, by and between Endo Pharmaceuticals Holdings Inc. and David P. Holveck (incorporated herein by reference to Exhibit 10.30 of the Current Report on Form 8-K dated March 12, 2008)

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No.	Title
10.31	License and Supply Agreement by and by and among Novartis, AG, Novartis Consumer Health, Inc. and Endo Pharmaceuticals Inc. dated as of March 4, 2008 (incorporated herein by reference to Exhibit 10.31 of the Form 10-Q for the Quarter ended March 31, 2008 filed with the Commission on May 2, 2008)
10.31.1	Amendment No. 1 to the License and Supply Agreement by and by and among Novartis, AG, Novartis Consumer Health, Inc. and Endo Pharmaceuticals Inc. dated as of March 28, 2008 (incorporated herein by reference to Exhibit 10.31.1 of the Form 10-Q for the Quarter ended March 31, 2008 filed with the Commission on May 2, 2008)
10.32	Sales and Marketing Services Agreement, dated as of May 15, 2008 between Endo Pharmaceuticals and Ventiv Commercial Services, LLC (incorporated herein by reference to Exhibit 10.32 of the Form 10-Q for the Quarter ended June 30, 2008 filed with the Commission on August 1, 2008)
10.32.1	Amendment to the Sales and Marketing Services Agreement, dated as of January 29, 2009 between Endo Pharmaceuticals and Ventiv Commercial Services, LLC (incorporated herein by reference to Exhibit 10.32.1 to the Form 10-K for the year ended December 31, 2008 filed with the Commission on March 2, 2009)
10.32.2	Amendment to the Sales Representative Service Agreement, dated as of April 1, 2009 between Endo Pharmaceuticals Inc. and Ventiv Commercial Services, LLC (incorporated herein by reference to Exhibit 10.32.2 of the Current Report on Form 8-K dated April 7, 2009)
10.32.3*	Amendment to the Sales Representative Services Agreement, dated as of May 11, 2009 between Endo Pharmaceuticals Inc. and Ventiv Commercial Services, LLC (incorporated herein by reference to Exhibit 10.32.3 of the Current Report on Form 8-K dated May 11, 2009)
10.33	[Intentionally Omitted.]
10.34	Lease Agreement, dated as of May 5, 2000, by and between Endo Pharmaceuticals and Painters Crossing One Associates, L.P. (incorporated herein by reference to Exhibit 10.34 of the Registration Statement filed with the Commission on June 9, 2000)
10.34.1	Amendment to Lease Agreement, dated as of November 6, 2006, by and between Endo Pharmaceuticals and Painters Crossing One Associates, L.P. (incorporated herein by reference to Exhibit 10.34.1 of the Form 10-Q for the quarter ended September 30, 2006 filed with the Commission on November 9, 2006)
10.35	Amended and Restated Employment Agreement, dated as of December 19, 2007, by and between the Company and Caroline B. Manogue (incorporated herein by reference to Exhibit 10.29 of the Form 10-K for the year ended December 31, 2007 filed with the Commission on February 26, 2008)
10.36	Employment Agreement between Endo Pharmaceuticals Holdings Inc. and Julie McHugh (incorporated herein by reference to Exhibit 10.2 of the Current Report on Form 8-K, dated March 12, 2010).
10.37	Endo Pharmaceuticals Holdings Inc. 2004 Stock Incentive Plan (incorporated herein by reference to Exhibit 10.37 of the Form 10-Q for the Quarter ended June 30, 2004 filed with the Commission on August 9, 2004)
10.38	Endo Pharmaceuticals Holdings Inc. 2007 Stock Incentive Plan (incorporated herein by reference to Exhibit D of the Definitive Proxy Statement on Schedule 14A filed with the Commission on April 30, 2007)
10.39	Master Development and Toll Manufacturing Agreement, dated as of May 3, 2001, by and between Novartis Consumer Health, Inc. and Endo Pharmaceuticals (incorporated herein by reference to Exhibit 10.39 of the Form 10-Q for the Quarter Ended June 30, 2001 filed with the Commission on August 14, 2001)
10.39.1	First Amendment, effective February 1, 2003, to the Master Development and Toll Manufacturing Agreement between Endo Pharmaceuticals and Novartis Consumer Health, Inc. (incorporated herein by reference to Exhibit 10.39.1 of the Form 10-Q for the Quarter Ended June 30, 2005 filed with the Commission on August 8, 2005)
10.39.2	Second Amendment, effective as of December 1, 2004, to the Master Development and Toll Manufacturing Agreement between Endo Pharmaceuticals and Novartis Consumer Health, Inc. (incorporated herein by reference to Exhibit 10.39.2 of the Form 10-Q for the Quarter Ended June 30, 2005 filed with the Commission on August 8, 2005)
10.40	Lease Agreement between Painters Crossing Three Associates, L.P. and Endo Pharmaceuticals Inc. dated January 19, 2007 (incorporated herein by reference to Exhibit 10.40 of the Annual Report on Form 10-K for the Year Ended December 31, 2006 filed with the Commission on March 1, 2007)

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- 10.40.1 First Amendment to Lease Agreement, dated as of March 3, 2008 by and between Partners Crossing Three Associates, L.P. and Endo Pharmaceuticals Inc. (incorporated herein by reference to Exhibit 10.40.1 of the Form 10-Q for the Quarter ended March 31, 2008 filed with the Commission on May 2, 2008)

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No.	Title
10.41	Policy of Endo Pharmaceuticals Holdings Inc. Relating to Insider Trading in Company Securities and Confidentiality of Information (incorporated herein by reference to Exhibit 10.41 of the Form 10-Q for the Quarter ended March 31, 2005 filed with the Commission on May 10, 2005)
10.42	Form of Indemnification Agreement (incorporated herein by reference to Exhibit 10.1 of the Current Report on Form 8-K, dated May 8, 2009).
10.43	Employment Agreement between Endo Pharmaceuticals Holdings Inc. and Alan G. Levin (incorporated herein by reference to Exhibit 10.2 of the Current Report on Form 8-K, dated May 8, 2009).
10.44	Lease Agreement, dated as of January 6, 2003, by and between Endo Pharmaceuticals and Dawson Holding Company (incorporated by reference to Exhibit 10.44 of the Annual Report on Form 10-K for the Year Ended December 31, 2002 filed with the Commission on March 27, 2003)
10.45	Lease Agreement, dated as of November 13, 2003, by and between Endo Pharmaceuticals and Painters Crossing Two Associates, L.P. (incorporated herein by reference to Exhibit 10.45 of the Annual Report on Form 10-K for the Year Ended December 31, 2003 filed with the Commission on March 15, 2004)
10.45.1	Amendment to Lease Agreement, dated as of February 16, 2005, by and between Endo Pharmaceuticals and Painters Crossing Two Associates, L.P. (incorporated herein by reference to Exhibit 10.45.1 of the Current Report on Form 8-K dated February 18, 2005)
10.45.2	Amendment to Lease Agreement, dated as of November 6, 2006, by and between Endo Pharmaceuticals and Painters Crossing Two Associates, L.P. (incorporated herein by reference to Exhibit 10.34.1 of the Form 10-Q for the quarter ended September 30, 2006 filed with the Commission on November 9, 2006)
10.46	License Agreement, dated as of February 25, 2004, by and between Endo Pharmaceuticals and Noven Pharmaceuticals, Inc. (incorporated herein by reference to Exhibit 10.46 of Amendment No. 2 to the Annual Report on Form 10-K for the Year Ended December 31, 2003 filed with the Commission on June 25, 2004)
10.46.1	Termination Agreement, dated as of February 24, 2006, by and between Noven Pharmaceuticals, Inc. and Endo Pharmaceuticals (incorporated herein by reference to Exhibit 10.46.1 of the Annual Report on Form 10-K for the Year Ended December 31, 2005 filed with the Commission on March 8, 2006)
10.47	Supply Agreement, dated as of February 25, 2004, by and between Endo Pharmaceuticals and Noven Pharmaceuticals, Inc. (incorporated herein by reference to Exhibit 10.47 of Amendment No. 2 to the Annual Report on Form 10-K for the Year Ended December 31, 2003 filed with the Commission on June 25, 2004)
10.48	License and Co-Promotion Rights Agreement, dated as of July 14, 2004, by and between Endo Pharmaceuticals and Vernalis Development Limited (incorporated herein by reference to Exhibit 10.48 of the Current Report on Form 8-K dated July 19, 2004)
10.48.1	Co-Promotion Agreement, dated as of July 1, 2005, by and between Endo Pharmaceuticals Inc. and Vernalis Development Limited (incorporated by reference to Exhibit 10.48.1 of the Current Report on Form 8-K dated July 8, 2005)
10.48.2	Second Amendment, dated as of December 12, 2005, to the License Agreement by and between Endo Pharmaceuticals Inc. and Vernalis Development Limited (incorporated by reference to Exhibit 10.48.2 of the Current Report on Form 8-K dated December 29, 2005)
10.48.3	First Amendment, dated as of December 12, 2005, to the Co-Promotion Agreement by and between Endo Pharmaceuticals Inc. and Vernalis Development Limited (incorporated by reference to Exhibit 10.48.3 of the Current Report on Form 8-K dated December 29, 2005)

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No.	Title
10.48.4	Third Amendment, dated as of July 23, 2007, to the License Agreement by and between Endo Pharmaceuticals Inc. and Vernalis Development Limited (incorporated by reference to Exhibit 10.48.4 of the Current Report on Form 8-K dated July 27, 2007)
10.48.5	Fourth Amendment, dated as of February 19, 2008, to the License Agreement by and between Endo Pharmaceuticals Inc. and Vernalis Development Limited (incorporated herein by reference to Exhibit 10.48.5 of the Form 10-K for the year ended December 31, 2007 filed with the Commission on February 26, 2008)
10.48.6	Agreement to Terminate the Co-Promotion Agreement by and between Endo Pharmaceuticals Inc. and Vernalis Development Limited, effective February 19, 2008 (incorporated herein by reference to Exhibit 10.48.6 of the Form 10-K for the year ended December 31, 2007 filed with the Commission on February 26, 2008)
10.49	Loan Agreement, dated as of July 14, 2004, by and between Endo Pharmaceuticals and Vernalis Development Limited (incorporated herein by reference to Exhibit 10.49 of the Current Report on Form 8-K dated July 19, 2004)
10.49.1	Agreement to Terminate the Loan Agreement by and between Endo Pharmaceuticals and Vernalis Development Limited, effective February 19, 2008 (incorporated herein by reference to Exhibit 10.49.1 of the Form 10-K for the year ended December 31, 2007 filed with the Commission on February 26, 2008)
10.50	Form of Stock Option Grant Agreement under the 2007 Stock Incentive Plan (incorporated herein by reference to Exhibit 10.50 of the Form 10-K for the year ended December 31, 2008 filed with the Commission on March 2, 2009)
10.51	Form of Restricted Stock Unit Grant Agreement under the 2007 Stock Incentive Plan (incorporated herein by reference to Exhibit 10.51 of the Form 10-K for the year ended December 31, 2008 filed with the Commission on March 2, 2009)
10.52	Agreement and Plan of Merger dated January 5, 2009, by and between Endo Pharmaceuticals Holdings Inc., BTB Purchaser, and Indevus Pharmaceuticals, Inc. (incorporated herein by reference to Exhibit 10.1 of the Current Report on Form 8-K dated January 5, 2009)
10.52.1	Amendment, dated January 7, 2009 to the Agreement and Plan of Merger, by and between Endo Pharmaceuticals Holdings Inc., BTB Purchaser, and Indevus Pharmaceuticals, Inc. (incorporated herein by reference to Exhibit 10.1 of the Current Report on Form 8-K dated January 7, 2009)
10.52.2	Amendment No. 2, dated February 4, 2009, to the Agreement and Plan of Merger, by and among Endo Pharmaceuticals Holdings, Inc., BTB Purchaser, and Indevus Pharmaceuticals, Inc. (incorporated herein by reference to Exhibit 2.1 of the Current Report on Form 8-K dated February 6, 2009)
10.53	Form of Stockholder Tender Agreement (incorporated herein by reference to Exhibit 10.2 of the Current Report on Form 8-K dated January 5, 2009)
10.54	Nebido® Contingent Cash Consideration Agreement, dated February 23, 2009, by and between Endo Pharmaceuticals Holdings Inc. and American Stock Transfer and Trust Company (incorporated herein by reference to Exhibit 10.54 of the Form 10-K for the year ended December 31, 2008 filed with the Commission on March 2, 2009)
10.55	Octreotide Contingent Cash Consideration Agreement, dated February 23, 2009, by and between Endo Pharmaceuticals Holdings Inc. and American Stock Transfer and Trust Company (incorporated herein by reference to Exhibit 10.55 of the Form 10-K for the year ended December 31, 2008 filed with the Commission on March 2, 2009)
10.56	Memorandum of Understanding, dated February 4, 2009, by and among (i) Wolf Popper LLP, counsel for Plaintiff Arthur Gober, CBM IRA Beneficiary Custodian, Beneficiary of Jerome Gober, (ii) Skadden, Arps, Slate, Meagher & Flom LLP, counsel for Defendants Endo Pharmaceuticals Holdings Inc. and BTB Purchaser Inc., (iii) The Weiser Law Firm, P.C., counsel for Plaintiff Martin Wexler, (iv) Young Conaway Stargatt & Taylor, LLP, counsel for Defendants Indevus Pharmaceuticals, Inc., Glenn L. Cooper, Andrew Ferrara, James C. Gale, Michael E. Hanson, Stephen C. McCluski, Cheryl P. Morley and Malcolm Morville, (v) Levi & Korsinsky LLP, counsel for Plaintiff Malena C. Schroeder and (vi) Johnson Bottini LLP, counsel for Plaintiff H. Steven Mishket (incorporated herein by reference to Exhibit 10.1 of the Current Report on Form 8-K dated February 6, 2009)
10.57	Amended and Restated License, Commercialization and Supply Agreement executed September 18, 2007 between Indevus and Esprit Pharma, Inc. (incorporated herein by reference to Exhibit 10.1 to the Indevus Current Report on Form 8-K dated September 21, 2007)
10.58	Lease Agreement between National Patent Development Corporation and Cedar Brook Corporate Center, L.P. dated October 6, 1997 (incorporated herein by reference to Exhibit 10.9 to the Valera Registration Statement on Form S-1 (File No. 333-123288)

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filed with the Commission on December 9, 2005)

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No.	Title
10.59	Amendment to Lease between Valera Pharmaceuticals, Inc. and Cedar Brook Corporate Center, L.P. dated January 7, 2004 (incorporated herein by reference to Exhibit 10.10 to the Valera Registration Statement on Form S-1 (File No. 333-123288) filed with the Commission on December 9, 2005)
10.60	Lease Agreement between Valera Pharmaceuticals, Inc. and Cedar Brook 7 Corporate Center, L.P. dated March 8, 2005 (incorporated herein by reference to Exhibit 10.11 to the Valera Registration Statement on Form S-1 (File No. 333-123288) filed with the Commission on December 9, 2005)
10.61	Agreement and Plan of merger, dated as of December 11, 2006, by and among Indevus, Hayden Merger Sub, Inc. and Valera Pharmaceuticals, Inc. (incorporated herein by reference to Exhibit 2.1 to the Indevus Current Report on Form 8-K, dated December 12, 2006)
10.62	License Agreement dated February 18, 1994 between Indevus and Rhone-Poulenc Rorer, S.A. (incorporated herein by reference to the Indevus Registration Statement on Form S-3 or Amendment I (File no. 33-75826))
10.63	Lease dated February 5, 1997 between Indevus and Ledgemont Realty Trust (incorporated herein by reference to Exhibit 10.87 to the Indevus Form 10-Q for the period ended December 31, 1996 filed with the Commission on February 14, 1997)
10.64	License Agreement effective as of November 26, 1999 between Madaus AG and Indevus (incorporated herein by reference to Exhibit 10.113 to the Indevus Form 10-K for the fiscal year ended September 30, 1999, filed with the Commission on December 28, 1999)
10.65	Indemnity and Release Agreement between American Home Products Corporation and Indevus dated as of May 30, 2001 (incorporated herein by reference to Exhibit 1.120 to the Indevus Form 10-Q for the period ended June 30, 2001, filed with the Commission on August 14, 2001)
10.66	Supply Agreement between Indevus and Madaus AG dated December 16, 2003 (incorporated herein by reference to Exhibit 10.129 to the Indevus Form 10-Q for the period ended December 31, 2002, filed with the Commission on February 14, 2003)
10.67	Development and License Agreement between Indevus and Shire Laboratories Inc. dated March 11, 2003 (incorporated herein by reference to Exhibit 10.130 to the Indevus Form 10-Q for the period ended March 31, 2003, filed with the Commission on April 13, 2003)
10.68	License, Commercialization and Supply Agreement dated April 6, 2004 between Indevus and Odyssey Pharmaceuticals Inc. (incorporated herein by reference to Exhibit 99.2 to the Indevus Current Report on Form 8-K dated April 19, 2004)
10.68.1	Amendment No. 1 to License, Commercialization and Supply Agreement dated April 30, 2005 between Indevus and Odyssey Pharmaceuticals, Inc. (incorporated herein by reference to Exhibit 10.143 to the Indevus Form 10-Q for the period ended March 31, 2005, filed with the Commission on May 10, 2005)
10.69	Indenture of Lease dated December 20, 2004 between Indevus and Mortimer B. Zuckerman and Edward H. Linde, Trustees of Hayden Office Trust (incorporated herein by reference to Exhibit 10.142 to the Indevus Form 10-Q for the period ended December 31, 2004, filed with the Commission on February 9, 2005)
10.70	Amendment and Consent Agreement dated May 14, 2005 between Indevus, Odyssey Pharmaceuticals, Inc., and Saturn Pharmaceuticals, Inc (incorporated herein by reference to Exhibit 10.1 to the Indevus Current Report on Form 8-K dated May 17, 2005)
10.71	License Agreement dated July 28, 2005 between Indevus and Schering Aktiengesellschaft (incorporated herein by reference to Exhibit 10.1 to the Indevus Current Report on Form 8-K dated August 2, 2005)
10.72	Manufacturing and Supply Agreement by and between Indevus and Schering AG, Germany dated on or about October 20, 2006 (incorporated herein by reference to Exhibit 10.158 to the Indevus Form 10-K for the fiscal year ended September 30, 2006, filed with the Commission on December 7, 2006)
10.73	License and Supply Agreement by and between Indevus and Madaus GmbH dated on or about November 3, 2006 (incorporated herein by reference to Exhibit 10.159 to the Indevus Form 10-K for the fiscal year ended September 30, 2006, filed with the Commission on December 7, 2006)
10.73.1	Amendment and Agreement by and between Indevus and Madaus GmbH dated on or about November 3, 2006 (incorporated herein by reference to Exhibit 10.160 to the Indevus Form 10-K for the fiscal year ended September 30, 2006, filed with the Commission on December 7, 2006)

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- 10.74 API Supply Agreement by and between Indevus and Helsinn Chemicals SA and Helsinn Advanced Synthesis SA dated on or about November 22, 2006 (incorporated herein by reference to Exhibit 10.162 to the Indevus Form 10-K for the fiscal year ended September 30, 2006, filed with the Commission on December 7, 2006)

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No.	Title
10.75	Supprelin Contingent Stock Rights Agreement, dated as of April 17, 2007, between Indevus and American Stock Transfer & Trust Company (incorporated herein by reference to Exhibit 10.1 to the Indevus Current Report on Form 8-K dated April 17, 2007)
10.75.1	Supplemental Supprelin CSR Agreement, dated as of March 23, 2009, by and between Endo American Stock Transfer & Trust (incorporated herein by reference to Exhibit 10.3 of the Current Report on Form 8-K dated March 23, 2009)
10.76	Stent Contingent Stock Rights Agreement, dated as of April 17, 2007, between Indevus and American Stock Transfer & Trust Company (incorporated herein by reference to Exhibit 10.2 to the Indevus Current Report on Form 8-K dated April 17, 2007)
10.76.1	Supplemental Stent CSR Agreement, dated as of March 23, 2009, by and between Endo American Stock Transfer & Trust (incorporated herein by reference to Exhibit 10.2 of the Current Report on Form 8-K dated March 23, 2009)
10.77	Octreotide Contingent Stock Rights Agreement, dated as of April 17, 2007, between Indevus and American Stock Transfer & Trust Company (incorporated herein by reference to Exhibit 10.3 to the Indevus Current Report on Form 8-K dated April 17, 2007)
10.77.1	Supplemental Octreotide CSR Agreement, dated as of March 23, 2009, by and between Endo American Stock Transfer & Trust (incorporated herein by reference to Exhibit 10.1 of the Current Report on Form 8-K dated March 23, 2009)
10.78	Supply Agreement by and between Valera Pharmaceuticals, Inc. and Plantex USA Inc. (incorporated herein by reference to Exhibit 10.1 to the Valera Form 10-Q for the period ended June 30, 2006, filed with the Commission on August 9, 2006)
10.79	Form of License, Supply and Distribution Agreement by and between Indevus Pharmaceuticals, Inc. and Orion Corporation dated April 2, 2008 (incorporated herein by reference to Exhibit 10.208 to the Indevus Form 10-K for the fiscal year ending September 30, 2008, filed with the Commission on December 11, 2008)
10.80	Form of Purchase and Sale Agreement by and between Ledgemont Royalty Sub LLC and Indevus dated August 26, 2008 (incorporated herein by reference to Exhibit 10.215 to the Indevus Form 10-K for the fiscal year ending September 30, 2008, filed with the Commission on December 11, 2008)
10.81	Form of Note Purchase Agreement by and among Ledgemont Royalty Sub LLC, Indevus and the purchasers named therein dated August 26, 2008 (incorporated herein by reference to Exhibit 10.216 to the Indevus Form 10-K for the fiscal year ending September 30, 2008, filed with the Commission on December 11, 2008)
10.82	Form of Indenture by and between Ledgemont Royalty Sub LLC and U.S. Bank National Association dated August 26, 2008 (incorporated herein by reference to Exhibit 10.217 to the Indevus Form 10-K for the fiscal year ending September 30, 2008, filed with the Commission on December 11, 2008)
10.83	Form of Pledge and Security Agreement made by Indevus to U.S. Bank National Association, as Trustee, dated August 26, 2008 (incorporated herein by reference to Exhibit 10.218 to the Indevus Form 10-K for the fiscal year ending September 30, 2008, filed with the Commission on December 11, 2008)
10.84	Form of Development, License and Commercialization Agreement made by and between Indevus and Teva Pharmaceutical Industries Ltd., dated September 25, 2008 (incorporated herein by reference to Exhibit 10.219 to the Indevus Form 10-K for the fiscal year ending September 30, 2008, filed with the Commission on December 11, 2008)
10.85	First Amendment to Amended and Restated License, Commercialization and Supply Agreement between Indevus Pharmaceuticals, Inc. and Allergan USA, Inc. dated as of January 9, 2009 (incorporated herein by reference to Exhibit 10.1 to the Indevus Current Report on Form 8-K, dated January 15, 2009)
10.86	Agreement between National Patent Development Corporation and Dento-Med Industries, Inc. dated November 30, 1989 (incorporated herein by reference to Exhibit 10.17 to the Valera Registration Statement on Form S-1 (File No. 333-123288) filed with the Commission on December 9, 2005).
10.87	Contribution Agreement between Hydro Med Sciences, Inc. and GP Strategies Corporation dated June 30, 2000 (incorporated herein by reference to Exhibit 10.12 to the Valera Registration Statement on Form S-1 (File No. 333-123288) filed with the Commission on December 9, 2005).
10.88	Termination of Agreement dated September 12, 1990 between National Patent Development Corporation and The Population Council, Inc. dated October 1, 1997 (incorporated herein by reference to Exhibit 10.6 to the Valera Registration Statement on Form S-1 (File No. 333-123288) filed with the Commission on December 9, 2005).

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Exhibit

No.	Title
10.88.1	Amendment to the Termination of the Joint Development Agreement between GP Strategies Corporation and The Population Council, Inc. dated November 29, 2001 (incorporated herein by reference to Exhibit 10.7 to the Valera Registration Statement on Form S-1 (File No. 333-123288) filed with the Commission on December 9, 2005).
10.88.2	Amendment No. 2 to Termination Agreement between Valera Pharmaceuticals, Inc. and The Population Council, Inc. dated August 31, 2004 (incorporated herein by reference to Exhibit 10.8 to the Valera Registration Statement on Form S-1 (File No. 333-123288) filed with the Commission on December 9, 2005).
21	Subsidiaries of the Registrant
31.1	Certification of the President and Chief Executive Officer of Endo pursuant to Section 302 of the Sarbanes-Oxley Act of 2002
31.2	Certification of the Chief Financial Officer of Endo pursuant to Section 302 of the Sarbanes-Oxley Act of 2002
32.1	Certification of the President and Chief Executive Officer of Endo pursuant to 18 U.S.C. Section 1350, as adopted pursuant to Section 906 of the Sarbanes-Oxley Act of 2002
32.2	Certification of the Chief Financial Officer of Endo pursuant to 18 U.S.C. Section 1350, as adopted pursuant to Section 906 of the Sarbanes-Oxley Act of 2002

* Confidential portions of this exhibit have been redacted and filed separately with the Commission pursuant to a confidential treatment request in accordance with Rule 24b-2 of the Securities Exchange Act of 1934, as amended.