U.S. SECURITIES AND EXCHANGE COMMISSION WASHINGTON, D.C. 20549 FORM 10-Q

(MARK ONE) [X] QUARTERLY REPORT UNDER SECTION 13 OR 15(d) OF THE SECURITIES EXCHANGE ACT OF 1934 For the quarterly period ended September 30, 1998 [] TRANSITION REPORT UNDER SECTION 13 OR 15(d) OF THE EXCHANGE ACT For the transition period from ______ to ___ Commission File Number 0-26918 CYTOCLONAL PHARMACEUTICS INC. (Exact Name of Small Business Issuer as Specified in Its Charter) DELAWARE
(State or Other jurisdication (I.R.S. Employer Identification Number) or Organization) 9000 Harry Hines Boulevard, Suite 330, Dallas, Texas 75235 (Address of Principal Executive Offices) (214)-353-2922 _____ (Issuer's Telephone Number, Including Area Code) (Former Name, Former Address and Former Fiscal Year, if changed since last report) Check whether the issuer: (1) filed all reports required to be filed by Section 13 or 15(d) of the Exchange Act during the past 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 X NO APPLICABLE ONLY TO CORPORATE ISSUERS State the number of shares outstanding of each of the issuer's classes of common equity, as of the latest practicable date: 10,208,441 SHARES OF COMMON STOCK, \$.01 PAR VALUE, OUTSTANDING AS OF NOVEMBER 9, 1998. CYTOCLONAL PHARMACEUTICS INC. TABLE OF CONTENTS <TABLE> <CAPTION> Page(s) PART I. FINANCIAL INFORMATION <C> <S>Item 1. -- Financial Statements:

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Item 1. Financial Statements				
CYTOCLONAL PHA	ARMACEUTICS INC.			
BALANCE SHE	ETS			
	EPTEMBER 30, DECEMBER 31, 1998 1997 UNAUDITED)			
ASSETS				
~~Current assets:~~				
Cash (principally money market)	\$ 7,801,000 \$ 1,849,000			
Prepaid expenses and other current a	108,000 35,000			
Total current assets	7,909,000 1,884,000			
Equipment, net	133,000 127,000			
Patent rights, less accumulated amortiz \$521,000 and \$463,000	zation of 729,000 787,000			
Other assets	4,000 4,000			
TOTAL	\$ 8,775,000 \$ 2,802,000			
LIABILITIES AND STOCKHOLDERS' EQUITY				
Current Liabilities:				
Accounts payable and accrued expenses 473,000 460,000 Deferred revenue from research and development collaborative contract 267,000				
Current portion of royalties payable				
Total current liabilities	881,000 554,000			

Royalties payable less current portion	1,031,000	1,125,000

Total liabilities 1,912,000 1,679,000

Stockholders' equity:

Preferred stock - \$.01 par value, 10,000,000 shares authorized; 764,003 and 934,563 shares of Series A convertible preferred issued and outstanding at September 30, 1998 and December 31, 1997, respectively (liquidation value \$1,910,000 and \$2,336,000 at September 30, 1998 and December 31, 1997, respectively) 8,000 9,000

Common Stock - \$.01 par value, 30,000,000 shares authorized: 10,191,252 and 8,793,998 shares issued and outstanding at September 30, 1998 and

December 31, 1997, respectively 102,000 88,000

23,792,000 Additional paid-in capital 16,130,000

Accumulated Deficit (17,039,000) (15,104,000)

Total Stockholders' Equity 6.863,000 1.123.000

TOTAL \$ 8,775,000 \$ 2,802,000

</TABLE>

CYTOCLONAL PHARMACEUTICS INC.

STATEMENTS OF OPERATIONS (UNAUDITED)

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	THREE MONTHS ENDED SEPTEMBER 30,			
-	1998	1997	1998	1997
<\$>	<c></c>	<c></c>	<c></c>	<c></c>
Revenue:				
Licensing & research collaborative				

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\$ 194,000 - \$ 983,000 agreement

Operating Expenses:

Research and development \$351,000 \$ 362,000 \$ 1,173,000 \$ 1,050,000 General and administrative 919,000 603,000 1,930,000 1,490,000

> 3,103,000 2,540,000 1,270,000 965,000

Operating (loss) (1,076,000) (965,000) (2,120,000) (2,540,000)

Other (Income) expenses:

(100,000)Interest (income) (20,000)(187,000)(80,000)

Interest expense 2,000 2,000

> (100,000)(20,000)(185,000)(78,000)

\$ (976,000) \$ (945,000) \$(1,935,000) \$(2,462,000) NET (LOSS)

Basic and diluted

loss per common share \$(0.10) \$(0.12) \$(0.22) (0.33)

Weighted average number of shares outstanding - basic and

diluted 10,172,000 8,261,000 9,585,000 8,136,000

</TABLE>

CYTOCLONAL PHARMACEUTICS INC.

STATEMENTS OF CASH FLOWS (UNAUDITED)

<TABLE> <CAPTION>

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NINE MONTHS ENDED SEPTEMBER 30,

Cash flows from operating activities:

Net (loss) \$(1,935,000) \$(2,462,000)

Adjustments to reconcile net (loss) to net cash (used in) operating activities:

Depreciation and amortization 93,000 86,000
Value assigned to warrants and options 197,000 133,000

Equity in loss of joint venture - 16,000

Changes in:

Other assets (73,000) 19,000 Deferred revenue 267,000 -

Accounts payable and accrued expenses 41,000 (2,000)

Net cash (used in) operating activities (1,410,000) (2,210,000)

Cash flows from investing activities:

Purchase of equipment (69,000) (41,000)

Net cash (used in) investing activities (69,000) (41,000)

Cash flows from financing activities:

Net proceeds from private placement 4,838,000

Proceeds from exercise of options and warrants 2,640,000 576,000

Payment of royalties (47,000) (24,000)

Net cash provided by financing activities 7,431,000 552,000

NET INCREASE (DECREASE) IN CASH 5,952,000 (1,699,000)

Cash at beginning of period 1,849,000 2,858,000

CASH AT END OF PERIOD \$ 7,801,000 \$ 1,159,000

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CYTOCLONAL PHARMACEUTICS INC. NOTES TO FINANCIAL STATEMENTS September 30, 1998 (unaudited)

(1) FINANCIAL STATEMENT PRESENTATION

The unaudited financial statements of Cytoclonal Pharmaceutics Inc., a Delaware corporation (the "Company"), included herein have been prepared in accordance with the rules and regulations promulgated by the Securities and Exchange Commission and, in the opinion of management, reflect all adjustments (consisting only of normal recurring accruals) necessary to present fairly the results of operations for the interim periods presented. Certain information and footnote disclosures normally included in financial statements, prepared in accordance with generally accepted accounting principles, have been condensed or omitted pursuant to such rules and regulations. However, management believes that the disclosures are adequate to make the information presented not misleading. These financial statements and the notes thereto should be read in conjunction with the financial statements and the notes thereto included in the Company's Annual Report on Form 10-KSB for the fiscal year ended December 31, 1997. The results for the interim periods are not necessarily indicative of the

results for the full fiscal year.

Through March 31, 1998, the Company was in the development stage and its efforts had been principally devoted to research and development, capital formation and organizational development.

(2) RESEARCH AND COLLABORATIVE AGREEMENT

In June 1998, the Company entered into a license and research agreement with Bristol-Myers Squibb ("BMS") on two technologies related to production of paclitaxel, the active ingredient in BMS's largest selling cancer product, Taxol-Registered Trademark-. The agreement includes fees, milestone payments, research and development support and minimum and sales based royalties.

(3) LOSS PER COMMON SHARE

In 1997, the Financial Accounting Standards Board issued Statement No. 128 "Earnings Per Share". Statement No. 128 replaced the calculation of primary and fully diluted earnings per share with basic and diluted earnings per share. Unlike primary earnings per share, basic earnings per share excludes any dilutive effects of option, warrants and convertible securities. Dilutive earnings per share is very similar to the previously reported fully diluted earnings per share. In accordance with Statement No. 128, which was adopted by the Company in 1997, basic and diluted net loss per common share is based on the net loss increased by dividends on preferred stock divided by the weighted average number of common shares outstanding during the year. No effect has been given to outstanding options, warrants or convertible preferred stock in the diluted computation as their effect would be antidilutive.

(4) PRIVATE PLACEMENT

In April and May 1998, the Company completed a private placement for an aggregate of 671,035 shares of Common Stock and 335,540 Class E Warrants and received net proceeds of \$4,838,000.

(5) REVENUE RECOGNITION

Revenue from licensing and research agreements is recognized as the expenses for research and development activities performed under the

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terms of the agreements are incurred. Revenues from nonrefundable licenses and up front fees is recognized upon signing the agreement. Revenue resulting from the achievement of milestones is recognized when the milestone is achieved. Amounts received in advance of services to be performed are recorded as deferred revenue.

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

THE FOLLOWING DISCUSSION SHOULD BE READ IN CONJUNCTION WITH, AND IS QUALIFIED IN ITS ENTIRETY BY, THE FINANCIAL STATEMENTS AND THE NOTES THERETO INCLUDED IN THIS REPORT. THIS DISCUSSION CONTAINS CERTAIN FORWARD-LOOKING STATEMENTS THAT INVOLVE SUBSTANTIAL RISKS AND UNCERTAINTIES. WHEN USED IN THIS REPORT, THE WORDS "ANTICIPATE," "BELIEVE," "ESTIMATE," "EXPECT" AND SIMILAR EXPRESSIONS AS THEY RELATE TO THE COMPANY OR ITS MANAGEMENT ARE INTENDED TO IDENTIFY SUCH FORWARD-LOOKING STATEMENTS. THE COMPANY'S ACTUAL RESULTS, PERFORMANCE OR ACHIEVEMENTS COULD DIFFER MATERIALLY FROM THOSE EXPRESSED IN, OR IMPLIED BY, THESE FORWARD-LOOKING STATEMENTS. HISTORICAL OPERATING RESULTS ARE NOT NECESSARILY INDICATIVE OF THE TRENDS IN OPERATING RESULTS FOR ANY FURTHER PERIOD.

Cytoclonal Pharmaceutics Inc., a Delaware corporation (the "Company"), was duly organized and commenced operations in September 1991. To date, the Company's efforts have been principally devoted to research and development activities and organizational efforts, including the development of products for the treatment of cancer and infectious diseases, recruiting its scientific and management personnel and advisors and raising capital.

The Company's plan of operation for the next 12 months will consist of research and development and related activities aimed at:

- continued collaboration with Bristol-Myers Squibb on the development of Paclitaxel production from Microbial Fermentation and Paclitaxelspecific genes.
- further development of the Paclitaxel treatment of polycystic kidney disease, a potential new Paclitaxel indication and establishing a strategic partnership.
- evaluation of potential new proprietary microbial anticancer drugs with Bristol-Myers Squibb.
- further development of a diagnostic test using the patented LCG gene and related MAb to test in vitro serum, tissue or respiratory aspirant material for the presence of cells which may indicate a predisposition to, or early sign of, lung or other cancers.
- further testing of peptide from UCLA for inhibition of breast cancer via steroid receptors.
- further analysis of TNF-PEG technology as an anti-cancer agent in animal studies.
- testing proprietary vectors which have been constructed for the expression of specific proteins that may be utilizable for vaccines for different diseases using Mycobacteria.
- further development and potential marketing of the anti-sense technology currently being conducted at the University of Texas at Dallas.

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- developing a humanized antibody or peptide specific for the protein associated with the LCG gene and, if successful, submission of an IND for clinical trials.
- making modest improvements to the Company's laboratory and corporate facilities.
- hiring additional research technicians and a financial vice president.
- seeking to establish strategic partnerships for the development, marketing, sales and manufacturing of the Company's proposed products.

The actual research and development and related activities of the Company may vary significantly from current plans depending on numerous factors, including changes in the cost of such activities from current estimates, the results of the Company's research and development programs, the results of clinical studies, the timing of regulatory submissions, technological advances, determinations as to commercial potential and the status of competitive products. The focus and direction of the Company's operations will also be dependent upon the establishment of collaborative arrangements with other companies, the availability of financing and other factors.

For the period from July 1, 1998 to September 30, 1998, the Company incurred a net loss of \$976,000 compared to a net loss of \$945,000 for the same period in 1997. For the period from January 1, 1998 to September 30, 1998, the Company incurred a net loss of \$1,935,000 compared to a net loss of \$2,462,000 for the same period in 1997. The decrease for the nine month period from the previous year was attributable to revenue received from licensing and research and development agreements and an increase in interest income The Company expects to incur additional losses in the foreseeable future.

The Company incurred general and administrative expenses of \$1,930,000 and \$1,490,000 for the nine months ended September 1998 and September 1997, respectively. The increase from the previous year was attributable to increased legal and professional fees, including

increased patent expenses, as well as, increased insurance costs, increased public relations and financial relations expenses and increased contract labor expenses, partially offset by a decrease in consulting fees and rent expenses.

The Company incurred research and development expenses of \$1,173,000 and \$1,050,000 for the nine months ended September 1998 and September 1997, respectively. The increase was attributable to increased funding for the research programs at Washington State University and Research & Development Institute, Inc. (RDI), an increase in laboratory rental costs, and an increase in royalty payable to RDI, partially offset by a decrease in laboratory supply costs.

The Company believes that the net proceeds from its initial public offering of November 1995, the exercise of the placement agent purchase options in February 1997, the net proceeds of approximately \$4,838,000 from the private placement in April and May 1998, and proceeds of approximately \$2,640,000 from the exercise of warrants and options through September 1998

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will be sufficient to finance the Company's plan of operation through the end of 1999. There can be no assurance that the Company will generate sufficient revenues to fund its operations after such period or that any required financings will be available, through bank borrowings, debt or equity offerings, or otherwise, on acceptable terms or at all.

PART II. OTHER INFORMATION

Item 2. CHANGES IN SECURITIES AND USE OF PROCEEDS

On August 7, 1998 the Company issued an option to acquire up to 75,000 shares of its Common Stock to Synergy Group International, Inc. in connection with the execution of a financial advisory agreement with the Company. The options are exercisable at prices ranging from \$7.00 per share to \$9.00 per share. Such issuance was made in reliance upon an exemption from registration provisions of the Securities Act of 1933 set forth in Section 4(2) thereof.

Item 6. EXHIBITS AND REPORTS ON FORM 8-K

- (a) Exhibit 11 Computation of net (loss) per share Exhibit 27 Financial Data Schedule
- (b) Reports on Form 8-K None

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SIGNATURES

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

CYTOCLONAL PHARMACEUTICS INC.

Date: November 13, 1998 /s/ Daniel M. Shusterman

Daniel M. Shusterman Vice President of Operations, Treasurer and Chief Financial Officer

EXHIBIT 11

CYTOCLONAL PHARMACEUTICS INC.

COMPUTATION OF NET (LOSS) PER COMMON SHARE (unaudited)

<TABLE> <CAPTION>

THREE MONTHS NINE MONTHS

ENDED SEPTEMBER 30, ENDED SEPTEMBER 30,

1998 1997 1998 1997

Net (loss) \$ (976,000) \$ (945,000) \$(1,935,000) \$(2,462,000)

Add cumulative preferred dividend (48,000) (69,000) (143,000) (207,000)

NET (LOSS) USED FOR COMPUTATION \$(1,024,000) \$(1,014,000) \$(2,078,000) \$(2,669,000)

Weighted average number of common

shares outstanding - basic and

diluted 10,172,000 8,261,000 9,585,000 8,136,000

Net (loss) per common share -

basic and diluted \$ (0.10) \$ (0.12) \$ (0.22) \$ (0.33)

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