UNITED STATES SECURITIES AND EXCHANGE COMMISSION Washington, D.C. 20549

FORM 10-Q

(Mark One)

ľ	OUARTERLY REPORT PURSUANT 7	O SECTION 13 OR 15(d) OF	THE SECURITIES EXCHAN	NCE ACT OF 1934
$I\Lambda$	OUAKIEKLI KEFOKI FUKSUANI	O SECTION 13 OK 13(u) OF	THE SECURITIES EACHAI	NGE ACT OF 1936

For the quarterly period ended May 31, 2012

or	
[] TRANSITION REPORT PURSUANT TO SECTION 13 OR 15(d) OF T	HE SECURITIES EXCHANGE ACT OF 1934
For the transition period from to	0
Commission File Number: <u>0-123</u>	<u>305</u>
REPRO-MED SYSTEMS	
(Exact name of registrant as specified in	its charter)
<u>New York</u> (State or other jurisdiction of incorporation or organization)	13-3044880 (I.R.S. Employer Identification No.)
24 Carpenter Road, Chester New York (Address of principal executive offices)	10918 (Zip Code)
(Registrant's telephone number, including	g area code)
(Former name, former address and former fiscal year, if	changed since last report)
Indicate by check mark whether the registrant (1) has filed all reports required to be Exchange Act of 1934 during the preceding 12 months (or for such shorter period reports), and (2) has been subject to such filing requirements for the past 90 days.	that the registrant was required to file such
Indicate by check mark whether the registrant has submitted electronically and po Interactive Data File required to be submitted and posted pursuant to Rule 405 of the preceding 12 months (or for such shorter period that the registrant was require	Regulation S-T (§232.405 of this chapter) during
Indicate by check mark whether the registrant is a large accelerated filer, an accelerated company. See the definitions of "large accelerated filer," "accelerated filer for the Exchange Act.	
Large accelerated filer []	Accelerated filer []
Non-accelerated filer [] (Do not check if a smaller reporting company)	Smaller reporting company [X]
Indicate by check mark whether the registrant is a shell company (as defined in Ru	ule 12b-2 of the Exchange Act). [] Yes [X] No

As of July 12, 2012, 35,196,667 shares of common stock, \$.01 par value per share, were outstanding.

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PART I – FINANCIAL INFORMATION

ITEM 1. FINANCIAL STATEMENTS.

REPRO-MED SYSTEMS, INC. BALANCE SHEETS

		May 31, 2012		February 29, 2012	
. commo	1	Unaudited			
ASSETS					
CURRENT ASSETS					
Cash and cash equivalents	\$	1,727,378	\$	1,757,223	
Certificates of deposit		255,356		255,228	
Accounts receivable less allowance for doubtful accounts of \$19,206 and \$17,718 for May 31, 2012 and February 29, 2012, respectively		736,990		884,727	
Inventory		1,317,444		1,167,456	
Prepaid expenses		132,172		188,902	
Total Current Assets		4,169,340		4,253,536	
PROPERTY & EQUIPMENT, net		581,395		498,940	
		,		, .	
OTHER ASSETS: Patents, net of accumulated amortization of \$108,736 and \$107,640 at May 31, 2012 and February					
29, 2012, respectively		23,417		24,513	
Security deposit		30,968		28,156	
Total Other Assets		54,385		52,669	
TOTAL ASSETS	\$	4,805,120	\$	4,805,145	
		, ,		, ,	
LIABILITIES AND STOCKHOLDERS' EQUITY					
CURRENT LIABILITIES					
Note payable - current portion		2,117		2,077	
Notes payable to related parties - current portion		42,041		41,417	
Deferred capital gain - current portion		22,481		22,481	
Accounts payable		226,649		199,527	
Accrued expenses		119,417		153,800	
Accrued payroll and related taxes		25,399		41,551	
Accrued tax liability		47,296		98,000	
Total Current Liabilities		485,400	_	558,853	
OTHER LIABILITIES					
Note payable - less current portion		930		1,474	
Note payable to related parties - less current portion		427,085		437,832	
Deferred capital gain less current portion		129,275		134,895	
Deferred tax liability		121,363		121,363	
Total Other Liabilities		678,653		695,564	
Total Liabilities		1,164,053	_	1,254,417	
STOCKHOLDERS' EQUITY					
Common Stock, \$0.01 par value, 50,000,000 shares authorized, 37,471,667 shares issued, and					
35,196,667 shares outstanding at May 31, 2012 and February 29, 2012		374,717		374,717	
Additional paid-in Capital		3,263,244		3,263,244	
Retained earnings		145,106		54,767	
Lace Transport Charle 2 275 000 shares at east of March 21 2012 and Education 20 2012		3,783,067		3,692,728	
Less: Treasury Stock, 2,275,000 shares at cost at May 31, 2012 and February 29, 2012 Total Stockholders' Equity		(142,000) 3,641,067		(142,000) 3,550,728	
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TOTAL LIABILITIES AND STOCKHOLDERS' EQUITY	\$	4,805,120	\$	4,805,145	

The accompanying notes are an integral part of these Financial Statements

REPRO-MED SYSTEMS, INC. STATEMENTS OF OPERATIONS (UNAUDITED)

	F	For the Three Months Ended May 31		
		2012		2011
NET SALES	\$	1,744,165	\$	1,493,970
COST AND EXPENSES				
Cost of goods sold		637,362		540,185
Selling, general and administrative		881,770		586,778
Research and development		38,375		12,727
Depreciation and amortization		40,537		22,319
TOTAL COSTS AND EXPENSES		1,598,044		1,162,009
NET OPERATING PROFIT		146,121		331,961
OTHER INCOME/(EXPENSES)				
Gain (Loss) currency exchange		(3,013)		9,891
Interest expense		(7,207)		(8,076)
Interest and other income		1,734		5,442
TOTAL OTHER INCOME/(EXPENSES)		(8,486)		7,257
NET PROFIT BEFORE TAXES		137,635		339,218
Provision for Income Taxes	_	(47,296)		(141,259)
NET INCOME	\$	90,339	\$	197,959
NET INCOME PER SHARE				
Basic	\$		\$	0.01
Diluted	\$		\$	0.01
WEIGHTED AVERAGE NUMBER OF COMMON SHARES OUTSTANDING				_
Basic		35,196,667		36,577,667
Diluted	_	35,287,576		37,327,204

The accompanying notes are an integral part of these Financial Statements

REPRO-MED SYSTEMS, INC STATEMENTS OF CASH FLOWS (UNAUDITED)

For the Three Months Ended May 31,

		May	/ J1,	
		2012		2011
CASH FLOWS FROM OPERATING ACTIVITIES				
Net income	\$	90,339	\$	197,959
Adjustments to reconcile net income to net cash from operating activities:				
Depreciation and amortization		40,537		22,319
Deferred capital gain - building lease		(5,620)		(5,620)
Changes in operating assets and liabilities:		, , ,		
Decrease in accounts receivable		147,737		101,711
Increase in inventory		(149,988)		(161,954)
Decrease in prepaid expense		56,730		28,329
Decrease in deferred tax asset		_		45,641
Increase (decrease) in accounts payable		27,122		(26,483)
Increase (decrease) in accrued payroll and related taxes		(16,152)		24,126
Increase (decrease) in accrued expense		(34,383)		28,098
Increase in security deposits		(2,812)		_
Increase (decrease) in accrued tax liability		(50,704)		92,493
NET CASH PROVIDED BY OPERATING ACTIVITIES		102,806		346,619
CASH FLOWS FROM INVESTING ACTIVITIES				
Payments for property and equipment		(121,896)		(31,460)
Purchase of certificates of deposit		(128)		
NET CASH USED IN INVESTING ACTIVITIES		(122,024)		(31,460)
CASH FLOWS FROM FINANCING ACTIVITIES				
Payments to note payable to related parties		(10,123)		(9,534)
Payments on notes payable		(504)		(469)
NET CASH USED IN FINANCING ACTIVITIES		(10,627)		(10,003)
NET INCREASE (DECREASE) IN CASH AND CASH EQUIVALENTS		(29,845)		305,156
CASH AND CASH EQUIVILENTS, BEGINNING OF PERIOD		1,757,223		1,322,250
CASH AND CASH EQUIVILENTS, END OF PERIOD	\$	1,727,378	\$	1,627,406
Supplemental Information				
Cash paid during the periods for:				
Interest	\$	7,207	\$	8,076
Taxes	\$	98,000	\$	3,125
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The accompanying notes are an integral part of these Financial Statements

REPRO-MED SYSTEMS, INC. NOTES TO THE UNAUDITED FINANCIAL STATEMENTS

NATURE OF OPERATIONS AND SUMMARY OF SIGNIFICANT ACCOUNTING POLICIES

THE NATURE OF OPERATIONS

Repro-Med Systems, Inc. (the "Company") designs, manufactures and markets proprietary medical devices primarily for the ambulatory infusion market and emergency medical applications. The FDA regulates these products.

BASIS OF PRESENTATION

The accompanying unaudited financial statements as of May 31, 2012 have been prepared in accordance with generally accepted accounting principles in accordance with instructions to regulation S-X. Accordingly, they do not include all of the information and disclosures required by accounting principles generally accepted in the United States of America for complete financial presentation.

In the opinion of the Company's management, the financial statements contain all adjustments (consisting of normal recurring accruals) necessary to present fairly the Company's financial position as of May 31, 2012 and the results of operations and cash flow for the three-month periods ended May 31, 2012 and 2011.

The results of operations for the three months ended May 31, 2012 and 2011 are not necessarily indicative of the results to be expected for the full year. These interim financial statements should be read in conjunction with the financial statements and notes thereto of the Company and management's discussion and analysis of financial condition and results of operations included in the Company's Annual Report for the year ended February 29, 2012, as filed with the Securities and Exchange Commission on Form 10-K

USE OF ESTIMATES IN THE FINANCIAL STATEMENTS

The preparation of financial statements in conformity with U.S. generally accepted accounting principles ("GAAP") requires management to make estimates and assumptions that affect the amounts reported in the consolidated financial statements and accompanying notes. Actual results could differ from those estimates. Important estimates include but are not limited to, asset lives, valuation allowances, inventory and accruals.

SUBSEQUENT EVENTS EVALUATION

The Company has evaluated subsequent events through July 16, 2012, the date on which the financial statements were issued.

EMERGING ACCOUNTING STANDARDS

Management does not believe that any of the standards adopted by the Financial Accounting Standards Board that have been adopted but are not yet effective will have a material effect on the Company's financial reporting.

LEASED AIRCRAFT

The Company leases an aircraft from a Company controlled by the president. The lease payments aggregated were \$5,375 for the three-months ended May 31, 2012 and 2011. The original lease agreement has expired and the Company is currently on a month-to-month basis for rental payments.

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

This Quarterly Report on Form 10-Q contains certain "forward-looking" statements (as such term is defined in the Private Securities Litigation Reform Act of 1995) and information relating to us that are based on the beliefs of the management, as well as assumptions made and information currently available. Our actual results may vary materially from the forward-looking statements made in this report due to important factors such as uncertainties associated with future operating results, unpredictability related to Food and Drug Administration regulations, introduction of competitive products, limited liquidity, reimbursement related risks, government regulation of the home health care industry, success of the research and development effort, market acceptance of FREEDOM60®, availability of sufficient capital to continue operations and dependence on key personnel. When used in this report, the words "estimate," "project," "believe," "anticipate," "intend," "expect" and similar expressions are intended to identify forwardlooking statements. Such statements reflect current views with respect to future events based on currently available information and are subject to risks and uncertainties that could cause actual results to differ materially from those contemplated in such forwardlooking statements. Readers are cautioned not to place undue reliance on these forward-looking statements, which speak only as of the date hereof. These statements involve risks and uncertainties with respect to the ability to raise capital to develop and market new products, acceptance in the marketplace of new and existing products, ability to penetrate new markets, our success in enforcing and obtaining patents, obtaining required Government approvals and attracting and maintaining key personnel that could cause the actual results to differ materially. Repro-Med does not undertake any obligation to release publicly any revision to these forward-looking statements to reflect events or circumstances after the date hereof or to reflect the occurrence of unanticipated events.

THREE MONTHS ENDED May 31, 2012 VS. May 31, 2011

Net sales increased 16.7% overall from \$1,493,970 in the quarter ended May 31, 2011 to \$1,744,165 in the quarter ended May 31, 2012. This was due in part to a substantial increase in sales of RMS High-FloTM Subcutaneous Safety Needle Sets, quarter over quarter. Available in Europe since late February, 2011, the new RMS High-Flo Subcutaneous Needle Sets were formally introduced to the US market in September, 2011, through an advertising campaign that included trade shows, mailings and a direct sales campaign. The company's sales of its FREEDOM60 and RMS High-FloTM needle set product lines improved in both domestic and international markets.

Net Operating Profit was \$146,121 for the quarter ended May 31, 2012 as compared to \$331,961 from the same period last year. This change is attributable to the increases in cost of goods sold, a bonus and salary increase authorized by the Board of Directors for the CEO, expansion of sales and marketing staffs, additional advertising and promotions, as well as increases in research and development associated with the development of new products and product enhancements. Accordingly, net income decreased 54.4% from \$197,959 to \$90,339.

Selling, General and Administrative costs increased 50.3% from \$586,778 in 2011 to \$881,770 in 2012 primarily as the result of hiring additional staff in the sales and marketing areas, a bonus and salary increase authorized by the Board of Directors for the CEO calculated to partially reimburse him for a federal tax payment due in connection with his exercise of stock options to acquire shares of our common stock during our prior fiscal year, increased payroll and increased marketing expenses for advertising and trade shows.

Cost of goods sold increased \$97,177, or 18.0%, from \$540,185 to \$637,362 due to an increase in sales, an expanded production payroll, and the addition of a cost differential for a night shift with related benefits. The gross profit margin remained nearly constant this quarter at 63.5% compared to 63.8% compared to the same quarter in 2011

Interest expense decreased by 10.8% to \$7,207 in 2012 from \$8,076 for the comparative quarter in 2011 as a result of lower interest payments on long term debt.

Research and Development expenses increased \$25,648 or 201.5% from \$12,727 in 2011 to \$38,375 primarily due to R & D expenses incurred on new product development, primarily associated with the new RMS High-Flo Subcutaneous Needle Sets.

Depreciation and amortization expenses increased by \$18,218 from \$22,319 in 2011 to \$40,537 in 2012 due to increased investment in capital equipment.

LIQUIDITY AND CAPITAL RESOURCES

Net Cash provided from Operations was \$102,806 for the three months ended May 31, 2012 as compared with net cash provided by operations of \$346,619 for the previous three months ended May 31, 2011. This change is primarily due to an increase in inventory to support the RMS High-Flo Subcutaneous Needle Set product line and lower net income for the recent quarter. As the result of improved collections, accounts receivables declined by 16.7% over the three months ended May 31, 2012 even though overall sales increased.

We continue to experience an increase in sales. With these increases and the capital we currently have at the end of this period, we will continue to meet or exceed the company's liquidity needs for the next twelve months.

SUBSEQUENT EVENTS

On June 5, 2012, we announced that the results of an Active Controlled Clinical Simulated Use Study confirmed that RMS High-FlowTM Subcutaneous Needle Sets are "safety sets." The sets' butterfly wing closures encase needles after use and help to protect against accidental needle stick injuries, an area of concern to the medical community. The sets were renamed to RMS High-FloTM Subcutaneous Safety Needle Sets to reflect the safety feature. This new name is used in this filing.

BRANDING AND RECOGNITION

We continue to enhance marketing effects with an expanded schedule of advertising for its product lines in appropriate industry publications on a monthly basis. The company also exhibited at several infusion and EMS trade shows in the first quarter of the fiscal year.

FREEDOM60®

The FREEDOM60® Syringe Infusion Pump is designed for ambulatory medication infusions. For the home care patient, FREEDOM60® is an easy-to-use lightweight mechanical pump using a 60cc syringe, completely portable, cost effective and maintenance free, with no batteries to replace and no cumbersome IV pole. For the infusion professional, FREEDOM60® delivers accurate infusion rates and uniform flow profiles providing consistent transfer of medication.

The FREEDOM60® is popular in the treatment of Primary Immune Deficiency by injecting immune globulin (IgG) under the skin as a subcutaneous administration (SCIg). This method has provided patients with vastly improved quality of life with much fewer unpleasant side effects over the traditional intravenous route. The FREEDOM60® is an ideal system for this administration since the patient is able to self-medicate at home. The pump is easily configured for this application, and the FREEDOM60® is the lowest cost infusion system available in a heavily cost constrained market. We have advertised to the IgG market that FREEDOM60® operates in "dynamic equilibrium", that is, the pump finds and maintains a balance between what a patient's subcutaneous tissues are able to manage and what the pump infuses. This balance is created by a safe, limited and controlled pressure which adjusts the flow rate automatically to the patient's needs providing a reliable, faster, and more comfortable administration with fewer side effects for those patients.

We have expanded the use of the FREEDOM60® to cover antibiotics including the widely used and somewhat difficult to administer Vancomycin and beta lactams with longer infusion times. We have also found a following for FREEDOM60® for use in treating thalissemia with the drug Desferal®. In Europe, we found success in using the FREEDOM60® for pain control, specifically post-operative epidural pain administration. Our European market also uses the FREEDOM60® for chemotherapy as well as subcutaneous immune globulin.

RMS HIGH-FLO™ SUBCUTANEOUS SAFETY NEEDLE SET ADDITION TO FREEDOM60® PRODUCT LINE

We received approval from the U.S. Food and Drug Administration (FDA) on May 20, 2011, for domestic marketing of our new subcutaneous needle administration set. Previously available internationally, the needle set is branded the RMS High-FloTM Subcutaneous Safety Needle Set.

The RMS High-FloTM Subcutaneous Safety Needle Set was developed as an improvement in performance and safety over similar devices. Our design permits drug flows which are the same or faster than those achieved with larger gauge needles currently on the market. Offered in needle lengths of 4mm, 6mm, 9mm, 12mm and 14mm, the sets are available in combinations for single, double, triple, and quadruple infusions. Using a Low Residual "Y" Connector, needle sets can deliver to as many as eight infusion sites.

THE MARKET FOR INFUSION PUMPS & DISPOSABLES

The ambulatory infusion market has been rapidly changing due to reimbursement issues. Insurance reimbursement has drastically reduced the market share of high-end electronic type delivery systems as well as high-cost disposable non-electric devices, providing an opportunity for the FREEDOM60®. We believe market pressures have moved providers to consider alternatives to expensive electronic systems especially for new subcutaneous administrations which usually cannot be done with gravity. Due to cost concerns, some patients have been trained to administer intravenous drugs through IV push where the drug is pushed into the vein directly from a syringe. This is a low-cost option but has been associated with complications and considered by many to be a high-risk procedure. Thus, the overall trend has been towards syringe pumps due to the low-cost of disposables.

IMPORTANCE OF INSURANCE REIMBURSEMENT TO FREEDOM60® SALES

In order to receive more favorable Medicare reimbursement for our FREEDOM60® Syringe Infusion System, we had submitted a formal request for a HCPCS coding verification with the Statistical Analysis Durable Medical Equipment Regional Carrier (SADMERC). It was the determination that the Medicare HCPCS code(s) to bill the four Durable Medical Regional Carries (DMERCs) should be: "E0779 Ambulatory infusion pump, mechanical, reusable, for infusion 8 hours or greater." The new code significantly increases the reimbursement for the FREEDOM60® for billable syringe pump application approved by Medicare. Current approved uses under Medicare include among others, subcutaneous immune globulin, antivirals, antifungals, and chemotherapeutics. In June 2007, Medicare issued a letter of clarification stating in part:

"The FREEDOM60® Syringe Infusion Pump is the only allowable pump to be billed with the Subcutaneous Immune Globulin (SCIg). The code for this pump for dates of service 1/1/00 - 5/16/07 is E0780. For dates of service on or after 5/17/07, the correct code is E0779 per SADMERC. The items being billed must be supported by corresponding documentation. All other pumps or modifiers will result in a denial."

COMPETITION FOR THE FREEDOM60®

Competition for the FREEDOM60® for IgG is consists mostly of electrically powered infusion devices which are more costly and can create high pressures during delivery which can cause complications for the administration of IgG. However, there can be no assurance that other companies with greater resources will not enter the market with competitive products which will have an adverse effect on our sales.

In expanded uses beyond SCIg, competition for FREEDOM60® would come from gravity bags and elastomeric pumps in addition to electric/electronic pumps.

There is the potential for new drugs to enter the market, such as using Hyaluronidase which can facilitate absorption of IgG, making multiple site infusions unnecessary and changing the market conditions for devices such as the FREEDOM60®. We believe the principle behind the FREEDOM60® is ideal for all these new drug combinations, but there can be no assurance that these newer drugs will have the same needs and requirements as the current drugs being used.

There can be no assurance that Medicare will continue to provide reimbursement for the FREEDOM60® or they may allow reimbursement for other infusion pumps that are currently in the market or new ones that may enter shortly, which could adversely affect our sales into this market.

RES-Q-VAC® PORTABLE MEDICAL SUCTION

The RES-Q-VAC® Emergency Airway Suction System is a lightweight, portable, hand-operated suction device that removes fluids from a patient's airway by attaching the RES-Q-VAC® pump to various proprietary sterile and non-sterile single-use catheters sized for adult and pediatric suctioning. The one-hand operation makes it extremely effective and the product is generally found in emergency vehicles, hospitals and wherever portable aspiration is a necessity, including backup support for powered suction systems. The Full Stop Protection® filter(FSP) and disposable features of the RES-Q-VAC® reduce the risk of exposing health professional to HIV or SARS when suctioning a patient or during post treatment cleanup. All of the parts that connect to the pump are disposable.

A critical component and advantage of the RES-Q-VAC® system is our Full Stop Protection® filter, a patented filtering system that both prevents leakage and overflow of the aspirated fluids, even at full capacity, and traps virtually all air and fluid borne pathogens and potentially infectious materials within the sealable container. This protects users from potential exposure to disease and contamination. The Full Stop Protection® meets the requirement of the Occupational Safety and Health Administration 'Occupational Exposure to Bloodborne Pathogens' CFR29 1910.1030. The Company has received a letter from OSHA confirming that the RES-Q-VAC® with the Full Stop Protection® falls under the engineering controls of the Bloodborne Pathogen regulation and that the Product's use would fulfill the regulatory requirements.

Recent concerns are for diseases that are easily transmitted by small aerosolized droplets such as Asian Bird Flu, Swine Flu, and resistant tuberculosis. Other concerns are hepatitis and HIV, among others.

One advantage of our RES-Q-VAC® airway suction system is versatility. With the addition of Full Stop Protection®, we created specific custom RES-Q-VAC® kits for various vertical markets:

Emergency Medicine - we make several special kits for emergency use, which contain all the catheters necessary to treat adults as well as infants or children. These first responder kits are generally non-sterile. We also have special attachments available for the advanced paramedic to treat patients who are intubated.

Respiratory - in-home care, long-term care, situations requiring frequent suctioning such as cystic fibrosis patients, patients with swallowing disorders, elderly, patients on ventilators and with tracheostomies all benefit from the portability, cost and performance of the RES-Q-VAC®. In hospitals, the RES-Q-VAC® provides emergency backup due to power loss or breakdown of the wall suction system.

Hospital Use - for crash carts, the emergency room, patients in isolation, patient transport (e.g., from ICU to Radiology) and backup for respiratory, RES-Q-VAC® is available sterile with Full Stop Protection® for the ultimate in performance and to meet all the OSHA regulations and CDC guidelines for use in treating patients in isolation, and in any location. Hospitals are required under the EMTALA regulations to provide emergency treatment to patients anywhere in the primary facility and up to 250 yards away. The RES-Q-VAC® ensures full compliance with these regulations and helps minimize unfavorable outcomes and potential lawsuits. We provide special hospital kits, which are fully stocked to meet all hospital applications for both adult and pediatric.

Nursing Homes, Hospice, Sub-acute - we provide special configurations for dining Areas and portable suctioning for outside events and travel. Chronic suction can be accommodated with RES-Q-VAC®, which can be left by the bedside for immediate use during critical times.

Dental Applications - we offer a version of the RES-Q-VAC®, called DENTAL-EVAC®, which addresses the needs of oral surgeons for emergency backup suction during a procedure. DENTAL-EVAC® is supplied with the dental suction attachments such as saliva ejector and high volume evacuator.

Military Applications - due to its lightweight, portability, and rapid deployment, we believe that the RES-Q-VAC® is ideal for any military situation. In addition, exposure to chemical weapons of mass destruction such as Sarin is best treated by rapid, aggressive, and repeated suctioning. We believe that the RES-Q-VAC®'s compact size, powerful pump, and full protection of the user from any contamination, gives us a competitive edge in this market.

We continue actively pursuing a direct sales effort into the hospital market and continue our effort into nursing homes working with direct sales and several regional distributors in the respiratory market. We also work with national regional distributors who are well represented in the hospital respiratory market.

As part of our sales efforts in the emergency medicine field, we exhibited at the EMS Today Show in Baltimore, March 5-9, 2012. This offered emergency medicine technicians, paramedics, firefighting and police professionals, and others the opportunity to test RES-Q-VAC® for themselves and helped to support the efforts of RES-Q-VAC distributors.

COMPETITION FOR THE RES-Q-VAC®

We believe that the RES-Q-VAC® is currently the performance leader for manual, portable suction instruments. In the emergency market, the primary competition is the V-VacTM from Laerdal. The V-VacTM is more difficult to use, cannot suction infants, and cannot be used while wearing heavy gloves such as in chemical warfare or in the extreme cold. Laerdal has more resources than Repro-Med Systems and had begun marketing the V-VacTM before RES-Q-VAC® entered the market. Another competitor is Ambu, with the Res-Cue brand pump, a product similar to our design, made in China. We believe that the product is not as well made or as versatile, and may not be purchased by the military segment of the market due to lines of supply concerns. We believe that the addition of Full Stop Protection® substantially separates the RES-Q-VAC® from competitive units, which tend to leak fluid when becoming full or could pass airborne pathogens during use. There is a heightened concern from healthcare professionals concerning exposure to disease and we believe the RES-Q-VAC® provides improved protection for these users.

PART I – ITEM 3. QUANTITATIVE AND QUALITATIVE DISCLOSURES ABOUT MARKET RISK.

Not Applicable

PART I – ITEM 4. CONTROLS AND PROCEDURES.

The Company's management, including the Company's Principal Executive Officer and Chief Financial Officer, have evaluated the effectiveness of the Company's "disclosure controls and procedures "as such is defined in Rule 13a-15(e) promulgated under the Securities Exchange Act of 1934, as amended (the "Exchange Act"). Based upon their evaluations, the Principal Executive Officer and Chief Financial Officer concluded that, as of the end of the period covered by this report, the Company's disclosure controls and procedures were effective for the purpose of ensuring that the information required to be disclosed in the reports that the Company files or submits under the Exchange Act with the Securities and Exchange Commission (the "SEC") (1) is recorded, processed, summarized and reported within the time periods specified in the SEC's rules and forms and (2) is accumulated and communicated to the Company's management, including its Principal Executive Officer and Chief Financial Officer, as appropriate to allow timely decisions regarding required disclosure.

There have been no changes in the Company's internal control over financial reporting during the quarter ended May 31, 2012 that have materially affected, or are reasonably likely to materially affect, the Company's internal control over financial reporting.

PART II – OTHER INFORMATION

ITEM 1. LEGAL PROCEEDINGS.

We are, from time to time, subject to claims and suits arising in the ordinary course of business, including claims for damages for personal injuries and employment related claims.

ITEM 1A. RISK FACTORS.

Not required for smaller reporting companies.

ITEM 2. UNREGISTERED SALES OF EQUITY SECURITIES AND USE OF PROCEEDS.

None.

ITEM 3. DEFAULTS UPON SENIOR SECURITIES.

None.

ITEM 4. MINE SAFETY DISCLOSURES.

Not applicable.

ITEM 5. OTHER INFORMATION.

None.

ITEM 6. EXHIBITS.

- 31.1 Certification of Principal Executive Officer Pursuant to Section 302 of Sarbanes-Oxley Act 2002
- 31.2 Certification of Chief Financial Officer Pursuant to Section 302 of Sarbanes-Oxley Act 2002
- 32.1 Certification of Principal Executive Officer Pursuant to Section 906 of the Sarbanes-Oxley Act 2002
- 32.2 Certification of Chief Financial Officer Pursuant to Section 906 of the Sarbanes-Oxley Act 2002
- 101* Interactive Data Files of Financial Statements and Notes.

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.

REPRO-MED SYSTEMS, INC.

July 16, 2012 /s/ Andrew I. Sealfon

Andrew I. Sealfon, President Chairman of the Board, Director,

Principal Executive Officer

July 16, 2012 /s/ Michael R. Boscher

Michael R. Boscher, Treasurer and Chief Financial Officer

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^{*} In accordance with Regulation S-T, the Interactive Data Files in Exhibit 101 to the Quarterly Report on Form 10-Q shall be deemed "furnished" and not "filed".

EXHIBIT 31.1

RULE 13A-14(A) / 15D-14(A) CERTIFICATION OF PRINCIPAL EXECUTIVE OFFICER

I, Andrew I. Sealfon, certify that:

- 1) I have reviewed Form 10-Q of Repro-Med Systems, Inc. (the "Report");
- 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) I am 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 me 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) 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:
 - (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 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 16, 2012

/s/ Andrew I. Sealfon Andrew I. Sealfon Principal Executive Officer

EXHIBIT 31.2

RULE 13A-14(A) / 15D-14(A) CERTIFICATION OF TREASURER / CHIEF FINANCIAL OFFICER

I, Michael R. Boscher, certify that:

- 1) I have reviewed Form 10-Q of Repro-Med Systems, Inc. (the "Report");
- 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) I am 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 me 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) 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:
 - (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 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 16, 2012

/s/ Michael R. Boscher Michael R. Boscher Treasurer and Chief Financial Officer

EXHIBIT 32.1

CERTIFICATION PURSUANT TO 18 U.S.C. SECTION 1350 AS ADDED BY SECTION 906 OF THE SARBANES-OXLEY ACT OF 2002

In connection with the Quarterly Report of Repro-Med Systems, Inc. (the "Company") on Form 10-Q (the "Report") for the period ending May 31, 2012 as filed with the Securities and Exchange Commission, I, Andrew I. Sealfon, certify, pursuant to 18 U.S.C. Section 1350, as adopted pursuant to Section 906 of the Sarbanes-Oxley Act of 2002, that:

- (1) the Report fully complies with the requirements of Section 13(a) or 15(d) of the Securities Exchange Act of 1934; and
- (2) the information contained in the Report fairly presents, in all material respects, the financial condition and results of operations of the Company as of the dates and for the periods expressed in this report.

Date: July 16, 2012

/s/ Andrew I. Sealfon Andrew I. Sealfon Principal Executive Officer

EXHIBIT 32.2

CERTIFICATION PURSUANT TO 18 U.S.C. SECTION 1350 AS ADDED BY SECTION 906 OF THE SARBANES-OXLEY ACT OF 2002

In connection with the Quarterly Report of Repro-Med Systems, Inc. (the "Company") on Form 10-Q (the "Report") for the period ending May 31, 2012 as filed with the Securities and Exchange Commission, I, Michael R. Boscher, certify, pursuant to 18 U.S.C. Section 1350, as adopted pursuant to Section 906 of the Sarbanes-Oxley Act of 2002, that:

- (1) the Report fully complies with the requirements of Section 13(a) or 15(d) of the Securities Exchange Act of 1934; and
- (2) the information contained in the Report fairly presents, in all material respects, the financial condition and results of operations of the Company as of the dates and for the periods expressed in this report.

Date: July 16, 2012

/s/ Michael R. Boscher Michael R. Boscher Treasurer and Chief Financial Officer