

SKINVISIBLE INC
Form 10-Q
May 12, 2014

UNITED STATES

SECURITIES AND EXCHANGE COMMISSION

Washington, DC 20549

FORM 10-Q

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

For the quarterly period ended **March 31, 2014**

Transition Report pursuant to 13 or 15(d) of the Securities Exchange Act of 1934

For the transition period from _____ to _____

Commission File Number: **000-25911**

Skinvisible, Inc.

(Exact name of Registrant as specified in its charter)

Nevada

(State or other jurisdiction of incorporation or organization)

88-0344219

(IRS Employer Identification No.)

6320 South Sandhill Road, Suite 10, Las Vegas, NV 89120

(Address of principal executive offices)

702.433.7154

(Registrant's telephone number)

(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 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

Edgar Filing: SKINVISIBLE INC - Form 10-Q

Yes No

Indicate by check mark whether the registrant has submitted electronically and posted on its corporate Web site, if any, every Interactive Data File required to be submitted and posted pursuant to Rule 405 of Regulation S-T (§ 229.405 of this chapter) during the preceding 12 months (or for such shorter period that the registrant was required to submit and post such files). Yes No

Indicate by check mark whether the registrant is a large accelerated filer, an accelerated filer, a non-accelerated filer, or a smaller reporting company.

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 No

State the number of shares outstanding of each of the issuer's classes of common stock, as of the latest practicable date:
111,089,969 common shares as of March 31, 2014

Table of Contents

TABLE OF CONTENTS

Page

PART I – FINANCIAL INFORMATION

Item 1: <u>Financial Statements</u>	3
Item 2: <u>Management’s Discussion and Analysis of Financial Condition and Results of Operations</u>	4
Item 3: <u>Quantitative and Qualitative Disclosures About Market Risk</u>	11
Item 4: <u>Controls and Procedures</u>	11

PART II – OTHER INFORMATION

Item 1: <u>Legal Proceedings</u>	12
Item 1A: <u>Risk Factors</u>	12
Item 2: <u>Unregistered Sales of Equity Securities and Use of Proceeds</u>	12
Item 3: <u>Defaults Upon Senior Securities</u>	12
Item 4: <u>Mine Safety Disclosure</u>	12
Item 5: <u>Other Information</u>	12
Item 6: <u>Exhibits</u>	12

Table of Contents

PART I - FINANCIAL INFORMATION

Item 1. Financial Statements

Our consolidated financial statements included in this Form 10-Q are as follows:

F-1 Consolidated Balance Sheets as of March 31, 2014 and December 31, 2013 (unaudited);

F-2 Consolidated Statements of Operations for the three months ended March 31, 2014 and 2013 (unaudited);

F-3 Consolidated Statements of Cash Flow for the three months ended March 31, 2014 and 2013 (unaudited);

F-4 Notes to Consolidated Financial Statements.

These consolidated financial statements have been prepared in accordance with accounting principles generally accepted in the United States of America for interim financial information and the SEC instructions to Form 10-Q. In the opinion of management, all adjustments considered necessary for a fair presentation have been included. Operating results for the interim period ended March 31, 2014 are not necessarily indicative of the results that can be expected for the full year.

Table of Contents

SKINVISIBLE, INC.

CONSOLIDATED BALANCE SHEETS

(Unaudited)

	March 31, 2014	December 31, 2013
ASSETS		
Current assets		
Cash	\$316,738	\$513,420
Accounts receivable	5,478	11,591
Inventory	35,563	22,437
Due from related party	1,145	1,145
Prepaid expense and other current assets	515	515
Total current assets	359,439	549,108
Fixed assets, net of accumulated depreciation of \$323,149 and \$322,813, respectively	3,219	3,555
Intangible and other assets:		
Patents and trademarks, net of accumulated amortization of \$257,574 and \$249,064, respectively	325,326	321,388
Total assets	\$687,984	\$874,051
LIABILITIES AND STOCKHOLDERS' DEFICIT		
Current liabilities		
Accounts payable and accrued liabilities	\$531,085	\$479,622
Accrued interest payable	191,696	165,197
Loans from related party	1,910	1,832
Loans payable	1,000,000	1,000,000
Convertible notes payable, net of unamortized debt discount of \$54,345 and \$66,490, respectively	1,098,241	1,091,097
Convertible notes payable related party, net of unamortized discount of \$881,075 and \$949,672, respectively	942,287	873,689
Total current liabilities	3,765,219	3,611,437
Total liabilities	3,765,219	3,611,437
Stockholders' deficit		
Common stock; \$0.001 par value; 200,000,000 shares authorized; 111,089,969 and 110,909,969 shares issued and outstanding at March 31, 2014 and December 31, 2013, respectively	111,090	110,910
Additional paid-in capital	21,071,732	21,066,512
Stock payable	—	—
Accumulated deficit	(24,260,057)	(23,914,808)
Total stockholders' deficit	(3,077,235)	(2,737,386)
Total liabilities and stockholders' deficit	\$687,984	\$874,051

See Accompanying Notes to Consolidated Financial Statements.

F-1

Table of Contents

SKINVISIBLE, INC.

CONSOLIDATED STATEMENT OF OPERATIONS

(Unaudited)

	Three Months Ending	
	March 31, 2014	March 31, 2013
Revenues	\$ 10,038	\$ 27,287
Cost of revenues	2,321	158
Gross profit	7,717	27,129
Operating expenses		
Depreciation and amortization	8,846	8,135
Selling general and administrative	173,111	151,518
Total operating expenses	181,957	159,653
Loss from operations	(174,240)	(132,524)
Other income and (expense)		
Other income	125	—
Gain on Sale of Equipment		—
Interest expense	(169,334)	(165,023)
Gain (loss) on extinguishment of debt	(1,800)	700
Total other expense	(171,009)	(164,323)
Net loss	\$(345,249)	\$(296,847)
Basic loss per common share	\$(0.00)	\$(0.00)
Basic weighted average common shares outstanding	110,415,949	109,982,330

See Accompanying Notes to Consolidated Financial Statements.

Table of Contents

SKINVISIBLE, INC.

CONSOLIDATED STATEMENT OF CASH FLOWS

(Unaudited)

	Three Months Ended	
	March 31, 2014	March 31, 2013
Cash flows from operating activities:		
Net loss	\$(345,249)	\$(296,847)
Adjustments to reconcile net loss to net cash used in operating activities:		
Depreciation and amortization	8,846	8,135
Stock-based compensation	—	9,800
Amortization of debt discount	80,742	73,266
Stock issued for interest expense		555
Gain (loss) on extinguishment of debt	1,800	(700)
Changes in operating assets and liabilities:		
(Increase) decrease in inventory	(13,126)	(2,611)
Decrease in accounts receivable	6,113	15,299
Increase in prepaid expenses and other current assets	—	—
Increase (decrease) in accounts payable and accrued liabilities	55,063	(68,811)
Increase in accrued interest	26,499	63,110
Decrease in unearned revenue	—	(19,792)
Net cash used in operating activities	(179,312)	(218,596)
Cash flows from investing activities:		
Purchase of fixed assets and intangible assets	(12,448)	(17,536)
Net cash used in investing activities	(12,448)	(17,536)
Cash flows from financing activities:		
Proceeds from, net of payments to, related parties for loans	78	(7,526)
Payments on related parties convertible notes payable	—	(12,475)
Proceeds from convertible notes payable	—	—
Payments on convertible notes payable	(5,000)	(5,000)
Payments on loans payable	—	—
Net cash provided by (used in) financing activities	(4,922)	(25,001)
Net change in cash	(196,682)	(261,133)
Cash, beginning of period	513,420	450,507
Cash, end of period	\$316,738	\$189,374
Supplemental disclosure of cash flow information:		
Cash paid for interest	\$47,422	\$3,696
Cash paid for tax	\$—	\$—
SUPPLEMENTAL DISCLOSURE OF CASH FLOW INFORMATION:		
Non-cash investing and financing activities:		
Common Stock issued on conversion of debts	\$—	\$16,277

Edgar Filing: SKINVISIBLE INC - Form 10-Q

Common stock issued on extinguishment of debts	\$5,400	\$5,700
--	---------	---------

See Accompanying Notes to Consolidation Financial Statements.

F-3

Table of Contents

SKINVISIBLE, INC.

NOTES TO CONSOLIDATED FINANCIAL STATEMENTS

(UNAUDITED)

1. DESCRIPTION OF BUSINESS, HISTORY AND SUMMARY OF SIGNIFICANT POLICIES

Description of business - Skinvisible, Inc., (referred to as the “Company”) is focused on the development and manufacture of innovative topical, transdermal and mucosal polymer-based delivery system technologies and formulations incorporating its patent-pending formula/process for combining hydrophilic and hydrophobic polymer emulsions. The technologies and formulations have broad industry applications within the pharmaceutical, over-the-counter, personal skincare and cosmetic arenas. Additionally, the Company’s non-dermatological formulations, offer solutions for a broad spectrum of markets women’s health, pain management, and others. The Company maintains executive and sales offices in Las Vegas, Nevada.

History - Skinvisible, Inc. (referred to as the “Company”) was incorporated in Nevada on March 6, 1998, under the name of Microbial Solutions, Inc. The Company underwent a name change on February 26, 1999, when it changed its name to Skinvisible, Inc. The Company’s subsidiary’s name of Manloe Labs, Inc. was also changed to Skinvisible Pharmaceuticals, Inc.

Skinvisible, Inc. together with its subsidiary shall herein be collectively referred to as the “Company”.

Going concern - The accompanying financial statements have been prepared on a going concern basis, which contemplates the realization of assets and the satisfaction of liabilities in the normal course of business. The Company has incurred cumulative net losses of \$24,260,057 since its inception and requires capital for its contemplated operational and marketing activities to take place. The Company’s ability to raise additional capital through the future issuances of common stock is unknown. The obtainment of additional financing, the successful development of the Company’s contemplated plan of operations, and its transition, ultimately, to the attainment of profitable operations are necessary for the Company to continue operations. The ability to successfully resolve these factors raise substantial doubt about the Company’s ability to continue as a going concern. The consolidated financial statements of the Company do not include any adjustments that may result from the outcome of these aforementioned uncertainties.

Principles of consolidation - The consolidated financial statements include the accounts of the Company and its subsidiary. All significant intercompany balances and transactions have been eliminated.

Use of estimates - The preparation of consolidated financial statements in conformity with accounting principles generally accepted in the United States requires management to make estimates and assumptions that affect the reported amounts of assets and liabilities and disclosure of contingent assets and liabilities at the date of the consolidated financial statements and the reported amounts of revenue and expenses during the reporting period. Actual results could differ from those estimates.

Cash and cash equivalents - For purposes of the statement of cash flows, the Company considers all highly liquid investments and short-term debt instruments with original maturities of three months or less to be cash equivalents. There are \$316,738 and \$513,420 in cash and no cash equivalents as of March 31, 2014 and December 31, 2013, respectively.

Fair Value of Financial Instruments - The carrying amounts reflected in the balance sheets for cash, accounts payable and accrued expenses approximate the respective fair values due to the short maturities of these items.

As required by the Fair Value Measurements and Disclosures Topic of the FASB ASC, fair value is measured based on a three-tier fair value hierarchy, which prioritizes the inputs used in measuring fair value as follows: (Level 1) observable inputs such as quoted prices in active markets; (Level 2) inputs, other than the quoted prices in active markets, that are observable either directly or indirectly; and (Level 3) unobservable inputs in which there is little or no market data, which require the reporting entity to develop its own assumptions.

The three levels of the fair value hierarchy are described below:

Level 1: Unadjusted quoted prices in active markets that are accessible at the measurement date for identical, unrestricted assets or liabilities;

Level 2: Quoted prices in markets that are not active, or inputs that are observable, either directly or indirectly, for substantially the full term of the asset or liability;

Level 3: Prices or valuation techniques that require inputs that are both significant to the fair value measurement and unobservable (supported by little or no market activity).

Table of Contents

Revenue recognition

Product sales – Revenues from the sale of products (Invisicare® polymers) are recognized when title to the products are transferred to the customer and only when no further contingencies or material performance obligations are warranted, and thereby have earned the right to receive reasonably assured payments for products sold and delivered.

Royalty sales – The Company also recognizes royalty revenue from licensing its patented product formulations only when earned, when no further contingencies or material performance obligations are warranted, and thereby have earned the right to receive and retain reasonably assured payments.

Distribution and license rights sales – The Company also recognizes revenue from distribution and license rights only when earned (and are amortized over a five year period), when no further contingencies or material performance obligations are warranted, and thereby have earned the right to receive and retain reasonably assured payments.

Costs of Revenue – Cost of revenue includes raw materials, component parts, and shipping supplies. Shipping and handling costs is not a significant portion of the cost of revenue.

Accounts Receivable - Accounts receivable is comprised of uncollateralized customer obligations due under normal trade terms requiring payment within 30 days from the invoice date. The carrying amount of accounts receivable is reviewed periodically for collectability. If management determines that collection is unlikely, an allowance that reflects management's best estimate of the amounts that will not be collected is recorded. Management reviews each accounts receivable balance that exceeds 30 days from the invoice date and, based on an assessment of creditworthiness, estimates the portion, if any, of the balance that will not be collected. As of March 31, 2014, the Company had not recorded a reserve for doubtful accounts. The Company has \$1,000,000 in convertible notes payable which are secured by the accounts receivable of a license agreement the Company has with Women's Choice Pharmaceuticals, LLC on its proprietary prescription product, ProCort®.

Inventory- Substantially all inventory consists of finished goods and are valued based upon first-in first-out ("FIFO") cost, not in excess of market. The determination of whether the carrying amount of inventory requires a write-down is based on an evaluation of inventory.

Goodwill and intangible assets - The Company follows Financial Accounting Standard Board's (FASB) Codification Topic 350-10 ("ASC 350-10"), "*Intangibles – Goodwill and Other*". According to this statement, goodwill and intangible assets with indefinite lives are no longer subject to amortization, but rather an annual assessment of impairment by applying a fair-value based test. Fair value for goodwill is based on discounted cash flows, market multiples and/or appraised values as appropriate. Under ASC 350-10, the carrying value of assets are calculated at the lowest level for

which there are identifiable cash flows.

Income taxes - The Company accounts for its income taxes in accordance with FASB Codification Topic ASC 740-10, “*Income Taxes*”, which requires recognition of deferred tax assets and liabilities for future tax consequences attributable to differences between the financial statement carrying amounts of existing assets and liabilities and their respective tax bases and tax credit carry-forwards. Deferred tax assets and liabilities are measured using enacted tax rates expected to apply to taxable income in the years in which those temporary differences are expected to be recovered or settled. The effect on deferred tax assets and liabilities of a change in tax rates is recognized in income in the period that includes the enactment date.

Stock-based compensation - The Company follows the guidelines in FASB Codification Topic ASC 718-10 “*Compensation-Stock Compensation*”, which requires the measurement and recognition of compensation expense for all share-based payment awards made to employees and directors including employee stock options and employee stock purchases related to a Employee Stock Purchase Plan based on the estimated fair values.

Stock based compensation expense recognized under ASC 718-10 for the three months ended March 31, 2014 and 2013 totaled \$0 and \$9,800, respectively.

Earnings (loss) per share - The Company reports earnings (loss) per share in accordance with FASB Codification Topic ASC 260-10 “*Earnings Per Share*”, Basic earnings (loss) per share is computed by dividing income (loss) available to common shareholders by the weighted average number of common shares available. Diluted earnings (loss) per share is computed similar to basic earnings (loss) per share except that the denominator is increased to include the number of additional common shares that would have been outstanding if the potential common shares had been issued and if the additional common shares were dilutive. Diluted earnings (loss) per share has not been presented since the effect of the assumed exercise of options and warrants to purchase common shares (common stock equivalents) would have an anti-dilutive effect.

Table of Contents

2. FIXED ASSETS

Fixed assets consist of the following as of March 31, 2014 and December 31, 2013:

	March 31, 2014	December 31, 2013
Machinery and equipment	\$48,163	\$48,163
Furniture and fixtures	113,635	113,635
Computers, equipment and software	38,540	38,540
Leasehold improvements	12,569	12,569
Lab equipment	113,461	113,461
Total	326,368	326,368
Less: accumulated depreciation	(323,149)	(322,813)
Fixed assets, net of accumulated depreciation	\$3,219	\$3,555

Depreciation expense for the three months ended March 31, 2014 and 2013 was \$336 and \$331, respectively.

3. INTANGIBLE AND OTHER ASSETS

Patents and trademarks are capitalized at their historical cost and are amortized over their estimated useful lives. As of March 31, 2014, patents and trademarks total \$325,326, net of \$257,574 of accumulated amortization. Amortization expense for the three months ended March 31, 2014 and 2013 was \$8,510 and \$7,804, respectively.

License and distributor rights (“agreement”) were acquired by the Company in January 1999 and provide exclusive use distribution of polymers and polymer based products. The Company has a non-expiring term on the license and distribution rights. Accordingly, the Company annually assesses this license and distribution rights for impairment and has determined that no impairment write-down is considered necessary as of March 31, 2014.

4. UNEARNED REVENUE

On January 16, 2013, the company terminated its licensing agreement with Panalab dated January 23, 2008. The agreement provided Panalab the right to distribute, market, sell and promote Skinvisible's proprietary formulas made with Invisicare and Adapalene through-out Panalabs assigned territory. Panalab had failed to sell or sub-license the products in the territory, thereby not fulfilling the conditions as set forth in the agreement and allowing for immediate termination of the agreement. As a result of this cancelation, unearned revenue of \$19,792 has been recognized as revenue during the year ended December 31, 2013.

5. STOCK OPTIONS AND WARRANTS

The following is a summary of option activity during the three months ended March 31, 2014.

	Number of Shares	Weighted Average Exercise Price
Balance, December 31, 2013	9,300,000	\$ 0.05
Options granted and assumed	—	\$ —
Options expired	—	\$ —
Options canceled	—	—
Options exercised	—	—
Balance, March 31, 2014	9,300,000	\$ 0.05

As of March 31, 2014, 9,300,000 stock options are exercisable.

Table of Contents

On September 23, 2013, the Company granted stock options for 300,000 shares of its common stock with a strike price of \$0.03. The stock options were exercisable upon grant and have a life of 5 years. The stock options were valued at \$8,400 using the Black-Scholes option pricing model based upon the following assumptions: term of 5 years, risk free interest rate of 1.33%, a dividend yield of 0% and volatility rates of 443%. The Company recorded an expense of \$8,400 for the year ended December 31, 2013.

Stock warrants -

The following is a summary of warrants activity during the three months ended March 31, 2014.

	Number of Shares	Weighted Average Exercise Price
Balance, December 31, 2013	3,818,780	\$ 0.05
Warrants granted and assumed	—	—
Warrants expired	387,500	0.06
Warrants canceled	—	—
Warrants exercised	—	—
Balance, March 31, 2014	3,631,280	\$ 0.05

All warrants outstanding as of March 31, 2014 are exercisable.

On February 11, 2013, the Company issued a warrant for 180,915 shares of common stock to the note holder with a exercise price of \$0.05. The vesting period on these grant was immediate. The value of these warrants were estimated by using the Black-Scholes option pricing model with the following assumptions: expected life of 2 years; risk free interest rate of 0.27%; dividend yield of 0% and expected volatility of 178%. To account for such grants to non-employees, we recorded the issuance as consulting expense in the amount of \$1,683.

On February 11, 2013, the Company issued a warrant for 90,365 shares of common stock to a note holder with a exercise price of \$0.05. The vesting period on these grant was immediate. The value of these warrants were estimated by using the Black-Scholes option pricing model with the following assumptions: expected life of 2 years; risk free interest rate of 0.27%; dividend yield of 0% and expected volatility of 178%. To account for such grants to non-employees, we recorded the issuance as consulting expense in the amount of \$840.

On May 22, 2013, the Company issued a warrant for 200,000 shares of common stock to Retire Happy, LLC for Consulting services. The warrants have an exercise price of \$0.04 and the vesting period on these grants was immediate. The value of these warrants were estimated by using the Black-Scholes option pricing model with the following assumptions: expected life of 1 years; risk free interest rate of 0.11%; dividend yield of 0% and expected volatility of 167%. To account for such grants to non-employees, we recorded the issuance as consulting expense in the amount of \$3,000.

6. NOTES PAYABLE

On May 22, 2013 the Company approved a financing plan to offer accredited investors up to \$1,000,000 in secured promissory notes. During the nine months ended September 30, 2013 the Company entered into twenty-four 9% notes payable to investors and received total proceeds of \$1,000,000. The notes are due two years from the anniversary date of execution. The Notes are secured by the US Patent rights granted for the Company's Sunscreen Products: US patent number #8,128,913: "Sunscreen Composition with Enhanced UV-A Absorber Stability and Methods".

7. RELATED PARTY TRANSACTIONS

During the year ended 2014 various officers advanced funds to support the daily operations of the company. As of March 31, 2014, \$1,910 remained due to related parties as repayment for advanced monies, all related other party notes have been extinguished or re-negotiated as convertible notes. See note 8.

Table of Contents

8. CONVERTIBLE NOTES PAYABLE

Convertible Notes Payable at consists of the following:	March 31, 2014	December 31, 2013
\$52,476 face value, 10% unsecured note payable to an investor, note interest and payment are due on demand. The note could be converted to option rights for Skinvisible, Inc. shares at ten cents per share (\$0.10), these rights expired January 12, 2010. Note is currently in default, no penalties occur due to default.	\$36,476	\$36,476
\$27,000 face value 10% unsecured notes payable to investors, due October, 2012. At the written request of the investor's until the repayment date, the note may be converted at the investors option to shares of the Company's common stock at a fixed price of \$0.05 per share along with additional warrants to purchase one share for every two shares issued at the exercise price of \$0.07 per share for two years after the conversion date. The Company has determined the value associated with the beneficial conversion feature in connection with the notes to be \$19,385. The aggregate beneficial conversion feature was accreted and charged to general and administrative expenses as a financing expense in the amount of \$19,385 in the year ending December 31, 2012. The beneficial conversion feature is valued under the intrinsic value method. Interest due to lender can also be converted at a rate of (\$0.05) per share into warrants.	5,000	10,000
\$1,000,000 face value 9% unsecured notes payable to investors, due in 2015. At the investor's option until the repayment date, the note and related interest may be converted to shares of the Company's common stock a discount of 90% of the current share price after the first anniversary of the note. The Notes are secured by the accounts receivable of a license agreement the Company has with Womens Choice Pharmaceuticals, LLC on its proprietary prescription product, ProCort®. The Company has determined the value associated with the beneficial conversion feature in connection with the notes and interest to be \$111,110. The aggregate original issue discount feature has been accreted and charged to general and administrative expenses as a financing expense in the amount of \$12,144 during the three months ended March 31, 2014. The original issue discount feature is valued under the intrinsic value method.	1,000,000	1,000,000
Original issue discount	111,110	111,110
Unamortized debt discount	(54,345)	(66,490)
	\$1,098,241	\$1,091,097

Table of Contents

9. CONVERTIBLE NOTES PAYABLE RELATED PARTY

Convertible Notes Payable Related Party at consists of the following:	MArch 31, 2014	December 31, 2013
On December 31, 2011, the Company re-negotiated accrued salaries and interest for three employees. Under the terms of the agreements, the notes dated before December 31, 2010, and all salaries not previously converted were converted to promissory notes convertible into common stock with a warrant feature. The promissory notes are unsecured, due five years from issuance, and bear an interest rate of 10%. At the investor's option until the repayment date, the note may be converted to shares of the Company's common stock at a fixed price of \$0.04 per share along with additional warrants to purchase one share for every two shares issued at the exercise price of \$0.06 per share for three years after the conversion date. The Company has determined the value associated with the beneficial conversion feature in connection with the notes negotiated on December 31, 2011 to be \$1,123,078. The aggregate beneficial conversion feature has been accreted and charged to general and administrative expenses as a financing expense in the amount of \$41,972 during the three months ending March 31, 2014. The beneficial conversion feature is valued under the intrinsic value method. In the year ending December 2013, the Company made a \$51,485 in cash payments to reduce the note balance.	1,071,593	1,071,593
Unamortized debt discount	(458,021)	(499,993)
On June 30, 2012, the Company re-negotiated accrued salaries and interest for three employees. Under the terms of the agreements, the notes dated before July 1, 2011, and all salaries not previously converted were converted to promissory notes convertible into common stock with a warrant feature. The promissory notes are unsecured, due five years from issuance, and bear an interest rate of 10%. At the investor's option until the repayment date, the note may be converted to shares of the Company's common stock at a fixed price of \$0.04 per share along with additional warrants to purchase one share for every two shares issued at the exercise price of \$0.06 per share for three years after the conversion date. The Company has determined the value associated with the beneficial conversion feature in connection with the notes to be \$209,809. The aggregate beneficial conversion feature has been accreted and charged to general and administrative expenses as a financing expense in the amount of \$10,481 during the three months ending March 31, 2014. The beneficial conversion feature is valued under the intrinsic value method. On January 18, 2013, the Company made a \$3,990 cash payment to reduce the note balance.	321,032	321,032
Unamortized debt discount	(134,981)	(145,462)
On December 30 and 31, 2012, the Company re-negotiated accrued salaries and interest for three employees. Under the terms of the agreements, \$182,083 of related party notes accrued interest and salaries not previously converted were converted to promissory notes convertible into common stock with a warrant feature. The \$182,083 face value promissory notes are unsecured, due five years from issuance, and bear an interest rate of 10%. At the investor's option until the repayment date, the note may be converted to shares of the Company's common stock at a fixed price of \$0.03 per share along with additional warrants to purchase one share for every two shares issued at the exercise price of \$0.04 per share for three years after the conversion date. The Company has determined the value associated with the beneficial conversion feature in connection with the notes to be \$182,083. The	182,083	182,083

Edgar Filing: SKINVISIBLE INC - Form 10-Q

aggregate beneficial conversion feature has been accreted and charged to general and administrative expenses as a financing expense in the amount of \$9,175 during the three months ending March 31, 2014. The beneficial conversion feature is valued under the intrinsic value method.

Unamortized debt discount

(136,497) (145,672)

F-9

Table of Contents

On June 30, 2013, the Company re-negotiated accrued salaries and interest for two employees. Under the terms of the agreements, \$106,153 of accrued interest and salaries were converted to promissory notes convertible into common stock with a warrant feature. The \$106,153 face value promissory notes are unsecured, due five years from issuance, and bear an interest rate of 10%. At the investor's option until the repayment date, the note may be converted to shares of the Company's common stock at a fixed price of \$0.03 per share along with additional warrants to purchase one share for every two shares issued at the exercise price of \$0.04 per share for three years after the conversion date. The Company has determined the value associated with the beneficial conversion feature in connection with the notes to be \$70,768. The aggregate beneficial conversion feature has been accreted and charged to general and administrative expenses as a financing expense in the amount of \$3,566 during the three months ending March 31, 2014. The beneficial conversion feature is valued under the intrinsic value method.	106,152	106,152
Unamortized debt discount	(63,636)	(63,636)
On December 31, 2013, the Company re-negotiated accrued salaries and interest for three employees. Under the terms of the agreements, \$142,501 of accrued interest and salaries not previously converted were converted to promissory notes convertible into common stock with a warrant feature. The \$142,501 face value promissory notes are unsecured, due five years from issuance, and bear an interest rate of 10%. At the investor's option until the repayment date, the note may be converted to shares of the Company's common stock at a fixed price of \$0.03 per share along with additional warrants to purchase one share for every two shares issued at the exercise price of \$0.04 per share for three years after the conversion date. The Company has determined the value associated with the beneficial conversion feature in connection with the notes to be \$94,909. The aggregate beneficial conversion feature has been accreted and charged to general and administrative expenses as a financing expense in the amount of \$3,403 during the three months ending March 31, 2014. The beneficial conversion feature is valued under the intrinsic value method.	142,501	142,501
Unamortized debt discount	(91,506)	(94,909)
	\$942,286	\$873,689

Table of Contents

10. STOCKHOLDERS' DEFICIT

The Company is authorized to issue 200,000,000 shares of \$0.001 par value common stock. The Company had 111,089,969 and 110,909,969 issued and outstanding shares of common stock as of March 31, 2014 and December 31, 2013, respectively.

On January 7, 2014 the Company issued 180,000 shares with a fair value of \$5,400 to settle \$3,600 in outstanding accounts payable. As a result of this transaction a loss on extinguishment of debt of \$1,800 was recognized.

During the year ended December 31, 2013, the Company issued a total of 1,402,560 shares of common stock, with a fair value of \$44,554 for the conversion of outstanding debts of \$37,567, related interest of \$1,277 and settlement of stock payable \$1,800. The Company recorded a gain of \$3,910 on extinguishment of debts.

11. COMMITMENTS AND CONTINGENCIES

Lease obligations – The Company has operating leases for its offices. Future minimum lease payments under the operating leases for the facilities as of March 31, 2014, are as follows:

2014 \$24,868
2015 \$5,526

Rental expense, resulting from operating lease agreements, approximated \$10,367 and \$12,280 for the three months ended March 31, 2014 and 2013, respectively.

12. DEFINITIVE AGREEMENTS

On January 16, 2013, the company terminated its licensing agreement with Panalab dated January 23, 2008. The agreement provided Panalab the right to distribute, market, sell and promote the Skinvisible's proprietary formulas made with Invisicare and Adapalene through-out Panalabs assigned territory. Panalab had failed to sell or sub-license the products in the territory, thereby not fulfilling the conditions as set forth in the agreement and allowing for immediate termination of the agreement. . As a result of this cancelation the deferred revenue of \$19,792 was

recognized as revenue in the in the period ended March 31, 2013.

13. SUBSEQUENT EVENTS

The Company has evaluated events subsequent to the balance sheet through the issuance date of these financial statements in accordance with FASB ASC 855 and has determined that there are no such events that would require adjustment to, or disclosure in, the financial statements.

F-11

Table of Contents

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

Forward-Looking Statements

Certain statements, other than purely historical information, including estimates, projections, statements relating to our business plans, objectives, and expected operating results, and the assumptions upon which those statements are based, are “forward-looking statements” within the meaning of the Private Securities Litigation Reform Act of 1995, Section 27A of the Securities Act of 1933 and Section 21E of the Securities Exchange Act of 1934. These forward-looking statements generally are identified by the words “believes,” “project,” “expects,” “anticipates,” “estimates,” “intends,” “strategy,” “plan,” “may,” “will,” “would,” “will be,” “will continue,” “will likely result,” and similar expressions. Such forward-looking statements to be covered by the safe-harbor provisions for forward-looking statements contained in the Private Securities Litigation Reform Act of 1995, and are including this statement for purposes of complying with those safe-harbor provisions. Forward-looking statements are based on current expectations and assumptions that are subject to risks and uncertainties which may cause actual results to differ materially from the forward-looking statements. Our ability to predict results or the actual effect of future plans or strategies is inherently uncertain. Factors which could have a material adverse effect on our operations and future prospects on a consolidated basis include, but are not limited to: changes in economic conditions, legislative/regulatory changes, availability of capital, interest rates, competition, and generally accepted accounting principles. These risks and uncertainties should also be considered in evaluating forward-looking statements and undue reliance should not be placed on such statements. We undertake no obligation to update or revise publicly any forward-looking statements, whether as a result of new information, future events or otherwise. Further information concerning our business, including additional factors that could materially affect our financial results, is included herein and in our other filings with the SEC.

Company Overview

We, through our wholly owned subsidiary Skinvisible Pharmaceuticals Inc., are a pharmaceutical research and development (“R&D”) company that has developed and patented an innovative polymer delivery system, Invisicare® and formulated over forty topical skin products, which we out-license globally. We were incorporated in 1998, and target an estimated \$80 billion global skincare and dermatology market and a \$30 billion global over-the-counter market as well as other healthcare / medical and consumer goods markets.

With the research and development complete on forty products and numerous patents issued (technology and product patents), we are ready to monetize our investment. Our business model is to out-license our patented prescription, over-the-counter (“OTC”) and cosmeceutical products featuring Invisicare to established manufacturers and marketers of brands internationally and to maximize profits from the products we have already out-licensed. We have also recently developed a product for Netherton syndrome, for which we are seeking “orphan drug” status in both the United States and Europe. This designation has the potential to be highly lucrative, with more global companies seeing the

value of an orphan drug.

The opportunity for us to license our products has recently increased due to improving market conditions and the need for pharmaceutical companies to access external R&D companies for new products due to their own down-sizing or elimination of internal R&D departments. The demand for our products is enhanced due to the granting of key US and international patents and the completed development of a number of unique products.

Table of Contents

Our Plan for the Next Twelve Months

Our growth strategy is to:

1. Capitalize on the success of current licensees;
2. Increase the value of our current pipeline; and
3. Boost licensing revenues by securing additional licensees globally and develop a robust royalty revenue stream that will finance our future growth.

1. Capitalize On Current Licensees:

We have licensees around the globe. Three of these licensees are currently in the marketplace: Avon Products globally, Women's Choice Pharmaceuticals in the United States and Alto Pharmaceuticals in Canada.

We continue to work diligently with our licensees to ensure they have a smooth manufacturing process, ongoing R&D support and marketing feedback.

Avon Products, Inc:

Product: We have a long-term contract with Avon globally for over ten years to provide Invisicare polymer for their long-lasting lipsticks.

Sales: Invisicare polymers are purchased directly from Skinvisible.

Alto Pharmaceuticals:

Product: DermSafe®, long lasting hand sanitizer lotion launched in Canada in Q4 of 2011 for commercial / industrial use

Sales and Royalties: Alto has received Health Canada marketing approval for DermSafe and is currently marketing the product directly and seeking distributors in the commercial / healthcare marketplace.

Women's Choice Pharmaceuticals:

Product: ProCort®, long lasting prescription hemorrhoid cream launched in the United States August 2011

Sales and Royalties: ProCort continues to increase sales every quarter. Skinvisible receives a royalty based on net sales. This past year Women's Choice Pharmaceuticals LLC partnered with Advanced Medical Enterprises, LLC to market ProCort® in Puerto Rico. With over thirty pharmaceutical sales reps calling on OBGYNs in the US, Women's Choice has been successfully growing their sales of ProCort® and we look forward to increased growth in 2014. Women's Choice is seeking to form other strategic alliances in 2014 in order to increase its sales efforts by targeting new territories and targeting medical specialists which previously were not called upon.

Table of Contents

Product Launches for 2014:

We have additional products which are anticipated to be in the market in 2014.

Triclosan Hand Sanitizer & First Aid Antiseptic

Previously licensed as Safe4Hours® Antibacterial/Antimicrobial Hand Sanitizers (1% Triclosan) and Safe4Hours® First Aid Antiseptic & Skin Protectant for North America to Dermal Defense, Skinvisible has received the rights back for both products. Skinvisible anticipates setting up distributors for these products on an international basis.

DermSafe® Hand Sanitizer

Skinvisible's hand sanitizer formulated with Invisicare® and chlorhexidine gluconate has received registration in Belgium on behalf of Skinvisible. This registration allows Skinvisible to make DermSafe® available in most of Europe through a simple registration process. The Company is currently seeking licensees and/or distributors to begin the sale of DermSafe in the EU.

Mayquest Pharmaceuticals PTY

Licensed DermSafe chlorhexidine hand sanitizer for Singapore, Taiwan, Thailand, Indonesia and the Philippines for a license fee and royalty.

Received importation approval for DermSafe from Canada to Singapore. Launch pending.

Currently seeking distribution partner or sub- licensee to launch the product.

Table of Contents

2. Increasing The Value Of Skinvisible's Pipeline: Clinical Enhancement Of Pipeline

We have a pipeline of over forty products which are available for licensing. Testing is conducted in-house generating proof of concept including release of the active ingredient as well as long term shelf life (stability). Additional studies conducted on specific products including skin sensitivity, toxicity and product efficacy are outsourced to FDA compliant laboratories. These studies are critical in attracting potential licensees. Our clinical strategy is to:

(1) Add new studies for our prescription products. Our clinical strategy is to increase the amount of outsourced studies, specifically for our prescription products. Additional studies including skin penetration and skin irritation studies will add to the integrity and value of our products available for licensing.

(2) Add new long-term efficacy studies for our DermSafe® hand sanitizer. Last year we commissioned an independent laboratory to further analyze the long-term effectiveness of DermSafe® when put in contact with two bacteria; the "super bug" MRSA and E. coli, the "restaurant bug" since it is often transmitted by food and food handlers. The long-term effectiveness of two bacteria; Methicillin-resistant Staphylococcus aureus or MRSA (ATCC #33591) and Escherichia coli or E. coli (ATCC #43888") were tested up to four hours after application. The results showed that the individual arms of subjects which had DermSafe® applied and were even rinsed prior to each bacteria challenge, showed a 95.83% reduction at the 4 hour time point for MRSA and 99.38% for E. coli. Late last year we obtained the registration rights for DermSafe® in Belgium. This designation allows for the sale and/ or registration of DermSafe in most EU countries. A strategy is being developed along with a larger global strategy to bring DermSafe to the EU and Asia in 2014

(3) Obtain orphan drug status for our Netherton syndrome product. Along with our research and development of products to treat common skin conditions, we have also developed a patent pending product to treat a rare skin condition called Netherton syndrome. This disease is caused by a genetic defect which causes the skin to continually exfoliate, never forming a skin bond. This leaves the patient highly susceptible to infection and dealing with a life-long condition that has no cure.

Our product has shown excellent results in lab studies blocking the enzyme that breaks down the skin and we are seeking "Orphan Drug" designation in both the US (FDA) and Europe (EMA).

The advantages of obtaining Orphan Drug designation is that it provides various incentives including a reduction or elimination of registration and market authorization fees, protocol assistance, and seven years of market exclusivity for the product in the US and ten years in Europe. These incentives are highly attractive to pharmaceutical companies targeting this market. We are currently in discussions with potential licensees and we are implementing a case report to study the moisturization effects of the product. This process began at the beginning of 2014 and we are committed in ensuring that the product is given a proper opportunity to be effective. The initial outcome of the case study was unfortunately invalidated and we are seeking alternatives in order to generate additional data to substantiate our claims for orphan drug status. If successful, this will add greatly to our request for Orphan Drug designation with both the

EMA and FDA. There can be no assurances that our project will be successful.

Seek clinical partnerships which will result in FDA approvals of our prescription products. There are three “Phases” involved in obtaining FDA approval. The completion of Phase 1 and/or Phase 2 will increase the value of the (4) license and royalty fees of our products significantly. We are also seeking partnerships with Clinical Research Organizations (CROs) in order to define and begin the regulatory pathway for one or more of our prescription products.

3. **Secure Additional Licensees:**

We are in discussions and undergoing internal studies with various global pharmaceutical and Consumer Goods companies for licenses.

We continue to focus on completing the (confidential) development projects that are in progress, continuing our negotiations with multi-national companies on a number of potential OTC product license agreements and advancing discussions with interested international clients regarding joint ventures.

To facilitate further expansion, we have entered discussions with potential partners that are dermatology service providers and knowledgeable and connected in the dermatology market.

Table of Contents

4. New Products Available for Exclusive Licensing:

During the past few months Skinvisible has developed a new sunless tanning mousse / foam which uses a unique foam with Invisicare®, developed specifically for its foaming properties. This adds to Skinvisible’s line of sunless tanning products which includes sunless tanning lotions (light, medium and dark), pre-sun moisturizer and after-sun moisturizer along with sunless tanning spray products for commercial use. The addition of a sunless tanning mousse enhances this line of products which are available for exclusive licensing.

Skinvisible completed its first production of its patented broad spectrum sunscreen. Skinvisible’s sunscreens with SPF 15, 30 and 50 (the highest SPF allowed by the FDA) are formulated with Avobenzone, the only UVA sun filter allowed under the US FDA monograph. This UVA/UVB sunscreen was granted a patent from the United States patent office last year. Avobenzone is known for breaking down in the sun after only two hours – thus the requirement to reapply every 2 hours. Skinvisible’s patent was granted based on Invisicare’s® minimum 8 hour photo stability. For countries outside the United States, Skinvisible has additionally patented UVA/UVB sunscreens formulated with Tinosorb S that are available for licensing.

In March 25, 2014, we announced that we have successfully completed independent testing to validate our broad spectrum sunscreen claims according to new labeling guidelines by the FDA which are designed to help reduce the incidents of skin cancer in the U.S.

Skinvisible sunscreens can be labeled with the following claims:

Claim # 1 – Broad-Spectrum: According to the FDA, in order for a sunscreen to be labeled “broad spectrum” it must prove it protects against both UVA and UVB rays by having an SPF (Sun Protection Factor) of at least 15 and a critical wave length of at least 370 nm. Skinvisible’s sunscreen has surpassed both of these criteria, allowing Skinvisible’s broad spectrum sunscreen label to also state “prevents sunburn, skin cancer and aging due to the sun.”

Claim # 2 – Water-Resistant 80 Minutes: The FDA sunscreen water resistant claim requires that a sunscreen must have the same SPF after being in water or sweating for 40 or 80 minutes. Skinvisible’s testing was conducted at an independent laboratory specializing in sunscreen testing. The test involved human subjects that applied sunscreen to their arm, followed by the immersion of the arm into a Jacuzzi for 80 minutes (10 minutes in / 10 minutes out). Skinvisible’s sunscreen successfully completed this testing and is allowed to use “Water-resistant for 80 Minutes” on its sunscreen label, the longest length of time allowed by the FDA.

Claim # 3 – Unique Patented Technology / Eight-Hour Photostability: As previously announced, Skinvisible was granted a patent from the United States Patent and Trademark Office entitled “Sunscreen Composition with Enhanced UVA Absorber Stability and Methods”, which provides protection until November 2029. Skinvisible successfully formulated a unique Invisicare® delivery system specifically for stabilizing avobenzone; the key sunscreen used in the USA. Data submitted to the US patent office proved that Skinvisible’s sunscreen provides a minimum of eight hours of photostability.

Results of Operations for the Three Months Ended March 31, 2014 and 2013

Revenues

Our revenue from product sales, royalties on patent licenses and license fees for the three months ended March 31, 2014 was \$10,038, a decrease from \$27,287 for the same period ended March 31, 2013. The decrease was mainly due to a decrease in product sales.

Cost of Revenues

Our cost of revenues for the three months ended March 31, 2014 increased to \$2,321 from the prior year period when cost of revenues was \$158.

Gross Profit

Gross profit for the three months ended March 31, 2014 was \$7,717, or approximately 76% of sales. Gross profit for three months ended March 31, 2013 was \$27,129, or approximately 99% of sales.

Table of Contents

Operating Expenses

Operating expenses increased to \$181,957 for the three months ended March 31, 2014 from \$159,653 for the same period ended March 31, 2013. Our operating expenses for the three months ended March 31, 2014 consisted mainly of salaries and wages of \$34,315, accrued salaries and wages of \$59,418, accounting and audit expenses of \$10,800, depreciation and amortization expenses of \$8,846, rent of \$10,367, consulting fees of \$10,200 and research and development of \$9,707. In comparison, our operating expenses for the three months ended March 31, 2013 consisted mainly of salaries and wages of \$33,408, accrued salaries and wages of \$32,985, consulting expenses of \$12,100, rent of \$12,280, depreciation and amortization expenses of \$8,135, stock-based compensation of \$9,800, and edgarization fees of \$6,898.

Other Expenses

We had other expenses of \$171,009 for the three months ended March 31, 2014, compared with other expenses of \$164,323 for the three months ended March 31, 2013. This was largely the result of \$169,334 we paid in interest expenses for the three months ended March 31, 2014 from \$165,023 in the prior period ended March 31, 2013.

Net Loss

We recorded a net loss of \$345,249 for the three months ended March 31, 2014, as compared with a net loss of \$296,847 for the three months ended March 31, 2013.

Liquidity and Capital Resources

As of March 31, 2014, we had total current assets of \$316,738 and total assets in the amount of \$687,984. Our total current liabilities as of March 31, 2014 were \$3,765,219. We had a working capital deficit of \$3,405,780 as of March 31, 2014.

Operating activities used \$179,312 in cash for the three months ended March 31, 2014. Our net loss of \$345,249 was the main component of our negative operating cash flow, offset mainly by amortization of debt discount of \$80,742, an increase of accrued interest of \$26,499 and an increase in accounts payable and accrued liabilities of \$55,063.

Cash flows used by investing activities during the three months ended March 31, 2014 was \$12,448 as a result of the purchase of fixed and intangible assets.

Cash flows used by financing activities during the three months ended March 31, 2014 amounted to \$4,922 and consisted mainly of \$5,000 in payments on convertible promissory notes.

During the year ended December 31, 2013, we executed Promissory Notes (the “Notes”) in the aggregate principal amount of \$1,000,000 to several investors. The proceeds of the Notes are to be used for our general working capital purposes. The Notes bear interest at the rate of 9% per annum and mature at various times from May to August of 2015. We received net proceeds of \$900,000 from the Notes. We expect to service the Notes with income derived from license fees. If we are unable to generate revenue from license fees, we would be unable to service the Notes in the long term or we would be forced to try and renegotiate terms with the lenders. There can be no assurance, however, that we would be able to renegotiate any terms with the lenders. For the short term, however, we have sufficient capital available to service the Notes.

The Notes are secured by US Patent rights granted for our Sunscreen Products: US patent number #8,128,913: “Sunscreen Composition with Enhanced UV-A Absorber Stability and Methods.”

Based upon our current financial condition, we do not have sufficient cash to operate our business at the current level for the next twelve months. We intend to fund operations through increased sales and debt and/or equity financing arrangements, which may be insufficient to fund expenditures or other cash requirements. We plan to seek additional financing in a private equity offering to secure funding for operations. There can be no assurance that we will be successful in raising additional funding. If we are not able to secure additional funding, the implementation of our business plan will be impaired. There can be no assurance that such additional financing will be available to us on acceptable terms or at all.

Table of Contents

Off Balance Sheet Arrangements

As of March 31, 2014, there were no off balance sheet arrangements.

Critical Accounting Policies

In December 2001, the SEC requested that all registrants list their most “critical accounting policies” in the Management Discussion and Analysis. The SEC indicated that a “critical accounting policy” is one which is both important to the portrayal of a company’s financial condition and results, and requires management’s most difficult, subjective or complex judgments, often as a result of the need to make estimates about the effect of matters that are inherently uncertain.

Going concern – The accompanying financial statements have been prepared on a going concern basis, which contemplates the realization of assets and the satisfaction of liabilities in the normal course of business. We have incurred cumulative net losses of \$24,260,057 since our inception and require capital for our contemplated operational and marketing activities to take place. Our ability to raise additional capital through the future issuances of common stock is unknown. The obtainment of additional financing, the successful development of our contemplated plan of operations, and our transition, ultimately, to the attainment of profitable operations are necessary for us to continue operations. The ability to successfully resolve these factors raise substantial doubt about our ability to continue as a going concern. These consolidated financial statements do not include any adjustments that may result from the outcome of these aforementioned uncertainties.

Product sales – Revenues from the sale of products (Invisicare® polymers) are recognized when title to the products are transferred to the customer and only when no further contingencies or material performance obligations are warranted, and thereby have earned the right to receive reasonably assured payments for products sold and delivered.

Royalty sales – We also recognize royalty revenue from licensing our patented product formulations only when earned, with no further contingencies or material performance obligations are warranted, and thereby have earned the right to receive and retain reasonably assured payments.

Distribution and license rights sales – We also recognize revenue from distribution and license rights only when earned (and are amortized over a five year period), with no further contingencies or material performance obligations are warranted, and thereby have earned the right to receive and retain reasonably assured payments.

Costs of Revenue – Cost of revenue includes raw materials, component parts, and shipping supplies. Shipping and handling costs is not a significant portion of the cost of revenue.

Accounts Receivable – Accounts receivable is comprised of uncollateralized customer obligations due under normal trade terms requiring payment within 30 days from the invoice date. The carrying amount of accounts receivable is reviewed periodically for collectability. If management determines that collection is unlikely, an allowance that reflects management’s best estimate of the amounts that will not be collected is recorded. Management reviews each accounts receivable balance that exceeds 30 days from the invoice date and, based on an assessment of creditworthiness, estimates the portion, if any, of the balance that will not be collected. As of March 31, 2014, we had not recorded a reserve for doubtful accounts.

Table of Contents

Recently Issued Accounting Pronouncements

We do not expect the adoption of recently issued accounting pronouncements to have a significant impact on our results of operations, financial position or cash flow.

Item 3. Quantitative and Qualitative Disclosures About Market Risk

A smaller reporting company is not required to provide the information required by this Item.

Item 4. Controls and Procedures

Disclosure Controls and Procedures

We carried out an evaluation of the effectiveness of the design and operation of our disclosure controls and procedures (as defined in Exchange Act Rules 13a-15(e) and 15d-15(e)) as of March 31, 2014. This evaluation was carried out under the supervision and with the participation of our Chief Executive Officer and our Chief Financial Officer. Based upon that evaluation, our Chief Executive Officer and Chief Financial Officer concluded that, as of March 31, 2014, our disclosure controls and procedures were not effective due to the presence of material weaknesses in internal control over financial reporting.

A material weakness is a deficiency, or a combination of deficiencies, in internal control over financial reporting, such that there is a reasonable possibility that a material misstatement of the company's annual or interim financial statements will not be prevented or detected on a timely basis. Management has identified the following material weaknesses which have caused management to conclude that, as of March 31, 2014, our disclosure controls and procedures were not effective: (i) inadequate segregation of duties and effective risk assessment; and (ii) insufficient written policies and procedures for accounting and financial reporting with respect to the requirements and application of both US GAAP and SEC guidelines.

Remediation Plan to Address the Material Weaknesses in Internal Control over Financial Reporting

Our company plans to take steps to enhance and improve the design of our internal controls over financial reporting. During the period covered by this quarterly report on Form 10-Q, we have not been able to remediate the material weaknesses identified above. To remediate such weaknesses, we plan to implement the following changes during our fiscal year ending December 31, 2014: (i) appoint additional qualified personnel to address inadequate segregation of duties and ineffective risk management; and (ii) adopt sufficient written policies and procedures for accounting and financial reporting. The remediation efforts set out are largely dependent upon our securing additional financing to cover the costs of implementing the changes required. If we are unsuccessful in securing such funds, remediation efforts may be adversely affected in a material manner.

Changes in Internal Control over Financial Reporting

There were no changes in our internal control over financial reporting during the three months ended March 31, 2014 that have materially affected, or are reasonable likely to materially affect, our internal control over financial reporting.

Table of Contents

PART II – OTHER INFORMATION

Item 1. Legal Proceedings

We are not a party to any pending legal proceeding. We are not aware of any pending legal proceeding to which any of our officers, directors, or any beneficial holders of 5% or more of our voting securities are adverse to us or have a material interest adverse to us.

Item 1A: Risk Factors

A smaller reporting company is not required to provide the information required by this Item.

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

The information set forth below relates to our issuances of securities without registration under the Securities Act of 1933 during the reporting period which were not previously included in a Quarterly Report on Form 10-Q or Current Report on Form 8-K.

On January 7, 2014, we issued 180,000 shares to settle \$3,600 in outstanding accounts payable.

These securities were issued pursuant to Section 4(2) of the Securities Act and/or Rule 506 promulgated thereunder. The holders represented their intention to acquire the securities for investment only and not with a view towards distribution. The investors were given adequate information about us to make an informed investment decision. We did not engage in any general solicitation or advertising. We directed our transfer agent to issue the stock certificates with the appropriate restrictive legend affixed to the restricted stock.

Item 3. Defaults upon Senior Securities

None

Item 4. Mine Safety Disclosures

Not applicable.

Item 5. Other Information

None

Item 6. Exhibits

<u>Exhibit Number</u>	<u>Description of Exhibit</u>
31.1	<u>Certification of Chief Executive Officer pursuant to 18 U.S.C. Section 1350, as adopted pursuant to Section 302 of the Sarbanes-Oxley Act of 2002</u>
31.2	<u>Certification of Chief Financial Officer pursuant to 18 U.S.C. Section 1350, as adopted pursuant to Section 302 of the Sarbanes-Oxley Act of 2002</u>
32.1	<u>Certification of Chief Executive Officer and Chief Financial Officer pursuant to 18 U.S.C. Section 1350, as adopted pursuant to Section 906 of the Sarbanes-Oxley Act of 2002</u>
101**	The following materials from the Company's Quarterly Report on Form 10-Q for the quarter ended March 31, 2014 formatted in Extensible Business Reporting Language (XBRL).

**Provided herewith

12

Table of Contents

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.

Skinvisible, Inc.

Date: May 12, 2014

By: /s/ Terry Howlett

Terry Howlett

Title: Chief Executive Officer, Chief Financial Officer and Director

