

ENDO PHARMACEUTICALS HOLDINGS INC

Form 10-Q

May 14, 2002

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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,
2002 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: 39040

ENDO PHARMACEUTICALS HOLDINGS INC.

(Exact Name of Registrant as Specified in Its Charter)

Delaware
(State or other jurisdiction of
incorporation or organization)

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

100 Painters Drive
Chadds Ford, Pennsylvania 19317
(Address of Principal Executive Offices)

(610) 558-9800
(Registrant's Telephone Number, Including Area Code)

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

The aggregate number of shares of the Registrant's common stock outstanding as of May 14, 2002 was 102,064,450.

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ENDO PHARMACEUTICALS HOLDINGS INC.

**REPORT ON FORM 10-Q
FOR THE QUARTERLY PERIOD ENDED MARCH 31, 2002**

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Forward Looking Statements

We have made forward-looking statements in this document within the meaning of Section 27A of the Securities Act of 1933 and Section 21E of the Securities Exchange Act of 1934, as amended. These statements, including estimates of future net sales and consolidated EBITDA contained in the section titled Management's Discussion and Analysis of Financial Condition and Results of Operations, 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, or similar expressions are forward-looking statements. We have based 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 in Management's Discussion and Analysis of Financial Condition and Results of Operations, Business and elsewhere in this Report could affect our future financial results and could cause our actual results to differ materially from those expressed in forward-looking statements contained in this Report. Important factors that could cause our actual results to differ materially from the expectations reflected in the forward-looking statements in this Report include, among others:

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

results of clinical trials on new 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 our products;

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

new regulatory action or lawsuits relating to the 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;

our ability to successfully implement our acquisition strategy;

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; and

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

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.

Table of Contents**PART I****FINANCIAL INFORMATION****Item 1. Financial Statements**

**ENDO PHARMACEUTICALS HOLDINGS INC.
CONSOLIDATED BALANCE SHEETS (UNAUDITED)
(In thousands, except share data)**

	March 31, 2002	December 31, 2001
<u>ASSETS</u>		
CURRENT ASSETS:		
Cash and cash equivalents	\$117,695	\$95,357
Accounts receivable, net	69,953	85,329
Inventories	30,565	27,766
Prepaid expenses	3,999	5,527
Deferred income taxes	28,067	26,946
<hr/>		
<hr/>		
Total current assets	250,279	240,925
<hr/>		
<hr/>		
PROPERTY AND EQUIPMENT, Net	9,364	9,883
GOODWILL	181,079	182,318
OTHER INTANGIBLES, Net	12,311	12,495
DEFERRED INCOME TAXES	20,418	23,420

RESTRICTED CASH

150 150

OTHER ASSETS

1,874 1,804

TOTAL ASSETS

\$475,475 \$470,995

**LIABILITIES AND
STOCKHOLDERS EQUITY**

CURRENT LIABILITIES:

Accounts payable

\$28,595 \$30,705

Accrued expenses

59,693 50,176

Income taxes payable

233 3,526

Current portion of long-term debt

92,968 91,259

Total current liabilities

181,489 175,666

OTHER LIABILITIES

218 207

COMMITMENTS AND
CONTINGENCIES

STOCKHOLDERS EQUITY

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Preferred Stock, \$.01 par value;
40,000,000 shares authorized; none
issued

Common Stock, \$.01 par value;
175,000,000 shares authorized; and
102,063,950 issued and outstanding
at March 31, 2002 and
December 31, 2001
1,021 1,021

Additional paid-in capital
512,586 519,316

Accumulated deficit
(219,839) (225,215)

Total Stockholders' Equity
293,768 295,122

TOTAL LIABILITIES AND
STOCKHOLDERS' EQUITY
\$475,475 \$470,995

See Notes to Consolidated Financial Statements

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ENDO PHARMACEUTICALS HOLDINGS INC.
CONSOLIDATED STATEMENTS OF OPERATIONS (UNAUDITED)
(In thousands, except per share data)

	Three Months Ended March 31,	
	2002	2001
NET SALES	\$ 67,026	\$ 39,382
COST OF SALES		
18,891 12,649		
GROSS PROFIT		
48,135 26,733		
COSTS AND EXPENSES:		
Selling, general and administrative		
23,583 15,890		
Research and development		
13,396 9,174		
Depreciation and amortization		
785 12,399		
OPERATING INCOME (LOSS)		
10,371 (10,730)		
INTEREST EXPENSE, Net of interest income of \$307 and \$1,089, respectively		
1,622 3,540		

INCOME (LOSS) BEFORE INCOME TAX
(BENEFIT)

8,749 (14,270)

INCOME TAX (BENEFIT)

3,373 (32)

NET INCOME (LOSS)

\$5,376 \$(14,238)

NET INCOME (LOSS) PER SHARE:

Basic

\$.05 \$(.16)

Diluted

\$.05 \$(.16)

NET INCOME (LOSS) PRO FORMA TO
EXCLUDE AMORTIZATION OF
GOODWILL AND WORKFORCE-IN-PLACE:

\$5,376 \$(1,572)

NET INCOME (LOSS) PER SHARE PRO
FORMA TO EXCLUDE AMORTIZATION OF
GOODWILL AND WORKFORCE-IN-PLACE:

Basic

\$.05 \$(.02)

Diluted
\$.05 \$(.02)

WEIGHTED AVERAGE SHARES:

Basic
102,064 89,139
Diluted
102,281 89,139

See Notes to Consolidated Financial Statements.

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ENDO PHARMACEUTICALS HOLDINGS INC.
CONSOLIDATED STATEMENTS OF CASH FLOWS (UNAUDITED)
(In thousands)

	Three Months Ended March 31,	
	2002	2001
OPERATING ACTIVITIES:		
Net Income (Loss)	\$5,376	\$(14,238)
Adjustments to reconcile net income (loss) to net cash provided by operating activities:		
Depreciation and amortization	785	12,399
Amortization of deferred financing costs	80	547
Accretion of promissory notes	1,708	1,178
Deferred income taxes	3,120	(35)
Changes in assets and liabilities which provided (used) cash:		
Accounts receivable	15,376	26,715
Inventories	(2,799)	(174)
Other assets	1,378	541
Accounts payable	(2,110)	864
Accrued expenses	9,516	(5,463)
Income taxes payable	(3,293)	(2,549)
Other liabilities	11	5,441
Net cash provided by operating activities	29,148	25,226

INVESTING ACTIVITIES:

Purchase of property and equipment
(80) (1,240)

Net cash used in investing activities
(80) (1,240)

FINANCING ACTIVITIES:

Repayments of long-term debt
(4,127)
Repurchase of Class A Transferable
and Class B Non-Transferable
Warrants
(6,730)

Net cash used in financing activities
(6,730) (4,127)

NET INCREASE IN CASH AND
CASH EQUIVALENTS
22,338 19,859
CASH AND CASH
EQUIVALENTS, BEGINNING OF
PERIOD
95,357 59,196

CASH AND CASH
EQUIVALENTS, END OF PERIOD
\$117,695 \$79,055

SUPPLEMENTAL
INFORMATION:

Interest Paid
\$20 \$2,861
Income Taxes Paid
\$3,293 \$2,786

See Notes to Consolidated Financial Statements.

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**ENDO PHARMACEUTICALS HOLDINGS INC.
NOTES TO CONSOLIDATED FINANCIAL STATEMENTS
(UNAUDITED)
FOR THE THREE MONTHS ENDED MARCH 31, 2002**

1. CONSOLIDATED FINANCIAL STATEMENTS

In the opinion of management, the accompanying condensed consolidated financial statements of Endo Pharmaceuticals Holdings Inc. (the Company or we) and its subsidiaries, which are unaudited, include all normal and recurring adjustments necessary to present fairly the Company's financial position as of March 31, 2002 and the results of operations and cash flows for the periods presented. The accompanying consolidated balance sheet as of December 31, 2001 is derived from the Company's audited financial statements. Certain information and footnote disclosure normally included in financial statements prepared in accordance with accounting principles generally accepted in the United States of America have been condensed or omitted as promulgated by APB Opinion No. 28 and Rule 10.01 of Regulation S-X. These condensed consolidated financial statements should be read in conjunction with the consolidated financial statements and notes thereto as of and for the year ended December 31, 2001 contained in the Company's Annual Report on Form 10-K. Certain reclassifications have been made to the prior period's financial statements to conform with the classifications used in 2002.

2. RECENT ACCOUNTING PRONOUNCEMENTS

In June 1998, the FASB issued SFAS No. 133, Accounting for Derivative Instruments and Hedging Activities, which was effective for all fiscal years beginning after June 15, 2000. SFAS 133, as amended by SFAS 137 and SFAS 138, establishes accounting and reporting standards for derivative instruments, including certain derivative instruments embedded in other contracts and for hedging activities. All derivatives, whether designated in hedging relationships or not, are required to be recorded on the balance sheet at fair value. If the derivative is designated in a fair value hedge, the changes in the fair value of the derivative and the hedged item are recognized in earnings. If the derivative is designated as a cash flow hedge, changes in the fair value of the derivative are recorded in other comprehensive income (OCI) and are recognized in the income statement when the hedged item affects earnings. SFAS 133 defines new requirements for designation and documentation of hedging relationships as well as ongoing effectiveness assessments in order to use hedge accounting. A derivative that does not qualify as a hedge is marked to fair value through earnings.

At January 1, 2001, we recorded \$228,000 as an accumulated transition adjustment as a reduction to earnings.

In December 1999, the SEC issued SAB 101, entitled Revenue Recognition in Financial Statements, as amended, effective as of October 1, 2000, which summarizes the SEC's views in applying generally accepted accounting principles to revenue recognition. The adoption of this guideline had no effect on our financial statements.

In March 2000, the FASB issued Financial Accounting Series Interpretation No. 44 entitled Accounting for Certain Transactions Involving Stock Compensation, which provides clarification to Accounting Principles Board Opinion No. 25 (APB No. 25), Accounting for Stock Issued to Employees. The adoption of this interpretation had no effect on our financial statements.

In June 2001, the FASB issued SFAS No. 141, Business Combinations, and SFAS No. 142, Goodwill and Other Intangible Assets. SFAS No. 141 was effective for all business combinations completed after June 30, 2001. SFAS No. 142 is effective for fiscal years beginning after December 15, 2001. SFAS No. 141 requires that all business combinations be accounted for under the purchase method only and that certain acquired intangible assets in a business combination be recognized as assets apart from goodwill. SFAS No. 142 establishes revised reporting requirements for goodwill and other intangible assets. See note 3 to the consolidated financial statements.

3. GOODWILL AND OTHER INTANGIBLES

Effective January 1, 2002, we adopted the provisions of SFAS No. 142, Goodwill and Other Intangible Assets.

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Goodwill and other intangible assets consist of the following (in thousands):

	March 31, 2002	December 31, 2001
Goodwill	\$ 181,079	\$ 182,318
Amortizable Intangibles:		
Licenses		
\$11,000	\$11,000	
Patents		
3,200	3,200	
<hr/>		
<hr/>		
14,200	14,200	
Less accumulated amortization		
(1,889)	(1,705)	
<hr/>		
<hr/>		
Other Intangibles, net		
\$12,311	\$12,495	
<hr/>		
<hr/>		

We have one reportable segment, pharmaceutical products. Goodwill arose as a result of the August 26, 1997 acquisition of certain branded and generic pharmaceutical products, related rights and certain assets of DuPont Pharmaceuticals Company (DuPont , formerly The DuPont Merck Pharmaceutical Company, DuPont Merck Pharma and Endo Laboratories, L.L.C.) and the July 17, 2000 acquisition of Algos Pharmaceutical Corporation (Algos). Although goodwill arose in two separate transactions, the components of our operating segment have been integrated and are managed as one reporting unit. Our components extensively share assets and other resources with the other components of our business.. In addition, our components do not maintain discrete financial information. Accordingly, the components of our business have been aggregated into one reporting unit and will be evaluated as such for goodwill impairment. Goodwill will be evaluated for impairment on an annual basis on January 1st of each year unless events or circumstances indicate that an impairment has occurred between annual dates. Goodwill has been evaluated for impairment upon the adoption of SFAS No. 142 and no impairment has been identified.

Effective January 1, 2002, the carrying amount of workforce-in-place was reclassified as goodwill. The cost of license fees is capitalized and is being amortized on a straight-line basis over their estimated useful life of twenty years. The cost of acquired patents is capitalized and is being amortized on a straight-line basis over their estimated useful life of seventeen years.

The pro forma effect of the adoption of SFAS No. 141 and SFAS No. 142 is as follows:

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(Unaudited)
Three Months Ended
March 31,

	2002	2001
	(in thousands, except per share data)	
Reported net income (loss)	\$5,376	\$(14,238)
Add back: Goodwill amortization		
10,225		
Add back: Amortization of workforce-in-place		
1,487		
Pro forma income tax benefit		
954		

Adjusted net income (loss)
\$5,376 \$(1,572)

Basic earnings (loss) per share:

Reported net income (loss)	\$.05	\$(-.16)
Add back: Goodwill amortization		
.11		
Add back: Amortization of workforce-in-place		
.02		
Pro forma income tax benefit		
.01		

Adjusted net income (loss)
\$.05 \$(-.02)

Diluted earnings (loss) per share:

Reported net income (loss)

\$.05 \$(.16)

Add back: Goodwill
amortization

.11

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	(Unaudited) Three Months Ended March 31,	
	2002	2001
Add back: Amortization of workforce-in-place		
.02		
Pro forma income tax benefit		
.01		
<hr/>		
<hr/>		
Adjusted net income (loss)		
\$.05 \$(.02)		
<hr/>		
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(in thousands, except per share data)

Estimated amortization of intangibles for the five fiscal years subsequent to December 31, 2001 is as follows (in thousands):

2002	\$741
2003	
741	
2004	
741	
2005	
741	
2006	
741	

4. COMPENSATION RELATED TO STOCK OPTIONS**Endo Pharma LLC 1997 Executive and Employee Stock Option Plans**

On November 25, 1997, the Company established the 1997 Employee Stock Option Plan and the 1997 Executive Stock Option Plan (collectively, the 1997 Stock Option Plans). Pursuant to the Recapitalization of the Company on July 17, 2000, the 1997 Stock Option Plans were amended and restated. The Endo Pharma LLC Amended and Restated 1997 Employee Stock Option Plan and the Endo Pharma LLC Amended and Restated 1997 Executive Stock Option Plan (collectively, the Endo Pharma LLC 1997 Stock Option Plans) reserve an aggregate of 25,615,339 shares of Common Stock of the Company held by Endo Pharma LLC for issuance. Stock options granted under the Endo Pharma LLC 1997 Stock Option Plans expire no later than December 31, 2012 unless an initial public offering of the Company Common Stock held by Endo Pharma LLC occurs, in which case the stock options granted will expire on August 26, 2007. The effect of the Recapitalization has been reflected in the accompanying financial statements. Subsequent to the Merger, the exercise of stock options pursuant to the Endo Pharma LLC 1997 Stock Option Plans does not result in the issuance of additional shares in the Company.

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The Class C stock options vest in four discrete tranches contingent upon (i) the Common Stock of the Company exceeding a defined closing price threshold for ninety consecutive trading days, (ii) the closing price of the Common Stock of the Company on the last trading day of such ninety consecutive trading day period being greater than or equal to 85% of the defined closing price and (iii) the holder being a director, officer or employee of the Company or any of its subsidiaries on such date. The defined closing price thresholds are as follows:

Option Class	MorphiDex® is Approved On or Prior to December 31, 2002	MorphiDex® is Not Approved On or Prior to December 31, 2002
	Common Stock Closing Price Threshold	Common Stock Closing Price Threshold
C1A and C1B	\$6.06	\$4.28
C2 \$9.38 \$6.62		
C3 \$14.99 \$10.58		
C4 \$24.50 \$17.29		

If these share price targets are achieved resulting in the vesting of each tranche of options, the Company will record up to four non-cash compensation charges related to the vesting of certain of the options. Under performance-based options, the measurement of expense is calculated and recorded as a non-cash charge at the time performance is achieved as the difference between the market price of the stock and the exercise price of the options.

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If these charges are recorded by the Company in connection with the above options, they will be significant. They will, however, not result in the issuance of additional shares of Company Common Stock. The aforementioned conditions have been achieved for the Class C1A, Class C1B and Class C2 stock options and therefore these stock options have vested. Accordingly, a non-cash compensation charge of \$15.3 million was recorded in the fourth quarter of 2000 for the vesting of the Class C1A and Class C1B stock options and a non-cash compensation charge of \$37.3 million was recorded in the third quarter of 2001 for the vesting of the Class C2 stock options.

The Class C1A, C1B, C2, C3 and C4 stock options are generally exercisable, if vested, upon the earlier of (i) the occurrence of a sale, disposition or transfer (Transfer) of Common Stock, after which Kelso no longer owns any shares of Common Stock or (ii) January 1, 2006.

Endo Pharma LLC 2000 Supplemental Executive and Employee Stock Option Plans

Pursuant to the Merger and Recapitalization of the Company on July 17, 2000, the Endo Pharma LLC 2000 Supplemental Employee Stock Option Plan and the Endo Pharma LLC 2000 Supplemental Executive Stock Option Plan (collectively, the Endo Pharma LLC 2000 Supplemental Stock Option Plans) were established. The Endo Pharma LLC 2000 Supplemental Stock Option Plans reserve an aggregate of 10,672,314 shares of Common Stock of the Company held by Endo Pharma LLC for issuance. The Endo Pharma LLC 2000 Supplemental Stock Option Plans are only effective on January 1, 2003 in the event that we have not received the approval from the U.S. Food and Drug Administration for Morphidex® for the treatment of pain by December 31, 2002. Stock options granted under the Endo Pharma LLC 2000 Supplemental Stock Option Plans expire no later than December 31, 2012 unless an initial public offering of the Company Common Stock held by Endo Pharma LLC occurs, in which case the stock options granted will expire on August 26, 2007. The exercise of stock options pursuant to the Endo Pharma LLC 2000 Supplemental Stock Option Plans does not result in the issuance of additional shares in the Company, however, may result in additional non-cash compensation charges upon issuance and/or attainment of defined common stock price targets. These charges may be substantial. The Endo Pharma LLC 2000 Supplemental Stock Option Plans are not currently effective, therefore no options have been granted.

5. WARRANTS

Class A Transferable Warrants and Class B Non-Transferable Warrants

The Class A Transferable Warrants and Class B Non-Transferable Warrants are exercisable at an exercise price of \$.01 per share into a specified number of shares of Common Stock depending on the timing of the FDA's approval of Morphidex® for one or more pain indications. These warrants become exercisable on the fifth business day following the date on which we receive approval from the FDA with respect to Morphidex® for the treatment of one or more pain indications. These warrants will remain exercisable for a period of six months after the exercisability date, at which time they will expire. If the FDA does not approve Morphidex® by March 31, 2003, each of these warrants expires without any payment therefor.

If the FDA approves Morphidex® on or prior to September 30, 2002, then upon exercise of these warrants, each warrant will be exercisable into 0.633803 shares of Common Stock. If the FDA approves Morphidex® after September 30, 2002 and prior to March 31, 2003, then upon exercise of these warrants, each warrant will be exercisable into 0.263158 shares of Common Stock. If the FDA does not approve Morphidex® before March 31, 2003, each of these warrants becomes void and all rights in respect of these warrants will cease.

On December 5, 2001, we commenced a tender offer to purchase up to 13,500,000 of our outstanding Class A Transferable Warrants and any and all of our outstanding Class B Non-Transferable Warrants. This tender offer expired at midnight on January 25, 2002. As of December 31, 2001, there were outstanding 17,810,526 of these warrants. We accepted an aggregate of 8,585,262 Class A Transferable Warrants and Class B Non-Transferable Warrants for payment at a purchase price of \$0.75 per warrant. We used cash on hand to finance the purchase of the tendered warrants. Following the purchase by us, there were outstanding 9,225,264 of these warrants.

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Pre-Merger Endo Warrants

The Pre-Merger Endo Warrants are exercisable at an exercise price of \$.01 per share into a specified number of shares of Common Stock if the FDA does not approve MorphiDex® for any pain indication prior to December 31, 2002. As of March 31, 2002, there were outstanding 71,328,424 of these warrants. If the FDA does not approve MorphiDex® before December 31, 2002, then these warrants become exercisable and upon exercise, each warrant will be exercisable into 0.416667 shares of Common Stock for a total of 29,720,177 shares of Common Stock.

6. RELATED PARTY TRANSACTIONS

On July 14, 2000, Endo Pharma LLC was formed to ensure that the stock options granted pursuant to the 1997 Employee Stock Option Plan and the 1997 Executive Stock Option Plan (collectively, as amended and restated, the Endo Pharma LLC 1997 Stock Option Plans) diluted only the pre-Merger holders of Endo Common Stock (see Note 4). Subsequent to the Merger, only currently outstanding shares of Common Stock of the Company held by Endo Pharma LLC will be issued upon the exercise of these stock options. Because Endo Pharma LLC, and not the Company, will provide the shares issued upon the exercise of the options, the Company has entered into a tax sharing agreement with Endo Pharma LLC under which the Company will pay to Endo Pharma LLC the amount of the tax benefits it receives as a result of the exercise of these stock options into shares of Common Stock held by Endo Pharma LLC for the years in which these tax benefits arise. As of March 31, 2002, no payments have been made or accrued.

7. COMMITMENTS AND CONTINGENCIES

We have entered into employment agreements with certain members of management.

We have entered into certain collaboration agreements with third parties for the development of pain management products. These agreements require us to share in the development costs of such products and grant marketing rights to us for such products. If our third party partners are unable to fund their portion of the collaboration project with us, this may adversely affect our results of operations and cash flows in the foreseeable future.

We are, and may in the future be, subject to various claims or legal proceedings arising out of the normal course of business with respect to commercial matters, including product liabilities, patent infringement matters, governmental regulation and other actions. We cannot predict the timing or outcome of these claims or proceedings. Currently, the Company is not involved in any claim and/or legal proceeding with respect to which the amount of ultimate liability will, in the opinion of management, materially affect our financial position, results of operations or liquidity.

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Item 2. *Management's Discussion and Analysis of Financial Condition and Results of Operations.*

Except for the historical information contained in this Report, this Report, including the following discussion, contains forward-looking statements that involve risks and uncertainties.

Overview

We, through our wholly owned subsidiary, Endo Pharmaceuticals Inc., are engaged in the research, development, sales and marketing of branded and generic prescription pharmaceuticals used primarily for the treatment and management of pain. Branded products comprised approximately 76%, 67% and 58% of net sales for the years ended December 31, 2000, 2001 and the three months ended March 31, 2002. On August 26, 1997, an affiliate of Kelso & Company and the then members of management entered into an asset purchase agreement with the then DuPont Merck Pharmaceutical Company to acquire certain branded and generic pharmaceutical products and exclusive worldwide rights to a number of new chemical entities in the DuPont research and development pipeline from DuPont Merck through the newly-formed Endo Pharmaceuticals Inc.

On July 17, 2000, we completed our merger with Algos. In the merger, we issued to the former Algos stockholders, in the aggregate, 17,810,526 shares of our common stock and 17,810,526 warrants to purchase in the aggregate up to 20,575,507 additional shares of our common stock in certain circumstances as more fully described under note 5 to the consolidated financial statements in this report. In the merger, we also issued to our pre-merger stockholders, in the aggregate, 71,328,424 warrants to purchase in the aggregate up to 29,720,177 additional shares of common stock in certain other circumstances as more fully described under note 5 to the consolidated financial statements in this report.

The stock of Endo Pharmaceuticals Inc. is our only asset and we have no other operations or business.

In May 2001, we entered into a long-term manufacturing and development agreement with Novartis Consumer Health, Inc., whereby Novartis has agreed to manufacture certain of our commercial products and products in development. We have incurred and expect to continue to incur significant costs associated with the preparation of Novartis' manufacturing operations under this agreement. These costs primarily relate to the preparation of test batches of drug product for FDA approval and our own quality assessment and administrative costs relating to the shifting of existing production to Novartis.

Our quarterly results have fluctuated in the past, and may continue to fluctuate. These fluctuations are primarily due to the timing of new product launches, purchasing patterns of our customers, market acceptance of our products and the impact of competitive products and pricing.

Critical Accounting Policies

To understand our financial statements, it is important to understand our accounting policies. The preparation of our financial statements in conformity with accounting principles generally accepted in the United States (generally accepted accounting principles) requires us to make estimates and assumptions that affect the reported amounts of assets and liabilities and disclosure of contingent assets and liabilities at the date of the financial statements and the reported amounts of revenues and expenses during the reporting period. Some of these judgments can be subjective and complex, and, consequently, actual results may differ from these estimates. For any given individual estimate or assumption made by us, there may also be other estimates or assumptions that are reasonable. We believe, however, that given current facts and circumstances, it is unlikely that applying any such other reasonable judgment would cause a material adverse effect on our consolidated results of operations, financial position or cash flows for the periods represented in this Report.

Our most critical accounting policies include the determination of sales deductions for estimated chargebacks, rebates, sales incentives and allowances, royalties and returns and losses, the utilization of deferred tax assets and the assessment of impairment of goodwill and other intangible assets. Note 2 to our consolidated financial statements contained in our annual report on Form 10K describes our significant accounting policies.

Table of Contents**Results of Operations***Goodwill and Other Intangibles*

Effective January 1, 2002, we adopted the provisions of SFAS No. 142, Goodwill and Other Intangible Assets.

Goodwill represents a significant portion of our assets and stockholders' equity. As of March 31, 2002, goodwill comprised approximately 38% of our total assets and 62% of our stockholders' equity. We assess the potential impairment of goodwill by comparing the fair value of goodwill to its carrying value for our one reporting unit. An impairment loss would be recognized when the estimated fair value is less than its carrying amount. As a result of the significance of goodwill, our results of operations and financial position in a future period could be negatively impacted should an impairment of goodwill occur.

We have one reportable segment, pharmaceutical products. Goodwill arose as a result of the August 26, 1997 acquisition of certain branded and generic pharmaceutical products, related rights and certain assets of DuPont Pharmaceuticals Company (DuPont , formerly The DuPont Merck Pharmaceutical Company, DuPont Merck Pharma and Endo Laboratories, L.L.C.) and the July 17, 2000 acquisition of Algos Pharmaceutical Corporation (Algos). Although goodwill arose in two separate transactions, the components of our operating segment have been integrated and are managed as one reporting unit. Our components extensively share assets and other resources with the other components of our business and have similar economic characteristics. In addition, our components do not maintain discrete financial information. Accordingly, the components of our business have been aggregated into one reporting unit and will be evaluated as such for goodwill impairment. Goodwill will be evaluated for impairment on an annual basis on January 1st of each year unless events or circumstances indicate that an impairment has occurred between annual dates. Goodwill has been evaluated for impairment upon the adoption of SFAS No. 142 on January 1, 2002 and, based on the fair value of our reporting unit, no impairment has been identified.

Goodwill and other intangible assets consist of the following (in thousands):

	March 31, 2002	December 31, 2001
Goodwill	\$ 181,079	\$ 182,318
Amortizable Intangibles:		
Licenses	\$11,000	\$11,000
Patents	3,200	3,200
	14,200	14,200
Less accumulated amortization	(1,889)	(1,705)
	12,311	12,495
Other Intangibles, net	\$12,311	\$12,495

Effective January 1, 2002, the carrying amount of workforce-in-place was reclassified as goodwill. The cost of license fees is capitalized and is being amortized on a straight-line basis over their estimated useful life of twenty years. The cost of acquired patents is capitalized and is being amortized on a straight-line basis over their estimated useful life of seventeen years.

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The pro forma effect of the adoption of SFAS No. 141 and SFAS No. 142 is as follows:

	(Unaudited) Three Months Ended March 31,	
	2002	2001
	(in thousands except per share data)	
Reported net income (loss)	\$ 5,376	\$ (14,238)
Add back: Goodwill amortization		
10,225		
Add back: Amortization of workforce-in-place		
1,487		
Pro forma income tax benefit		
954		
<hr/>		
<hr/>		
Adjusted net income (loss)	\$5,376	\$(1,572)
<hr/>		
<hr/>		
Basic earnings (loss) per share:		
Reported net income (loss)		
\$.05	\$ (.16)	
Add back: Goodwill amortization		
.11		
Add back: Amortization of workforce-in-place		
.02		
Pro forma income tax benefit		
.01		
<hr/>		
<hr/>		
Adjusted net income (loss)	\$.05	\$ (.02)

Diluted earnings (loss) per share:

Reported net income (loss)
 \$.05 \$(.16)
 Add back: Goodwill
 amortization
 .11
 Add back: Amortization of
 workforce-in-place
 .02
 Pro forma income tax benefit
 .01

Adjusted net income (loss)
 \$.05 \$(.02)

Estimated amortization of intangibles for the five fiscal years subsequent to December 31, 2001 is as follows (in thousands):

2002	\$ 741
2003	741
2004	741
2005	741
2006	741

Compensation Related to Stock Options

During our fourth quarter ended December 31, 2000 we incurred a non-cash charge of \$15.3 million, and during our third quarter ended September 30, 2001, we recorded a non-cash charge of \$37.3 million, in each case for stock-based compensation relating to the vesting of options that were issued under the Endo Pharma LLC stock option plans. Under these plans, tranches of options vest when we attain certain stock price targets. As each tranche vests, we incur a non-cash charge representing the difference between the market price of the shares underlying the options and the exercise price of such options. We may in the future incur up to two additional charges in relation to the Endo Pharma LLC options as a result of the attainment of these common stock price targets. These charges may be substantial. These options are exercisable into shares of common stock that are presently held by Endo Pharma LLC. As a result, the exercise of these options will not result in

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the issuance of additional shares of common stock.

In connection with the Algos merger and our related recapitalization on July 17, 2000, the Endo Pharma LLC 2000 Supplemental Employee Stock Option Plan and the Endo Pharma LLC 2000 Supplemental Executive Stock Option Plan (collectively, the Endo Pharma LLC 2000 Supplemental Stock Option Plans) were established. The Endo Pharma LLC 2000 Supplemental Stock Option Plans reserve an aggregate of 10,672,314 shares of our common stock that is held by Endo Pharma LLC for issuance.

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The Endo Pharma LLC 2000 Supplemental Stock Option Plans are only effective on January 1, 2003 in the event that we have not received the approval from the FDA of MorphiDex® for the treatment of pain by December 31, 2002. The exercise of stock options pursuant to the Endo Pharma LLC 2000 Supplemental Stock Option Plans does not result in the issuance of additional shares in the Company, however, may result in additional non-cash compensation charges upon issuance and/or attainment of defined common stock price targets. These charges may be substantial. The Endo Pharma LLC 2000 Supplemental Stock Option Plans are not currently effective, therefore no options have been granted.

All the options we have granted pursuant to the Endo Pharmaceuticals Holdings Inc. 2000 Stock Incentive Plan have exercise prices equal to the market price of our stock on the date granted and, under generally accepted accounting principles, a measurement date had occurred on the date of grant. Consequently, we do not expect to incur a charge upon the vesting or exercise of those options.

Net Sales

Our net sales consist of revenues from sales of our pharmaceutical products, less estimates for certain chargebacks, rebates, sales incentives and allowances, royalties and the cost of returns and losses. We estimate the accrual for sales deductions based on historical data, estimated future trends and other competitive factors. Net sales are recognized when products are shipped.

The following table presents our unaudited net sales by product category for the three months ended March 31, 2002 and 2001.

	Three Months Ended March 31,	
	2002	2001
	(in thousands, unaudited)	
Percocet®	\$ 23,468	\$ 20,159
Lidoderm®		
10,002	3,643	
Other brands		
5,326	5,002	
<hr/>		
<hr/>		
Total brands		
38,796	28,804	
Total generics		
28,230	10,578	
<hr/>		
<hr/>		
Total net sales		
\$67,026	\$39,382	
<hr/>		
<hr/>		

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The following table presents our unaudited net sales as a percentage of total net sales for select products for the three months ended March 31, 2002 and 2001.

	Three Months Ended March 31,	
	2002	2001
	(unaudited)	
Percocet®	35%	51%
Lidoderm®		
15 9		
Other brands		
8 13		

Total brands		
58 73		
Total generics		
42 27		

Total net sales		
100% 100%		

Three Months Ended March 31, 2002 Compared to the Three Months Ended March 31, 2001

Net sales for the three months ended March 31, 2002 increased by 70% to \$67.0 million from \$39.4 million in the comparable 2001 period. This increase in net sales was primarily due to the increase in net sales of Lidoderm®, the first FDA-approved product for the treatment of the pain of post-herpetic neuralgia, certain generic products and the new strengths of Percocet®.

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In September 1999, we launched Lidoderm®, which continues to gain market share due to our ongoing promotional and educational efforts. Net sales of Lidoderm® increased to \$10.0 million from \$3.6 million in the comparable 2001 period. Percocet® net sales increased 16% to \$23.5 million from \$20.2 million in the comparable 2001 period due to the new strengths of Percocet® launched in November 2001. In April 2001, generic equivalents of Percocet® 7.5/500 and Percocet® 10.0/650 were introduced. In November 2001, we launched Percocet® 7.5/325 and Percocet® 10.0/325 which do not currently have generic equivalents. Generic products increased to \$28.2 million from \$10.6 million in the comparable 2001 period primarily due to the growth of our generic morphine sulfate extended release tablets and Endocet®. In November 1998, we launched the 15mg, 30mg and 60mg strengths, in May 2001, we launched the 100mg strength and in September 2001, we launched the 200mg strength of our generic morphine sulfate extended release tablets. These products continue to gain market share. In April 2001, we launched two new strengths of our generic product Endocet®. Generic competition with our products may have a material impact on our results of operations and cash flows in the future.

Gross profit for the three months ended March 31, 2002 increased by 80% to \$48.1 million from \$26.7 million in the comparable 2001 period. Gross profit margins increased to 72% from 68% due to a more favorable mix of higher margin brand and generic products resulting from the product launches discussed above, and the discontinuation of some lower margin non-core products. In addition, the increase in gross profit margins was also due to the existing fixed cost nature of our manufacturing relationship with Bristol-Myers Squibb Pharma Company (formerly DuPont Pharmaceuticals), currently our most significant contract manufacturing relationship. If we achieve our forecast for revenue and product mix, we expect the increase in gross profits and gross profit margins to continue.

Selling, general and administrative expenses for the three months ended March 31, 2002 increased by 48% to \$23.6 million from \$15.9 million in the comparable 2001 period. This increase was due to a \$4.2 million increase in sales and promotional efforts in 2002 over the comparable 2001 period to support Lidoderm® and Percocet®. In addition, we experienced an increase in personnel-related costs in the general and administrative functions in order to support our new product marketing and new product development.

Research and development expenses for the three months ended March 31, 2002 increased by 46% to \$13.4 million from \$9.2 million in the comparable 2001 period. This increase was due to our increased spending on new products under development that are focused in pain management. We are currently conducting Phase III clinical trials on MorphiDex® and on both an oral extended-release and oral immediate-release version of oxymorphone.

Depreciation and amortization for the three months ended March 31, 2002 decreased to \$.8 million from \$12.4 million in the comparable 2001 period. Effective January 1, 2002, we have adopted the provisions of SFAS 142, Goodwill and Other Intangible Assets and will no longer amortize goodwill unless evidence of an impairment exists. If SFAS 142 had been adopted as of January 1, 2001, depreciation and amortization for the three months ended March 31, 2001 would have been \$.7 million.

Interest expense, net for the three months ended March 31, 2002 decreased by 54% to \$1.6 million from \$3.5 million in the comparable 2001 period. This decrease is substantially due to the repayment on October 29, 2001 of the term loans outstanding on our credit facility. Interest expense for the three months ended March 31, 2002 substantially represents the accretion of promissory notes issued to Bristol-Myers Squibb which bear no interest and therefore have been discounted in the accompanying financial statements. For the three months ended March 31, 2001, due to the adoption of SFAS 133 on January 1, 2001, the Company recorded a \$.2 million charge for the accumulated transition adjustment relating to derivative instruments that do not qualify as a hedge under SFAS 133.

Income tax (benefit) for the three months ended March 31, 2002 increase to \$3.4 million from nil in the comparable 2001 period. This increase is due to the increase in income before income tax for the three months ended March 31, 2002. For the three months ended March 31, 2001, we recorded a valuation allowance on our existing deferred tax assets due to the uncertainty of the utilization of such amounts in the foreseeable future. During the fourth quarter of 2001, we evaluated our anticipated future taxable income based upon the repayment of our outstanding term loans, new product approvals and other existing and estimated future product performance and determined that it is more likely than not that we will utilize our deferred tax benefits. Accordingly, we reversed our valuation reserves that had been recorded against those deferred tax assets.

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Liquidity and Capital Resources

Net cash provided by operating activities increased by \$3.9 million to \$29.1 million for the three months ended March 31, 2002 from \$25.2 million for the three months ended March 31, 2001. This increase was due to the cash provided by the increase in net sales and gross profit for the three months ended March 31, 2002 compared to the three months ended March 31, 2001 offset by an increase in selling, general and administrative expenses and research and development expenses for the three months ended March 31, 2002 as compared to the three months ended March 31, 2001.

Net cash utilized in investing activities decreased by \$1.1 million to \$.1 million for the three months ended March 31, 2002 from \$1.2 million for the three months ended March 31, 2001 due to the purchase in 2001 of leasehold improvements and other furniture and fixtures related to our principal executive offices and the implementation of an electronic document management system during 2001.

Net cash utilized in financing activities increased by \$2.6 million to \$6.7 million for the three months ended March 31, 2002 from \$4.1 million for the three months ended March 31, 2001. During the three months ended March 31, 2002, we utilized \$6.7 million of cash, including fees, to repurchase 8.6 million Class A Transferable Warrants and Class B Non-Transferable Warrants. During the three months ended March 31, 2001, we made a scheduled principal payment on our term loans which were repaid in full on October 29, 2001.

In its annual report filed on Form 10K for the year ended December 31, 2001, Penwest Pharmaceuticals Co., a collaboration partner of Endo with which Endo has an alliance agreement and with which Endo is developing one of its pipeline projects, stated that its existing capital resources, will enable Penwest to maintain currently planned operations at least through March 31, 2003. If Penwest is unable to fund their portion of the collaboration project with Endo, this may adversely affect our results of operations and cash flows in the foreseeable future.

Our cash and cash equivalents totaled \$117.7 million at March 31, 2002. We believe that our (a) cash and cash equivalents, (b) cash flow from operations and (c) our credit facility (which has an available unused line of credit of \$75 million) will be sufficient to meet our normal operating, investing and financing requirements in the foreseeable future, including the funding of our pipeline projects in the event that our collaboration partners are unable to fund their portion of any particular project. We may use a portion of our cash and cash equivalents to repay all or a portion of the notes that we have issued to Bristol-Myers Squibb Company (formerly DuPont Pharmaceuticals) or for possible acquisitions.

In December 2001, we amended and restated our senior secured credit facility with a number of lenders, including affiliates of certain of the underwriters of our recent public offering. This amended and restated credit facility provides for a line of credit of \$75.0 million and a delayed draw term loan of \$25.0 million. The line of credit and delayed draw term loan mature December 21, 2006. Any loans outstanding under the credit facility are secured by a first priority security interest in substantially all of our assets. The credit facility contains representations and warranties, covenants, events of default and other provisions customarily found in similar agreements.

Item 3. Quantitative and Qualitative Disclosures about Market Risk.

During the fourth quarter of 2001, we repaid the remaining outstanding balance of our variable rate term loans. Prior to the repayment of our variable rate term loans, our primary market risk exposure was to changes in interest rates (LIBOR) on our variable rate borrowings. As of March 31, 2002, we only have outstanding fixed rate borrowings. These fixed rate borrowings are substantially comprised of promissory notes payable to Bristol-Myers Squibb issued in consideration for manufacturing and supply services provided under the Manufacturing and Supply Agreement. The promissory notes issued annually over the initial five year term of the Manufacturing and Supply Agreement have an aggregate face value of \$23.0 million per year and are payable on August 26, 2002. The promissory notes bear no interest and therefore have been discounted in the accompanying financial statements using our borrowing rate for similar instruments at the time of borrowing. We also financed a portion of the purchase price of the Acquisition through the issuance of a promissory note to DuPont. The note has a face value of \$3.9 million and is payable on August 26, 2002. The promissory note bears no interest and therefore has been discounted in the

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accompanying financial statements using a rate which approximates our borrowing rate for similar instruments at the time of borrowing. On December 21, 2001, we entered into a new credit facility that provides for a line of credit of \$75.0 million and a delayed draw term loan of \$25.0 million. Borrowings under the new credit facility are variable rate borrowings. There are no amounts outstanding under the new credit facility. We do not utilize financial instruments for trading purposes and hold no derivative financial instruments that could expose us to significant market risk. We monitor interest rates and enter into interest rate agreements as considered appropriate.

PART II

OTHER INFORMATION

Item 1. *Legal Proceedings.*

On October 20, 2000, The Purdue Frederick Company and related companies (Purdue Frederick) filed suit against us and our subsidiary, Endo Pharmaceuticals Inc. (EPI), in the U.S. District Court for the Southern District of New York alleging that EPI's bioequivalent version of Purdue Frederick's OxyContin® (oxycodone hydrochloride extended-release tablets), 40mg strength, infringes three of its patents. This suit arose after EPI provided the plaintiffs with notice that its ANDA submission for a bioequivalent version of Purdue Frederick's OxyContin®, 40mg strength, challenged the listed patents for OxyContin® 40mg tablets. On March 13, 2001, Purdue Frederick filed a second suit against us and EPI in the U.S. District Court for the Southern District of New York alleging that EPI's bioequivalent versions of Purdue Frederick's OxyContin®, 10mg and 20mg strengths, infringe the same three patents. This suit arose from EPI having amended its earlier ANDA on February 9, 2001 to add bioequivalent versions of the 10mg and 20mg strengths of OxyContin®. On August 30, 2001, Purdue Frederick filed a third suit against us and EPI in the U.S. District Court for the Southern District of New York alleging that EPI's bioequivalent version of Purdue Frederick's OxyContin®, 80mg strength, infringes the same three patents. This suit arose from EPI having amended its earlier ANDA on July 30, 2001 to add the bioequivalent version of the 80mg strength of OxyContin®.

For each of the 10mg, 20mg, 40mg and 80mg strengths of this product, EPI made the required Paragraph IV certification against the patents listed in the FDA's Orange Book as covering these strengths of OxyContin®. EPI has pleaded counterclaims that the patents asserted by Purdue Frederick are invalid, unenforceable and/or not infringed by EPI's formulation of oxycodone hydrochloride extended-release tablets, 10mg, 20mg, 40mg and 80mg strengths. EPI has also counterclaimed for antitrust damages based on allegations that Purdue Frederick obtained the patents through fraud on the United States Patent and Trademark Office and is asserting them while aware of their invalidity and unenforceability. However, we cannot make any assurances as to the outcome of this patent challenge. Purdue Frederick was granted a preliminary injunction (Purdue Pharma L.P. v. Boehringer Ingelheim GmbH, 98 F. Supp. 2d 362 (SDNY 2000)), which decision was affirmed on appeal (Purdue Pharma L.P. v. Boehringer Ingelheim GmbH, 237 F.3d 1359 (Fed. Cir. 2001)), against a different manufacturer based on the same patents that are being asserted against us and EPI, and in the same court in which Purdue Frederick sued. We believe the defenses rejected in the preliminary injunction decision and in the appellate decision do not substantially impact the principal defenses raised by us and EPI.

On November 15, 2001, SmithKline Beecham Corporation (and related companies) filed suit against EPI in the U.S. District Court for the Eastern District of Pennsylvania alleging that EPI's bioequivalent version of SmithKline's Paxil®, 40 mg strength, infringes five of its patents. The FDA accepted EPI's ANDA submission for a bioequivalent version of SmithKline's Paxil®, 40 mg strength, earlier in 2001. In this ANDA, EPI made the required Paragraph IV certification against all of the SmithKline patents listed in the FDA's Orange Book as covering Paxil®. Paxil® is indicated for the treatment of depression, obsessive compulsive disorder and panic disorder. Although we believe the patents asserted by SmithKline Beecham are invalid and/or not infringed, no assurance can be given as to the outcome of this patent challenge process.

Litigation similar to that described above may also result from products we currently have in development, as well as those that we may develop in the future. We, however, cannot predict the timing or outcome of any such litigation, or whether any such litigation will be brought against us.

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On November 15, 2001, EPI was named, along with ten other pharmaceutical companies, as a defendant in a class action lawsuit filed by Bennie Toombs in the United States District Court for the Western District of Louisiana. According to the complaint, each of the defendant pharmaceutical companies had allegedly manufactured and sold products containing phenylpropanolamine (PPA). The complaint alleges that the defendants failed to adequately warn plaintiff of the hazards of the use of the subject products containing PPA and that as a result of this failure to warn, plaintiffs suffered injury. The action has been transferred by order of the United States Judicial Panel on Multidistrict Litigation to the Western District of Washington, where it has been consolidated for pretrial proceedings with other cases involving claims against manufacturers of PPA-containing products. EPI is not a party to any of these other actions and intends to vigorously defend itself in the *Toombs* litigation.

In addition to the above, the Company is involved in, or has been involved in, arbitrations or legal proceedings that arise from the normal course of its business. The Company cannot predict the timing or outcome of these claims and proceedings. Currently, the Company is not involved in any arbitration and/or legal proceeding that it expects to have a material effect on its business, financial condition or results of operations and cash flows.

Item 2. *Changes in Securities and Use of Proceeds.*

None.

Item 3. *Defaults Upon Senior Securities.*

None.

Item 4. *Submission of Matters to a Vote of Security Holders.*

None.

Item 5. *Other Information.*

None.

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Item 6. Exhibits and Reports on Form 8-K.

(a) *Exhibits.*

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

(b) *Reports on Form 8-K.*

We filed the following Form 8-Ks in the quarter ended March 31, 2002:

	<u>Dates</u>	<u>Items</u>
January 17, 2002 Items 5 & 7	January 8, 2002	Items 7 & 9
February 21, 2002 Items 7 & 9		
March 6, 2002 Items 7 & 9		

No financial statements were filed in connection with any such Form 8-K.

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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)

/s/ Carol A. Ammon

Name: Carol A. Ammon
Title: *Chairman and Chief Executive
Officer*

/s/ Jeffrey R. Black

Name: Jeffrey R. Black
Title: *Senior Vice President and Chief
Financial Officer*

Date: May 14, 2002

Table of Contents**Exhibit Index**

Exhibit No.	Title
2.1	Amended and Restated Agreement and Plan of Merger, dated as of March 3, 2000 (the Merger Agreement), by and among Endo Pharmaceuticals Holdings Inc. (Endo), Endo Inc. and Algos Pharmaceutical Corporation (Algos) (incorporated herein by reference to Exhibit 2.1 of the Registration Statement on Form S-4 of the Registrant (Registration No. 333-39040) (the Registration Statement), filed with the Securities and Exchange Commission (the Commission) on June 9, 2000)
2.2	Amendment, dated as of April 17, 2000, to the Merger Agreement, by and between Endo, Endo Inc. and Algos (incorporated herein by reference to Exhibit 2.2 of the Registration Statement filed with the Commission on June 9, 2000)
2.3	Asset Purchase Agreement, dated as of August 27, 1997, by and between Endo Pharmaceuticals Inc. (Endo Pharmaceuticals) and The DuPont Merck Pharmaceutical Company (DuPont Merck Pharmaceutical) (incorporated herein by reference to Exhibit 2.3 of the Registration Statement filed with the Commission on June 9, 2000)
3.1	Amended and Restated Certificate of Incorporation of Endo (incorporated herein by reference to Exhibit 3.1 of the Form 10-Q for the Quarter ended June 30, 2000 filed with the Commission on August 15, 2000)
3.2	Amended and

Restated By-laws of
Endo (incorporated
herein by reference
to Exhibit 3.2 of the
Form 10-Q for the
Quarter ended
June 30, 2000 filed
with the
Commission on
August 15,
2000) 4.1
Amended and
Restated Executive
Stockholders
Agreement, dated
as of July 14, 2000,
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, 2000 filed
with the
Commission on
August 15,
2000) 4.2
Amended and
Restated Employee
Stockholders
Agreement, dated
as of July 14, 2000,
by and among
Endo, Endo LLC,
KIA V, KEP V and
the Employee
Stockholders (as
defined therein)
(incorporated
herein by reference
to Exhibit 4.2 of the
Form 10-Q for the
Quarter ended
June 30, 2000 filed
with the
Commission on
August 15,
2000) 4.3 Form of
Stock Certificate of
Endo Common
Stock (incorporated
herein by reference

to Exhibit 4.3 of the Form 10-Q for the Quarter ended June 30, 2000 filed with the Commission on August 15, 2000) 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) 10.1 Endo Warrant Agreement, dated as of July 17, 2000, by and between Endo and United States Trust Company of New York (incorporated herein by reference to Exhibit 10.1 of the Form 10-Q for the Quarter ended June 30, 2000 filed with the Commission on August 15, 2000)

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Exhibit No.	Title
10.2	Algos Warrant Agreement, dated as of July 17, 2000, by and between Endo and United States Trust Company of New York (incorporated herein by reference to Exhibit 10.2 of the Form 10-Q for the Quarter ended June 30, 2000 filed with the Commission on August 15, 2000)
10.3	Form of Series A Warrant to Purchase Shares of Common Stock and Warrants of Endo (incorporated herein by reference to Exhibit 10.3 of the Registration Statement filed with the Commission on June 9, 2000)
10.4	Letter Agreement, dated as of November 26, 1999, by and among Algos, Endo, KIA V and KEP V (incorporated herein by reference to Exhibit 10.4 of the Registration Statement filed with the Commission on June 9, 2000)
10.5	Tax Sharing Agreement, dated as of July 17, 2000, by and among Endo, Endo Inc. and Endo LLC (incorporated herein by reference to Exhibit 10.5 of the Form 10-Q for the Quarter ended June 30, 2000 filed

with the
Commission on
August 15,
2000) 10.6
[Intentionally
Omitted.] 10.7
Amended and
Restated Credit
Agreement, dated
as of December 21,
2001, by and
between Endo,
Endo
Pharmaceuticals,
the Lenders Party
There to and
JPMorgan Chase
Bank (incorporated
by reference to
Exhibit 10.7 of the
Annual Report on
Form 10-K for the
Year Ended
December 31, 2001
filed with the
Commission on
March 29,
2002) 10.8
[Intentionally
Omitted.] 10.9
[Intentionally
Omitted.] 10.10
Sole and Exclusive
License Agreement,
dated as of
November 23,
1998, by and
between Endo
Pharmaceuticals
and Hind Health
Care, Inc.
(incorporated
herein by reference
to Exhibit 10.10 of
the Registration
Statement filed with
the Commission on
June 9,
2000) 10.11
Analgesic License
Agreement, dated
as of October 27,
1997, by and
among Endo
Pharmaceuticals,
Endo Laboratories,
LLC and DuPont
Merck
Pharmaceutical
(incorporated
herein by reference
to Exhibit 10.11 of

the Registration Statement filed with the Commission on June 9, 2000) 10.12 Anti-Epileptic License Agreement, dated as of October 27, 1997, by and among Endo Pharmaceuticals, Endo Laboratories, LLC and DuPont Merck Pharmaceutical (incorporated herein by reference to Exhibit 10.12 of the Registration Statement filed with the Commission on June 9, 2000) 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) 10.15 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)

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Exhibit No.	Title
10.17	Manufacture and Supply Agreement, dated as of August 26, 1997, by and among Endo Pharmaceuticals, DuPont Merck Pharmaceutical and DuPont Merck Pharma (n/k/a Bristol-Myers Squibb Pharma Company) (incorporated herein by reference to Exhibit 10.17 of the Registration Statement filed with the Commission on June 9, 2000)
10.18	Amended and Restated Strategic Alliance Agreement, dated as of April 2, 2002, by and between Endo Pharmaceuticals and Penwest Pharmaceuticals Co.**
10.19	Agreement, dated as of February 1, 2000, by and between Endo Pharmaceuticals and UPS Supply Chain Management, Inc. (f/k/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
Employment
Agreement, dated
as of July 17, 2000,
by and between
Endo and John W.
Lyle (incorporated
herein by reference
to Exhibit 10.26 of
the Form 10-Q for
the Quarter ended
June 30, 2000 filed
with the
Commission on
August 14,
2000) *10.27
Amended and
Restated
Employment
Agreement, dated
as of September 1,
2001, by and
between Endo
Pharmaceuticals
and Carol A.
Ammon

(incorporated herein by reference to Exhibit 10.27 of the Current Report on Form 8-K dated August 31, 2001) *10.28 Amended and Restated Employment Agreement, dated as of September 1, 2001, by and between Endo Pharmaceuticals and Jeffrey R. Black (incorporated herein by reference to Exhibit 10.28 of the Current Report on Form 8-K dated August 31, 2001)

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Exhibit No.	Title
<p>*10.29 Amended and Restated Employment Agreement, dated as of September 1, 2001, by and between Endo Pharmaceuticals and David Allen Harvey Lee, MD, Ph.D. (incorporated herein by reference to Exhibit 10.29 of the Current Report on Form 8-K dated August 31, 2001)</p> <p>*10.30 Amended and Restated Employment Agreement, dated as September 1, 2001, by and between Endo Pharmaceuticals and Mariann T. MacDonald (incorporated herein by reference to Exhibit 10.30 of the Current Report on Form 8-K dated August 31, 2001)</p> <p>10.31 Separation and Release Agreement, dated as of March 22, 2000, by and between Endo Pharmaceuticals, Endo and Osagie O. Imasogie (incorporated herein by reference to Exhibit 10.31 of the Registration Statement filed with the Commission on June 9, 2000)</p> <p>10.32 Separation and Release Agreement, dated as of April 20, 2000, by and between Endo</p>	

Pharmaceuticals,
Endo and Louis J.
Vollmer
(incorporated
herein by reference
to Exhibit 10.32 of
the Registration
Statement filed with
the Commission on
June 9,
2000) 10.33
Office Lease, dated
as of August 26,
1997, by and
between Endo
Pharmaceuticals
and Northstar
Development
Company
(incorporated
herein by reference
to Exhibit 10.33 of
the Registration
Statement filed with
the Commission on
June 9,
2000) 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) *10.35
Amended and
Restated
Employment
Agreement, dated
as of September 1,
2001, by and
between Endo and
Caroline B.
Manogue (formerly
Berry)
(incorporated
herein by reference
to Exhibit 10.35 of
the Current Report
on Form 8-K dated
August 31,
2001) *10.36
Amended and
Restated

Employment Agreement, dated as of September 1, 2001, by and between Endo and Peter A. Lankau (incorporated herein by reference to Exhibit 10.36 of the Current Report on Form 8-K dated August 31, 2001) 10.37

License Agreement, dated as of August 16, 1993, by and between Endo Pharmaceuticals (as successor in interest to Algos Pharmaceutical Corporation) and The Medical College of Virginia (incorporated herein by reference to Exhibit 10.4.1 of the registration statement on Form S-1 of Algos Pharmaceutical Corporation declared effective on September 25, 1996) 10.38

[Intentionally Omitted.] 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) 10.40

[Intentionally Omitted.]

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Exhibit No.	Title
10.41	Service Agreement, dated as of February 1, 2001, by and between Endo Pharmaceuticals and Ventiv Health U.S. Sales Inc. (incorporated herein by reference to Exhibit 10.41 of the Current Report on Form 8-K dated August 31, 2001)11 Statement Regarding Computation of per Share Earnings

* A management contract or compensatory plan or arrangement required to be filed as an Exhibit pursuant to Item 14(c) of Form 10-K.

** Confidential portions of this exhibit have been redacted and filed separately with the Commission pursuant to a confidential treatment request in accordance with Rule 406 of the Securities Act of 1933.