

PROVECTUS PHARMACEUTICALS INC
Form 10-Q
August 14, 2008
United States

Securities And Exchange Commission
Washington, DC 20549

FORM 10-Q

(Mark One)

Quarterly Report under Section 13 or 15(d) of the Securities Exchange Act of 1934

For the quarterly period ended June 30, 2008

OR

Transition Report under Section 13 or 15(d) of the Securities Exchange Act of 1934

For the transition period from _____ to _____

Commission file number: 0-9410

Provectus Pharmaceuticals, Inc.
(Exact Name of Small Business Issuer as Specified in Its Charter)

Nevada
(State or other jurisdiction of incorporation or organization)

90-0031917
(I.R.S. Employer Identification Number)

7327 Oak Ridge Highway Suite A, Knoxville, TN 37931
(Address of Principal Executive Offices)

866/594-5999
(Issuer's Telephone Number, Including Area Code)

N/A
(Former Name, Former Address and Former Fiscal Year, if Changed Since Last Report)

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

Indicate by check mark whether the registrant is a large accelerated filer, an accelerated filer, or a non-accelerated filer. (Check one):

Large Accelerated Filer Accelerated Filer

Non-Accelerated Filer Smaller reporting
company

Indicate by check mark whether the registrant is a shell company (as defined in Rule 12b-2 of the Exchange Act). Yes
o No x

The number of shares outstanding of the issuer's stock, \$0.001 par value per share, as of June 30, 2008 was
51,352,855.

Transitional Small Business Disclosure Format (check one): Yes o No x

Item 1. Financial Statements

PROVECTUS PHARMACEUTICALS, INC.
(A Development-Stage Company)

CONSOLIDATED BALANCE SHEETS

	June 30, 2008 (Unaudited)	December 31, 2007 (Audited)
Assets		
Current Assets		
Cash and cash equivalents	\$ 5,169	352,389
United States Treasury Notes, total face value \$2,806,568 and \$6,910,157, respectively	2,805,308	6,907,837
Cash in escrow	2,000,000	--
Prepaid expenses and other current assets	46,014	99,460
Total Current Assets	4,856,491	7,359,686
Equipment and furnishings, less accumulated depreciation of \$386,605 and \$381,977, respectively	38,318	42,946
Patents, net of amortization of \$3,769,457 and \$3,433,897, respectively	7,945,988	8,281,548
Other assets	27,000	27,000
	\$ 12,867,797	\$ 15,711,180
Liabilities and Stockholders' Equity		
Current Liabilities		
Accounts payable – trade	\$ 143,730	\$ 455,192
Accrued compensation and payroll taxes	109,526	274,011
Accrued consulting expense	70,000	102,037
Other accrued expenses	39,500	48,430
Total Current Liabilities	362,756	879,670
Stockholders' Equity		
Preferred stock; par value \$.001 per share; 25,000,000 shares authorized; no shares issued and outstanding	--	--
Common stock; par value \$.001 per share; 100,000,000 shares authorized; 51,352,855 and 49,399,281 shares issued and outstanding, respectively	51,353	49,399
Paid in capital	62,711,598	59,988,147

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Deficit accumulated during the development stage	(50,257,910)	(45,206,036)
Total Stockholders' Equity	12,505,041	14,831,510
	\$ 12,867,797	\$ 15,711,180

See accompanying notes to consolidated financial statements.

PROVECTUS PHARMACEUTICALS, INC.
(A Development-Stage Company)
CONSOLIDATED STATEMENTS OF OPERATIONS
(Unaudited)

	Three Months Ended June 30, 2008	Three Months Ended June 30, 2007	Six Months Ended June 30, 2008	Six Months Ended June 30, 2007	Cumulative Amounts from January 17, 2002 (Inception) Through June 30, 2008
Revenues					
OTC product revenue	\$ --	\$ --	\$ --	\$ --	\$ 25,648
Medical device revenue	--	--	--	--	14,109
Total revenues	--	--	--	--	39,757
Cost of sales	--	--	--	--	15,216
Gross profit	--	--	--	--	24,541
Operating expenses					
Research and development	1,248,668	1,063,282	2,311,784	2,152,585	13,844,949
General and administrative	1,298,446	1,160,777	2,463,440	2,366,008	24,429,469
Amortization	167,780	167,780	335,560	335,560	3,769,457
Total operating loss	(2,714,894)	(2,391,839)	(5,110,784)	(4,854,153)	(42,019,334)
Gain on sale of fixed assets	--	--	--	--	55,075
Loss on extinguishment of debt	--	--	--	--	(825,867)
Investment income	19,005	84,159	58,910	169,748	630,220
Interest expense	--	--	--	(11,409)	(8,098,004)
Net loss	\$ (2,695,889)	\$ (2,307,680)	\$ (5,051,874)	\$ (4,695,814)	\$ (50,257,910)
Basic and diluted loss per common share	\$ (0.05)	\$ (0.05)	\$ (0.10)	\$ (0.10)	
	50,981,758	45,597,872			

Weighted average number of common shares outstanding – basic and diluted	50,433,460	44,929,819
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See accompanying notes to consolidated financial statements.

PROVECTUS PHARMACEUTICALS, INC.
(A Development-Stage Company)
CONSOLIDATED STATEMENTS OF STOCKHOLDERS' EQUITY
(Unaudited)

	Common Stock				
	Number of shares	Par value	Paid-in capital	Accumulated deficit	Total
Balance, at January 17, 2002	--	\$ --	\$ --	\$ --	\$ --
Issuance to founding shareholders	6,000,000	6,000	(6,000)	--	--
Sale of stock	50,000	50	24,950	--	25,000
Issuance of stock to employees	510,000	510	931,490	--	932,000
Issuance of stock for services	120,000	120	359,880	--	360,000
Net loss for the period from January 17, 2002 (inception) to April 23, 2002 (date of reverse merger)	--	--	--	(1,316,198)	(1,316,198)
Balance, at April 23, 2002	6,680,000	\$ 6,680	\$ 1,310,320	\$ (1,316,198)	\$ 802
Shares issued in reverse merger	265,763	266	(3,911)	--	(3,645)
Issuance of stock for services	1,900,000	1,900	5,142,100	--	5,144,000
Purchase and retirement of stock	(400,000)	(400)	(47,600)	--	(48,000)
Stock issued for acquisition of Valley Pharmaceuticals	500,007	500	12,225,820	--	12,226,320
Exercise of warrants	452,919	453	--	--	453
Warrants issued in connection with convertible debt	--	--	126,587	--	126,587
Stock and warrants issued for acquisition of Pure-ific	25,000	25	26,975	--	27,000
Net loss for the period from April 23, 2002 (date of reverse merger) to December 31, 2002	--	--	--	(5,749,937)	(5,749,937)
Balance, at December 31, 2002	9,423,689	\$ 9,424	\$ 18,780,291	\$ (7,066,135)	\$ 11,723,580
Issuance of stock for services	764,000	764	239,036	--	239,800
Issuance of warrants for services	--	--	145,479	--	145,479
Stock to be issued for services	--	--	281,500	--	281,500
Employee compensation from stock options	--	--	34,659	--	34,659
Issuance of stock pursuant to Regulation S	679,820	680	379,667	--	380,347
Beneficial conversion related to convertible debt	--	--	601,000	--	601,000
Net loss for the year ended December 31, 2003	--	--	--	(3,155,313)	(3,155,313)
Balance, at December 31, 2003	10,867,509	\$ 10,868	\$ 20,461,632	\$ (10,221,448)	\$ 10,251,052
Issuance of stock for services	733,872	734	449,190	--	449,923
Issuance of warrants for services	--	--	495,480	--	495,480
Exercise of warrants	132,608	133	4,867	--	5,000
Employee compensation from stock options	--	--	15,612	--	15,612

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Issuance of stock pursuant to Regulation S	2,469,723	2,469	790,668	--	793,137
Issuance of stock pursuant to Regulation D	1,930,164	1,930	1,286,930	--	1,288,861
Beneficial conversion related to convertible debt	--	--	360,256	--	360,256
Issuance of convertible debt with warrants	--	--	105,250	--	105,250
Repurchase of beneficial conversion feature	--	--	(258,345)	--	(258,345)
Net loss for the year ended December 31, 2004	--	--	--	(4,344,525)	(4,344,525)
Balance, at December 31, 2004	16,133,876	\$ 16,134	\$ 23,711,540	\$ (14,565,973)	\$ 9,161,701
Issuance of stock for services	226,733	227	152,058	--	152,285
Issuance of stock for interest payable	263,721	264	195,767	--	196,031
Issuance of warrants for services	--	--	1,534,405	--	1,534,405
Issuance of warrants for contractual obligations	--	--	985,010	--	985,010
Exercise of warrants and stock options	1,571,849	1,572	1,438,223	--	1,439,795
Employee compensation from stock options	--	--	15,752	--	15,752
Issuance of stock pursuant to Regulation D	6,221,257	6,221	6,506,955	--	6,513,176
Debt conversion to common stock	3,405,541	3,405	3,045,957	--	3,049,795
Issuance of warrants with convertible debt	--	--	1,574,900	--	1,574,900
Beneficial conversion related to convertible debt	--	--	1,633,176	--	1,633,176
Beneficial conversion related to interest expense	--	--	39,259	--	39,529
Repurchase of beneficial conversion feature	--	--	(144,128)	--	(144,128)
Net loss for the year ended 2005	--	--	--	(11,763,853)	(11,763,853)
Balance, at December 31, 2005	27,822,977	\$ 27,823	\$ 40,689,144	\$ (26,329,826)	\$ 14,387,141
Issuance of stock for services	719,246	719	676,024	--	676,743
Issuance of stock for interest payable	194,327	195	183,401	--	183,596
Issuance of warrants for services	--	--	370,023	--	370,023
Exercise of warrants and stock options	1,245,809	1,246	1,188,570	--	1,189,816
Employee compensation from stock options	--	--	1,862,456	--	1,862,456
Issuance of stock pursuant to Regulation D	10,092,495	10,092	4,120,329	--	4,130,421
Debt conversion to common stock	2,377,512	2,377	1,573,959	--	1,576,336
Beneficial conversion related to interest expense	--	--	16,447	--	16,447
Net loss for the year ended 2006	--	--	--	(8,870,579)	(8,870,579)
Balance, at December 31, 2006	42,452,366	\$ 42,452	\$ 50,680,353	\$ (35,200,405)	\$ 15,522,400

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Issuance of stock for services	150,000	150	298,800	--	298,950
Issuance of stock for interest payable	1,141	1	1,257	--	1,258
Issuance of warrants for services	--	--	472,635	--	472,635
Exercise of warrants and stock options	3,928,957	3,929	3,981,712	--	3,985,641
Employee compensation from stock options	--	--	2,340,619	--	2,340,619
Issuance of stock pursuant to Regulation D	2,376,817	2,377	1,845,761	--	1,848,138
Debt conversion to common stock	490,000	490	367,010	--	367,500
Net loss for the year ended 2007	--	--	--	(10,005,631)	(10,005,631)
Balance, at December 31, 2007	49,399,281	\$ 49,399	\$ 59,988,147	\$ (45,206,036)	\$ 14,831,510
Issuance of stock for services	112,500	113	135,387	--	135,500
Issuance of warrants for services	--	--	111,333	--	111,333
Exercise of warrants and stock options	1,841,074	1,841	1,321,500	--	1,323,341
Employee compensation from stock options	--	--	1,155,231	--	1,155,231
Net loss for the six months ended June 30, 2008	--	--	--	(5,051,874)	(5,051,874)
Balance, at June 30, 2008	51,352,855	\$ 51,353	\$ 62,711,598	\$ (50,257,910)	\$ 12,505,041

See accompanying notes to consolidated financial statements.

PROVECTUS PHARMACEUTICALS, INC.
(A Development-Stage Company)
CONSOLIDATED STATEMENTS OF CASH FLOW
(Unaudited)

	Six Months Ended June 30, 2008	Six Months Ended June 30, 2007	Cumulative Amounts from January 17, 2002 (Inception) through June 30, 2008
Cash Flows From Operating Activities			
Net loss	\$ (5,051,874)	\$ (4,695,814)	\$ (50,257,910)
Adjustments to reconcile net loss to net cash used in operating activities			
Depreciation	4,628	4,628	409,606
Amortization of patents	335,560	335,560	3,769,457
Amortization of original issue discount	--	2,797	3,845,721
Amortization of commitment fee	--	--	310,866
Amortization of prepaid consultant expense	--	84,019	1,295,226
Amortization of deferred loan costs	--	3,713	2,261,584
Accretion of United States Treasury Notes	(16,451)	(101,167)	(373,295)
Loss on extinguishment of debt	--	--	825,867
Loss on exercise of warrants	--	--	236,146
Beneficial conversion of convertible interest	--	--	55,976
Convertible interest	--	1,258	389,950
Compensation through issuance of stock options	1,155,231	1,426,190	5,424,329
Compensation through issuance of stock	--	--	932,000
Issuance of stock for services	135,500	110,667	6,458,148
Issuance of warrants for services	111,333	174,118	1,127,137
Issuance of warrants for contractual obligations	--	--	985,010
Gain on sale of equipment	--	--	(55,075)
(Increase) decrease in assets			
Prepaid expenses and other current assets	53,446	47,870	(46,014)
Increase (decrease) in liabilities			
Accounts payable	(311,462)	18,461	140,085
Accrued expenses	(205,452)	8,717	368,656
Net cash used in operating activities	(3,789,541)	(2,578,983)	(21,896,530)
Cash Flows from Investing activities			
Proceeds from sale of fixed asset	--	--	180,075
Capital expenditures	--	(22,127)	(62,049)
Increase in Cash in escrow	(2,000,000)	--	(2,000,000)
Proceeds from investments	9,230,000	10,005,000	39,711,644
Purchase of investments	(5,111,020)	(9,972,814)	(42,143,657)

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Net cash provided by (used in) investing activities	2,118,980	10,059	(4,313,987)
Cash Flows from Financing Activities			
Net proceeds from loans from stockholder	--	--	174,000
Proceeds from convertible debt	--	--	6,706,795
Net proceeds from sale of common stock	--	1,830,588	14,979,081
Proceeds from exercise of warrants and stock options	1,323,341	332,912	7,707,900
Cash paid to retire convertible debt	--	--	(2,385,959)
Cash paid for deferred loan costs	--	--	(747,612)
Premium paid on extinguishments of debt	--	--	(170,519)
Purchase and retirement of common stock	--	--	(48,000)

6

Net cash provided by financing activities	1,323,341	2,163,500	26,215,686
Net change in cash and cash equivalents	\$ (347,220)	\$ (405,424)	\$ 5,169
Cash and cash equivalents, at beginning of period	\$ 352,389	\$ 638,334	\$ --
Cash and cash equivalents, at end of period	\$ 5,169	\$ 232,910	\$ 5,169

Supplemental Disclosure of Noncash Investing and Financing Activities:

June 30, 2008

None

June 30, 2007

1. Debt converted to common stock of \$367,500
2. Payment of accrued interest through the issuance of stock of \$1,258
3. Issuance of stock for stock issuance costs of \$17,550 incurred in 2006
4. Stock committed to be issued for services of \$26,667 accrued at June 30, 2007

See accompanying notes to consolidated financial statements.

NOTES TO CONSOLIDATED FINANCIAL STATEMENTS
(unaudited)

1. Basis of Presentation

The accompanying unaudited condensed consolidated financial statements have been prepared in accordance with accounting principles generally accepted in the United States of America for interim financial information pursuant to Regulation S-X. Accordingly, they do not include all of the information and footnotes required by accounting principles generally accepted in the United States of America for complete financial statements. In the opinion of management, all adjustments (consisting of normal recurring accruals) considered necessary for a fair presentation have been included. Operating results for the three and six months ended June 30, 2008 are not necessarily indicative of the results that may be expected for the year ended December 31, 2008.

2. Recapitalization and Merger

Provectus Pharmaceuticals, Inc., formerly known as "Provectus Pharmaceutical, Inc." and "SPM Group, Inc.," was incorporated under Colorado law on May 1, 1978. SPM Group ceased operations in 1991, and became a development-stage company effective January 1, 1992, with the new corporate purpose of seeking out acquisitions of properties, businesses, or merger candidates, without limitation as to the nature of the business operations or geographic location of the acquisition candidate.

On April 1, 2002, SPM Group changed its name to "Provectus Pharmaceutical, Inc." and reincorporated in Nevada in preparation for a transaction with Provectus Pharmaceuticals, Inc., a privately-held Tennessee corporation ("PPI"). On April 23, 2002, an Agreement and Plan of Reorganization between Provectus Pharmaceutical and PPI was approved by the written consent of a majority of the outstanding shares of Provectus Pharmaceutical. As a result, Provectus Pharmaceuticals, Inc. issued 6,680,000 shares of common stock in exchange for all of the issued and outstanding shares of PPI. As part of the acquisition, Provectus Pharmaceutical changed its name to "Provectus Pharmaceuticals, Inc." and PPI became a wholly-owned subsidiary of Provectus. This transaction was recorded as a recapitalization of PPI.

On November 19, 2002, the Company acquired Valley Pharmaceuticals, Inc., a privately-held Tennessee corporation formerly known as Photogen, Inc., by merging PPI with and into Valley and naming the surviving corporation "Xantech Pharmaceuticals, Inc." Photogen, Inc. was separated from Photogen Technologies, Inc. in a non-pro rata split-off to some of its shareholders. The assets of Photogen, Inc. consisted primarily of the equipment and intangibles related to its therapeutic activity and were recorded at their fair value. The majority shareholders of Valley were also the majority shareholders of Provectus. Valley had no revenues prior to the transaction with the Company. By acquiring Valley, the Company acquired its intellectual property, including issued U.S. patents and patentable inventions.

3. Basic and Diluted Loss Per Common Share

Basic and diluted loss per common share is computed based on the weighted average number of common shares outstanding. Loss per share excludes the impact of outstanding options, warrants, and convertible debt as they are antidilutive. Potential common shares excluded from the calculation at June 30, 2008 and 2007 relate to 21,655,672 and 26,563,081 of warrants, and 8,915,093 and 9,084,419 of options.

4. Equity and Debt Transactions

(a) During the three months ended March 31, 2008, the Company issued 100,000 shares of common stock to consultants in exchange for services. Consulting costs charged to operations were \$122,500. During the three months ended June 30, 2008, the Company issued 12,500 shares of common stock to consultants in exchange for services. Consulting costs charged to operations were \$13,000.

(b) During the three months ended March 31, 2008, the Company issued 60,000 warrants to consultants in exchange for services. Consulting costs charged to operations were \$40,657. During the three months ended March 31, 2008, 197,013 warrants were exercised for \$184,402 resulting in 197,013 shares being issued. 24,050 of the warrants exercised had an exercise price of \$1.00 that was reduced to \$0.80. Additional consulting costs of \$4,810 were charged to operations as a result of the reduction of the exercise price of the 24,050 warrants. During the three months ended March 31, 2008, 143,999 warrants were forfeited. Additionally, 330,881 shares committed to be issued as of December 31, 2007 were issued January 2, 2008. During the three months ended June 30, 2008, the Company issued 12,000 warrants to consultants in exchange for services. Consulting costs charged to operations were \$5,254. During the three months ended June 30, 2008, 1,075,104 warrants were exercised for \$980,064 resulting in 1,075,104 shares being issued. 576,012 of the warrants exercised had an exercise price of \$1.00 that was reduced to \$0.90. Additional consulting costs of \$57,602 were charged to operations as a result of the reduction of the exercise price of the 576,012 warrants. 15,050 of the warrants exercised had an exercise price of \$1.00 that was reduced to \$0.80. Additional consulting costs of \$3,010 were charged to operations as a result of the reduction of the exercise price of the 15,050 warrants.

5. Stock-Based Compensation

One employee of the Company exercised a total of 193,281 options during the three months ended June 30, 2008 at exercise prices ranging from \$0.32 to \$1.02 per share of common stock for \$109,600. Another employee of the Company exercised a total of 44,795 options during the three months ended June 30, 2008 at an exercise price of \$1.10 per share of common stock for \$49,275. On June 3, 2008, the Company issued 50,000 stock options to a newly appointed Member of the Board. The options vested on the date of grant. The exercise price is the fair market price on the date of issuance, and all options were outstanding at June 30, 2008. On June 27, 2008, the Company issued 200,000 stock options to its re-elected Members of the Board. The options vested on the date of grant. The exercise price is the fair market price on the date of issuance, and all options were outstanding at June 30, 2008.

Effective January 1, 2006, the Company adopted FASB 123R. This change in accounting replaces existing requirements under FASB 123 and eliminates the ability to account for share-based compensation transaction using APB 25. The compensation cost relating to share-based payment transactions is measured based on the fair value of the equity or liability instruments issued. For purposes of estimating the fair value of each stock option on the date of grant, the Company utilized the Black-Scholes option-pricing model. The Black-Scholes option valuation model was developed for use in estimating the fair value of traded options, which have no vesting restrictions and are fully transferable. In addition, option valuation models require the input of highly subjective assumptions including the expected volatility factor of the market price of the company's common stock (as determined by reviewing its historical public market closing prices). Because the Company's employee stock options have characteristics significantly different from those of traded options and because changes in the subjective input assumptions can materially affect the fair value estimate, in management's opinion, the existing models do not necessarily provide a reliable single measure of the fair value of its employee stock options. Included in the results for the three and six months ended June 30, 2008, is \$661,000 and \$1,155,231, respectively, of stock-based compensation expense which relates to the fair value of stock options. Included in the results for the three and six months ended June 30, 2007, is \$852,795 and \$1,426,190, respectively, of stock-based compensation expense which relates to the fair value of stock options.

6. Cash in Escrow

The Company invested the Cash held in Escrow at June 30, 2008 into preferred units on July 1, 2008 and the investment was liquidated on August 7, 2008.

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

The following discussion is intended to assist in the understanding and assessment of significant changes and trends related to our results of operations and our financial condition together with our consolidated subsidiaries. This discussion and analysis should be read in conjunction with the consolidated financial statements and notes thereto included elsewhere in this Quarterly Report on Form 10-Q. Historical results and percentage relationships set forth in the statement of operations, including trends which might appear, are not necessarily indicative of future operations.

Capital Structure

Our ability to continue as a going concern is reasonably assured due to our financing completed during 2006 and warrants exercised in 2007 and 2008. At the current rate of expenditures, we do not plan to raise additional capital until late 2008, although our existing funds are sufficient to meet anticipated needs throughout 2008 and well into 2009.

We have implemented our integrated business plan, including execution of the current and next phases in clinical development of our pharmaceutical products and continued execution of research programs for new research initiatives.

We intend to proceed as rapidly as possible with the asset sale and licensure of our OTC products that can be sold with a minimum of regulatory compliance and with the further development of revenue sources through licensing of our existing medical device and biotech intellectual property portfolio. Although we believe that there is a reasonable basis for our expectation that we will become profitable due to the asset sale and licensure of our OTC products, we cannot assure you that we will be able to achieve, or maintain, a level of profitability sufficient to meet our operating expenses.

Our current plans include continuing to operate with our four employees during the immediate future, but we have added additional consultants and anticipate adding more consultants in the next 12 months. Our current plans also include minimal purchases of new property, plant and equipment, and increased research and development for additional clinical trials.

Plan of Operation

With the reorganization of Provectus and PPI and the acquisition and integration into the Company of Valley and Pure-ific, we believe we have obtained a unique combination of core intellectual properties and OTC and other non-core products. This combination represents the foundation for an operating company that we believe will provide both profitability and long-term growth. In 2007 and the first six months of 2008, we continued to carefully control expenditures in preparation for the asset sale and licensure or spin out of our OTC products, medical device and biotech technologies, and we will issue equity only when it makes sense and primarily for purposes of attracting strategic investors.

In the short term, we intend to develop our business by selling the OTC assets and licensing our existing OTC products, principally Pure-Stick, GloveAid and Pure-ific. We are also now considering a spin out of the wholly-owned subsidiary that contains the OTC assets. We will also sell and/or license our medical device and biotech technologies and consider a spin out of those non-core wholly-owned subsidiaries. In the longer term, we expect to continue the process of developing, testing and obtaining the approval of the U. S. Food and Drug Administration for prescription drugs in particular. Additionally, we have restarted our research programs that will identify additional conditions that our intellectual properties may be used to treat as well as additional treatments for those and other conditions.

We have continued to make significant progress with the major research and development projects expected to be ongoing in the next 12 months. Our expanded Phase 1 metastatic melanoma clinical trial and the second group of our expanded Phase 1 breast carcinoma clinical trial was completed in April 2007 for approximately \$1,000,000 in the aggregate, most of which has been expended in 2005 and 2006. The planning phase for the expected Phase 2 trial in metastatic melanoma has been completed which will cost approximately \$3,000,000 through 2008. This includes expenditures in 2008 to significantly advance the Phase 2 trial in metastatic melanoma that commenced in August 2007 and which may provide pivotal efficacy. Additionally, we planned \$1,000,000 of expenditures in 2007 and 2008 to substantially advance our work with other oncology indications which included the initiation of the third group of our expanded Phase 1 breast carcinoma clinical trial. Our Phase 2 psoriasis trial commenced in November 2007 and will cost approximately \$1,500,000 over 12 months. Our Phase 2 atopic dermatitis trial commenced in May 2008 and the cost is included in the psoriasis trial budget. Our Phase 1- 2 liver cancer trial is expected to cost approximately \$500,000 in total and is expected to commence shortly in 2008. Total research and development project expense in 2007 was approximately \$3,000,000. We anticipate expending the same amount in 2008. The remaining research and development expense in 2007 does not specifically relate to the above project expense in 2007.

Comparison of Three and Six Months Ended June 30, 2008 and June 30, 2007.

Revenues

OTC Product Revenue was \$-0- in both the three and six months ended June 30, 2008 and 2007. We discontinued our proof-of-concept program in November 2006 and have therefore ceased selling our OTC products. There was no medical device revenue in both the three and six months ended June 30, 2008 and 2007. The lack of medical device revenue resulted due to no emphasis on selling. The Company has designated the OTC and medical device products as non-core and is considering the sale of the underlying assets in conjunction with the planned spin-out of the respective wholly-owned subsidiaries.

Research and development

Research and development costs of \$1,248,668 for the three months ended June 30, 2008 included depreciation expense of \$2,314, consulting and contract labor of \$355,049, lab supplies and pharmaceutical preparations of \$45,889, insurance of \$18,456, legal of \$50,618, payroll of \$757,808, and rent and utilities of \$18,534. Research and development costs of \$1,063,282 for the three months ended June 30, 2007 included depreciation expense of \$2,314, consulting and contract labor of \$166,755, lab supplies and pharmaceutical preparations of \$84,263, insurance of \$24,088, legal of \$90,925, payroll of \$680,016, and rent and utilities of \$14,921. The increase in consulting and contract labor is primarily the result of expense for the Phase 2 metastatic melanoma clinical trial and related melanoma study expenditures. The increase in payroll is the result of bonuses.

Research and development costs of \$2,311,784 for the six months ended June 30, 2008 included depreciation expense of \$4,628, consulting and contract labor of \$531,094, lab supplies and pharmaceutical preparations of \$65,381, insurance of \$36,978, legal of \$142,566, payroll of \$1,491,809, and rent and utilities of \$39,328. Research and development costs of \$2,152,585 for the six months ended June 30, 2007 included depreciation expense of \$4,628, consulting and contract labor of \$304,493, lab supplies and pharmaceutical preparations of \$99,011, insurance of \$43,656, legal of \$143,755, payroll of \$1,525,984, and rent and utilities of \$31,058. The increase in consulting and contract labor is primarily the result of expense for the Phase 2 metastatic melanoma clinical trial and related melanoma study expenditures.

General and administrative

General and administrative expenses increased by \$137,669 in the three months ended June 30, 2008 to \$1,298,446 from \$1,160,777 for the three months ended June 30, 2007. The increase resulted primarily from higher payroll expenses for general corporate purposes as a result of raises and bonuses.

General and administrative expenses increased by \$97,432 in the six months ended June 30, 2008 to \$2,463,440 from \$2,366,008 for the six months ended June 30, 2007. The components of general and administrative expenses are consistent and comparable during the six months ended June 30, 2008 and 2007.

Cash Flow

As of June 30, 2008, we held approximately \$4,800,000 in cash and short-term United States Treasury Notes. At our current cash expenditure rate, the Company's current funds will be sufficient to meet our current and planned needs in 2008 and well into 2009. We have been increasing our expenditure rate by accelerating some of our research programs for new research initiatives; in addition, we are seeking to improve our cash flow through the asset sale and licensure of our OTC products as well as other non-core assets. However, we cannot assure you that we will be successful in selling the OTC and other non-core assets and licensing our existing OTC products. Moreover, even if we are successful in improving our current cash flow position, we nonetheless plan to need additional funds to meet our long-term requirements in 2009 and beyond. We anticipate that these funds will come from the proceeds of private

placements, the exercise of existing warrants outstanding, or public offerings of debt or equity securities.

Capital Resources

As noted above, our present cash flow is currently sufficient to meet our short-term operating needs. The Company's cash will be used to finance the current and next phases in clinical development of our pharmaceutical products. We anticipate that any required funds for our operating and development needs beyond 2008 will come from the proceeds of private placements, the exercise of existing warrants outstanding, or public offerings of debt or equity securities. While we believe that we have a reasonable basis for our expectation that we will be able to raise additional funds, we cannot assure you that we will be able to complete additional financing in a timely manner. In addition, any such financing may result in significant dilution to shareholders. For further information on funding sources, please see the notes to our financial statements included in this report.

Critical Accounting Policies

Long-Lived Assets

We review the carrying values of our long-lived assets for possible impairment whenever an event or change in circumstances indicates that the carrying amount of the assets may not be recoverable. Any long-lived assets held for disposal are reported at the lower of their carrying amounts or fair value less cost to sell.

Patent Costs

Internal patent costs are expensed in the period incurred. Patents purchased are capitalized and amortized over the remaining life of the patent. The patents are being amortized over the remaining lives of the patents, which range from 11-14 years. Annual amortization of the patents is expected to be approximately \$671,000 per year for the next five years.

Stock-Based Compensation

We adopted Financial Accounting Standards Board (“FASB”) Statement No. 123 (revised 2004), “Share-Based Payment” (FASB 123R), effective January 1, 2006 under the modified prospective method, which recognizes compensation cost beginning with the effective date (a) based on the requirements of FASB 123R for all share-based payments granted after the effective date and to awards modified, repurchased, or cancelled after that date and (b) based on the requirements of FASB 123 for all awards granted to employees prior to the effective date of FASB 123R that remain unvested on the effective date. There was no cumulative effect of our initially applying this Statement. At June 30, 2008 we have estimated that an additional \$150,836 will be expensed over the applicable remaining vesting periods for all share-based payments granted to employees on or before December 31, 2005 which remained unvested on January 1, 2006.

The compensation cost relating to share-based payment transactions is measured based on the fair value of the equity or liability instruments issued and is expensed on a straight-line basis. For purposes of estimating the fair value of each stock option on the date of grant, we utilized the Black-Scholes option-pricing model. The Black-Scholes option valuation model was developed for use in estimating the fair value of traded options, which have no vesting restrictions and are fully transferable. In addition, option valuation models require the input of highly subjective assumptions including the expected volatility factor of the market price of the company’s common stock (as determined by reviewing its historical public market closing prices). Because our employee stock options have characteristics significantly different from those of traded options and because changes in the subjective input assumptions can materially affect the fair value estimate, in management’s opinion, the existing models do not necessarily provide a reliable single measure of the fair value of its employee stock options.

Research and Development

Research and development costs are charged to expense when incurred. An allocation of payroll expenses is made based on a percentage estimate of time spent. The research and development costs include the following: depreciation expense, consulting and contract labor, lab supplies and pharmaceutical preparations, insurance, legal, payroll, and rent and utilities.

Accounting Pronouncement

In September of 2006, the FASB issued Statement No. 157, Fair Value Measurements, or SFAS 157. SFAS 157 establishes a standard framework for measuring fair value in generally accepted accounting principles (GAAP), clarifies the definition of “fair value” within that framework, and expands disclosures about the use of fair value measurements. The Company adopted SFAS 157 which had no impact on our financial statements.

Contractual Obligations - Leases

We lease office and laboratory space in Knoxville, Tennessee, on an annual basis, renewable for one year at our option. Our current lease term expired at June 30, 2008 and it was subsequently renewed in August 2008 with an expiration of June 30, 2009.

Forward-Looking Statements

This Quarterly Report on Form 10-Q contains forward-looking statements regarding, among other things, our anticipated financial and operating results. Forward-looking statements reflect our management's current assumptions, beliefs, and expectations. Words such as "anticipate," "believe," "estimate," "expect," "intend," "plan," and similar expressions are intended to identify forward-looking statements. While we believe that the expectations reflected in our forward-looking statements are reasonable, we can give no assurance that such expectations will prove correct. Forward-looking statements are subject to risks and uncertainties that could cause our actual results to differ materially from the future results, performance, or achievements expressed in or implied by any forward-looking statement we make. Some of the relevant risks and uncertainties that could cause our actual performance to differ materially from the forward-looking statements contained in this report are discussed below under the heading "Risk Factors" and elsewhere in this Quarterly Report on Form 10-Q. We caution investors that these discussions of important risks and uncertainties are not exclusive, and our business may be subject to other risks and uncertainties which are not detailed there.

Investors are cautioned not to place undue reliance on our forward-looking statements. We make forward-looking statements as of the date on which this Quarterly Report on Form 10-Q is filed with the SEC, and we assume no obligation to update the forward-looking statements after the date hereof whether as a result of new information or events, changed circumstances, or otherwise, except as required by law.

Item 3. Quantitative and Qualitative Disclosures About Market Risk.

None.

Item 4T. Controls and Procedures.

(a) Evaluation of Disclosure Controls and Procedures. Our chief executive officer and chief financial officer have evaluated the effectiveness of the design and operation of our "disclosure controls and procedures" (as that term is defined in Rule 13a-15(e) under the Exchange Act) as of June 30, 2008, the end of the fiscal quarter covered by this Quarterly Report on Form 10-Q. Based on that evaluation, the chief executive officer and chief financial officer have concluded that our disclosure controls and procedures are effective to ensure that material information relating to the Company and the Company's consolidated subsidiaries is made known to such officers by others within these entities, particularly during the period this Quarterly Report on Form 10-Q was prepared, in order to allow timely decisions regarding required disclosure.

(b) Changes in Internal Controls. There has been no change in our internal control over financial reporting that occurred during the fiscal quarter covered by this Quarterly Report on Form 10-Q that has materially affected, or is reasonably likely to materially affect, our internal control over financial reporting.

PART II. OTHER INFORMATION

Item 1. Legal Proceedings.

The Company was not involved in any legal proceedings during the fiscal quarter covered by this Quarterly Report of Form 10-Q.

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

Recent Sales of Unregistered Securities

None.

Item 3. Defaults Upon Senior Securities.

None.

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

(a) Our annual meeting of shareholders was held on June 19, 2008.

(b) The following is a list of all nominees for Director of the Company who were elected at the annual meeting and whose term of office continued after the annual meeting:

H. Craig Dees
 Timothy D. Scott
 Eric A. Wachter
 Stuart R. Fuchs

(c) There were present at the annual meeting in person or by proxy 26,375,922 (representing 52%) shares of our common stock out of a total of 50,930,931 shares of our common stock issued and outstanding and entitled to vote at the annual meeting.

(d) The results of the vote of the shareholders taken at the annual meeting by ballot and by proxy as solicited by us on behalf of the board of directors for the election of the nominees for our board of directors were as follows:

Nominee	For	Against	Withheld Authority
H. Craig Dees	26,337,022	400	38,500
Timothy C. Scott	26,337,022	400	38,500
Eric A. Wachter	26,337,022	400	38,500
Stuart Fuchs	26,337,022	400	38,500

Item 5. Other Information.

None.

Item 6. Exhibits

31.1 Certification Pursuant to Rule 13a-14(a) (Section 302 Certification), dated August 14, 2008, executed by H. Craig Dees, Ph.D., Chief Executive Officer of the Company.

31.2 Certification Pursuant to Rule 13a-14(a) (Section 302 Certification), dated August 14, 2008, executed by Peter R. Culpepper, Chief Financial Officer of the Company.

32.1 Certification Pursuant to 18 U.S.C. ss. 1350 (Section 906 Certification), dated August 14, 2008, executed by H. Craig Dees, Ph.D., Chief Executive Officer of the Company, and Peter R. Culpepper, Chief Financial Officer of the Company.

Signatures

In accordance with Section 13 or 15(d) of the Exchange Act, the Registrant caused this report to be signed on its behalf by the undersigned, thereunto duly authorized.

Provectus Pharmaceuticals, Inc.

Date: August 14, 2008

By: /s/ H. Craig Dees, Ph.D.
H. Craig Dees, Ph.D.
Chief Executive Officer

EXHIBIT INDEX

Exhibit No. Description

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