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PART I. FINANCIAL INFORMATION

ITEM 1. FINANCIAL STATEMENTS

QUESTCOR PHARMACEUTICALS, INC.

CONSOLIDATED BALANCE SHEETS

(In thousands)
(unaudited)

	June 30, 2011	December 31, 2010
ASSETS		
Current assets:		
Cash and cash equivalents	\$ 65,277	\$ 41,508
Short-term investments	63,849	73,324
Total cash, cash equivalents and short-term investments	129,126	114,832
Accounts receivable, net of allowances of \$22 and \$25 at June 30, 2011 and December 31, 2010, respectively	23,714	11,128
Inventories, net of allowances of \$160 and \$158 at June 30, 2011 and December 31, 2010, respectively	3,998	3,726
Prepaid income taxes	4,532	3,532
Prepaid expenses and other current assets	1,492	1,864
Deferred tax assets	8,237	8,417
Total current assets	171,099	143,499
Property and equipment, net	1,930	872
Purchased technology, net	2,927	3,074
Goodwill	—	299
Deposits and other assets	59	65
Deferred tax assets	4,184	4,184
Total assets	<u>\$ 180,199</u>	<u>\$ 151,993</u>
LIABILITIES AND SHAREHOLDERS' EQUITY		
Current liabilities:		
Accounts payable	\$ 2,780	\$ 3,869
Accrued compensation	5,270	4,158
Sales-related reserves	27,066	21,511
Other accrued liabilities	1,586	1,973
Total current liabilities	36,702	31,511
Lease termination, deferred rent and other non-current liabilities	192	355
Total liabilities	36,894	31,866
Shareholders' equity:		
Preferred stock, no par value, 7,500,000 shares authorized; none outstanding	—	—
Common stock, no par value, 105,000,000 shares authorized, 62,317,624 and 62,418,464 shares issued and outstanding at June 30, 2011 and December 31, 2010, respectively	72,887	74,809
Retained earnings	70,393	45,295
Accumulated other comprehensive income	25	23
Total shareholders' equity	143,305	120,127
Total liabilities and shareholders' equity	<u>\$ 180,199</u>	<u>\$ 151,993</u>

See accompanying notes.

QUESTCOR PHARMACEUTICALS, INC.
CONSOLIDATED STATEMENTS OF INCOME
(In thousands, except per share data)
(unaudited)

	Three Months Ended		Six Months Ended	
	June 30,		June 30,	
	2011	2010	2011	2010
Revenue				
Net sales	\$45,980	\$28,316	\$82,813	\$54,560
Cost of sales (exclusive of amortization of purchased technology)	<u>2,856</u>	<u>2,000</u>	<u>4,728</u>	<u>3,998</u>
Gross profit	43,124	26,316	78,085	50,562
Operating expenses:				
Selling and marketing	14,746	6,028	25,998	12,678
General and administrative	3,791	2,943	7,663	5,669
Research and development	3,891	2,943	6,872	5,690
Depreciation and amortization	273	130	471	255
Impairment of goodwill	—	—	299	—
Total operating expenses	<u>22,701</u>	<u>12,044</u>	<u>41,303</u>	<u>24,292</u>
Income from operations	20,423	14,272	36,782	26,270
Interest and other income, net	<u>120</u>	<u>119</u>	<u>384</u>	<u>215</u>
Income before income taxes	20,543	14,391	37,166	26,485
Income tax expense	<u>6,669</u>	<u>5,109</u>	<u>12,068</u>	<u>9,351</u>
Net income	<u>\$13,874</u>	<u>\$ 9,282</u>	<u>\$25,098</u>	<u>\$17,134</u>
Net income per share:				
Basic	<u>\$ 0.22</u>	<u>\$ 0.15</u>	<u>\$ 0.40</u>	<u>\$ 0.28</u>
Diluted	<u>\$ 0.21</u>	<u>\$ 0.14</u>	<u>\$ 0.38</u>	<u>\$ 0.27</u>
Shares used in computing net income per share:				
Basic	<u>62,034</u>	<u>62,022</u>	<u>62,126</u>	<u>61,957</u>
Diluted	<u>65,464</u>	<u>64,543</u>	<u>65,483</u>	<u>64,057</u>

See accompanying notes.

QUESTCOR PHARMACEUTICALS, INC.
CONSOLIDATED STATEMENTS OF CASH FLOWS
(In thousands)
(unaudited)

	Six Months Ended June 30,	
	2011	2010
OPERATING ACTIVITIES		
Net income	\$ 25,098	\$ 17,134
Adjustments to reconcile net income to net cash provided by operating activities:		
Share-based compensation expense	3,528	1,908
Deferred income taxes	180	41
Amortization of investments	376	329
Depreciation and amortization	471	255
Impairment of goodwill	299	—
Loss on disposal of property and equipment	11	—
Changes in operating assets and liabilities:		
Accounts receivable	(12,586)	2,904
Inventories	(272)	70
Prepaid income taxes	(1,000)	—
Prepaid expenses and other current assets	372	(4)
Accounts payable	(1,089)	(9,448)
Accrued compensation	1,112	665
Sales-related reserves	5,555	2,237
Income taxes payable	—	590
Other accrued liabilities	(387)	(265)
Other non-current liabilities	(163)	(171)
Net cash flows provided by operating activities	<u>21,505</u>	<u>16,245</u>
INVESTING ACTIVITIES		
Purchase of property and equipment	(1,393)	(208)
Purchase of short-term investments	(53,859)	(54,065)
Proceeds from maturities of short-term investments	62,960	14,880
Deposits and other assets	6	—
Net cash flows provided by / (used in) investing activities	<u>7,714</u>	<u>(39,393)</u>
FINANCING ACTIVITIES		
Income tax benefit realized from share-based compensation plans	3,735	320
Issuance of common stock, net	2,268	823
Repurchase of common stock	(11,453)	—
Net cash flows (used in) / provided by financing activities	<u>(5,450)</u>	<u>1,143</u>
Increase (decrease) in cash and cash equivalents	<u>23,769</u>	<u>(22,005)</u>
Cash and cash equivalents at beginning of period	41,508	45,829
Cash and cash equivalents at end of period	<u>\$ 65,277</u>	<u>\$ 23,824</u>
Supplemental Disclosures of Cash Flow Information:		
Cash paid for interest	<u>\$ 7</u>	<u>\$ 2</u>
Cash paid for income taxes	<u>\$ 3,120</u>	<u>\$ 8,400</u>

See accompanying notes.

QUESTCOR PHARMACEUTICALS, INC.
NOTES TO CONSOLIDATED FINANCIAL STATEMENTS
(unaudited)

1. The Company

Questcor Pharmaceuticals, Inc. is a biopharmaceutical company whose primary product helps patients with serious, difficult-to-treat medical conditions. Our primary product is H.P. Acthar® Gel (repository corticotropin injection), or Acthar, an injectable drug that is approved by the U.S. Food and Drug Administration, or FDA, for the treatment of 19 indications. Of these 19 indications, we currently generate substantially all of our net sales from three indications: (i) the treatment of acute exacerbations of multiple sclerosis, or MS, in adults, (ii) the treatment of nephrotic syndrome, or NS, and (iii) the treatment of infantile spasms, or IS, in infants and children under two years of age. With respect to NS, Acthar is approved to induce a diuresis or a remission of proteinuria in the nephrotic syndrome without uremia of the idiopathic type or due to lupus erythematosus. Acthar is approved both as maintenance therapy and to treat exacerbations in selected cases of systemic lupus erythematosus, or SLE, and in July 2011, we announced that we are exploring SLE as our next targeted therapeutic area for Acthar. We continue to explore the possibility of developing markets for other on-label indications, and the possibility of pursuing FDA approval of additional indications not currently on the Acthar label, where there is high unmet medical need.

Our other product is Doral® (quazepam), which is indicated for the treatment of insomnia characterized by difficulty in falling asleep, frequent nocturnal awakenings, and/or early morning awakenings. We own the U.S. rights to and have immaterial sales of Doral.

2. Summary of Significant Accounting Policies

Basis of Presentation

The consolidated financial statements include the accounts of Questcor and our wholly-owned subsidiary, Ribogene, Inc.

Use of Estimates

The preparation of financial statements in conformity with United States generally accepted accounting principles requires us to make estimates and assumptions that affect the amounts reported in the financial statements and accompanying notes. Actual results could differ significantly from those estimates.

Revenue Recognition

We recognize revenue in accordance with Accounting Standards Codification 605, "Revenue Recognition-Products," or ASC 605, from sales of Acthar and Doral. Pursuant to ASC 605, we recognize revenue when we have persuasive evidence that an arrangement, agreement or contract exists, when title for our product and risk of loss have passed to our customer, the price we charge for our product is fixed or is readily determinable, and we are reasonably assured of collecting the amounts owed under the resulting receivable. For sales of both of our products, we do not require collateral from our customers. We also support Acthar patient assistance programs administered by the National Organization of Rare Diseases, or NORDD, and the Chronic Disease Fund. These and other patient-oriented support programs have now provided free drug with a commercial value of over \$90 million to patients since September 2007 through June 30, 2011. We do not recognize any revenue from our free drug program.

In the United States, our exclusive customer for Acthar is CuraScript Specialty Distributor, or CuraScript SD. For our sales to CuraScript SD, a sale of Acthar occurs when CuraScript SD accepts a shipment of Acthar. We sell Acthar at a discount from our list price to CuraScript SD, which then sells Acthar primarily to approximately 12 specialty pharmacies, including CuraScript Specialty Pharmacy, or CuraScript SP, and to many hospitals. We sell Doral to pharmaceutical wholesalers, which in turn sell Doral primarily to retail pharmacies and hospitals.

International sales of our products are immaterial.

Net Sales

We record net sales after establishing reserves for the following:

- Medicaid rebates;
- Tricare retail program rebates;

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- Medicare Part D Coverage Gap Discount Program rebates;
- Chargebacks due to other government programs;
- Questcor-sponsored co-pay assistance programs; and
- Other deductions such as payment discounts.

We currently provide our products to Medicaid participants under an agreement with the Center for Medicare and Medicaid Services, or CMS. Under this agreement, states are eligible to receive rebates from us for Medicaid patients in accordance with CMS's regulations. For the quarters ended June 30, 2011 and 2010, the rebate amount equaled 100% of the Average Manufacturers' Price, or AMP, which approximates the amount we charge to CuraScript SD. States have historically provided us with rebate invoices for their Medicaid Fee for Service reimbursements between 60 to 90 days after the end of the calendar quarter in which our products were provided. Certain states are taking longer to submit their initial rebate invoices for the Medicaid Managed Care utilization that became rebate eligible as a result of Health Care Reform Acts on March 23, 2010. We estimate the end of period liability and the sales reserve needed for these Medicaid rebates based on the following multi-step process:

- Using a predictive model, we review national Medicaid statistics as well as internal information received from our Acthar reimbursement support center and from CuraScript SP for the most recent completed quarter to develop an estimate of future Medicaid rebate invoices that we expect to receive. This includes an estimate for both future Medicaid Fee for Service and Medicaid Managed Care Organization rebate invoices.
- We review the Medicaid rebate invoices received during the last 90 days and compare those invoices to the reserve that we had previously set at the end of the prior quarter. Based on this comparison and using the predictive model and other available information, which is updated quarterly, we estimate the remaining liability that we believe is still outstanding for periods prior to the most recently completed quarter.
- Based on estimated end-of-quarter inventory held at CuraScript SD, all specialty pharmacies and hospitals, we calculate the expected future rebate liability for that portion of the inventory which will eventually be used to fill prescriptions for Medicaid patients.

Using similar processes, we estimate the end of period liability and the sales reserve needed for Tricare retail program rebates, Medicare Part D Coverage Gap Discount Program rebates, or Coverage Gap Discount rebates (commonly referred to as the Medicare Part D "donut hole"), and chargebacks due to other government programs. The Medicare Part D Coverage Gap Discount Program took effect on January 1, 2011. Approximately 25% of our sales for MS are to Medicare insureds. As of March 31, 2011, we established a reserve for our estimated obligation with respect to Coverage Gap Discount rebates in the amount of \$120,000, and in April 2011 we received our first quarter bill, for \$71,000. As of June 30, 2011, we continued to reserve the remaining \$49,000 for our obligation with respect to Coverage Gap Discount rebates. We do not believe this program will have a material effect on our cash flows or results of operations.

Our resulting total sales reserve for the quarter includes the sum of the Medicaid sales reserve, the Tricare sales reserve, the Coverage Gap Discount reserves, the chargeback sales reserve, co-pay assistance payments, and payment discounts provided.

Significant judgment is inherent in the selection of assumptions and the interpretation of historical experience as well as the identification of external and internal factors affecting the determination of our reserves for Medicaid rebates and other government program rebates and chargebacks. We believe that the assumptions used to determine these sales reserves are reasonable considering known facts and circumstances. However, our Medicaid rebates and other government program rebates and chargebacks could differ significantly from our reserve amounts because of unanticipated changes in prescription trends or patterns in the states' submissions of Medicaid claims, adjustments to the amount of product in the distribution channel, or if our estimates of the number of Medicaid patients with IS, MS and NS are incorrect. We have greater visibility on the future submission of Medicaid claims and the amount of product in the distribution channel for Acthar distributed to CuraScript SP (which is owned by CuraScript SD) than we have with respect to Acthar distributed through other specialty pharmacies. If actual Medicaid rebates, or other government program rebates and chargebacks are materially different from our estimates, we would account for such differences as a change in estimate in the period in which they become known. If actual future payments for such reserves exceed the estimates we made at the time of sale, our financial position, results of operations and cash flows may be negatively impacted.

Medicaid Rebates and the New National Health Care Legislation

In March 2010, Congress passed, and the President signed into law, health care legislation entitled the Patient Protection and Affordable Care Act of 2010 and the Health Care and Education Affordability Reconciliation Act of 2010, and subsequent related legislation passed during the third quarter of 2010, which we refer to collectively as the Health Care

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Reform Acts. The Health Care Reform Acts contain a number of provisions that have impacted, both positively and negatively, our financial position, results of operations and cash flows. The provisions of the Health Care Reform Acts have reduced our rebate provided to states for prescriptions filled for Medicaid patients to 100% of the AMP, which approximates the amount we charge to CuraScript SD. Before the passage of the Health Care Reform Acts, the formula used to calculate the per vial rebate required us to rebate 110% of our AMP for Acthar. Effective March 23, 2010, the Health Care Reform Acts extended Medicaid rebates to Medicaid Managed Care plans. Medicaid Managed Care plans provide for the delivery of Medicaid health benefits and additional services through an arrangement between a state Medicaid agency and managed care organizations. Our provision for expected Medicaid rebate liability and our quarterly sales reserves have included an estimate for Medicaid Managed Care usage since March 23, 2010.

Other Impacts from Health Care Reform Acts

In addition to the impact on our required Medicaid rebates, the Health Care Reform Acts contain a number of provisions that we expect to continue to impact, both positively and negatively, our financial position, results of operations and cash flows.

- *Positive Impact.* The Health Care Reform Acts contain provisions that create a national high-risk insurance pool, temporarily extend health coverage to individuals with pre-existing medical conditions, prohibit the denial of health coverage to children with pre-existing conditions, prohibit the denial of health coverage to adults with pre-existing conditions and place limits on insurers with respect to lifetime and annual caps on health coverage, and increase the number of patients with private insurance.
- *Negative Impact.* The Health Care Reform Acts contain the following provisions that we have identified as having a negative or potentially negative impact on our overall financial position, results of operations and cash flows:
 - Effective January 1, 2011, pharmaceutical companies, including the Company, must provide Coverage Gap Discount rebates, as described above.
 - Effective January 1, 2011, the U.S. federal government allocates an annual fee among manufacturers of branded prescription drugs based on market share, in the aggregate, for specified government programs. The Health Care Reform Acts determine an individual manufacturer's market share as the ratio of its aggregate sales of branded prescription drugs during the preceding calendar year as a percentage of the aggregate branded prescription drug sales for all covered manufacturers.
 - We expect the number of Medicaid patients to increase gradually through 2014, and that this expansion will more likely impact the number of adults in Medicaid because many states have already set their eligibility criteria for children at or above the level designated in the Health Care Reform Acts. An increase in the proportion of patients who receive Acthar and who are covered by Medicaid could adversely affect our net sales.

Many of the provisions of the Health Care Reform Acts require rulemaking action by governmental agencies to implement. As various agencies implement these rules and regulations, our business may be negatively impacted other than as described above. In addition, Congress and the President may make additional refinements to the Health Care Reform Acts which may have additional, potentially negative impacts on our overall financial position, results of operations and cash flows. Furthermore, there continues to be active debate in Congress and the courts ranging from repeal of the Health Care Reform Acts to no change in the law. At this time, we cannot predict the full impact of the Health Care Reform Acts, or the timing and impact of any future rules or regulations promulgated to implement the Health Care Reform Acts. It is possible that the Health Care Reform Acts and related rulemaking actions may have an overall negative effect on our net sales over time.

TRICARE Retail Pharmacy Programs

The Department of Defense, or DoD, Tricare Retail Pharmacy program became effective on May 26, 2009 pursuant to section 703 of the National Defense Authorization Act of 2008. This program and its regulations require manufacturers to pay rebates, retroactive to January 28, 2008, to the DoD on products distributed to Tricare beneficiaries through retail pharmacies. The regulations further require that pharmaceutical products paid for by the DoD through the Tricare Retail Pharmacy program be subject to the Federal Ceiling Price program, which requires manufacturers to provide the DoD with a refund on pharmaceutical products utilized through the Tricare Retail Pharmacy program. As a result, we established a sales reserve of \$3.5 million for Tricare rebates as of the year ended December 31, 2009, which covered 100% of our estimated liability for the time period January 28, 2008 through December 31, 2009. The regulations upon which the reserve is based are currently being challenged by the pharmaceutical industry. As it is unclear whether this challenge will be successful, our sales reserve related to the period from January 28, 2008 through December 31, 2009 remained at \$3.5 million at June 30, 2011.

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Effective January 1, 2010, we entered into a new pricing agreement with the Veterans Administration, resulting in a rebate for pharmaceutical products utilized through the Tricare Retail Pharmacy program during 2010 of \$5,670 per vial, or a reduction of \$14,865 from the previous rebates of \$20,535. Effective January 1, 2011, our rebate decreased to \$5,528 per vial. We recorded sales reserves of \$0.6 million for each of the six month periods ended June 30, 2011 and 2010, respectively.

Government Chargebacks

We permit certain other government-supported entities, such as those covered by our contract with the Veterans Administration or eligible Public Health Service, or PHS, 340(B) entities, to purchase Acthar from CuraScript SD based on a contractual amount. Because our payment terms with CuraScript SD are approximately 30 days, we include actual chargebacks taken plus an estimate applied to the units in channel when estimating the sales reserve related to government chargebacks. Sales to the Veterans Administration and PHS 340(B) entities are generally immaterial to our financial position as a whole.

Co-Pay Assistance Programs

We sponsor co-pay assistance programs for Acthar patients which are administered by NORD and the Chronic Disease Fund. We account for these co-pay assistance program payments as a reduction to our revenue.

Total Sales-related Reserves

At June 30, 2011 and December 31, 2010, sales-related reserves included in the accompanying Consolidated Balance Sheets were as follows (in thousands):

	<u>June 30, 2011</u>	<u>December 31, 2010</u>
Medicaid rebates	\$22,856	\$ 17,384
Tricare rebates	4,087	4,125
Medicare Part D Coverage Gap Discount Program rebates	49	—
Government chargebacks	45	2
Other discounts	29	—
Total	<u>\$27,066</u>	<u>\$ 21,511</u>

Product Exchanges

On a limited basis, we generally authorize Acthar exchanges for expiring and expired product in accordance with our stated return policy, which allows CuraScript SD to return product within one month of its expiration date and for a period up to three months after such product has reached its expiration date. We exchange returns for replacement product and we include in the cost of sales the estimated costs for such exchanges, which include actual product material costs and related shipping charges. Product exchanges have been insignificant since we began utilizing the services of CuraScript SD to distribute Acthar.

Concentration of Credit Risk

Financial instruments which subject us to a significant concentration of credit risk principally consist of cash and cash equivalents, short-term investments and accounts receivable. We invest our cash in high credit quality government and corporate debt instruments and believe the financial risks associated with these instruments are minimal.

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Cash and cash equivalents are maintained at financial institutions and, at times, balances may exceed federally insured limits. We have never experienced any losses related to these balances. All of our non-interest bearing cash balances were fully insured at June 30, 2011 due to a temporary federal program in effect from December 31, 2010 through December 31, 2012. Under the program, there is no limit to the amount of insurance for eligible accounts. Beginning in 2013, insurance coverage will revert to \$250,000 per depositor at each financial institution, and our non-interest bearing cash balances may again exceed federally insured limits. Interest-bearing amounts on deposit in excess of federally insured limits at June 30, 2011 would have approximated \$1.7 million.

We extend credit to our customer, CuraScript SD, which accounts for nearly 100% of our gross product sales and accounts receivable. We have not experienced significant credit losses on our customer accounts.

Inventories

We state inventories, net of allowances, at the lower of cost or market value. Cost is determined by the first-in, first-to-expire method.

We review inventory periodically for slow-moving or obsolete status. We adjust our inventory if we do not expect to recover the cost of inventory. We would record a reserve to adjust inventory to its net realizable value: (i) when a product is close to expiration and we do not expect it to be sold, (ii) when a product has reached its expiration date or (iii) when we do not expect a product to be saleable. In determining the reserves for our products, we consider factors such as the amount of inventory on hand and its remaining shelf life, and current and expected market conditions. We have evaluated the current level of inventory and based on our evaluation have recorded adjustments to reflect inventory at its net realizable value. These adjustments are estimates, which could vary significantly from actual results if future economic conditions, customer demand, competition or other relevant factors differ from expectations. These estimates require us to assess the future demand for our products in order to categorize the status of such inventory items as slow-moving, obsolete or in excess-of-need. These future estimates are subject to the ongoing accuracy of our forecasts of market conditions, industry trends, competition and other factors. Differences between our estimated reserves and actual inventory adjustments have been immaterial, and we account for such adjustments in the current period as a change in estimate.

Property and Equipment

Equipment and leasehold improvements and related accumulated depreciation and amortization are as follows (in thousands):

	<u>June 30,</u> <u>2011</u>	<u>December 31,</u> <u>2010</u>
Laboratory equipment	\$ 8	\$ 8
Manufacturing equipment	733	692
Office equipment, furniture and fixtures	1,895	1,971
Leasehold improvements	946	420
	<u>3,582</u>	<u>3,091</u>
Less accumulated depreciation and amortization	(1,652)	(2,219)
	<u>\$ 1,930</u>	<u>\$ 872</u>

Total depreciation and amortization expense amounted to \$0.3 million for the six months ended June 30, 2011 and \$0.2 million for the year ended December 31, 2010.

Supply Concentration Risks

We obtain some materials used in our products from a single source. We have a supply agreement with BioVectra dcl, our sole source supplier for the active pharmaceutical ingredient, or API, in Acthar. We also have a supply agreement with Cangene bioPharma, Inc., or Cangene, pursuant to which Cangene manufactures Acthar final product. Cangene is our sole source for Acthar final product. Additionally, we use a sole source provider for potency testing.

Cash Equivalents and Short-Term Investments

We consider highly liquid investments with maturities from the date of purchase of three months or less to be cash equivalents. We classify available-for-sale debt instruments with maturities at the date of purchase of greater than three months as short-term investments.

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We carry available-for-sale securities at fair value, with the unrealized gains and losses, if any, reported in a separate component of shareholders' equity. If we deem the decline in value to be other-than-temporary and we intend to sell such securities before their full cost can be recovered, we write down such securities to fair value and we charge the loss to net realized losses on investments. We use significant judgment in the determination of when an other-than-temporary decline in value has occurred. We evaluate our investment securities for other-than-temporary declines based on quantitative and qualitative factors. As of June 30, 2011, none of our investments had an other-than-temporary decline in valuation, and no other-than-temporary losses were recognized during the year ended December 31, 2010. We base the cost of securities sold on the specific identification method. We include realized gains and losses, if any, in the accompanying Consolidated Statements of Income, in Other Income.

A summary of cash and cash equivalents and short-term investments, classified as available-for-sale, and carried at fair value is as follows (in thousands):

	<u>Amortized Cost</u>	<u>Gross Unrealized Gain</u>	<u>Gross Unrealized (Loss)</u>	<u>Estimated Fair Value</u>
June 30, 2011				
Cash and cash equivalents	\$ 65,277	\$ —	\$ —	\$ 65,277
Short-term investments:				
Certificates of deposit	\$ 9,320	\$ 24	\$ (3)	\$ 9,341
Corporate bonds	48,103	34	(27)	48,110
Government-sponsored enterprises	4,395	—	(1)	4,394
Municipal bonds	2,003	1	—	2,004
	<u>\$ 63,821</u>	<u>\$ 59</u>	<u>\$ (31)</u>	<u>\$ 63,849</u>
December 31, 2010				
Cash and cash equivalents	\$ 41,508	\$ —	\$ —	\$ 41,508
Short-term investments:				
Certificates of deposit	\$ 9,080	\$ 39	\$ (4)	\$ 9,115
Corporate bonds	15,427	9	(5)	15,431
Government-sponsored enterprises	41,983	12	(27)	41,968
Municipal bonds	6,808	3	(1)	6,810
	<u>\$ 73,298</u>	<u>\$ 63</u>	<u>\$ (37)</u>	<u>\$ 73,324</u>

The amortized cost and fair value of short-term investment securities at June 30, 2011, by contractual maturity, are as follows (in thousands):

	<u>Amortized Cost</u>	<u>Estimated Fair Value</u>
Due in one year or less	\$ 37,739	\$ 37,739
Due after one through two years	26,082	26,110
Total short-term investments	<u>\$ 63,821</u>	<u>\$ 63,849</u>

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As of June 30, 2011, the average contractual maturity of our short-term investments was approximately 14 months.

As of June 30, 2011, we had the following available-for-sale securities that were in an unrealized loss position but were not deemed to be other-than-temporarily impaired (in thousands):

	<u>Less Than 12 Months</u>		<u>12 Months or Greater</u>	
	<u>Gross Unrealized Losses</u>	<u>Estimated Fair Value</u>	<u>Gross Unrealized Losses</u>	<u>Estimated Fair Value</u>
Corporate bonds	\$ (12)	\$ 5,846	\$ (15)	\$ 5,846
Government-sponsored enterprises	(1)	1,043	—	—
Certificates of deposit	—	—	(3)	477
Total	<u>\$ (13)</u>	<u>\$ 6,889</u>	<u>\$ (18)</u>	<u>\$ 6,323</u>

The gross unrealized losses reported above for June 30, 2011 were caused by general fluctuations in market interest rates from the respective purchase date of these securities through June 30, 2011. No significant facts or circumstances have occurred to indicate that these unrealized losses are related to any deterioration in the creditworthiness of the issuers of the marketable securities we own. Based on our review of these securities, including our assessment of the duration and severity of the related unrealized losses, we have not recorded any other-than-temporary impairments on these investments.

Fair Value of Financial Instruments

Our financial instruments include cash and cash equivalents, short-term investments, accounts receivable, accounts payable and accrued liabilities. We believe that the fair value of these financial instruments approximate the reported carrying amounts.

Fair Value Measurements

We account for fair value measurements under Accounting Standards Codification 820 "Fair Value Measurements and Disclosures," or ASC 820, which defines fair value as the exchange price that would be received for an asset or paid to transfer a liability (an exit price) in the principal or most advantageous market for the asset or liability in an orderly transaction between market participants at the measurement date. ASC 820 establishes a three-level fair value hierarchy that prioritizes the inputs used to measure fair value. This hierarchy requires entities to maximize the use of observable inputs and minimize the use of unobservable inputs. The three levels of inputs used to measure fair value are as follows:

- Level 1 – Quoted prices in active markets for identical assets or liabilities.
- Level 2 – Observable inputs other than quoted prices included in Level 1, such as quoted prices for similar assets and liabilities in active markets; quoted prices for identical or similar assets and liabilities in markets that are not active; or other inputs that are observable or can be corroborated by observable market data.
- 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. This includes certain pricing models, discounted cash flow methodologies and similar techniques that use significant unobservable inputs.

We have segregated all assets and liabilities measured at fair value on a recurring basis (at least annually) into the most appropriate level within the fair value hierarchy based on the inputs used to determine the fair value at the measurement date in the table below. As of June 30, 2011, all of our assets and liabilities are valued using Level 1 inputs except for our short-term investments which are valued using Level 2 inputs.

Assets measured at fair value on a recurring basis are summarized below (in thousands):

	<u>Basis of Fair Value Measurements</u>			
	<u>Balance at June 30, 2011</u>	<u>Level 1</u>	<u>Level 2</u>	<u>Level 3</u>
Cash and cash equivalents	\$ 65,277	\$65,277	\$ —	\$ —
Certificates of deposit	9,341	9,341	—	—
Corporate bonds	48,110	—	48,110	—
Government-sponsored enterprises	4,394	—	4,394	—
Municipal bonds	2,004	—	2,004	—
Total	<u>\$129,126</u>	<u>\$74,618</u>	<u>\$54,508</u>	<u>\$ —</u>

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Investment securities are exposed to various risk factors, such as interest rate, market and credit risk. Due to the level of risk associated with certain investment securities and the level of uncertainty related to changes in the value of investment securities, it is possible that changes in these risk factors in the near term could have an adverse material impact on our results of operations or shareholders' equity.

Certain assets and liabilities are measured at fair value on a nonrecurring basis. In other words, the instruments are not measured at fair value on an ongoing basis but are subject to fair value adjustments only in certain circumstances (for example, when there is evidence of impairment). There were no assets or liabilities measured at fair value on a nonrecurring basis during the periods ended June 30, 2011 and December 31, 2010, other than goodwill associated with a 1999 transaction, which was impaired during the six months ended June 30, 2011 resulting in a net realizable value of zero.

Comprehensive Income

Accounting Standards Codification 220 "Comprehensive Income," or ASC 220, requires reporting and displaying comprehensive income (loss) and its components, which includes net income and unrealized gains and losses on investments and foreign currency translation gains and losses. The following table summarizes comprehensive income (in thousands):

	Three Months Ended June 30,		Six Months Ended June 30,	
	2011	2010	2011	2010
Net income	\$13,874	\$9,282	\$25,098	\$17,134
Change in unrealized gains or losses on available-for-sale securities, net of related tax effects	(15)	54	2	77
Comprehensive income	<u>\$13,859</u>	<u>\$9,336</u>	<u>\$25,100</u>	<u>\$17,211</u>

Share-based Compensation

We recognize compensation expense for all share-based awards made to employees and directors. The fair value of share-based awards is estimated at grant date using an option pricing model and the portion that is ultimately expected to vest is recognized as compensation cost over either (1) the requisite service period or (2) the performance period.

Since share-based compensation is recognized only for those awards that are ultimately expected to vest, we have applied an estimated forfeiture rate to unvested awards for the purpose of calculating compensation cost. These estimates will be revised, if necessary, in future periods if actual forfeitures differ from estimates. Changes in forfeiture estimates impact compensation cost in the period in which the change in estimate occurs.

We use the Black-Scholes option-pricing model to estimate the fair value of share-based awards. The determination of fair value using the Black-Scholes option-pricing model is affected by our stock price as well as assumptions regarding a number of complex and subjective variables, including expected stock price volatility, risk-free interest rate, expected dividends and projected employee stock option behaviors. We estimate the expected term based on the contractual term of the awards and employees' exercise and expected post-vesting termination behavior.

Additionally, we are required to disclose in our consolidated statement of cash flows the income tax effects resulting from share-based payment arrangements. We adopted the simplified method to calculate the beginning balance of the additional paid-in capital, or APIC, pool of excess tax benefits, and to determine the subsequent effect on the APIC pool and consolidated statements of cash flows of the tax effects of employee share-based compensation awards.

At June 30, 2011, there was \$14.1 million of total unrecognized compensation cost related to unvested stock options, which is expected to be recognized over a remaining weighted average vesting period of approximately 2.7 years.

Share-based compensation cost is summarized below (in thousands):

	Three Months Ended June 30,		Six Months Ended June 30,	
	2011	2010	2011	2010
Selling and marketing	\$ 415	\$ 238	\$ 789	\$ 470
General and administrative	1,041	442	2,167	1,017
Research and development	260	240	572	462
Total	<u>\$ 1,716</u>	<u>\$ 920</u>	<u>\$3,528</u>	<u>\$1,949</u>

[Table of Contents](#)**Net Income Per Share**

We compute basic net income per share by dividing the net income for the period by the weighted average number of common shares outstanding during the period. We compute diluted net income per share by dividing the net income for the period by the weighted-average number of common and common equivalent shares, such as stock options and restricted shares outstanding during the period. Diluted earnings for holders of our common stock per share consider the impact of potentially dilutive securities. Dilutive potential common shares resulting from the assumed exercise of outstanding stock options are determined based on the treasury stock method. Under the treasury stock method, the dilutive impact of a stock option that is "in-the money" is based on the difference between that stock option's exercise price and the Company's stock price at the time of measurement. The more the stock price exceeds the exercise price, the greater the number of potential common shares and thus the greater the dilutive impact of the stock option.

Basic and diluted net income per share was calculated as follows (in thousands):

	Three Months Ended June 30,		Six Months Ended June 30,	
	2011	2010	2011	2010
Net income applicable to common shareholders	\$13,874	\$ 9,282	\$25,098	\$17,134
Shares used in computing net income per share applicable to common shareholders:				
Basic	62,034	62,022	62,126	61,957
Effect of dilutive potential common shares:				
Stock options	3,418	2,504	3,344	2,086
Restricted stock	12	17	13	14
ESPP	—	—	—	—
Diluted	65,464	64,543	65,483	64,057
Net income per share applicable to common shareholders:				
Basic	\$ 0.22	\$ 0.15	\$ 0.40	\$ 0.28
Diluted	\$ 0.21	\$ 0.14	\$ 0.38	\$ 0.27

The following table presents the amounts excluded from the computation of diluted net income per share applicable to common shareholders for the periods ended June 30, 2011 and 2010 as the inclusion of these securities would have been anti-dilutive (in thousands):

	Three Months Ended June 30,		Six Months Ended June 30,	
	2011	2010	2011	2010
Stock options	348	180	846	2,029

Purchased Technology and Goodwill

Purchased technology consists of the following (in thousands):

	June 30, 2011	December 31, 2010
Purchased technology	\$ 4,386	\$ 4,386
Less accumulated amortization	(1,459)	(1,312)
	\$ 2,927	\$ 3,074

Purchased technology at June 30, 2011 and December 31, 2010 consists of our acquisition costs for Doral. Amortization expense for purchased technology totaled \$0.1 million for each of the six months ended June 30, 2011 and June 30, 2010, respectively. As of June 30, 2011 and December 31, 2010, we determined that purchased technology was not impaired and will continue to monitor the carrying value of the remaining purchased technology.

Goodwill consists of the following (in thousands):

	June 30, 2011	December 31, 2010
Goodwill	\$ 1,023	\$ 1,023
Less accumulated amortization	(1,023)	(724)
	\$ —	\$ 299

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During the six months ended June 30, 2011, we determined the carrying value of the remaining goodwill impaired and, therefore, charged the remaining balance to impairment of goodwill. At December 31, 2010, we had not yet made this determination.

Indemnification, Commitments and Contingencies

As permitted under California law and in accordance with our Amended and Restated Bylaws, we indemnify our officers and directors for certain events or occurrences while the officer or director is or was serving at our request in such capacity. The potential future indemnification limit is to the fullest extent permissible under California law. However, we have a director and officer insurance policy that limits our exposure and may enable us to recover a portion of any future amounts paid. We believe the fair value of these indemnification agreements in excess of applicable insurance coverage is minimal. Accordingly, we have no liabilities recorded for these agreements as of June 30, 2011 and December 31, 2010.

Segment Reporting

We have determined that we operate in one business segment.

Income Taxes

We account for income taxes under the provisions of Accounting Standards Codification 740, "Income Taxes," or ASC 740. We make certain estimates and judgments in determining income tax expense for financial statement purposes. These estimates and judgments occur in the calculation of certain tax assets and liabilities, which arise from differences in the timing of recognition of revenue and expense for tax and financial statement purposes.

As part of the process of preparing our consolidated financial statements, we are required to estimate income taxes in each of the jurisdictions in which we operate. This process involves estimating our tax exposure under the most current tax laws and assessing temporary differences resulting from differing treatment of items for tax and accounting purposes. These differences result in deferred tax assets and liabilities, which are included in our consolidated balance sheets.

We regularly assess the likelihood that we will be able to recover our deferred tax assets, which is ultimately dependent on us generating future taxable income. We consider all available evidence, both positive and negative, including historical levels of income, expectations and risks associated with estimates of future taxable income and ongoing prudent and feasible tax planning strategies in assessing the need for a valuation allowance. If it is not considered "more likely than not" that we will recover our deferred tax assets, we will increase our provision for taxes by recording a valuation allowance against the deferred tax assets that we estimate will not ultimately be recoverable. Changes in the valuation allowance based on our assessment will result in an income tax benefit if the valuation allowance is decreased and an income tax expense if the valuation allowance is increased.

Equity Transactions

On February 29, 2008, our Board of Directors approved a stock repurchase plan that provides for the Company's repurchase of up to 7 million shares of our common stock. Stock repurchases under this plan may be made through either open market or privately negotiated transactions in accordance with all applicable laws, rules and regulations. On May 29, 2009, our Board of Directors increased the stock repurchase plan authorization by an additional 6.5 million shares.

During the six months ended June 30, 2011, we used \$11.5 million to repurchase 884,300 shares of our common stock. Under this stock repurchase plan, we have repurchased a total of 9.2 million shares of our common stock for \$48.1 million through June 30, 2011, at an average price of \$5.21 per share. As of June 30, 2011, there are 4.3 million shares authorized remaining under our stock repurchase plan.

Recent Accounting Pronouncements

In April 2011, the Financial Accounting Standards Board, or FASB, issued Accounting Standards Update No. 2011-04 "Fair Value Measurement (Topic 820) – Amendments to Achieve Common Fair Value Measurement and Disclosure Requirements in U.S. GAAP and IFRS," or ASU No. 2011-04. ASU No. 2011-04 is the result of the continuing convergence projects between the FASB, and the International Accounting Standards Board, or IASB, to create a common set of high quality global accounting standards. The amendments in ASU No. 2011-04 explain how to measure fair value. They do not require additional fair value measurements and are not intended to establish standards or affect valuation practices outside of financial reporting. The amendment will be effective for interim and annual periods beginning after December 15, 2011. We plan to adopt ASU No. 2011-04 and do not anticipate a material effect on our financial position or results of operation.

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In June 2011, the FASB issued Accounting Standards Update No. 2011-05 “Presentation of Comprehensive Income,” or ASU No. 2011-05. ASU No. 2011-05 improves the comparability, consistency, and transparency of financial reporting and increases the prominence of items reported in other comprehensive income, or OCI, by eliminating the option to present OCI as part of the statement of changes in shareholders’ equity. The amendments in this standard require that all non-owner changes in shareholders’ equity be presented either in a single continuous statement of comprehensive income or in two separate but consecutive statements. Under either method, adjustments must be displayed for items that are reclassified from OCI to net income, in both net income and OCI. The standard does not change the current option for presenting components of OCI gross or net of the effect of income taxes, provided that such tax effects are presented in the statement in which OCI is presented or disclosed in the notes to the financial statements. Additionally, the standard does not affect the calculation or reporting of earnings per share. For public entities, the amendments in this ASU are effective for fiscal years, and interim periods within those years, beginning after December 15, 2011 and are to be applied retrospectively, with early adoption permitted. We plan to adopt ASU No. 2011-05 and do not anticipate a material effect on our financial position or results of operation.

Subsequent Events

We evaluated subsequent events that have occurred after June 30, 2011, and determined that there were no events or transactions occurring during this reporting period which require recognition or disclosure in our consolidated financial statements.

On July 26, 2011, we announced that we are exploring systemic lupus erythematosus, or SLE, as our next targeted therapeutic area for Acthar. Acthar is approved both as maintenance therapy and to treat exacerbations in selected cases of SLE.

ITEM 2. MANAGEMENT'S DISCUSSION AND ANALYSIS OF FINANCIAL CONDITION AND RESULTS OF OPERATIONS

Except for the historical information contained herein, the following discussion contains forward-looking statements that involve risks and uncertainties. Our actual results could differ materially from those discussed herein. Factors that could cause or contribute to such differences include, but are not limited to, those discussed in this section, in Item 1A "Risk Factors" of Part II of this Quarterly Report, those discussed in our Annual Report on Form 10-K for the year ended December 31, 2010, including Item 1 "Business of Questcor," and Item 1A "Risk Factors" of Part I of that Annual Report, as well as factors discussed in any documents incorporated by reference herein or therein.

Overview

Questcor Pharmaceuticals, Inc. is a biopharmaceutical company whose primary product helps patients with serious, difficult-to-treat medical conditions. Our primary product is H.P. Acthar® Gel (repository corticotropin injection), or Acthar, an injectable drug that is approved by the U.S. Food and Drug Administration, or FDA, for the treatment of 19 indications. Of these 19 indications, we currently generate substantially all of our net sales from three indications: (i) the treatment of acute exacerbations of multiple sclerosis, or MS, in adults, (ii) the treatment of nephrotic syndrome, or NS, and (iii) the treatment of infantile spasms, or IS, in infants and children under two years of age. With respect to NS, Acthar is approved to induce a diuresis or a remission of proteinuria in the nephrotic syndrome without uremia of the idiopathic type or due to lupus erythematosus. Acthar is approved both as maintenance therapy and to treat exacerbations in selected cases of systemic lupus erythematosus, or SLE, and in July 2011, we announced that we are exploring SLE as our next targeted therapeutic area for Acthar. We continue to explore the possibility of developing markets for other on-label indications, and the possibility of pursuing FDA approval of additional indications not currently on the Acthar label, where there is high unmet medical need.

While there are 19 indications on the Acthar label, we currently target the following four on-label therapeutic areas:

Therapeutic Area	Labeled Indication	Commercial Efforts
MS Exacerbations	Multiple Sclerosis: H.P. Acthar Gel (repository corticotropin injection) is indicated for the treatment of acute exacerbations of multiple sclerosis in adults. Controlled clinical trials have shown H.P. Acthar Gel to be effective in speeding the resolution of acute exacerbations of multiple sclerosis. However, there is no evidence that it affects the ultimate outcome or natural history of the disease.	Sales force expanded from 38 to 77, effective November 1, 2010; 751 paid prescriptions in the second quarter ended June 30, 2011, an increase of 147% over the second quarter ended June 30, 2010.
Nephrotic Syndrome	Edematous State: To induce a diuresis or a remission of proteinuria in the nephrotic syndrome without uremia of the idiopathic type or that due to lupus erythematosus.	Initial sales force of five specialists commenced marketing activities in March 2011, and generated 45 paid prescriptions in the second quarter ended June 30, 2011. We are in the process of increasing our NS selling effort, with the goal of calling on over 3,000 nephrologists.
Infantile Spasms	Infantile spasms: H.P. Acthar Gel (repository corticotropin injection) is indicated as monotherapy for the treatment of infantile spasms in infants and children under 2 years of age.	Acthar has been the standard of care for this condition for many years. Second quarter paid prescriptions remained within the normal historic range.
Systemic Lupus Erythematosus	Collagen Diseases: During an exacerbation or as maintenance therapy in selected cases of systemic lupus erythematosus.	Announced as Questcor's next targeted therapeutic area in July 2011.

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Acthar has also been used to treat other conditions not on the label of approved indications for Acthar. Since January 1, 2010, we believe physicians have written prescriptions for Acthar for a variety of conditions that are not on its label of approved indications, including the following: Adrenal Insufficiency, Encephalopathy, Landau-Kleffner Syndrome, Myasthenia Gravis, Neurosarcoidosis, Opsoclonus Myoclonus, Peripheral Neuropathy, Scleroderma and Ulcerative Colitis. We do not promote Acthar for these indications.

We also maintain a research and development program focused on gathering data to: (i) evaluate the proper use of Acthar for on-label indications; (ii) investigate other potential uses of Acthar that are not currently FDA approved indications; and (iii) improve our understanding of how Acthar works in the human body (pharmacology), and ultimately, its mechanism(s) of action in the disease states for which it is currently used, or may be used in the future:

- On-Label Development. On-label clinical development efforts include the following:
 - Nephrotic Syndrome. We are the sponsor of a Phase IV clinical trial evaluating Acthar for the treatment of proteinuria associated with treatment-resistant idiopathic membranous nephropathy (IMN), with enrollment scheduled to begin in the third quarter of 2011; we are also supporting clinical nephrology Investigator Initiated Studies (IIS) evaluating: (i) safety and efficacy of Acthar in IMN; (ii) safety and efficacy of Acthar in focal segmental glomerular sclerosis (FSGS); and (iii) safety and efficacy of Acthar in treatment-resistant nephrotic syndrome (including IMN, FSGS, IgA nephropathy and minimal change disease).
 - Infantile Spasms (IS). We are supporting an IIS aimed at establishing quality of care indicators for IS.
 - Systemic Lupus Erythematosus (SLE). We are considering conducting clinical studies evaluating Acthar for treatment of SLE.
- Other Indications, Not On-Label. Research and development efforts with respect to the use of Acthar to treat conditions that are not on the label of approved indications for Acthar include the following:
 - Diabetic Nephropathy (DN). We are developing a clinical protocol for a small Company-sponsored study to evaluate the safety and efficacy of Acthar in treating Diabetic Nephropathy. We are also supporting a clinical IIS evaluating safety and efficacy of Acthar in treatment of DN.
 - Multiple Sclerosis (MS) - Pulse Therapy. We are supporting a clinical IIS examining pulse administration of Acthar in MS in conjunction with disease-modifying therapy to evaluate possible disease modifying effects of Acthar.
 - Cognitive Protection/Autism. We are supporting a preclinical IIS, investigating whether Acthar has protective effects in an animal model of epilepsy with concomitant autism-related cognitive dysfunction.
 - Traumatic Brain Injury (TBI): We are supporting a preclinical IIS investigating whether Acthar has protective effects in an animal model of TBI.
 - Amyotrophic Lateral Sclerosis (ALS): We are supporting a preclinical IIS investigating whether Acthar has protective effects in an animal model of ALS (commonly referred to as Lou Gehrig's disease).
- Pharmacology. We are conducting non-clinical and clinical pharmacology studies in the following areas:
 - General. We seek to gain a better understanding of the mechanism of action of Acthar as well as the pharmacology of Acthar across and within each indication. We are also conducting studies to understand why Acthar acts differently than steroids or potentially other melanocortin peptides.
 - Multiple Sclerosis. We are supporting an IIS evaluating immune modulating effects of Acthar applied to serum from MS patients and an IIS evaluating neuroprotective properties of adrenocorticotropic hormone that are relevant to MS.

Net sales of Acthar are derived from our sales of vials to CuraScript SD, which in turn sells Acthar primarily to specialty pharmacies. These specialty pharmacies place orders to CuraScript SD based on their respective levels of sales and inventory practices. End-user demand for Acthar results from physicians writing prescriptions to patients for the treatment of

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MS exacerbations, NS, IS and various other conditions. Recommended treatment regimens among physicians prescribing Acthar vary within each therapeutic area. Due to various factors, including inventory levels at both the specialty pharmacies and at CuraScript SD, the duration of treatment regimens and the timing of the placement of re-fill prescription orders, there is typically a delay between changes in prescription levels and changes in the levels of orders we receive from CuraScript SD. Additionally, treatment regimens and patient compliance with physician recommended treatment regimens, may vary over time. We experienced significant prescription growth in the quarter ended March 31, 2011 and the quarter ended June 30, 2011, and believe this growth has resulted in higher net sales in the quarter ended June 30, 2011.

Set forth below is a table which provides information for new (non-refill) paid Acthar prescriptions for our three primary therapeutic area and vials shipped to CuraScript SD for the three months ended June 30, 2011 and 2010:

	Three Months Ended		Increase	% Change
	2011	June 30, 2010		
New Paid Prescriptions:				
MS	751	304	447	147%
NS	45	4	41	1025%
IS	106	95	11	12%
Vials Shipped to CuraScript SD	2,430	1,680	750	45%

Important Notes Regarding Prescription Data

(1) Because Acthar prescriptions are filled at specialty pharmacies, we do not receive complete information regarding either the number of prescriptions or the number of vials by therapeutic area for all of the patients being treated with Acthar. However, we are able to monitor trends in payer mix and areas of therapeutic use for new (non-refill) Acthar prescriptions based on data we receive from our reimbursement support center. We estimate that over 90% of new Acthar prescriptions are processed by this support center, but believe that very few refill prescriptions are processed there.

(2) Prescription figures include related conditions for each therapeutic area. Related conditions are diagnoses that are either alternative descriptions of the medical condition or are closely related to the medical condition which is the focus of the table. For example, a prescription for "Demyelinating disease of the central nervous system" would be included as an MS related condition for purpose of this table. About 5% of the prescriptions in the tables are for related conditions.

(3) A new prescription may or may not represent a new patient or a new therapy for the patient receiving the prescription. We use business rules to determine whether a prescription should be classified as new for inclusion in this table. From time to time, we may modify these rules, which could cause some changes to the historic figures in the tables above.

We have increased the size of our Specialty Sales Force, which calls on neurologists who treat patients for MS or IS, several times since the beginning of 2008, most recently when we expanded from 38 representatives to 77 representatives effective November 2010. Additionally, in March 2011, we assembled a Nephrology Sales Force which promotes Acthar exclusively to nephrologists for use in treating NS. Our initial Nephrology Sales Force was comprised of just five representatives and, based on the results of their efforts, we are significantly expanding our NS selling effort. Specifically, we are in the process of hiring approximately 23 additional representatives for our Nephrology Sales Force and we are giving a limited supportive selling role to our 77 representative Specialty Sales Force. Based on this expansion, we expect the total number of target nephrologists that we call on will increase from less than 400 pre-expansion to over 3,000.

Our other product is Doral® (quazepam), which is indicated for the treatment of insomnia characterized by difficulty in falling asleep, frequent nocturnal awakenings, and/or early morning awakenings. We own the U.S. rights to and have immaterial sales of Doral.

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Results of Operations

Three months ended June 30, 2011 compared to the three months ended June 30, 2010:

Recorded Net Sales

	Three Months Ended June 30,		Increase/ (Decrease)	% Change
	2011	2010		
	(in \$000's)			
Revenue	\$60,087	\$38,837	\$ 21,250	55%
Less sales reserves:				
Provision for Medicaid rebates	13,135	9,749	3,386	35%
Provision for chargebacks	16	—	16	100%
Provision for Coverage Gap Discount	—	—	—	—
Provision for Tricare	337	369	(32)	-9%
Co-payment assistance and other	619	403	216	54%
Total sales reserves	14,107	10,521	3,586	34%
Net sales	<u>\$45,980</u>	<u>\$28,316</u>	<u>\$ 17,664</u>	62%

Net sales for the three months ended June 30, 2011 and 2010 were comprised of net sales of our products Acthar and Doral. Net sales of Acthar for the three months ended June 30, 2011 totaled \$45.8 million as compared to \$28.2 million during the same period in 2010. Net sales for the three months ended June 30, 2011 were positively affected by increased unit demand from CuraScript SD, our distributor for Acthar. We shipped 2,430 vials for the three months ended June 30, 2011 as compared to 1,680 vials shipped for the three months ended June 30, 2010. Net sales also increased due to changes in the price we charge CuraScript SD for Acthar. We increased the price we charge CuraScript SD for Acthar by 5% in January 2011 and June 2011.

While we do not receive complete information regarding prescriptions by therapeutic area, we believe increased demand from CuraScript SD was driven by strong prescription growth in each of our primary therapeutic areas: MS, NS and IS. During the three months ended June 30, 2011, the number of prescriptions for Acthar to treat MS exacerbations increased to 751 from 304 in the quarter ended June 30, 2010, which was attributable to our expanded sales force calling on physicians who treat patients with MS. In addition, during the first half of 2011, we hired, trained and deployed five sales representatives who in March 2011 started promoting Acthar exclusively to nephrologists for use in treating NS. In the quarter ended June 30, 2011, this pilot selling effort resulted in 45 new, paid NS prescriptions, a significant increase over the four new, paid NS prescriptions in the quarter ended June 30, 2010. We also experienced a strong quarter with respect to IS prescriptions, though the 106 paid, new IS prescriptions in the second quarter was within the normal historic range. These prescription figures are based on internal company estimates and are subject to change as discussed in the "Important Notes Regarding Prescription Data" to the prescription table on page 19 of this report. As Acthar is already considered by most child neurologists to be the treatment of choice in IS, we will be reducing the number of sales calls to child neurologist, in order to make time available for our Specialty Sales Force to have a limited supportive role in our expanded nephrology effort.

Total sales reserves decreased as a percentage of gross revenue to 23.5% for the quarter ended June 30, 2011 from 27.1% for the quarter ended June 30, 2010. We utilize a multi-step approach to determine our sales reserves each quarter, which includes an analysis of a predictive model, a review of Medicaid and other invoices received during the quarter and an estimate of in channel inventory. We believe the decrease in total sales reserves as a percent of gross revenue was primarily due to the increased percentage of our business attributable to MS and NS, as IS prescriptions have a higher Medicaid incidence rate than MS or NS prescriptions.

On a sequential basis, net sales increased by \$9.2 million to \$46.0 million in the three months ended June 30, 2011, compared to \$36.8 million in the three months ended March 31, 2011.

While we believe that we have not significantly penetrated our MS market, we cannot assure you that the November 2010 expansion of our Specialty Sales Force will continue to generate incremental MS prescriptions. Additionally, it is unclear whether our strategy to expand our Nephrology selling effort will be successful. Acthar orders may be affected by several factors, including inventory levels at specialty pharmacies and hospitals, greater use of patient assistance programs, the overall pattern of usage by the health care community, including Medicaid and government-supported entities, the use of alternative therapies, and the reimbursement policies of insurance companies. Our specialty distributor ships Acthar to specialty pharmacies and hospitals to meet end user demand. We track our own Acthar shipments daily, but those shipments vary compared to end user demand because of changes in inventory levels at specialty pharmacies and hospitals.

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Cost of Sales and Gross Profit

	Three Months Ended June 30,		Increase/ (Decrease)	% Change
	2011	2010		
	(in \$000's)			
Cost of sales	\$ 2,856	\$ 2,000	\$ 856	43%
Gross profit	\$43,124	\$26,316	\$ 16,808	64%
Gross margin	94%	93%		

Cost of sales was \$2.9 million for the three months ended June 30, 2011, as compared to \$2.0 million for the three months ended June 30, 2010. We include in cost of sales material costs, packaging, warehousing and distribution, product liability insurance, royalties, quality control (which primarily includes product stability testing), quality assurance and reserves for excess or obsolete inventory. Our gross margin was 94%, or \$43.1 million, for the three months ended June 30, 2011, as compared to 93%, or \$26.3 million for the three months ended June 30, 2010. The increase in gross margin in 2011 as compared to 2010 is primarily the result of the 5% price increase that was taken in January and June 2011, continued growth in paid prescriptions for patients with MS and a reduction in direct material costs, offset by an increase in royalties on Acthar net sales. The manufacturing process for Acthar is complex and problems may arise during manufacturing for a variety of reasons, including equipment malfunction, failure to follow specific protocols and procedures, problems with raw materials, natural disasters, and environmental factors.

Selling and Marketing

	Three Months Ended June 30,		Increase/ (Decrease)	% Change
	2011	2010		
	(in \$000's)			
Selling and marketing expense	\$14,746	\$6,028	\$ 8,718	145%

Selling and marketing expenses were \$14.7 million for the three months ended June 30, 2011, as compared to \$6.0 million for the three months ended June 30, 2010. The increase of \$8.7 million in 2011 as compared to 2010 is due primarily to increases in headcount-related costs and costs associated with our expanded sales and marketing effort. During the latter part of 2010, to further build on positive prescription trends, we increased the size of our Specialty Sales Force, which calls on neurologists who treat patients for MS or IS, from 38 representatives to 77 representatives effective November 2010. Additionally, in March 2011, we assembled a Nephrology Sales Force which promotes Acthar exclusively to nephrologists for use in treating NS. Our initial Nephrology Sales Force was comprised of just five representatives and, based on the results of their efforts we are significantly expanding our NS selling effort. Specifically, we are in the process of hiring approximately 23 additional representatives for our Nephrology Sales Force and we are giving a limited supportive selling role to our 77 representative Specialty Sales Force.

We cannot guarantee you that this prescription growth trend will continue or that our strategy to expand our Nephrology selling effort will be successful, and, even if it is successful in the long-term, our sales force expansion may impact negatively our short-term financial results.

General and Administrative

	Three Months Ended June 30,		Increase/ (Decrease)	% Change
	2011	2010		
	(in \$000's)			
General and administrative expense	\$ 3,791	\$ 2,943	\$ 848	29%

General and administrative expenses were \$3.8 million for the three months ended June 30, 2011, as compared to \$2.9 million for the three months ended June 30, 2010. The increase of \$0.8 million in 2011 as compared to 2010 is due primarily to increases in costs to support the increased sales and marketing teams and the increase in share-based compensation which is directly correlated to the increase in our stock price year over year.

Research and Development

	Three Months Ended June 30,		Increase/ (Decrease)	% Change
	2011	2010		
	(in \$000's)			
Research and development	\$ 3,891	\$ 2,943	\$ 948	32%

Research and development expenses were \$3.9 million in the three months ended June 30, 2011, as compared to \$2.9 million for the three months ended June 30, 2010. The increase in research and development expenses in 2011 as compared to 2010 was primarily due to increases in headcount related costs to support our efforts to explore the use of Acthar as a therapeutic alternative for the treatment of NS, costs incurred associated with the initiation of our Phase IV dose response clinical trial for idiopathic membranous nephropathy, offsetting a reduction in costs which occurred during the three months ended June 30, 2010 associated with the IS supplemental New Drug Application. Costs included in research and development also include costs associated with the funding of medical research projects to better understand the therapeutic benefit of Acthar in current and new therapeutic applications, product development efforts and compliance activities.

We plan to continue our research and development efforts to explore the use of Acthar as a therapeutic alternative for the treatment of NS. In 2010, we supported investigator-initiated studies in patients with idiopathic membranous nephropathy (on-label) and diabetic nephropathy (not on-label). Based on the results of these investigations, we have started a Phase IV dose response clinical trial for idiopathic membranous nephropathy and are developing a clinical protocol for a Company-sponsored study to evaluate the safety and efficacy of Acthar in treating diabetic nephropathy. These clinical trials will result in a significant increase in research and development expenses in the second half of 2011 through 2013. We may also pursue clinical trials to evaluate the use of Acthar to treat other therapeutic uses, including conditions that are not currently on the label of approved indications for Acthar.

Share-based compensation costs. Total share-based compensation costs for the three months ended June 30, 2011 and 2010 were \$1.7 million and \$0.9 million, respectively. For the three months ended June 30, 2011, we granted options to employees and non-employee directors to purchase 43,000 shares of our common stock at a weighted average exercise price of \$21.90 per share, which was equal to the fair market value of our common stock on the date of the grant. During the first quarter of 2011, we issued 274,000 performance-based options. These performance-based options include a one-time performance achievement, followed by a time-based vesting of an additional 12 months, should the performance be achieved. During the quarter ended June 30, 2011, we determined that achievement of the one-time performance milestone was reasonably estimated and probable. As such, we recorded share-based compensation costs related to these performance-based options.

In addition to stock options, we may also issue restricted stock awards to certain employees. No restricted stock awards were issued for the three months ended June 30, 2011. The total share-based compensation costs for the three months ended June 31, 2011 and 2010 included \$9,000 and \$14,000, respectively, related to these restricted stock awards. The following table sets forth our share-based compensation costs for the three months ended June 30, 2011 and 2010, respectively:

	Three Months Ended June 30,	
	2011	2010
Selling and marketing	\$ 415	\$ 238
General and administrative	1,041	442
Research and development	260	240
Total	<u>\$ 1,716</u>	<u>\$ 920</u>

Income tax expense. Income tax expense for the three months ended June 30, 2011 was \$6.7 million, as compared to \$5.1 million for the three months ended June 30, 2010. The increase in income tax expense of \$1.6 million in 2011 as compared to 2010 was due to an increase in net sales (resulting in a higher basis for income taxes) offset by a reduction in our effective tax rate. For the three months ended June 30, 2011, we used the single sales factor methodology for California, which has resulted in tax savings of approximately \$0.5 million. Because most of our sales are sourced outside of California, we do not expect to continue to pay significant income taxes in California.

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Six months ended June 30, 2011 compared to the six months ended June 30, 2010:

Recorded Net Sales

	Six Months Ended June 30,		Increase/ (Decrease)	% Change
	2011	2010		
	(in \$000's)			
Revenue	\$108,717	\$72,298	\$ 36,419	50%
Less sales reserves:				
Provision for Medicaid rebates	24,124	16,350	7,774	48%
Provision for chargebacks	120	—	120	100%
Provision for Coverage Gap Discount	120	—	120	100%
Provision for Tricare	604	561	43	8%
Co-payment assistance and other	936	827	109	13%
Total sales reserves	25,904	17,738	8,166	46%
Net sales	\$ 82,813	\$54,560	\$ 28,253	52%

Net sales for the six months ended June 30, 2011 and 2010 were comprised of net sales of our products Acthar and Doral. Net sales of Acthar for the six months ended June 30, 2011 totaled \$82.6 million as compared to \$54.3 million during the same period in 2010. Net sales for the six months ended June 30, 2011 were positively affected by increased unit demand from CuraScript SD, our distributor for Acthar. We shipped 4,440 vials for the six months ended June 30, 2011 as compared to 3,126 vials shipped for the six months ended June 30, 2010. Effective January 3, 2011, we increased the price we charge CuraScript SD for Acthar by 5%. We increased the price we charge CuraScript SD for Acthar by another 5% on June 1, 2011.

During the six months ended June 30, 2011, we achieved a significant increase in the number of prescriptions for Acthar to treat MS exacerbations, which was attributable to our expanded sales force calling on physicians who treat patients with MS. While there can be a delay between changes in end-user demand and the impact of such changes on the level of orders we receive from our distributor, MS paid prescriptions increased by approximately 135% in the first six months, from 535 to 1,259 prescriptions and we believe that this increase resulted in higher net sales in the period.

In addition, during the first half of 2011, we hired, trained and deployed five sales representatives who in March 2011 started marketing Acthar exclusively to nephrologists for use in treating NS. This pilot selling effort proved to be successful by showing a significant increase in paid NS prescriptions from 15 to 63 prescriptions, or a 320% increase, for the six months ended June 30, 2011.

Total sales reserves decreased as a percentage of gross revenue to 23.8% for the six months ended June 30, 2011 from 24.5% for the quarter ended June 30, 2010, primarily due to a proportionate decrease in our Medicaid reserve, as IS prescriptions have a higher Medicaid incidence rate than MS or NS prescriptions. This percentage decrease is offset by the addition of reserves related to Medicaid Managed Care Organizations, which we began accruing in connection with the adoption of the Health Care Reform Acts on March 23, 2010.

While we believe that we have not significantly penetrated our MS market, we cannot assure you that the November 2010 expansion of our Specialty Sales Force will continue to generate incremental MS prescriptions. Additionally, it is unclear whether our strategy to expand our Nephrology selling effort will be successful. Acthar orders may be affected by several factors, including inventory levels at specialty pharmacies and hospitals, greater use of patient assistance programs, the overall pattern of usage by the health care community, including Medicaid and government-supported entities, the use of alternative therapies, and the reimbursement policies of insurance companies. Our specialty distributor ships Acthar to specialty pharmacies and hospitals to meet end user demand. We track our own Acthar shipments daily, but those shipments vary compared to end user demand because of changes in inventory levels at specialty pharmacies and hospitals.

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Cost of Sales and Gross Profit

	Six Months Ended June 30,		Increase/ (Decrease)	% Change
	2011	2010		
Cost of sales	\$ 4,728	\$ 3,998	\$ 730	18%
Gross profit	\$78,085	\$50,562	\$ 27,523	54%
Gross margin	94%	93%		

Cost of sales was \$4.7 million for the six months ended June 30, 2011, as compared to \$4.0 million for the six months ended June 30, 2010. We include in cost of sales material costs, packaging, warehousing and distribution, product liability insurance, royalties, quality control (which primarily includes product stability testing), quality assurance and reserves for excess or obsolete inventory. Our gross margin was 94%, or \$78.1 million, for the six months ended June 30, 2011, as compared to 93%, or \$50.6 million for the six months ended June 30, 2010. The increase in gross margin in 2011 as compared to 2010 is primarily the result of two 5% price increases that were taken on January 1, 2011 and June 1, 2011, continued growth in paid prescriptions for patients with MS and a reduction in direct material costs, offset by an increase in royalties on Acthar net sales. The manufacturing process for Acthar is complex and problems may arise during manufacturing for a variety of reasons, including equipment malfunction, failure to follow specific protocols and procedures, problems with raw materials, natural disasters, and environmental factors.

Selling and Marketing

	Six Months Ended June 30,		Increase/ (Decrease)	% Change
	2011	2010		
Selling and marketing expense	\$25,998	\$12,678	\$ 13,320	105%

Selling and marketing expenses were \$26.0 million for the six months ended June 30, 2011, as compared to \$12.7 million for the six months ended June 30, 2010. The increase of \$13.3 million in 2011 as compared to 2010 is due primarily to increases in headcount-related costs and costs associated with our expanded sales and marketing effort. During the latter part of 2010, to further build on positive prescription trends, we doubled the size of our sales organization, increasing the sales force to 77 Acthar specialists and an additional five nephrology sales representatives.

We cannot guarantee you that this prescription growth trend will continue or that our sales force expansion will be successful, and, even if it is successful in the long-term, our sales force expansion may impact negatively our short-term financial results.

General and Administrative

	Six Months Ended June 30,		Increase/ (Decrease)	% Change
	2011	2010		
General and administrative expense	\$7,663	\$5,669	\$ 1,994	35%

General and administrative expenses were \$7.7 million for the six months ended June 30, 2011, as compared to \$5.7 million for the six months ended June 30, 2010. The increase of \$2.0 million in 2011 as compared to 2010 is due primarily to increases in headcount related costs to support the increased sales and marketing teams and the increase in share-based compensation which is directly correlated to the increase in our stock price year over year.

Research and Development

	Six Months Ended June 30,		Increase/ (Decrease)	% Change
	2011	2010		
Research and development	\$6,872	\$5,690	\$ 1,182	21%

Research and development expenses were \$6.9 million in the six months ended June 30, 2011, as compared to \$5.7 million for the six months ended June 30, 2010. The increase in research and development expenses in 2011 as compared to 2010 was primarily due to increases in headcount related costs to support our efforts to explore the use of Acthar as a therapeutic alternative for the treatment of NS, costs incurred associated with the initiation of our Phase IV dose response clinical trial for

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idiopathic membranous nephropathy, offsetting a reduction in costs which occurred during the three months ended June 30, 2010 associated with the IS supplemental New Drug Application. Costs included in research and development also include costs associated with the funding of medical research projects to better understand the therapeutic benefit of Acthar in current and new therapeutic applications, product development efforts and compliance activities.

We plan to continue our research and development efforts to explore the use of Acthar as a therapeutic alternative for the treatment of NS. In 2010, we supported investigator-initiated studies in patients with idiopathic membranous nephropathy (on-label) and diabetic nephropathy (not on-label). Based on the results of these investigations, we have started a Phase IV dose response clinical trial for idiopathic membranous nephropathy and are developing a clinical protocol for a Company-sponsored study to evaluate the safety and efficacy of Acthar in treating diabetic nephropathy. These clinical trials will result in a significant increase in research and development expenses in the second half of 2011 through 2013. We may also pursue clinical trials to evaluate the use of Acthar to treat other therapeutic uses, including conditions that are not currently on the label of approved indications for Acthar.

Share-based compensation costs. Total share-based compensation costs for the six months ended June 30, 2011 and 2010 were \$3.5 million and \$1.9 million, respectively. For the six months ended June 30, 2011, we granted options to employees and non-employee directors to purchase 1.3 million shares of our common stock at a weighted average exercise price of \$14.80 per share, which was equal to the fair market value of our common stock on the date of the grant. Included in the 1.3 million options granted during the quarter, were 274,000 performance-based options. These performance-based options include a one-time performance achievement, followed by a time-based vesting of an additional 12 months, should the performance be achieved. During the quarter ended June 30, 2011, we determined that achievement of the one-time performance milestone was reasonably estimated and probable. As such, we recorded share-based compensation costs related to these performance-based options.

In addition to stock options, we may also issue restricted stock awards to certain employees. The total share-based compensation costs for the six months ended June 30, 2011 and 2010 included \$19,000 and \$27,000, respectively, related to these restricted stock awards. The following table sets forth our share-based compensation costs for the six months ended June 30, 2011 and 2010, respectively:

	Six Months Ended	
	June 30,	
	2011	2010
Selling and marketing	\$ 789	\$ 470
General and administrative	2,167	1,017
Research and development	572	462
Total	<u>\$3,528</u>	<u>\$1,949</u>

Income tax expense. Income tax expense for the six months ended June 30, 2011 was \$12.1 million, as compared to \$9.4 million for the six months ended June 30, 2010. The increase in income tax expense of \$2.7 million in 2011 as compared to 2010 was due to an increase in net sales (resulting in a higher basis for income taxes) offset by a reduction in our effective tax rate. Beginning in 2011, we intend to use the single sales factor methodology for California, which has resulted in tax savings for the six months ended June 30, 2011 by approximately \$0.9 million. Because most of our sales are sourced outside of California, we do not expect to continue to pay significant income taxes in California.

Liquidity and Capital Resources

Cash and cash equivalents, short term investments and working capital as of June 30, 2011 and December 31, 2010 were as follows:

Financial Assets:

	June 30, 2011	December 31, 2010
Cash and cash equivalents	\$ 65,277	\$ 41,508
Short term investments	63,849	73,324
Cash, cash equivalents and short term investments	<u>\$ 129,126</u>	<u>\$ 114,832</u>

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Select measures of liquidity and capital resources:

	June 30, 2011	December 31, 2010
Current assets	\$ 179,785	\$ 143,499
Current liabilities	(45,388)	(31,511)
Working Capital	<u>\$ 134,397</u>	<u>\$ 111,988</u>
Current ratio	<u>3.96</u>	<u>4.55</u>

Until required for use in our business, we invest our cash reserves in money market funds and high quality commercial, corporate and U.S. government and agency bonds in accordance with our investment policy. The objective of our investment policy is to preserve capital, provide liquidity consistent with forecasted cash flow requirements, maintain appropriate diversification and generate returns relative to these investment objectives and prevailing market conditions.

The increase in cash, cash equivalents and short term investments was primarily due to the increase in net sales and the related cash generated from operations, the proceeds from maturities of short term investments, offset by the repurchase of shares of our common stock through our approved stock repurchase plan. The increase in our working capital was primarily due to increases in our cash, cash equivalents and short term investments, accounts receivable and inventories, offset primarily by increases in our sales-related reserves and our income taxes payable.

We expect continued growth in our research and development expenses, particularly those related to clinical trials associated with our on-label indication for NS. However, we anticipate that cash generated from operations and our existing cash, cash equivalents and short term investments should provide us adequate resources to fund our operations as currently planned for the foreseeable future.

Cash Flows

Change in cash and cash equivalents:

	Six Months Ended		Increase/ (Decrease)
	June 30,		
	2011	2010	
Net cash flows provided by operating activities	\$21,505	\$ 16,245	\$ 5,260
Net cash flows provided by / (used in) investing activities	7,714	(39,393)	47,107
Net cash flows (used in) / provided by financing activities	(5,450)	1,143	(6,593)
Net change in cash and cash equivalents	<u>\$23,769</u>	<u>(\$ 22,005)</u>	<u>\$ 45,774</u>

Operating Activities

The increase in net cash and cash equivalents from June 30, 2010 is primarily due to the net income achieved in 2011 versus the net income achieved in the same period in 2010. The components of cash flows from operating activities, as reported on our Consolidated Statement of Cash Flows, are as follows:

- Our reported net income, adjusted for non-cash items, including share-based compensation expense, deferred income taxes, amortization of investments, depreciation and amortization and loss on disposal of property and equipment was \$30.0 million and \$19.7 million for the six months ended June 30, 2011 and 2010, respectively.
- Net cash outflow due to changes in operating assets and liabilities was (\$8.5) million for the six months ended June 30, 2011 and net cash outflow was (\$3.4) million for the six months ended June 30, 2010. The (\$8.5) million change in operating assets and liabilities primarily relates to an increase in our accounts receivable of \$12.6 million due to an increase in net sales, offset by an increase in sales-related reserves of \$5.6 million, which relates to an increase in Acthar gross sales.

Investing Activities

The components of cash flows from investing activities consisted of the following:

- Purchases of property and equipment of \$1.3 million;
- Purchases of short term investments of \$53.9 million; and
- Maturities of short term investments of \$63.0 million.

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Financing Activities

Net cash flows from financing activities reflected the following:

- the income tax benefit realized on our share-based compensation plans of \$3.7 million;
- the issuance of common stock related to the exercise of stock options for \$2.3 million; offset by
- the repurchase of shares of our common stock of \$11.5 million to repurchase 884,300 shares of our common stock. Under this stock repurchase plan, we have repurchased a total of 9.2 million shares of our common stock for \$48.1 million through June 30, 2011, at an average price of \$5.21 per share. Additionally, we have purchased 6.2 million shares of our common stock outside of our stock repurchase plan for a total of \$30.4 million through June 30, 2011 at an average price of \$4.93 per share for a total repurchase value of \$78.5 million. As of June 30, 2011, there are 4.3 million shares authorized remaining under our stock repurchase plan.

We do not currently intend to conduct business development activities which would utilize a material portion of our liquidity. We review our level of liquidity and anticipated cash needs for the business on an ongoing basis, and consider whether to return additional capital to our shareholders as well as alternative methods to return capital.

Recent Accounting Pronouncements

In April 2011, the FASB issued Accounting Standards Update No. 2011-04 “Fair Value Measurement (Topic 820) – Amendments to Achieve Common Fair Value Measurement and Disclosure Requirements in U.S. GAAP and IFRS,” or ASU No. 2011-04. ASU No. 2011-04 is the result of the continuing convergence projects between the FASB, and the International Accounting Standards Board, or IASB, to create a common set of high quality global accounting standards. The amendments in ASU No. 2011-04 explain how to measure fair value. They do not require additional fair value measurements and are not intended to establish standards or affect valuation practices outside of financial reporting. The amendment will be effective for interim and annual periods beginning after December 15, 2011. We plan to adopt ASU No. 2011-04 and do not anticipate a material effect on our financial position or results of operation.

In June 2011, the FASB issued Accounting Standards Update No. 2011-05 “Presentation of Comprehensive Income,” or ASU No. 2011-05. ASU No. 2011-05 improves the comparability, consistency, and transparency of financial reporting and increases the prominence of items reported in other comprehensive income, or OCI, by eliminating the option to present OCI as part of the statement of changes in shareholders’ equity. The amendments in this standard require that all non-owner changes in stockholders’ equity be presented either in a single continuous statement of comprehensive income or in two separate but consecutive statements. Under either method, adjustments must be displayed for items that are reclassified from OCI to net income, in both net income and OCI. The standard does not change the current option for presenting components of OCI gross or net of the effect of income taxes, provided that such tax effects are presented in the statement in which OCI is presented or disclosed in the notes to the financial statements. Additionally, the standard does not affect the calculation or reporting of earnings per share. For public entities, the amendments in this ASU are effective for fiscal years, and interim periods within those years, beginning after December 15, 2011 and are to be applied retrospectively, with early adoption permitted. We plan to adopt ASU No. 2011-05 and do not anticipate a material effect on our financial position or results of operation.

ITEM 3. QUANTITATIVE AND QUALITATIVE DISCLOSURES ABOUT MARKET RISK

Our exposure to market risk at June 30, 2011 has not changed materially from December 31, 2010, and reference is made to the more detailed disclosures of market risk included in our Annual Report on Form 10-K for the year ended December 31, 2010, as filed with the Securities and Exchange Commission, or SEC, on February 23, 2011.

ITEM 4. CONTROLS AND PROCEDURES

(a) Evaluation of Disclosure Controls and Procedures

We maintain disclosure controls and procedures that are designed to ensure that information required to be disclosed in our reports filed pursuant to the Securities Exchange Act of 1934, as amended, or Exchange Act, is recorded, processed, summarized and reported within the time periods specified in the SEC’s rules and forms and that such information is accumulated and communicated to our management, including our Chief Executive Officer and Principal Accounting Officer (principal executive officer) and Chief Financial Officer (principal financial officer), as appropriate, to allow for timely decisions regarding required disclosure.

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In designing and evaluating the disclosure controls and procedures, management recognizes that any controls and procedures, no matter how well designed and operated, can provide only reasonable assurance of achieving the desired control objectives, and management is required to apply its judgment in evaluating the cost-benefit relationship of possible controls and procedures. Our disclosure controls and procedures were designed to provide reasonable assurance that the controls and procedures would meet their objectives.

As required by SEC Rule 13a-15(b), we carried out an evaluation, under the supervision and with the participation of our management, including our Chief Executive Officer (principal executive officer) and Chief Financial Officer (principal financial officer), of the effectiveness of the design and operation of our disclosure controls and procedures as of the end of the quarter covered by this quarterly report on Form 10-Q. Based on the foregoing, our Chief Executive Officer (principal executive officer) and Chief Financial Officer (principal financial officer) concluded that our disclosure controls and procedures were effective as of June 30, 2011.

There has been no change in our internal controls over financial reporting during our most recent fiscal quarter that has materially affected, or is reasonably likely to affect materially, our internal controls over financial reporting.

PART II. OTHER INFORMATION

ITEM 1. LEGAL PROCEEDINGS

We operate in a highly regulated industry. We are subject to the regulatory authority of the SEC, the FDA and numerous other federal and state governmental agencies including state attorney general offices, which have become more active in investigating the business practices of pharmaceutical companies. From time to time, we receive requests for information from various governmental agencies. In addition, from time to time, we may become involved in litigation relating to claims arising from our ordinary course of business. In June 2011, Glenridge Pharmaceuticals LLC filed a lawsuit against us in Superior Court of California, Santa Clara County, alleging that we had underpaid royalties to Glenridge, in connection with the timing of the impact of various offsets in the calculation of net sales. We intend to defend this lawsuit vigorously. We are not aware of any claims or actions pending or threatened against us, the ultimate disposition of which we believe would have a material adverse effect on us.

ITEM 1A. RISK FACTORS

Information about material risks related to our business, financial condition and results of operations for the quarterly period ended June 30, 2011 does not materially differ from that described in Part I, Item 1A of our Annual Report on Form 10-K for the year ended December 31, 2010, as filed with the SEC on February 23, 2011, other than the following additional risk factor:

Changes in treatment regimens or patient compliance may adversely affect our business.

Recommended treatment regimens among physicians prescribing Acthar for use in treating MS exacerbations, NS and IS vary within each therapeutic area. If physicians prescribe a lower number of vials for the treatment of MS exacerbations, NS or IS, our net sales from the sale of Acthar would decline. Additionally, we are aware that some prescriptions are initially for a lower number of vials than is necessary to complete the physician's recommended treatment regimen, and allow for one or more prescription refills. If patients do not obtain their refill prescriptions in order to complete their recommended treatment regimen, our net sales from the sale of Acthar would decline. There can be no assurance that we would be able to increase prescription levels by enough to offset any such decline in vials per prescription.

Forward Looking Statements

This Quarterly Report on Form 10-Q contains forward-looking statements that involve risks and uncertainties. Our actual results could differ materially from those discussed herein. Factors that could cause or contribute to such differences include, but are not limited to:

- our reliance on Acthar for substantially all of our net sales and profits;
- reductions in vials used per prescription resulting from changes in physician recommended treatment regimens or patient compliance with physician recommendations;
- the complex nature of our manufacturing process and the potential for supply disruptions or other business disruptions;
- the lack of patent protection for Acthar, and the possible FDA approval and market introduction of competitive products;
- our ability to generate revenue from sales of Acthar to treat on-label indications associated with NS, and our ability to develop other therapeutic uses for Acthar, including SLE;
- research and development risks, including risks associated with our clinical trials with respect to NS and potential clinical trials with respect to SLE, and our reliance on third-parties to conduct research and development and the ability of research and development to generate successful results;
- regulatory changes or other policy actions by governmental authorities and other third parties as the Health Care Reform Acts are implemented or efforts to reduce federal and/or state government deficits;
- our ability to receive high reimbursement levels from third party payers;
- an increase in the proportion of our Acthar unit sales comprised of Medicaid-eligible patients and government entities;
- our ability to estimate reserves required for Acthar used by government entities and Medicaid-eligible patients and the impact that unforeseen invoicing of historical Medicaid prescriptions may have upon our results;
- our ability to operate within an industry that is highly regulated at both the Federal and state level;

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- our ability to effectively manage our growth, including the expansion of our NS selling effort, and our reliance on key personnel;
- the impact to our business caused by economic conditions;
- our ability to protect our proprietary rights;
- our ability to maintain effective controls over financial reporting;
- the risk of product liability lawsuits;
- unforeseen business interruptions;
- volatility in our monthly and quarterly Acthar shipments and end-user demand, as well as volatility in our stock price; and
- other risks discussed in our Annual Report on Form 10-K for the year ended December 31, 2010, as filed with the SEC on February 23, 2011, and other documents filed with the SEC.

ITEM 2-5.

Not applicable.

ITEM 6. EXHIBITS

<u>Exhibit No</u>	<u>Description</u>
10.1	Amended and Restated 2006 Equity Incentive Plan.
10.2	Amended and Restated Employee Stock Purchase Plan.
31.1	Certification of Principal Executive Officer Pursuant to Rule 13a-14(a)/15d-14(a) of the Securities Exchange Act of 1934.
31.2	Certification of Principal Financial Officer Pursuant to Rule 13a-14(a)/15d-14(a) of the Securities Exchange Act of 1934.
32.1	Certification of Principal Executive Officer Pursuant to Rule 13a-14(b)/15d-14(b) of the Securities Exchange Act of 1934 and 18 U.S.C. Section 1350.
32.2	Certification of Principal Financial Officer Pursuant to Rule 13a-14(b)/15d-14(b) of the Securities Exchange Act of 1934 and 18 U.S.C. Section 1350.
101	The following financial statements are from Questcor Pharmaceutical, Inc.'s Quarterly Report on Form 10-Q for the quarter ended June 30, 2011, are furnished herewith, formatted in Extensible Business Reporting Language ("XBRL"): (i) the Condensed Consolidated Balance Sheets, (ii) the Condensed Consolidated Statements of Income, (iii) the Condensed Consolidated Statements of Cash Flows, and (iv) the Notes to the Condensed Consolidated Financial Statements, tagged as blocks of text.

SIGNATURES

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

Date: July 29, 2011

QUESTCOR PHARMACEUTICALS, INC.

By: /s/ Don M. Bailey
Don M. Bailey
President and Chief Executive Officer
(Principal Executive Officer)

By: /s/ Michael H. Mulroy
Michael H. Mulroy
Chief Financial Officer and General Counsel
(Principal Financial Officer)

By: /s/ Kristine Engelke
Kristine Engelke
Controller
(Principal Accounting Officer)

Exhibit Index

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**QUESTCOR PHARMACEUTICALS, INC.
2006 EQUITY INCENTIVE AWARD PLAN**

ARTICLE I

PURPOSE

The purpose of the Questcor Pharmaceuticals, Inc. 2006 Equity Incentive Award Plan (the “**Plan**”) is to promote the success and enhance the value of Questcor Pharmaceuticals, Inc., a California corporation (the “**Company**”), by linking the personal interests of the members of the Board, Employees, and Consultants to those of Company shareholders and by providing such individuals with an incentive for performance to generate returns to Company shareholders. The Plan is further intended to provide flexibility to the Company in its ability to motivate, attract, and retain the services of members of the Board, Employees, and Consultants upon whose judgment, interest, and special effort the successful conduct of the Company’s operation is largely dependent.

ARTICLE II

DEFINITIONS AND CONSTRUCTION

Wherever the following terms are used in the Plan they shall have the meanings specified below, unless the context clearly indicates otherwise. The singular pronoun shall include the plural where the context so indicates.

2.1 “**Administrator**” means the entity that conducts the general administration of the Plan as provided herein. With reference to the administration of the Plan with respect to Awards granted to Independent Directors, the term “**Administrator**” shall refer to the Board. With reference to the administration of the Plan with respect to any other Award, the term “Administrator” shall refer to the Committee unless the Board has assumed the authority for administration of the Plan generally as provided in Section 13.1. With reference to the duties of the Committee under the Plan which have been delegated to one or more persons pursuant to Section 13.5, the term “**Administrator**” shall refer to such person(s) unless the Committee or the Board has revoked such delegation.

2.2 “**Award**” means an Option, a Restricted Stock award, a Stock Appreciation Right award, a Dividend Equivalents award, a Stock Payment award, a Restricted Stock Unit award or a Performance-Based Award granted to a Participant pursuant to the Plan.

2.3 “**Award Agreement**” means any written or electronic agreement, contract, or other instrument or document evidencing an Award.

2.4 “**Board**” means the Board of Directors of the Company.

2.5 “**Cause,**” unless otherwise defined in an employment or services agreement between the Participant and the Company or any Parent or Subsidiary or Successor Entity, means:

- (a) a Participant’s conviction or no contest plea to a felony, or a crime involving moral turpitude, under any federal or state criminal law;

(b) the commission of a fraud by a Participant against the Company or any Parent or Subsidiary or Successor Entity;

(c) with respect to a Participant who is an Employee, the repeated, unexplained or unjustified absence by an Employee from the Company or any Parent or Subsidiary or Successor Entity; or

(d) the gross negligence or willful misconduct of a Participant where such gross negligence or willful misconduct has resulted or is likely to result in substantial and material damage to the Company or any Parent or Subsidiary or Successor Entity.

The existence of “Cause” shall be determined by the Administrator, in its sole discretion. The foregoing definition shall not in any way preclude or restrict the right of the Company or any Parent or Subsidiary or Successor Entity to discharge or dismiss any Participant or other person in the service of the Company or any Parent or Subsidiary or Successor Entity for any other acts or omissions, but such other acts or omissions shall not be deemed, for purposes of the Plan, to constitute grounds for termination for Cause.

2.6 “Change in Control” means and includes each of the following:

(a) the acquisition, directly or indirectly, by any “person” or “group” (as those terms are defined in Sections 3(a)(9), 13(d) and 14(d) of the Exchange Act and the rules thereunder) of “beneficial ownership” (as determined pursuant to Rule 13d-3 under the Exchange Act) of securities entitled to vote generally in the election of directors (“voting securities”) of the Company that represent 50% or more of the combined voting power of the Company’s then outstanding voting securities, other than:

(i) an acquisition by a trustee or other fiduciary holding securities under any employee benefit plan (or related trust) sponsored or maintained by the Company or any person controlled by the Company or by any employee benefit plan (or related trust) sponsored or maintained by the Company or any person controlled by the Company, or

(ii) an acquisition of voting securities by the Company or a corporation owned, directly or indirectly by the shareholders of the Company in substantially the same proportions as their ownership of the stock of the Company.

(b) individuals who, as of the Effective Date, constitute the Board together with any new director(s) whose appointment or election by the Board or nomination for election by the Company’s shareholders was approved by a vote of at a majority of the directors then still in office who either were directors on the Effective Date or whose election or nomination for election was previously so approved (other than any director designated by a person who shall have entered into an agreement with the Company to effect a transaction described in Section 2.6(a) or Section 2.6(c)), cease for any reason to constitute a majority thereof;

(c) the consummation by the Company (whether directly involving the Company or indirectly involving the Company through one or more intermediaries) of (x) a merger, consolidation, reorganization, or business combination or (y) a sale or other disposition of all or substantially all of the Company’s assets, in each case other than a transaction which results in the Company’s voting securities outstanding immediately before the transaction continuing to represent

(either by remaining outstanding or by being converted into voting securities of the Company or the person that, as a result of the transaction, controls, directly or indirectly, the Company or owns, directly or indirectly, all or substantially all of the Company's assets or otherwise succeeds to the business of the Company (the Company or such person, the "**Successor Entity**") directly or indirectly, at least a majority of the combined voting power of the Successor Entity's outstanding voting securities immediately after the transaction; and

(d) the liquidation or dissolution of the Company.

For purposes of subsection (a) above, the calculation of voting power shall be made as if the date of the acquisition were a record date for a vote of the Company's shareholders, and for purposes of subsection (c) above, the calculation of voting power shall be made as if the date of the consummation of the transaction were a record date for a vote of the Company's shareholders.

The Administrator shall have full and final authority, which shall be exercised in its discretion, to determine conclusively whether a Change in Control of the Company has occurred pursuant to the above definition, and the date of the occurrence of such Change in Control and any incidental matters relating thereto.

2.7 "**Code**" means the Internal Revenue Code of 1986, as amended from time to time, and the regulations issued thereunder.

2.8 "**Committee**" means the committee of the Board described in Article 13.

2.9 "**Consultant**" means any consultant or adviser if:

(a) The consultant or adviser renders bona fide services to the Company or any Parent or Subsidiary;

(b) The services rendered by the consultant or adviser are not in connection with the offer or sale of securities in a capital-raising transaction and do not directly or indirectly promote or maintain a market for the Company's securities; and

(c) The consultant or adviser is a natural person who has contracted directly with the Company or any Parent or Subsidiary to render such services.

2.10 "**Covered Employee**" means an Employee who is, or is likely to become, a "covered employee" within the meaning of Section 162(m)(3) of the Code.

2.11 "**Disability**" means a permanent and total disability within the meaning of Section 22(e)(3) of the Code, as it may be amended from time to time.

2.12 "**Dividend Equivalents**" means a right granted to a Participant pursuant to Article 8 to receive the equivalent value (in cash or Stock) of dividends paid on Stock.

2.13 "**Effective Date**" has the meaning set forth in Section 14.1.

2.14 "**Eligible Individual**" means any person who is a member of the Board, a Consultant or an Employee, as determined by the Administrator.

2.15 “**Employee**” means any officer or other employee (as defined in accordance with Section 3401(c) of the Code) of the Company or any Parent or Subsidiary.

2.16 “**Exchange Act**” means the Securities Exchange Act of 1934, as amended from time to time.

2.17 “**Expiration Date**” has the meaning set forth in Section 14.2.

2.18 “**Fair Market Value**” means, as of any date, the value of Stock determined as follows:

(a) If the Stock is listed on any established stock exchange or a national market system, including without limitation The Nasdaq Global Market or The Nasdaq Capital Market of The Nasdaq Stock Market, its Fair Market Value shall be the closing sales price for such stock as quoted on such exchange or system for the last market trading day prior to the date of determination for which a closing sales price is reported, as reported in The Wall Street Journal or such other source as the Administrator deems reliable;

(b) If the Stock is regularly quoted by a recognized securities dealer but selling prices are not reported, its Fair Market Value shall be the mean of the closing bid and asked prices for the Stock on the date prior to the date of determination as reported in The Wall Street Journal or such other source as the Administrator deems reliable; or

(c) In the absence of an established market for the Stock, the Fair Market Value thereof shall be determined in good faith by the Administrator.

2.19 “**Good Reason**” means, without a Participant’s express written consent, the occurrence of any of the following actions in connection with a Change in Control:

(a) a material reduction in job responsibilities given the Participant’s position with the Company or any Parent or Subsidiary or Successor Entity, and the Participant’s prior responsibilities with the Company or any Parent or Subsidiary or Successor Entity;

(b) any reduction in the Participant’s annual base compensation from the Company or any Parent or Subsidiary or Successor Entity as in effect immediately prior to such reduction; or

(c) a relocation of the Participant’s workplace for the Company or any Parent or Subsidiary or Successor Entity to a facility or location more than twenty-five (25) miles from the Participant’s workplace immediately prior to such relocation; provided, however, that the new workplace also increases the Participant’s commuting distance from the Participant’s primary residence.

A Participant shall provide thirty (30) day’s written notice to the Company or any Parent or Subsidiary or Successor Entity (whichever entity is the Participant’s employer) of any resignation for “**Good Reason.**”

2.20 “**Incentive Stock Option**” means an Option that is intended to be an incentive stock option and meets the requirements of Section 422 of the Code or any successor provision thereto.

2.21 “**Independent Director**” means a member of the Board who is not an Employee.

2.22 “**Minimum Vesting Awards**” has the meaning set forth in Section 11.7.

2.23 “**Non-Employee Director**” means a member of the Board who qualifies as a “Non-Employee Director” as defined in Rule 16b-3(b)(3) of the Exchange Act, or any successor rule.

2.24 “**Non-Qualified Stock Option**” means an Option that is not intended to be or otherwise does not qualify as an Incentive Stock Option.

2.25 “**Option**” means a right granted to a Participant pursuant to Article 5 of the Plan to purchase a specified number of shares of Stock at a specified price during specified time periods. An Option may be either an Incentive Stock Option or a Non-Qualified Stock Option.

2.26 “**Parent**” means any “**parent corporation**” as defined in Section 424(e) of the Code and any applicable regulations promulgated thereunder of the Company or any other entity which beneficially owns, directly or indirectly, a majority of the outstanding voting stock or voting power of the Company.

2.27 “**Participant**” means any Eligible Individual who, as a member of the Board, a Consultant or an Employee, has been granted an Award pursuant to the Plan.

2.28 “**Performance-Based Award**” means an Award granted to selected Covered Employees pursuant to Articles 6 and 8, but which is subject to the terms and conditions set forth in Article 9.

2.29 “**Performance Criteria**” means the criteria that the Administrator selects for purposes of establishing the Performance Goal or Performance Goals for a Participant for a Performance Period. The Performance Criteria that will be used to establish Performance Goals are limited to the following: net earnings (either before or after interest, taxes, depreciation and amortization), sales or revenue, net income (either before or after taxes), operating earnings, cash flow (including, but not limited to, operating cash flow and free cash flow), return on net assets, return on stockholders’ equity, return on sales, gross or net profit margin, working capital, earnings per share and price per share of Stock, the achievement of certain scientific milestones, customer retention rates, licensing, partnership or other strategic transactions, execution of a corporate collaboration agreement relating to a product candidate of the Company, acceptance by the U.S. Food and Drug Administration (“**FDA**”) or a comparable foreign regulatory authority of a final New Drug Application or similar document, approval for marketing of a product candidate of the Company by the FDA or a comparable foreign regulatory authority, obtaining a specified level of financing for the Company, as determined by the Committee, including through government grants (or similar awards) and the issuance of securities, commencement of a particular stage of clinical trials for a product candidate of the Company, or the achievement of one or more corporate, divisional or individual scientific or inventive measures. Any of the criteria identified above may be measured either in absolute terms or as compared to any incremental increase or as compared to results of a peer group. The Administrator shall, within the time prescribed by Section 162(m) of the Code, define in an objective fashion the manner of calculating the Performance Criteria it selects to use for such Performance Period for such Participant.

2.30 “**Performance Goals**” means, for a Performance Period, the goals established in writing by the Administrator for the Performance Period based upon the Performance Criteria. Depending on the Performance Criteria used to establish such Performance Goals, the Performance Goals may be expressed in terms of overall Company performance or the performance of a Subsidiary, division or other operational unit, or an individual. The Administrator, in its discretion, may, within the time prescribed by Section 162(m) of the Code, adjust or modify the calculation of Performance Goals for such Performance Period in order to prevent the dilution or enlargement of the rights of Participants (i) in the event of, or in anticipation of, any unusual or extraordinary corporate item, transaction, event, or development, or (ii) in recognition of, or in anticipation of, any other unusual or nonrecurring events affecting the Company, or the financial statements of the Company, or in response to, or in anticipation of, changes in applicable laws, regulations, accounting principles, or business conditions.

2.31 “**Performance Period**” means the one or more periods of time, which may be of varying and overlapping durations, as the Administrator may select, over which the attainment of one or more Performance Goals will be measured for the purpose of determining a Participant’s right to, and the payment of, a Performance-Based Award.

2.32 “**Plan**” means this Questcor Pharmaceuticals, Inc. 2006 Equity Incentive Award Plan, as it may be amended from time to time.

2.33 “**Qualified Performance-Based Compensation**” means any compensation that is intended to qualify as “qualified performance-based compensation” as described in Section 162(m)(4)(C) of the Code.

2.34 “**Restricted Stock**” means Stock awarded to a Participant pursuant to Article 6 that is subject to certain restrictions and may be subject to risk of forfeiture or repurchase.

2.35 “**Restricted Stock Unit**” means a right to receive a share of Stock during specified time periods granted pursuant to Section 8.3.

2.36 “**Securities Act**” means the Securities Act of 1933, as amended from time to time.

2.37 “**Section 409A Award**” has the meaning set forth in Section 10.1.

2.38 “**Stock**” means the common stock of the Company and such other securities of the Company that may be substituted for Stock pursuant to Article 12.

2.39 “**Stock Appreciation Right**” or “**SAR**” means a right granted pursuant to Article 7 to receive a payment equal to the excess of the Fair Market Value of a specified number of shares of Stock on the date the SAR is exercised over the Fair Market Value of such number of shares of Stock on the date the SAR was granted as set forth in the applicable Award Agreement.

2.40 “**Stock Payment**” means (a) a payment in the form of shares of Stock, or (b) an option or other right to purchase shares of Stock, as part of any bonus, deferred compensation or other arrangement, made in lieu of all or any portion of the compensation, granted pursuant to Section 8.2.

2.41 “**Subsidiary**” means any “subsidiary corporation” as defined in Section 424(f) of the Code and any applicable regulations promulgated thereunder of the Company or any other entity of which a majority of the outstanding voting stock or voting power is beneficially owned directly or indirectly by the Company.

2.42 “**Successor Entity**” has the meaning set forth in Section 2.6.

2.43 “**Termination of Consultancy**” means the time when the engagement of a Participant as a Consultant to the Company or a Parent or Subsidiary is terminated for any reason, with or without cause, including, but not by way of limitation, by resignation, discharge, death or retirement, but excluding terminations where there is a simultaneous commencement of employment with the Company or any Parent or Subsidiary. The Administrator, in its absolute discretion, shall determine the effect of all matters and questions relating to Termination of Consultancy, including, but not by way of limitation, the question of whether a Termination of Consultancy resulted from a discharge for good cause, and all questions of whether a particular leave of absence constitutes a Termination of Consultancy. Notwithstanding any other provision of the Plan, the Company or any Parent or Subsidiary has an absolute and unrestricted right to terminate a Consultant’s service at any time for any reason whatsoever, with or without cause, except to the extent expressly provided otherwise in writing.

2.44 “**Termination of Directorship**” shall mean the time when a Participant who is a Non-Employee Director ceases to be a member of the Board for any reason, including, but not by way of limitation, a termination by resignation, failure to be elected, death or retirement. The Board, in its sole and absolute discretion, shall determine the effect of all matters and questions relating to Termination of Directorship with respect to Non-Employee Directors.

2.45 “**Termination of Employment**” shall mean the time when the employee-employer relationship between a Participant and the Company or any Parent or Subsidiary is terminated for any reason, with or without cause, including, but not by way of limitation, a termination by resignation, discharge, death, disability or retirement; but excluding: (a) terminations where there is a simultaneous reemployment or continuing employment of a Participant by the Company or any Parent or Subsidiary, (b) at the discretion of the Administrator, terminations which result in a temporary severance of the employee-employer relationship, and (c) at the discretion of the Administrator, terminations which are followed by the simultaneous establishment of a consulting relationship by the Company or a Parent or Subsidiary with the former employee. The Administrator, in its absolute discretion, shall determine the effect of all matters and questions relating to Termination of Employment, including, but not by way of limitation, the question of whether a Termination of Employment resulted from a discharge for good cause, and all questions of whether a particular leave of absence constitutes a Termination of Employment; provided, however, that, with respect to Incentive Stock Options, unless otherwise determined by the Administrator in its discretion, a leave of absence, change in status from an employee to an independent contractor or other change in the employee-employer relationship shall constitute a Termination of Employment if, and to the extent that, such leave of absence, change in status or other change interrupts employment for the purposes of Section 422(a)(2) of the Code and the then applicable regulations and revenue rulings under said Section.

ARTICLE III

SHARES SUBJECT TO THE PLAN

3.1 Number of Shares.

(a) Subject to Article 12 and Section 3.1(b), the aggregate number of shares of Stock which may be issued or transferred pursuant to Awards under the Plan shall be 9,750,000 shares as of the Effective Date.

(b) To the extent that an Award under Plan terminates, expires, or lapses for any reason, any shares of Stock subject to the Award shall again be available for the grant of an Award pursuant to the Plan. To the extent permitted by applicable law or any exchange rule, shares of Stock issued in assumption of, or in substitution for, any outstanding awards of any entity acquired in any form of combination by the Company or any Parent or Subsidiary shall not be counted against shares of Stock available for grant pursuant to this Plan. If any shares of Restricted Stock are forfeited by a Participant or repurchased by the Company pursuant to Section 6.3 hereof, such shares shall again be available for the grant of an Award pursuant to the Plan. The payment of Dividend Equivalents in cash in conjunction with any outstanding Awards shall not be counted against the shares available for issuance under the Plan.

(c) Notwithstanding the provisions of this Section 3.1, no shares of Stock may again be optioned, granted or awarded if such action would cause an Incentive Stock Option to fail to qualify as an Incentive Stock Option under Section 422 of the Code.

3.2 Stock Distributed. Any Stock distributed pursuant to an Award may consist, in whole or in part, of authorized and unissued Stock, treasury stock or Stock purchased on the open market.

3.3 Limitation on Number of Shares Subject to Incentive Stock Options. The maximum number of shares of Stock that may be issued pursuant to Options that are intended to be Incentive Stock Options shall be 9,750,000 shares.

3.4 Individual Participant Limitations. Notwithstanding any provision in the Plan to the contrary, and subject to Article 12, the maximum number of shares of Stock with respect to one or more Awards that may be granted to any one Participant during any calendar year shall be 600,000, except for an employee serving as Chief Executive Officer of the Company, who is eligible to be granted options covering an aggregate number of shares of up to 1,500,000 in a calendar year. Notwithstanding the foregoing, in connection with his or her initial service to the Company, the aggregate number of shares of Stock with respect to which Awards may be granted to any Participant shall not exceed 1,000,000 shares of Stock during the calendar year which includes such individual's initial service to the Company.

ARTICLE IV

ELIGIBILITY AND PARTICIPATION

4.1 Eligibility. Persons eligible to participate in this Plan include Employees, Consultants and members of the Board, as determined by the Administrator.

4.2 Participation. Subject to the provisions of the Plan, the Administrator may, from time to time, select from among all Eligible Individuals those to whom Awards shall be granted and shall determine the nature and amount of each Award. No individual shall have any right to be granted an Award pursuant to this Plan.

ARTICLE V

STOCK OPTIONS

5.1 General. The Administrator is authorized to grant Options to Eligible Individuals on the following terms and conditions:

(a) Exercise Price. The exercise price per share of Stock subject to an Option shall be determined by the Administrator and set forth in the Award Agreement; provided that the exercise price per share for any Option shall not be less than 100% of the Fair Market Value of a share of Stock on the date of grant.

(b) Time and Conditions of Exercise. The Administrator shall determine the time or times at which an Option may be exercised in whole or in part; provided that the term of any Option granted under the Plan shall not exceed ten years. The Administrator shall also determine the performance or other conditions, if any, that must be satisfied before all or part of an Option may be exercised.

(c) Payment. The Administrator shall determine the methods, terms and conditions by which the exercise price of an Option may be paid, and the form and manner of payment, including, without limitation, payment in the form of cash, a promissory note bearing interest at no less than such rate as shall then preclude the imputation of interest under the Code, shares of Stock, or other property acceptable to the Administrator and payment through the delivery of a notice that the Participant has placed a market sell order with a broker with respect to shares of Stock then issuable upon exercise of the Option, and that the broker has been directed to pay a sufficient portion of the net proceeds of the sale to the Company in satisfaction of the Option exercise price; provided that payment of such proceeds is then made to the Company upon settlement of such sale, and the methods by which shares of Stock shall be delivered or deemed to be delivered to Participants. Notwithstanding any other provision of the Plan to the contrary, no Participant who is a member of the Board or an "executive officer" of the Company within the meaning of Section 13(k) of the Exchange Act shall be permitted to pay the exercise price of an Option, or continue any extension of credit with respect to the exercise price of an Option with a loan from the Company or a loan arranged by the Company, in any method which would violate Section 13(k) of the Exchange Act.

(d) Evidence of Grant. All Options shall be evidenced by an Award Agreement between the Company and the Participant. The Award Agreement shall include such additional provisions as may be specified by the Administrator.

5.2 Incentive Stock Options. Incentive Stock Options may be granted only to employees (as defined in accordance with Section 3401(c) of the Code) of the Company or a Subsidiary which constitutes a “subsidiary corporation” of the Company within Section 424(f) of the Code or a Parent which constitutes a “parent corporation” of the Company within the meaning of Section 424(e) of the Code, and the terms of any Incentive Stock Options granted pursuant to the Plan must comply with the following additional provisions of this Section 5.2 in addition to the requirements of Section 5.1:

(a) Ten Percent Owners. An Incentive Stock Option shall be granted to any individual who, at the date of grant, owns stock possessing more than ten percent of the total combined voting power of all classes of Stock of the Company or any “subsidiary corporation” of the Company or “parent corporation” of the Company (each within the meaning of Section 424 of the Code) only if such Option is granted at an exercise price per share that is not less than 110% of the Fair Market Value per share of the Stock on the date of the grant and the Option is exercisable for no more than five years from the date of grant.

(b) Transfer Restriction. An Incentive Stock Option shall not be transferable by the Participant other than by will or by the laws of descent or distribution.

(c) Right to Exercise. During a Participant’s lifetime, an Incentive Stock Option may be exercised only by the Participant.

(d) Failure to Meet Requirements. Any Option (or portion thereof) purported to be an Incentive Stock Option which, for any reason, fails to meet the requirements of Section 422 of the Code shall be considered a Non-Qualified Stock Option.

5.3 Substitution of Stock Appreciation Rights. The Administrator may provide in the Award Agreement evidencing the grant of an Option that the Administrator, in its sole discretion, shall have the right to substitute a Stock Appreciation Right for such Option at any time prior to or upon exercise of such Option; provided that such Stock Appreciation Right shall be exercisable with respect to the same number of shares of Stock for which such substituted Option would have been exercisable.

ARTICLE VI

RESTRICTED STOCK AWARDS

6.1 Grant of Restricted Stock. The Administrator is authorized to make Awards of Restricted Stock to any Eligible Individual selected by the Administrator in such amounts and subject to such terms and conditions as determined by the Administrator. All Awards of Restricted Stock shall be evidenced by an Award Agreement.

6.2 Issuance and Restrictions. Restricted Stock shall be subject to such repurchase restrictions, forfeiture restrictions, restrictions on transferability and other restrictions as the Administrator may impose (including, without limitation, limitations on the right to vote Restricted Stock or the right to receive dividends on the Restricted Stock). These restrictions may lapse separately or in combination at such times, pursuant to such circumstances or installments or otherwise as the Administrator determines at the time of the grant of the Award or thereafter. Alternatively, these restrictions may lapse pursuant to the satisfaction of one or more Performance Goals or other specific performance goals as the Administrator determines to be appropriate at the time of the grant of the Award or thereafter, in each case on a specified date or dates or over any period or periods determined by the Administrator.

6.3 Repurchase or Forfeiture. Except as otherwise determined by the Administrator at the time of the grant of the Award or thereafter, upon a Participant's Termination of Employment, Termination of Directorship or Termination of Consultancy during the applicable restriction period, Restricted Stock that is at that time subject to restrictions shall be forfeited or subject to repurchase by the Company (or its assignee) under such terms as the Administrator shall determine; provided, however, that the Administrator may (a) provide in any Restricted Stock Award Agreement that restrictions or forfeiture conditions relating to Restricted Stock will be waived in whole or in part in the event of a Participant's Termination of Employment, Termination of Directorship or Termination of Consultancy under certain circumstances, and (b) in other cases waive in whole or in part restrictions or forfeiture conditions relating to Restricted Stock.

6.4 Certificates for Restricted Stock. Restricted Stock granted pursuant to the Plan may be evidenced in such manner as the Administrator shall determine. If certificates representing shares of Restricted Stock are registered in the name of the Participant, certificates must bear an appropriate legend referring to the terms, conditions, and restrictions applicable to such Restricted Stock, and the Company may, at its discretion, retain physical possession of the certificate until such time as all applicable restrictions lapse or the Award Agreement may provide that the shares shall be held in escrow by an escrow agent designated by the Company.

ARTICLE VII

STOCK APPRECIATION RIGHTS

7.1 Grant of Stock Appreciation Rights. A Stock Appreciation Right may be granted to any Eligible Individual selected by the Administrator. A Stock Appreciation Right shall be subject to such terms and conditions not inconsistent with the Plan as the Administrator shall impose and shall be evidenced by an Award Agreement.

7.2 Terms of Stock Appreciation Rights.

(a) A Stock Appreciation Right shall have a term set by the Administrator. A Stock Appreciation Right shall be exercisable in such installments as the Administrator may determine. A Stock Appreciation Right shall cover such number of shares of Stock as the Administrator may determine. The exercise price per share of Stock subject to each Stock Appreciation Right shall be set by the Administrator and set forth in the Award Agreement, except that in no event shall the exercise price be less than 100% of the Fair Market Value of the Stock underlying the Stock Appreciation Right at the time the Stock Appreciation Right is granted.

(b) A Stock Appreciation Right shall entitle the Participant (or other person entitled to exercise the Stock Appreciation Right pursuant to the Plan) to exercise all or a specified portion of the Stock Appreciation Right (to the extent then exercisable pursuant to its terms) and to receive from the Company an amount determined by multiplying (i) the amount (if any) by which the Fair Market Value of a share of Stock on the date of exercise of the Stock Appreciation Right exceeds the exercise price per share of the Stock Appreciation Right, by (ii) the number of shares of Stock with respect to which the Stock Appreciation Right shall have been exercised, subject to any limitations the Administrator may impose.

7.3 Payment and Limitations on Exercise.

(a) Subject to Sections 7.3(b) and (c), payment of the amounts determined under Sections 7.2(b) above shall be in cash, in Stock (based on its Fair Market Value as of the date the Stock Appreciation Right is exercised) or a combination of both, as determined by the Administrator.

(b) To the extent payment for a Stock Appreciation Right is to be made in cash, the Award Agreement shall specify the date of payment, which may be different than the date of exercise of the Stock Appreciation Right. If the date of payment for a Stock Appreciation Right is later than the date of exercise, the Award Agreement may specify that the Participant be entitled to earnings on such amount until paid.

(c) To the extent any payment under Section 7.2(b) is effected in Stock, it shall be made subject to satisfaction of all provisions of Article 5 above pertaining to Options.

7.4 Compliance with Code Section 409A. Notwithstanding anything in this Article 7 to the contrary, all Awards of Stock Appreciation Rights shall be structured to satisfy the requirements of Code Section 409A, as provided in Section 10 below.

ARTICLE VIII

OTHER TYPES OF AWARDS

8.1 Dividend Equivalents.

(a) Any Eligible Individual selected by the Administrator may be granted Dividend Equivalents based on the dividends on the shares of Stock that are subject to any Award, to be credited as of dividend payment dates, during the period between the date the Award is granted and the date the Award is exercised, vests or expires, as determined by the Administrator. Such Dividend Equivalents shall be converted to cash or additional shares of Stock by such formula and at such time and subject to such limitations as may be determined by the Administrator.

(b) Dividend Equivalents granted with respect to Options or SARs that are intended to be Qualified Performance-Based Compensation shall be payable, with respect to pre-exercise periods, regardless of whether such Option or SAR is subsequently exercised.

8.2 Stock Payments. Any Eligible Individual selected by the Administrator may receive Stock Payments in the manner determined from time to time by the Administrator; provided, that unless otherwise determined by the Administrator such Stock Payments, which may be subject to vesting, shall be made in lieu of base salary, bonus, or other cash compensation otherwise payable to such Eligible Individual. The number of shares shall be determined by the Administrator and may be based upon the Performance Goals or other specific performance goals determined appropriate by the Administrator.

8.3 Restricted Stock Units. The Administrator is authorized to make Awards of Restricted Stock Units to any Eligible Individual selected by the Administrator in such amounts and subject to such terms and conditions as determined by the Administrator. At the time of grant, the Administrator shall specify the date or dates on which the Restricted Stock Units shall become fully vested and nonforfeitable, and may specify such conditions to vesting as it deems appropriate. Alternatively, Restricted Stock Units may become fully vested and nonforfeitable pursuant to the satisfaction of one or more Performance Goals or other specific performance goals as the

Administrator determines to be appropriate at the time of the grant of the Restricted Stock Units or thereafter, in each case on a specified date or dates or over any period or periods determined by the Administrator. At the time of grant, the Administrator shall specify the maturity date applicable to each grant of Restricted Stock Units which shall be no earlier than the vesting date or dates of the Award and may be determined at the election of the Eligible Individual to whom the Award is granted. On the maturity date, the Company shall transfer to the Participant one unrestricted, fully transferable share of Stock for each Restricted Stock Unit that is vested and scheduled to be distributed on such date and not previously forfeited. The Administrator shall specify the purchase price, if any, to be paid by the Participant to the Company for such shares of Stock.

8.4 Term. Except as otherwise provided herein, the term of any Award of Dividend Equivalents, Stock Payments or Restricted Stock Units shall be set by the Administrator in its discretion.

8.5 Exercise or Purchase Price. The Administrator may establish the exercise or purchase price, if any, of any Award of Stock Payments or Restricted Stock Units; provided, however, that such price shall not be less than the par value of a share of Stock on the date of grant, unless otherwise permitted by applicable state law.

8.6 Form of Payment. Payments with respect to any Awards granted under Sections 8.1, 8.2 or 8.3 shall be made in cash, in Stock or a combination of both, as determined by the Administrator.

8.7 Award Agreement. All Awards under this Article 8 shall be subject to such additional terms and conditions as determined by the Administrator and shall be evidenced by a written Award Agreement.

8.8 Compliance with Code Section 409A. Notwithstanding anything in this Article 8 to the contrary, all Awards of Dividend Equivalents, Stock Payments, and Restricted Stock Units shall be structured to satisfy the requirements of Code Section 409A, as provided in Section 10 below.

ARTICLE IX

PERFORMANCE-BASED AWARDS

9.1 Purpose. The purpose of this Article 9 is to provide the Administrator the ability to qualify Awards other than Options and SARs and that are granted pursuant to Articles 6 and 8 as Qualified Performance-Based Compensation. If the Administrator, in its discretion, decides to grant a Performance-Based Award to a Covered Employee, the provisions of this Article 9 shall control over any contrary provision contained in Articles 6 or 8; provided, however, that the Administrator may in its discretion grant Awards to Covered Employees that are based on Performance Criteria or Performance Goals but that do not satisfy the requirements of this Article 9.

9.2 Applicability. This Article 9 shall apply only to those Covered Employees selected by the Administrator to receive Performance-Based Awards. The designation of a Covered Employee as a Participant for a Performance Period shall not in any manner entitle the Participant to receive an Award for the period. Moreover, designation of a Covered Employee as a Participant for a particular Performance Period shall not require designation of such Covered Employee as a Participant in any subsequent Performance Period and designation of one Covered Employee as a Participant shall not require designation of any other Covered Employees as a Participant in such period or in any other period.

9.3 Procedures with Respect to Performance-Based Awards. To the extent necessary to comply with the Qualified Performance-Based Compensation requirements of Section 162(m)(4)(C) of the Code, with respect to any Award granted under Articles 6 and 8 which may be granted to one or more Covered Employees, no later than ninety (90) days following the commencement of any fiscal year in question or any other designated fiscal period or period of service (or such other time as may be required or permitted by Section 162(m) of the Code), the Administrator shall, in writing, (a) designate one or more Covered Employees, (b) select the Performance Criteria applicable to the Performance Period, (c) establish the Performance Goals, and amounts of such Awards, as applicable, which may be earned for such Performance Period, and (d) specify the relationship between Performance Criteria and the Performance Goals and the amounts of such Awards, as applicable, to be earned by each Covered Employee for such Performance Period. Following the completion of each Performance Period, the Administrator shall certify in writing whether the applicable Performance Goals have been achieved for such Performance Period. In determining the amount earned by a Covered Employee, the Administrator shall have the right to reduce or eliminate (but not to increase) the amount payable at a given level of performance to take into account additional factors that the Administrator may deem relevant to the assessment of individual or corporate performance for the Performance Period.

9.4 Payment of Performance-Based Awards. Unless otherwise provided in the applicable Award Agreement, a Participant must be employed by the Company or a Parent or Subsidiary on the day a Performance-Based Award for such Performance Period is paid to the Participant. Furthermore, a Participant shall be eligible to receive payment pursuant to a Performance-Based Award for a Performance Period only if the Performance Goals for such period are achieved.

9.5 Additional Limitations. Notwithstanding any other provision of the Plan, any Award which is granted to a Covered Employee and is intended to constitute Qualified Performance-Based Compensation shall be subject to any additional limitations set forth in Section 162(m) of the Code (including any amendment to Section 162(m) of the Code) or any regulations or rulings issued thereunder that are requirements for qualification as qualified performance-based compensation as described in Section 162(m)(4)(C) of the Code, and the Plan shall be deemed amended to the extent necessary to conform to such requirements.

ARTICLE X

COMPLIANCE WITH SECTION 409A OF THE CODE

10.1 Awards subject to Code Section 409A. Any Award that constitutes, or provides for, a deferral of compensation subject to Section 409A of the Code (a “**Section 409A Award**”) shall satisfy the requirements of Section 409A of the Code and this Article 10, to the extent applicable. The Award Agreement with respect to a Section 409A Award shall incorporate the terms and conditions required by Section 409A of the Code and this Article 10. If any deferral of compensation is to be permitted in connection with a 409A Award, the Committee shall establish rules and procedures relating to such deferral in a manner intended to comply with the requirements of Section 409A of the Code, including, without limitation, the time when an election to defer may be made, the time period of the deferral and the events that would result in payment of the deferred amount, the interest or other earnings attributable to the deferral and the method of funding, if any, attributable to the deferred amount.

10.2 Distributions under a Section 409A Award. Any shares of Stock or other property or amounts to be paid or distributed upon the grant, issuance, vesting, exercise or payment of a Section 409A Award shall be distributed in accordance with the requirements of Section 409A(a)(2) of the Code, and, subject to any additional limitations in Section 409A and the Treasury Regulations issued thereunder, shall not be distributed earlier than:

- (a) the Participant's separation from service;
- (b) the date the Participant becomes disabled;
- (c) the Participant's death;
- (d) a specified time (or pursuant to a fixed schedule) specified under the Award Agreement at the date of the deferral compensation;
- (e) a change in the ownership or effective control of the Company or a Parent or Subsidiary, or in the ownership of a substantial portion of the assets of the Company or a Parent or Subsidiary; or
- (f) the occurrence of an unforeseeable emergency with respect to the Participant.

10.3 Prohibition on Acceleration of Benefits. The time or schedule of any distribution or payment of any shares of Stock or other property or amounts under a Section 409A Award shall not be accelerated, except as otherwise permitted under Section 409A(a)(3) of the Code and the Treasury Regulations thereunder.

10.4 Elections under Section 409A Awards. Any deferral election provided under or with respect to an Award to any Eligible Individual, or to the Participant holding a Section 409A Award, shall satisfy the requirements of Section 409A(a)(4)(B) of the Code, to the extent applicable, and any such deferral election with respect to compensation for services performed during a taxable year shall be made not later than the close of the preceding taxable year, or at such other time as provided in Treasury Regulations.

10.5 Compliance in Form and Operation. A Section 409A Award, and any election under or with respect to such Section 409A Award, shall comply in form and operation with the requirements of Section 409A of the Code and the Treasury Regulations thereunder.

ARTICLE XI

PROVISIONS APPLICABLE TO AWARDS

11.1 Stand-Alone and Tandem Awards. Awards granted pursuant to the Plan may, in the discretion of the Administrator, be granted either alone, in addition to, or in tandem with, any other Award granted pursuant to the Plan. Awards granted in addition to or in tandem with other Awards may be granted either at the same time as or at a different time from the grant of such other Awards.

11.2 Award Agreement. Awards under the Plan shall be evidenced by Award Agreements that set forth the terms, conditions and limitations for each Award which may include the term of an Award, the provisions applicable in the event of the Participant's Termination of Employment, Termination of Directorship or Termination of Consultancy, and the Company's authority to unilaterally or bilaterally amend, modify, suspend, cancel or rescind an Award.

11.3 Limits on Transfer.

(a) Except as otherwise provided by the Administrator pursuant to Section 11.3(b), no right or interest of a Participant in any Award may be pledged, encumbered, or hypothecated to or in favor of any party other than the Company or a Parent or Subsidiary, or shall be subject to any lien, obligation, or liability of such Participant to any other party other than the Company or a Parent or Subsidiary. Except as otherwise provided by the Administrator pursuant to Section 11.3(b), no Award shall be assigned, transferred, or otherwise disposed of by a Participant other than by will or the laws of descent and distribution, unless and until such Award has been exercised, or the shares underlying such Award have been issued, and all restrictions applicable to such shares have lapsed.

(b) Notwithstanding Section 11.3(a), the Administrator, in its sole discretion, may permit an Award (other than an Incentive Stock Option) to be transferred to, exercised by and paid to any one or more Permitted Transferees (as defined below), subject to the following terms and conditions: (i) an Award transferred to a Permitted Transferee shall not be assignable or transferable by the Permitted Transferee other than by will or the laws of descent and distribution; (ii) any Award which is transferred to a Permitted Transferee shall continue to be subject to all the terms and conditions of the Award as applicable to the original Participant (other than the ability to further transfer the Award); and (iii) the Participant and the Permitted Transferee shall execute any and all documents requested by the Administrator, including, without limitation documents to (A) confirm the status of the transferee as a Permitted Transferee, (B) satisfy any requirements for an exemption for the transfer under applicable federal and state securities laws and (C) evidence the transfer. For purposes of this Section 11.3(b), "**Permitted Transferee**" shall mean, with respect to a Participant, any child, stepchild, grandchild, parent, stepparent, grandparent, spouse, former spouse, sibling, niece, nephew, mother-in-law, father-in-law, son-in-law, daughter-in-law, brother-in-law, or sister-in-law, including adoptive relationships, any person sharing the Participant's household (other than a tenant or employee), a trust in which these persons (or the Participant) control the management of assets, and any other entity in which these persons (or the Participant) own more than fifty percent of the voting interests, or any other transferee specifically approved by the Administrator.

11.4 Beneficiaries. Notwithstanding Section 11.3, a Participant may, in the manner determined by the Administrator, designate a beneficiary to exercise the rights of the Participant and to receive any distribution with respect to any Award upon the Participant's death. A beneficiary, legal guardian, legal representative, or other person claiming any rights pursuant to the Plan is subject to all terms and conditions of the Plan and any Award Agreement applicable to the Participant, except to the extent the Plan and Award Agreement otherwise provide, and to any additional restrictions deemed necessary or appropriate by the Administrator. If the Participant is married and resides in a community property state, a designation of a person other than the Participant's spouse as his or her beneficiary with respect to more than 50% of the Participant's interest in the Award shall not be effective without the prior written consent of the Participant's spouse. If no beneficiary has been designated or survives the Participant, payment shall be made to the person entitled thereto pursuant to the Participant's will or the laws of descent and distribution. Subject to the foregoing, a beneficiary designation may be changed or revoked by a Participant at any time provided the change or revocation is filed with the Administrator.

11.5 Stock Certificates; Book-Entry Procedures.

(a) Notwithstanding anything herein to the contrary, the Company shall not be required to issue or deliver any certificates evidencing shares of Stock pursuant to the exercise of any Award, unless and until the Board has determined, with advice of counsel, that the issuance and delivery of such certificates is in compliance with all applicable laws, regulations of governmental authorities and, if applicable, the requirements of any exchange on which the shares of Stock are listed or traded. All Stock certificates delivered pursuant to the Plan are subject to any stop-transfer orders and other restrictions as the Administrator deems necessary or advisable to comply with federal, state, or foreign jurisdiction, securities or other laws, rules and regulations and the rules of any national securities exchange or automated quotation system on which the Stock is listed, quoted, or traded. The Administrator may place legends on any Stock certificate to reference restrictions applicable to the Stock. In addition to the terms and conditions provided herein, the Administrator may require that a Participant make such reasonable covenants, agreements, and representations as the Administrator, in its discretion, deems advisable in order to comply with any such laws, regulations, or requirements. The Administrator shall have the right to require any Participant to comply with any timing or other restrictions with respect to the settlement or exercise of any Award, including a window-period limitation, as may be imposed in the discretion of the Administrator.

(b) Notwithstanding any other provision of the Plan, unless otherwise determined by the Administrator or required by applicable law, rule or regulation, the Company shall not deliver to any Participant certificates evidencing shares of Stock issued in connection with any Award and instead such shares of Stock shall be recorded in the books of the Company (or, as applicable, its transfer agent or stock plan administrator).

11.6 Paperless Exercise. In the event that the Company establishes, for itself or using the services of a third party, an automated system for the exercise of Awards, such as a system using an internet website or interactive voice response, then the paperless exercise of Awards by a Participant may be permitted through the use of such an automated system.

11.7 Minimum Vesting Periods. Subject to Article XII hereof, any Awards granted under Article VI, Section 8.3 or Article IX hereof (the “**Minimum Vesting Awards**”), granted after July 25, 2011 that vest solely based on the passage of time may not provide for vesting any faster than the following schedule: (i) no more than one-third vested prior to the first anniversary of the award date, (ii) no more than two-thirds vested prior to the second anniversary of the award date and (iii) the balance shall vest at a rate no more than ratably over the period from the second anniversary of the award date to the third anniversary of the award date. Subject to Article XII hereof, any Minimum Vesting Awards granted after July 25, 2011 that do not vest solely based on the passage of time shall not vest any faster than ratably over the period from the award date to the first anniversary of the award date.

CHANGES IN CAPITAL STRUCTURE

12.1 Adjustments.

(a) In the event of any stock dividend, stock split, combination or exchange of shares, merger, consolidation, spin-off, recapitalization, distribution of Company assets to shareholders (other than normal cash dividends), or any other corporate event affecting the Stock or the share price of the Stock, the Administrator may make such proportionate adjustments, if any, as the Administrator in its discretion may deem appropriate to reflect such change with respect to (i) the aggregate number and type of shares that may be issued under the Plan (including, but not limited to, adjustments of the limitations in Sections 3.1 and 3.3); (ii) the terms and conditions of any outstanding Awards (including, without limitation, any applicable performance targets or criteria with respect thereto); and (iii) the grant, exercise or purchase price per share for any outstanding Awards under the Plan. Any adjustment affecting an Award intended as Qualified Performance-Based Compensation shall be made consistent with the requirements of Section 162(m) of the Code.

(b) In the event of any transaction or event described in Section 12.1(a) or any unusual or nonrecurring transactions or events affecting the Company, any affiliate of the Company, or the financial statements of the Company or any affiliate (including without limitation any Change in Control), or of changes in applicable laws, regulations or accounting principles, and whenever the Administrator determines that such action is appropriate in order to prevent the dilution or enlargement of the benefits or potential benefits intended to be made available under the Plan or with respect to any Award under the Plan, to facilitate such transactions or events or to give effect to such changes in laws, regulations or principles, the Administrator, in its sole discretion and on such terms and conditions as it deems appropriate, either by the terms of the Award or by action taken prior to the occurrence of such transaction or event and either automatically or upon the Participant's request, is hereby authorized to take any one or more of the following actions:

(i) To provide for either (A) termination of any such Award in exchange for an amount of cash, if any, equal to the amount that would have been received upon the exercise of such Award or realization of the Participant's rights (and, for the avoidance of doubt, if as of the date of the occurrence of the transaction or event described in this Section 12.1(b) the Administrator determines in good faith that no amount would have been attained upon the exercise of such Award or realization of the Participant's rights, then such Award may be terminated by the Company without payment) or (B) the replacement of such Award with other rights or property selected by the Administrator in its sole discretion;

(ii) To provide that such Award be assumed by the successor or survivor corporation, or a parent or subsidiary thereof, or shall be substituted for by similar options, rights or awards covering the stock of the successor or survivor corporation, or a parent or subsidiary thereof, with appropriate adjustments as to the number and kind of shares and prices; and

(iii) To make adjustments in the number and type of shares of Stock (or other securities or property) subject to outstanding Awards, and in the number and kind of outstanding Restricted Stock and/or in the terms and conditions of (including the grant or exercise price), and the criteria included in, outstanding options, rights and awards and options, rights and awards which may be granted in the future;

(iv) To provide that such Award shall be exercisable or payable or fully vested with respect to all shares covered thereby, notwithstanding anything to the contrary in the Plan or the applicable Award Agreement; and

(v) To provide that the Award cannot vest, be exercised or become payable after such event.

12.2 Acceleration Upon a Change in Control.

(a) Notwithstanding Section 12.1(b), if a Change in Control occurs and a Participant's Awards are not continued, converted, assumed, or replaced by (i) the Company or a Parent or Subsidiary of the Company, or (ii) a Successor Entity, such Awards shall become fully exercisable and/or payable, as applicable, and all forfeiture, repurchase and other restrictions on such Awards shall lapse immediately prior to such Change in Control. Upon, or in anticipation of, a Change in Control, the Administrator may cause any and all Awards outstanding hereunder to terminate at a specific time in the future, including but not limited to the date of such Change in Control, and shall give each Participant the right to exercise such Awards during a period of time as the Administrator, in its sole and absolute discretion, shall determine

(b) With respect to any Participant who was providing services as an Employee, member of the Board or Consultant, if such Participant has a Termination of Employment, Termination of Directorship or Termination of Consultancy in contemplation of a Change in Control, other than by reason of a discharge by the Company, any Parent or Subsidiary or any Successor Entity for Cause, a resignation by the Participant without Good Reason, or the Participant's death or Disability, within the sixty (60) days prior to the consummation of a Change in Control, any Awards held by such Participant shall become exercisable and/or payable, as applicable, and the forfeiture, repurchase and other restrictions on such Awards shall lapse in accordance with the schedule set forth in Section 12.2(f) below immediately prior to the date of such Change in Control and such Awards shall be exercisable for the longer of twelve (12) months following such Change in Control or the expiration of any applicable underwriters' lock-up agreements and thereafter shall terminate, but such period shall not extend beyond the expiration date of such Awards.

(c) With respect to any Participant who was providing services as an Employee, member of the Board or Consultant immediately prior to the consummation of a Change in Control, if such Participant has a Termination of Employment, Termination of Directorship or Termination of Consultancy other than by reason of the Participant's discharge by the Company, any Parent or Subsidiary or any Successor Entity for Cause, a resignation by the Participant without Good Reason, or the Participant's death or Disability, within the thirteen (13) months following such Change in Control, any Awards held by such Participant shall become exercisable and/or payable, as applicable, and the forfeiture, repurchase and other restrictions on such Awards shall lapse in accordance with the schedule set forth in Section 12.2(f) below on the date of such Termination of Employment, Termination of Directorship or Termination of Consultancy and such Awards shall be exercisable for the longer of twelve (12) months following such Change in Control or the expiration of any applicable underwriters' lock-up agreements and thereafter shall terminate, but such period shall not extend beyond the expiration date of such Awards.

(d) With respect to any Participant who was providing services as an Employee, member of the Board or Consultant immediately prior to the consummation of a Change in Control, if such Participant does not have a Termination of Employment, Termination of Directorship or

Termination of Consultancy prior to thirteen (13) months after such Change in Control, any Awards held by such Participant shall become exercisable and/or payable, as applicable, and the forfeiture, repurchase and other restrictions on such Awards shall lapse in accordance with the schedule set forth in Section 12.2(f) below at the end of such thirteen (13) month period and such Awards shall be exercisable for the longer of twelve (12) months following the end of such thirteen (13) month period or the expiration of any applicable underwriters' lock-up agreements and thereafter shall terminate, but such period shall not extend beyond the expiration date of such Awards.

(e) Notwithstanding anything to the contrary in this Article 12, an Award Agreement evidencing an Award may provide that a Participant shall have additional rights under such Award in the event of a Change in Control.

(f) For purposes of this Section 12.2, Awards shall become exercisable and/or payable, as applicable, and the forfeiture, repurchase and other restrictions on such Awards shall lapse pursuant to Sections 12.2(b), (c) and (d) in accordance with the number of years of service a Participant has with the Company, or any Parent or Subsidiary (or any predecessor organization, including, without limitation, RiboGene, Inc.), or any Successor Entity as of the date of determination (measured from the Participant's date of hire) as follows:

Length of Service	Percentage of Award to Become Exercisable and/or Payable and Percentage of Award as to Which Forfeiture, Repurchase and Other Restrictions Shall Lapse
0-180 days	0%
181 days to 1 year	25%
1 year and 1 day to 2 years	50%
Greater than 2 years	100%

12.3 No Other Rights. Except as expressly provided in the Plan, no Participant shall have any rights by reason of any subdivision or consolidation of shares of stock of any class, the payment of any dividend, any increase or decrease in the number of shares of stock of any class or any dissolution, liquidation, merger, or consolidation of the Company or any other corporation. Except as expressly provided in the Plan or pursuant to action of the Administrator under the Plan, no issuance by the Company of shares of stock of any class, or securities convertible into shares of stock of any class, shall affect, and no adjustment by reason thereof shall be made with respect to, the number of shares of Stock subject to an Award or the grant or exercise price of any Award.

ARTICLE XIII

ADMINISTRATION

13.1 Administrator. The Administrator of the Plan shall be the Compensation Committee of the Board (or another committee or a subcommittee of the Board to which the Board delegates administration of the Plan) (such committee, the "**Committee**"), which Committee shall consist solely of two or more members of the Board each of whom is both an "outside director," within the meaning of Section 162(m) of the Code and a Non-Employee Director. Notwithstanding the

foregoing: (a) the full Board, acting by a majority of its members in office, shall conduct the general administration of the Plan with respect to all Awards granted to Independent Directors, and for purposes of such Awards the term “**Administrator**” as used in this Plan shall be deemed to refer to the Board and (b) the Committee may delegate its authority hereunder to the extent permitted by Section 13.5. Appointment of Committee members shall be effective upon acceptance of appointment. In its sole discretion, the Board may at any time and from time to time exercise any and all rights and duties of the Administrator under the Plan except with respect to matters which under Rule 16b-3 under the Exchange Act or Section 162(m) of the Code, or any regulations or rules issued thereunder, are required to be determined in the sole discretion of the Committee. Committee members may resign at any time by delivering written notice to the Board. Vacancies in the Committee may only be filled by the Board.

13.2 Action by the Administrator. A majority of the Administrator shall constitute a quorum. The acts of a majority of the members present at any meeting at which a quorum is present, and, subject to applicable law, acts approved in writing by a majority of the Administrator in lieu of a meeting, shall be deemed the acts of the Administrator. Each member of the Administrator is entitled to, in good faith, rely or act upon any report or other information furnished to that member by any officer or other employee of the Company or any Parent or Subsidiary, the Company’s independent certified public accountants, or any executive compensation consultant or other professional retained by the Company to assist in the administration of the Plan.

13.3 Authority of Administrator. Subject to any specific designation in the Plan, the Administrator has the exclusive power, authority and discretion to:

- (a) Designate Participants to receive Awards;
- (b) Determine the type or types of Awards to be granted to each Participant;
- (c) Determine the number of Awards to be granted and the number of shares of Stock to which an Award will relate;

(d) Determine the terms and conditions of any Award granted pursuant to the Plan, including, but not limited to, the exercise price, grant price, or purchase price, any reload provision, any restrictions or limitations on the Award, any schedule for lapse of forfeiture restrictions or restrictions on the exercisability of an Award, and accelerations or waivers thereof, any provisions related to non-competition and recapture of gain on an Award, based in each case on such considerations as the Administrator in its sole discretion determines; provided, however, that the Administrator shall not have the authority to accelerate the vesting or waive the forfeiture of any Performance-Based Awards;

(e) Determine whether, to what extent, and pursuant to what circumstances an Award may be settled in, or the exercise price of an Award may be paid in, cash, Stock, other Awards, or other property, or an Award may be canceled, forfeited, or surrendered;

(f) Prescribe the form of each Award Agreement, which need not be identical for each Participant;

(g) Decide all other matters that must be determined in connection with an Award;

(h) Establish, adopt, or revise any rules and regulations as it may deem necessary or advisable to administer the Plan;

(i) Interpret the terms of, and any matter arising pursuant to, the Plan or any Award Agreement; and

(j) Make all other decisions and determinations that may be required pursuant to the Plan or as the Administrator deems necessary or advisable to administer the Plan.

13.4 Decisions Binding. The Administrator's interpretation of the Plan, any Awards granted pursuant to the Plan, any Award Agreement and all decisions and determinations by the Administrator with respect to the Plan are final, binding, and conclusive on all parties.

13.5 Delegation of Authority. To the extent permitted by applicable law, the Committee may from time to time delegate to a committee of one or more members of the Board or one or more officers of the Company the authority to grant or amend Awards to Participants other than (a) senior executives of the Company who are subject to Section 16 of the Exchange Act, (b) Covered Employees, or (c) officers of the Company (or members of the Board) to whom authority to grant or amend Awards has been delegated hereunder. Any delegation hereunder shall be subject to the restrictions and limits that the Committee specifies at the time of such delegation, and the Committee may at any time rescind the authority so delegated or appoint a new delegatee. At all times, the delegatee appointed under this Section 13.5 shall serve in such capacity at the pleasure of the Committee.

ARTICLE XIV

EFFECTIVE AND EXPIRATION DATES

14.1 Effective Date. The Plan will be effective as of the date on which the Plan is approved by the Company's shareholders (the "**Effective Date**").

14.2 Expiration Date. The Plan will expire on, and no Award may be granted pursuant to the Plan after, the earlier of the tenth anniversary of (i) the date this Plan is approved by the Board or (ii) the Effective Date (the "**Expiration Date**"). Any Awards that are outstanding on the Expiration Date shall remain in force according to the terms of the Plan and the applicable Award Agreement.

ARTICLE XV

AMENDMENT, MODIFICATION, AND TERMINATION

15.1 Amendment, Modification, and Termination. The Board may terminate, amend or modify the Plan at any time and from time to time; provided, however, that (a) to the extent necessary to comply with any applicable law, regulation, or stock exchange rule, the Company shall obtain shareholder approval of any Plan amendment in such a manner and to such a degree as required, and (b) shareholder approval is required for any amendment to the Plan that increases the number of shares available under the Plan (other than any adjustment as provided by Article 12).

15.2 Awards Previously Granted. No termination, amendment, or modification of the Plan shall adversely affect in any material way any Award previously granted pursuant to the Plan without the prior written consent of the Participant.

ARTICLE XVI

GENERAL PROVISIONS

16.1 No Rights to Awards. No Participant, Employee, or other person shall have any claim to be granted any Award pursuant to the Plan, and neither the Company nor the Administrator is obligated to treat Participants, Employees, and other persons uniformly.

16.2 No Shareholder Rights. Except as otherwise provided herein, a Participant shall have none of the rights of a shareholder with respect to shares of Stock covered by any Award until the Participant becomes the record owner of such shares of Stock.

16.3 Withholding. The Company or any Parent or Subsidiary shall have the authority and the right to deduct or withhold, or require a Participant to remit to the Company an amount sufficient to satisfy federal, state, local and foreign taxes (including the Participant's employment tax obligations) required by law to be withheld with respect to any taxable event concerning a Participant arising as a result of this Plan. The Administrator may in its discretion and in satisfaction of the foregoing requirement allow a Participant to elect to have the Company or a Parent or Subsidiary, as applicable, withhold shares of Stock otherwise issuable under an Award (or allow the return of shares of Stock) having a Fair Market Value equal to the sums required to be withheld. Notwithstanding any other provision of the Plan, the number of shares of Stock which may be withheld with respect to the issuance, vesting, exercise or payment of any Award (or which may be repurchased from the Participant of such Award within six months (or such other period as may be determined by the Administrator) after such shares of Stock were acquired by the Participant from the Company) in order to satisfy the Participant's federal, state, local and foreign income and payroll tax liabilities with respect to the issuance, vesting, exercise or payment of the Award shall be limited to the number of shares which have a Fair Market Value on the date of withholding or repurchase equal to the aggregate amount of such liabilities based on the minimum statutory withholding rates for federal, state, local and foreign income tax and payroll tax purposes that are applicable to such supplemental taxable income.

16.4 No Right to Employment or Services. Nothing in the Plan or any Award Agreement shall interfere with or limit in any way the right of the Company or any Parent or Subsidiary to terminate any Participant's employment or services at any time, nor confer upon any Participant any right to continue in the employ or service of the Company or any Parent or Subsidiary.

16.5 Unfunded Status of Awards. The Plan is intended to be an unfunded plan for incentive compensation. With respect to any payments not yet made to a Participant pursuant to an Award, nothing contained in the Plan or any Award Agreement shall give the Participant any rights that are greater than those of a general creditor of the Company or any Parent or Subsidiary.

16.6 Indemnification. To the extent allowable pursuant to applicable law, the Administrator (and each member thereof) shall be indemnified and held harmless by the Company from any loss, cost, liability, or expense that may be imposed upon or reasonably incurred by such member in connection with or resulting from any claim, action, suit, or proceeding to which he or she

may be a party or in which he or she may be involved by reason of any action or failure to act pursuant to the Plan and against and from any and all amounts paid by him or her in satisfaction of judgment in such action, suit, or proceeding against him or her; provided he or she gives the Company an opportunity, at its own expense, to handle and defend the same before he or she undertakes to handle and defend it on his or her own behalf. The foregoing right of indemnification shall not be exclusive of any other rights of indemnification to which such persons may be entitled pursuant to the Company's Certificate of Incorporation or Bylaws, as a matter of law, or otherwise, or any power that the Company may have to indemnify them or hold them harmless.

16.7 Relationship to other Benefits. No payment pursuant to the Plan shall be taken into account in determining any benefits pursuant to any pension, retirement, savings, profit sharing, group insurance, welfare or other benefit plan of the Company or any Parent or Subsidiary except to the extent otherwise expressly provided in writing in such other plan or an agreement thereunder.

16.8 Expenses. The expenses of administering the Plan shall be borne by the Company and its Subsidiaries.

16.9 Titles and Headings. The titles and headings of the Sections in the Plan are for convenience of reference only and, in the event of any conflict, the text of the Plan, rather than such titles or headings, shall control.

16.10 Fractional Shares. No fractional shares of Stock shall be issued and the Administrator shall determine, in its discretion, whether cash shall be given in lieu of fractional shares or whether such fractional shares shall be eliminated by rounding up or down as appropriate.

16.11 Limitations Applicable to Section 16 Persons. Notwithstanding any other provision of the Plan, the Plan, and any Award granted or awarded to any Participant who is then subject to Section 16 of the Exchange Act, shall be subject to any additional limitations set forth in any applicable exemptive rule under Section 16 of the Exchange Act (including any amendment to Rule 16b-3 under the Exchange Act) that are requirements for the application of such exemptive rule. To the extent permitted by applicable law, the Plan and Awards granted or awarded hereunder shall be deemed amended to the extent necessary to conform to such applicable exemptive rule.

16.12 Government and Other Regulations. The obligation of the Company to make payment of awards in Stock or otherwise shall be subject to all applicable laws, rules, and regulations, and to such approvals by government agencies as may be required. The Company shall be under no obligation to register pursuant to the Securities Act, any of the shares of Stock paid pursuant to the Plan. If the shares paid pursuant to the Plan may in certain circumstances be exempt from registration pursuant to the Securities Act, the Company may restrict the transfer of such shares in such manner as it deems advisable to ensure the availability of any such exemption.

16.13 Governing Law. The Plan and all Award Agreements shall be construed in accordance with and governed by the laws of the State of California, without regard to the conflicts of law principles thereof.

QUESTCOR PHARMACEUTICALS, INC.

Amended and Restated
Employee Stock Purchase Plan1. Purpose.

(a) The purpose of the 2003 Employee Stock Purchase Plan (the “**Plan**”) is to provide a means by which employees of Questcor Pharmaceuticals, Inc., a California corporation (the “**Company**”), and its Affiliates, as defined in subparagraph 1(b), which are designated as provided in subparagraph 2(b), may be given an opportunity to purchase stock of the Company.

(b) The word “**Affiliate**” as used in the Plan means any parent corporation or subsidiary corporation of the Company, as those terms are defined in Sections 424(e) and (f), respectively, of the Internal Revenue Code of 1986, as amended (the “**Code**”).

(c) The Company, by means of the Plan, seeks to retain the services of its employees, to secure and retain the services of new employees, and to provide incentives for such persons to exert maximum efforts for the success of the Company.

(d) The Company intends that the rights to purchase stock of the Company granted under the Plan be considered options issued under an “employee stock purchase plan” as that term is defined in Section 423(b) of the Code.

2. Administration.

(a) The Plan shall be administered by the Board of Directors (the “**Board**”) of the Company unless and until the Board delegates administration to a Committee, as provided in subparagraph 2(c). Whether or not the Board has delegated administration, the Board shall have the final power to determine all questions of policy and expediency that may arise in the administration of the Plan.

(b) The Board shall have the power, subject to, and within the limitations of, the express provisions of the Plan:

(i) To determine when and how rights to purchase stock of the Company shall be granted and the provisions of each offering of such rights (which need not be identical).

(ii) To designate from time to time which Affiliates of the Company shall be eligible to participate in the Plan.

(iii) To construe and interpret the Plan and rights granted under it, and to establish, amend and revoke rules and regulations for its administration. The Board, in the exercise of this power, may correct any defect, omission or inconsistency in the Plan, in a manner and to the extent it shall deem necessary or expedient to make the Plan fully effective.

(iv) To amend the Plan as provided in paragraph 13.

(v) Generally, to exercise such powers and to perform such acts as the Board deems necessary or expedient to promote the best interests of the Company and its Affiliates and to carry out the intent that the Plan be treated as an “employee stock purchase plan” within the meaning of Section 423 of the Code.

(c) The Board may delegate administration of the Plan to a Committee composed of two (2) or more members of the Board (the “**Committee**”). If administration is delegated to a Committee, the Committee shall have, in connection with the administration of the Plan, the powers theretofore possessed by the Board, subject, however, to such resolutions, not inconsistent with the provisions of the Plan, as may be adopted from time to time by the Board. The Board may abolish the Committee at any time and revert in the Board the administration of the Plan.

3. Shares Subject to the Plan.

(a) Subject to the provisions of paragraph 12 relating to adjustments upon changes in stock, the stock that may be sold pursuant to rights granted under the Plan shall not exceed in the aggregate Three Million Five Hundred Thousand (3,500,000) shares of the Company’s common stock, no par value (the “**Common Stock**”). If any right granted under the Plan shall for any reason terminate without having been exercised, the Common Stock not purchased under such right shall again become available for the Plan.

(b) The stock subject to the Plan may be unissued shares or reacquired shares, bought on the market or otherwise.

4. Grant of Rights; Offering.

(a) The Board or the Committee may from time to time grant or provide for the grant of rights to purchase Common Stock of the Company under the Plan to eligible employees (an “**Offering**”) on a date or dates (the “**Offering Date(s)**”) selected by the Board or the Committee. Each Offering shall be in such form and shall contain such terms and conditions as the Board or the Committee shall deem appropriate, which shall comply with the requirements of Section 423(b)(5) of the Code that all employees granted rights to purchase stock under the Plan shall have the same rights and privileges. The terms and conditions of an Offering shall be incorporated by reference into and made part of the Plan and shall be attached hereto as part of the Plan. The provisions of separate Offerings need not be identical, but each Offering shall include (through incorporation of the provisions of this Plan by reference in the document comprising the Offering or otherwise) the period during which the Offering shall be effective, which period shall not exceed six (6) months beginning with the Offering Date, and the substance of the provisions contained in paragraphs 5 through 8, inclusive.

(b) If an employee has more than one right outstanding under the Plan, unless he or she otherwise indicates in agreements or notices delivered hereunder: (1) each agreement or notice delivered by that employee will be deemed to apply to all of his or her rights under the Plan, and (2) a right with a lesser exercise price (or an earlier-granted right, if two rights have identical exercise prices), will be exercised to the fullest possible extent before a right with a higher exercise price (or a later-granted right, if two rights have identical exercise prices) will be exercised.

5. Eligibility.

(a) Rights may be granted only to employees of the Company or, as the Board or the Committee may designate as provided in subparagraph 2(b), to employees of any Affiliate of the

Company. Except as provided in subparagraph 5(b), an employee of the Company or any Affiliate shall not be eligible to be granted rights under the Plan, unless, on the Offering Date, such employee has been in the employ of the Company or any Affiliate for such continuous period preceding such grant as the Board or the Committee may require, but in no event shall the required period of continuous employment be equal to or greater than two (2) years. In addition, unless otherwise determined by the Board or the Committee and set forth in the terms of the applicable Offering, no employee of the Company or any Affiliate shall be eligible to be granted rights under the Plan, unless, on the Offering Date, such employee's customary employment with the Company or such Affiliate is for at least twenty (20) hours per week and at least five (5) months per calendar year.

(b) The Board or the Committee may provide that each person who, during the course of an Offering, first becomes an eligible employee of the Company or designated Affiliate will, on a date or dates specified in the Offering which coincides with the day on which such person becomes an eligible employee or occurs thereafter, receive a right under that Offering, which right shall thereafter be deemed to be a part of that Offering. Such right shall have the same characteristics as any rights originally granted under that Offering, as described herein, except that:

(i) the date on which such right is granted shall be the "Offering Date" of such right for all purposes, including determination of the exercise price of such right;

(ii) the period of the Offering with respect to such right shall begin on its Offering Date and end coincident with the end of such Offering;

and
(iii) the Board or the Committee may provide that if such person first becomes an eligible employee within a specified period of time before the end of the Offering, he or she will not receive any right under that Offering.

(c) No employee shall be eligible for the grant of any rights under the Plan if, immediately after any such rights are granted, such employee owns stock possessing five percent (5%) or more of the total combined voting power or value of all classes of stock of the Company or of any Affiliate. For purposes of this subparagraph 5(c), the rules of Section 424(d) of the Code shall apply in determining the stock ownership of any employee, and stock which such employee may purchase under all outstanding rights and options shall be treated as stock owned by such employee.

(d) An eligible employee may be granted rights under the Plan only if such rights, together with any other rights granted under "employee stock purchase plans" of the Company and any Affiliates, as specified by Section 423(b)(8) of the Code, do not permit such employee's rights to purchase stock of the Company or any Affiliate to accrue at a rate which exceeds twenty-five thousand dollars (\$25,000) of fair market value of such stock (determined at the time such rights are granted) for each calendar year in which such rights are outstanding at any time.

(e) Officers of the Company and any designated Affiliate shall be eligible to participate in Offerings under the Plan, provided, however, that the Board may provide in an Offering that certain employees who are highly compensated employees within the meaning of Section 423(b)(4)(D) of the Code shall not be eligible to participate.

6. Rights; Purchase Price.

(a) On each Offering Date, each eligible employee, pursuant to an Offering made under the Plan, shall be granted the right to purchase up to the number of shares of Common Stock of the Company purchasable with a percentage designated by the Board or the Committee (or fifteen percent (15%) in the absence of any designation) of such employee's Earnings (as defined in subparagraph 7(a)) during the period which begins on the Offering Date (or such later date as the Board or the Committee determines for a particular Offering) and ends on the date stated in the Offering, which date shall be no later than the end of the Offering. The Board or the Committee shall establish one or more dates during an Offering (the "**Purchase Date(s)**") on which rights granted under the Plan shall be exercised and purchases of Common Stock carried out in accordance with such Offering.

(b) In connection with each Offering made under the Plan, the Board or the Committee may specify a maximum number of shares that may be purchased by any employee as well as a maximum aggregate number of shares that may be purchased by all eligible employees pursuant to such Offering. In addition, in connection with each Offering that contains more than one Purchase Date, the Board or the Committee may specify a maximum aggregate number of shares which may be purchased by all eligible employees on any given Purchase Date under the Offering. If the aggregate purchase of shares upon exercise of rights granted under the Offering would exceed any such maximum aggregate number, the Board or the Committee shall make a pro rata allocation of the shares available in as nearly a uniform manner as shall be practicable and as it shall deem to be equitable.

(c) The purchase price of stock acquired pursuant to rights granted under the Plan shall be not less than the lesser of:

- (i) an amount equal to eighty-five percent (85%) of the fair market value of the stock on the Offering Date; or
- (ii) an amount equal to eighty-five percent (85%) of the fair market value of the stock on the Purchase Date.

7. Participation; Withdrawal; Termination.

(a) An eligible employee may become a participant in the Plan pursuant to an Offering by delivering a participation agreement to the Company within the time specified in the Offering, in such form as the Company provides. Each such agreement shall authorize payroll deductions of up to the maximum percentage specified by the Board or the Committee of such employee's Earnings during the Offering. "**Earnings**" is defined as an employee's regular salary or wages (including amounts thereof elected to be deferred by the employee, that would otherwise have been paid, under any arrangement established by the Company intended to comply with Section 401(k), Section 402(e)(3), Section 125, Section 402(h), or Section 403(b) of the Code, and also including any deferrals under a non-qualified deferred compensation plan or arrangement established by the Company), and may also include or exclude (as provided for each Offering) the following items of compensation: bonuses, commissions, overtime pay, incentive pay, profit sharing, other remuneration paid directly to the employee, the cost of employee benefits paid for by the Company or an Affiliate, education or tuition reimbursements, imputed income arising under any group insurance or benefit program, traveling expenses, business and moving expense reimbursements,

income received in connection with stock options, contributions made by the Company or an Affiliate under any employee benefit plan, and similar items of compensation, as determined by the Board or Committee. The payroll deductions made for each participant shall be credited to an account for such participant under the Plan and shall be deposited with the general funds of the Company. A participant may reduce (including to zero), but not increase, such payroll deductions, and an eligible employee may begin such payroll deductions, after the beginning of any Offering only as provided for in the Offering.

(b) At any time during an Offering, a participant may terminate his or her payroll deductions under the Plan and withdraw from the Offering by delivering to the Company a notice of withdrawal in such form as the Company provides. Such withdrawal may be elected at any time prior to the end of the Offering except as provided by the Board or the Committee in the Offering. Upon such withdrawal from the Offering by a participant, the Company shall distribute to such participant all of his or her accumulated payroll deductions (reduced to the extent, if any, such deductions have been used to acquire stock for the participant) under the Offering, without interest, and such participant's interest in that Offering shall be automatically terminated. A participant's withdrawal from an Offering will have no effect upon such participant's eligibility to participate in any other Offerings under the Plan but such participant will be required to deliver a new participation agreement in order to participate in subsequent Offerings under the Plan.

(c) Rights granted pursuant to any Offering under the Plan shall terminate immediately upon cessation of a participant's employment with the Company and any designated Affiliate, for any reason, and the Company shall distribute to such terminated employee all of his or her accumulated payroll deductions (reduced to the extent, if any, such deductions have been used to acquire stock for the terminated employee), under the Offering, without interest.

(d) Rights granted under the Plan shall not be transferable by a participant otherwise than by will or the laws of descent and distribution, or by a beneficiary designation as provided in paragraph 14 and, otherwise during his or her lifetime, shall be exercisable only by the person to whom such rights are granted.

8. Exercise.

(a) On each Purchase Date specified in the relevant Offering, each participant's accumulated payroll deductions and other additional payments specifically provided for in the Offering (without any increase for interest) will be applied to the purchase of whole shares of stock of the Company, up to the maximum number of shares permitted pursuant to the terms of the Plan and the applicable Offering, at the purchase price specified in the Offering. No fractional shares shall be issued upon the exercise of rights granted under the Plan, unless the Offering document specifically provides otherwise. The amount, if any, of accumulated payroll deductions remaining in each participant's account after the purchase of shares which is less than the amount required to purchase one share of stock on the final Purchase Date of an Offering shall be held in each such participant's account for the purchase of shares under the next Offering under the Plan, unless such participant withdraws from such next Offering, as provided in subparagraph 7(b), or is no longer eligible to be granted rights under the Plan, as provided in paragraph 5, in which case such amount shall be distributed to the participant after such final Purchase Date, without interest. The amount, if any, of accumulated payroll deductions remaining in any participant's account after the purchase of shares which is equal to the amount required to purchase whole shares of stock on the final Purchase Date of an Offering shall be distributed in full to the participant after such Purchase Date, without interest.

(b) No rights granted under the Plan may be exercised to any extent unless the shares to be issued upon such exercise under the Plan (including rights granted thereunder) are covered by an effective registration statement pursuant to the Securities Act of 1933, as amended (the “**Securities Act**”) and the Plan is in material compliance with all applicable state, foreign and other securities and other laws applicable to the Plan. If on a Purchase Date in any Offering hereunder the Plan is not so registered or in such compliance, no rights granted under the Plan or any Offering shall be exercised on such Purchase Date, and the Purchase Date shall be delayed until the Plan is subject to such an effective registration statement and such compliance, except that the Purchase Date shall not be delayed more than twelve (12) months and the Purchase Date shall in no event be more than six (6) months from the Offering Date. If on the Purchase Date of any Offering hereunder, as delayed to the maximum extent permissible, the Plan is not registered and in such compliance, no rights granted under the Plan or any Offering shall be exercised and all payroll deductions accumulated during the Offering (reduced to the extent, if any, such deductions have been used to acquire stock) shall be distributed to the participants, without interest.

9. Covenants of the Company.

(a) During the terms of the rights granted under the Plan, the Company shall keep available at all times the number of shares of stock required to satisfy such rights.

(b) The Company shall seek to obtain from each federal, state, foreign or other regulatory commission or agency having jurisdiction over the Plan such authority as may be required to issue and sell shares of stock upon exercise of the rights granted under the Plan. If, after reasonable efforts, the Company is unable to obtain from any such regulatory commission or agency the authority which counsel for the Company deems necessary for the lawful issuance and sale of stock under the Plan, the Company shall be relieved from any liability for failure to issue and sell stock upon exercise of such rights unless and until such authority is obtained.

10. Use of Proceeds from Stock.

Proceeds from the sale of stock pursuant to rights granted under the Plan shall constitute general funds of the Company.

11. Rights as a Stockholder.

A participant shall not be deemed to be the holder of, or to have any of the rights of a holder with respect to, any shares subject to rights granted under the Plan unless and until the participant’s shares acquired upon exercise of rights hereunder are recorded in the books of the Company.

12. Adjustments upon Changes in Stock.

(a) If any change is made in the stock subject to the Plan, or subject to any rights granted under the Plan (through merger, consolidation, reorganization, reclassification, recapitalization, stock dividend, dividend in property other than cash, stock split, reverse stock split, split-up, spin-off, repurchase, liquidating dividend, combination of shares, exchange of shares, liquidation, dissolution or sale, transfer, exchange or other disposition of all or substantially all of the

assets of the Company, or exchange of Common Stock or other securities of the Company, issuance of warrants or other rights to purchase securities of the Company, change in corporate structure or other transaction not involving the receipt of consideration by the Company), the Plan and outstanding rights will be appropriately adjusted in the class(es) and maximum number of shares subject to the Plan and the class(es) and number of shares and price per share of stock subject to outstanding rights. Such adjustments shall be made by the Board or the Committee, the determination of which shall be final, binding and conclusive. (The conversion of any convertible securities of the Company shall not be treated as a “transaction not involving the receipt of consideration by the Company.”)

(b) In the event of: (1) a dissolution or liquidation of the Company; (2) a merger or consolidation in which the Company is not the surviving corporation; (3) a reverse merger in which the Company is the surviving corporation, but the shares of the Company’s Common Stock outstanding immediately preceding the merger are converted by virtue of the merger into other property, whether in the form of securities, cash or otherwise; or (4) the acquisition by any person, entity or group within the meaning of Section 13(d) or 14(d) of the Securities Exchange Act of 1934, as amended (the “**Exchange Act**”) or any comparable successor provisions (excluding any employee benefit plan, or related trust, sponsored or maintained by the Company or any Affiliate of the Company) of the beneficial ownership (within the meaning of Rule 13d-3 promulgated under the Exchange Act, or comparable successor rule) of securities of the Company representing at least fifty percent (50%) of the combined voting power entitled to vote in the election of directors, then, as determined by the Board in its sole discretion (i) any surviving or acquiring corporation may assume outstanding rights or substitute similar rights for those under the Plan, (ii) such rights may continue in full force and effect, (iii) participants’ accumulated payroll deductions may be used to purchase Common Stock immediately prior to the transaction described above and the participants’ rights under the ongoing Offering terminated, (iv) all outstanding rights shall terminate without being exercised, or (v) all outstanding rights shall be purchased for an amount of cash equal to the amount that could have been obtained upon the exercise of such rights had such rights been currently exercisable, or the replacement of such rights with other rights or property selected by the Board.

(c) No adjustment or action described in this paragraph 12 or in any other provision of the Plan shall be authorized to the extent that such adjustment or action would cause the Plan to fail to satisfy the requirements of Section 423 of the Code.

13. Amendment of the Plan.

(a) The Board at any time, and from time to time, may amend the Plan. However, except as provided in paragraph 12 relating to adjustments upon changes in stock, no amendment shall be effective unless approved by the shareholders of the Company within twelve (12) months before or after the adoption of the amendment if such amendment requires stockholder approval in order for the Plan to obtain employee stock purchase plan treatment under Section 423 of the Code or to comply with the requirements of Rule 16b-3 promulgated under the Exchange Act or any NASDAQ or securities exchange requirements.

(b) The Board may amend the Plan in any respect the Board deems necessary or advisable to provide eligible employees with the maximum benefits provided or to be provided under the provisions of the Code and the regulations promulgated thereunder relating to employee stock purchase plans and/or to bring the Plan and/or rights granted under it into compliance therewith.

(c) Rights and obligations under any rights granted before amendment of the Plan shall not be altered or impaired by any amendment of the Plan, except with the consent of the person to whom such rights were granted, or except as necessary to comply with any laws or governmental regulations, or except as necessary to ensure that the Plan and/or rights granted under the Plan comply with the requirements of Section 423 of the Code.

14. Designation of Beneficiary.

(a) A participant may file a written designation of a beneficiary who is to receive any shares and cash, if any, from the participant's account under the Plan in the event of such participant's death subsequent to the end of an Offering but prior to delivery to the participant of such shares and cash. In addition, a participant may file a written designation of a beneficiary who is to receive any cash from the participant's account under the Plan in the event of such participant's death during an Offering.

(b) Such designation of beneficiary may be changed by the participant at any time by written notice. In the event of the death of a participant and in the absence of a beneficiary validly designated under the Plan who is living at the time of such participant's death, the Company shall deliver such shares and/or cash to the executor or administrator of the estate of the participant, or if no such executor or administrator has been appointed (to the knowledge of the Company), the Company, in its sole discretion, may deliver such shares and/or cash to the spouse or to any one or more dependents or relatives of the participant, or if no spouse, dependent or relative is known to the Company, then to such other person as the Company may designate.

15. Termination or Suspension of the Plan.

(a) The Board in its discretion, may suspend or terminate the Plan at any time. No rights may be granted under the Plan while the Plan is suspended or after it is terminated.

(b) Rights and obligations under any rights granted while the Plan is in effect shall not be altered or impaired by suspension or termination of the Plan, except as expressly provided in the Plan or with the consent of the person to whom such rights were granted, or except as necessary to comply with any laws or governmental regulation, or except as necessary to ensure that the Plan and/or rights granted under the Plan comply with the requirements of Section 423 of the Code.

16. Effective Date of Plan.

The Plan shall become effective as determined by the Board, but no rights granted under the Plan shall be exercised unless and until the Plan has been approved by the shareholders of the Company within twelve (12) months before or after the date the Plan is adopted by the Board.

CERTIFICATION

I, Don M. Bailey, certify that:

1. I have reviewed this quarterly report on Form 10-Q of Questcor Pharmaceuticals, Inc.;
2. Based on my knowledge, this report does not contain any untrue statement of a material fact or omit to state a material fact necessary to make the statements made, in light of the circumstances under which such statements were made, not misleading with respect to the period covered by this report;
3. Based on my knowledge, the financial statements, and other financial information included in this report, fairly present in all material respects the financial condition, results of operations and cash flows of the registrant as of, and for, the periods presented in this report;
4. The registrant's other certifying officer and I are responsible for establishing and maintaining disclosure controls and procedures (as defined in Exchange Act Rules 13a-15(e) and 15d-15(e)) and internal control over financial reporting (as defined in Exchange Act Rules 13a-15(f) and 15d-15(f)) for the registrant and have:
 - a. Designed such disclosure controls and procedures, or caused such disclosure controls and procedures to be designed under our supervision, to ensure that material information relating to the registrant, including its consolidated subsidiaries, is made known to us by others within those entities, particularly during the period in which this report is being prepared;
 - b. Designed such internal control over financial reporting, or caused such internal control over financial reporting to be designed under our supervision, to provide reasonable assurance regarding the reliability of financial reporting and the preparation of financial statements for external purposes in accordance with generally accepted accounting principles;
 - c. Evaluated the effectiveness of the registrant's disclosure controls and procedures and presented in this report our conclusions about the effectiveness of the disclosure controls and procedures, as of the end of the period covered by this report based on such evaluation; and
 - d. Disclosed in this report any change in the registrant's internal control over financial reporting that occurred during the registrant's most recent fiscal quarter (the registrant's fourth fiscal quarter in the case of an annual report) that has materially affected, or is reasonably likely to materially affect, the registrant's internal control over financial reporting; and
5. The registrant's other certifying officer and I have disclosed, based on our most recent evaluation of internal control over financial reporting, to the registrant's auditors and the audit committee of the registrant's board of directors (or persons performing the equivalent functions):
 - a. All significant deficiencies and material weaknesses in the design or operation of internal control over financial reporting which are reasonably likely to adversely affect the registrant's ability to record, process, summarize and report financial information; and
 - b. Any fraud, whether or not material, that involves management or other employees who have a significant role in the registrant's internal control over financial reporting.

Date: July 29, 2011

/s/ Don M. Bailey

Don M. Bailey

President and Chief Executive Officer

(Principal Executive Officer)

CERTIFICATION

I, Michael H. Mulroy, certify that:

1. I have reviewed this quarterly report on Form 10-Q of Questcor Pharmaceuticals, Inc.;
2. Based on my knowledge, this report does not contain any untrue statement of a material fact or omit to state a material fact necessary to make the statements made, in light of the circumstances under which such statements were made, not misleading with respect to the period covered by this report;
3. Based on my knowledge, the financial statements, and other financial information included in this report, fairly present in all material respects the financial condition, results of operations and cash flows of the registrant as of, and for, the periods presented in this report;
4. The registrant's other certifying officer and I are responsible for establishing and maintaining disclosure controls and procedures (as defined in Exchange Act Rules 13a-15(e) and 15d-15(e)) and internal control over financial reporting (as defined in Exchange Act Rules 13a-15(f) and 15d-15(f)) for the registrant and have:
 - a. Designed such disclosure controls and procedures, or caused such disclosure controls and procedures to be designed under our supervision, to ensure that material information relating to the registrant, including its consolidated subsidiaries, is made known to us by others within those entities, particularly during the period in which this report is being prepared;
 - b. Designed such internal control over financial reporting, or caused such internal control over financial reporting to be designed under our supervision, to provide reasonable assurance regarding the reliability of financial reporting and the preparation of financial statements for external purposes in accordance with generally accepted accounting principles;
 - c. Evaluated the effectiveness of the registrant's disclosure controls and procedures and presented in this report our conclusions about the effectiveness of the disclosure controls and procedures, as of the end of the period covered by this report based on such evaluation; and
 - d. Disclosed in this report any change in the registrant's internal control over financial reporting that occurred during the registrant's most recent fiscal quarter (the registrant's fourth fiscal quarter in the case of an annual report) that has materially affected, or is reasonably likely to materially affect, the registrant's internal control over financial reporting; and
5. The registrant's other certifying officer and I have disclosed, based on our most recent evaluation of internal control over financial reporting, to the registrant's auditors and the audit committee of the registrant's board of directors (or persons performing the equivalent functions):
 - a. All significant deficiencies and material weaknesses in the design or operation of internal control over financial reporting which are reasonably likely to adversely affect the registrant's ability to record, process, summarize and report financial information; and
 - b. Any fraud, whether or not material, that involves management or other employees who have a significant role in the registrant's internal control over financial reporting.

Date: July 29, 2011

/s/ Michael H. Mulroy

Michael H. Mulroy
Chief Financial Officer
(Principal Financial Officer)

CERTIFICATION

I, Don M. Bailey, hereby certify pursuant to Rule 13a-14(b) or Rule 15d-14(b) of the Securities Exchange Act of 1934 and 18 U.S.C. Section 1350, that the Quarterly Report of Questcor Pharmaceuticals, Inc. on Form 10-Q for the quarterly period ended June 30, 2011 fully complies with the requirements of Section 13(a) or 15(d) of the Securities Exchange Act of 1934 and that information contained in such Quarterly Report on Form 10-Q for the period ended June 30, 2011 fairly presents in all material respects the financial condition and results of operations of Questcor Pharmaceuticals, Inc.

Date: July 29, 2011

/s/ Don M. Bailey

Don M. Bailey
President and Chief Executive Officer
(Principal Executive Officer)

This certification accompanies the Quarterly Report on Form 10-Q pursuant to Rule 13a-14(b) or Rule 15d-14(b) under the Securities Exchange Act of 1934 and 18 U.S.C. Section 1350 and shall not be deemed filed by Questcor Pharmaceuticals, Inc. for purposes of Section 18 of the Securities Exchange Act of 1934.

CERTIFICATION

I, Michael H. Mulroy, hereby certify pursuant to Rule 13a-14(b) or Rule 15d-14(b) of the Securities Exchange Act of 1934 and 18 U.S.C. Section 1350, that the Quarterly Report of Questcor Pharmaceuticals, Inc. on Form 10-Q for the quarterly period ended June 30, 2011 fully complies with the requirements of Section 13(a) or 15(d) of the Securities Exchange Act of 1934 and that information contained in such Quarterly Report on Form 10-Q for the period ended June 30, 2011 fairly presents in all material respects the financial condition and results of operations of Questcor Pharmaceuticals, Inc.

Date: July 29, 2011

/s/ Michael H. Mulroy

Michael H. Mulroy
Chief Financial Officer
(Principal Financial Officer)

This certification accompanies the Quarterly Report on Form 10-Q pursuant to Rule 13a-14(b) or Rule 15d-14(b) under the Securities Exchange Act of 1934 and 18 U.S.C. Section 1350 and shall not be deemed filed by Questcor Pharmaceuticals, Inc. for purposes of Section 18 of the Securities Exchange Act of 1934.